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Ms. Jennifer Tufts
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Dear Mr. Mullins and Ms. Tufts:

TRANSMITTAL OF REPLACEMENT PAGES FOR METHODS FOR CONDUCTING RISK ASSESSMENTS AND RISK EVALUATIONS AT THE PADUCAH GASEOUS DIFFUSION PLANT, PADUCAH, KENTUCKY, VOLUME 1. HUMAN HEALTH, (DOE/LX/07-0107&D2/R4/V1)

Please find enclosed replacement pages for *Methods for Conducting Risk Assessments and Risk Evaluations at the Paducah Gaseous Diffusion Plant, Paducah, Kentucky, Volume 1. Human Health*, DOE/LX/07-0107&D2/R4/V1. These replacement pages for the Risk Methods Document for the Paducah Gaseous Diffusion Plant update the July 2014 version of the document numbered DOE/LX/07-0107&D2/R3/V1.

This change addresses the Kentucky Department for Environmental Protection's requirement for their granting of approval of the 2014 revision of the document.

The U.S. Department of Energy requests approval of the subject document no later than September 30, 2014.

If you have any questions or require additional information, please contact Rich Bonczek at (859) 219-4051.

Sincerely,

Jennifer Woodard
Paducah Site Lead

Portsmouth/Paducah Project Office

Enclosures:

1. Replacement Pages for *Methods for Conducting Risk Assessments and Risk Evaluations at the Paducah Gaseous Diffusion Plant, Paducah, Kentucky, Volume 1. Human Health, DOE/LX/07-0107&D2/R4/V1* (Clean)
2. Replacement Pages for *Methods for Conducting Risk Assessments and Risk Evaluations at the Paducah Gaseous Diffusion Plant, Paducah, Kentucky, Volume 1. Human Health, DOE/LX/07-0107&D2/R4/V1* (Redline)

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DOE/LX/07-0107&D2/R4/V1

**Methods for Conducting Risk Assessments
and Risk Evaluations
at the Paducah Gaseous Diffusion Plant
Paducah, Kentucky
Volume 1. Human Health**



CLEARED FOR PUBLIC RELEASE

**Methods for Conducting Risk Assessments
and Risk Evaluations
at the Paducah Gaseous Diffusion Plant
Paducah, Kentucky
Volume 1. Human Health**

Date Issued—September 2014

Prepared for the
U.S. DEPARTMENT OF ENERGY
Office of Environmental Management

Managed by
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Paducah Gaseous Diffusion Plant
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PREFACE

This *Methods for Conducting Human Health Risk Assessments and Risk Evaluations at the Paducah Gaseous Diffusion Plant, Paducah Kentucky*, DOE/LX/07-0107&D2/R4/V1 (previous versions issued as DOE/LX/07-0107&D2/R2/V1, DOE/LX/07-0107&D2/R1/V1, and DOE/OR/07-1506&D1/V1/R1), was prepared in accordance with the requirements under both the Comprehensive Environmental Response, Compensation, and Liability Act (CERCLA) and the Resource Conservation and Recovery Act (RCRA). This document is not meant to be prescriptive, rather it is meant to provide guidance for the completion of risk analyses beyond the guidance found in the most recent revision of *Site Management Plan, Paducah Gaseous Diffusion Plant, Paducah, Kentucky* (DOE 2014). Specifically, this document integrates results of comment resolution meetings and technical meetings between the regulatory agencies and the U.S. Department of Energy and provisions in the Federal Facility Agreement (FFA) for the Paducah Gaseous Diffusion Plant (PGDP) and provides methods that should be followed when completing risk analyses to ensure consistency in risk analyses. Risk analyses considered in this document are human health risk assessments and risk evaluations prepared for both informal and formal reports. This document and its appendices, including preliminary remediation goal values, are for use at PGDP and are not applicable to other sites within the Commonwealth of Kentucky.

In accordance with Section IV of the FFA for PGDP, this integrated technical document was developed to satisfy both CERCLA and RCRA corrective action requirements. The phases of the investigation process are referenced by CERCLA terminology within this document to reduce the potential for confusion.

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ACRONYMS

AF	adherence factor
ALM	Adult Lead Model
AOC	area of concern
ARAR	applicable or relevant and appropriate requirement
AT123D	Analytical Transient 1-, 2-, 3-Dimensional Simulation of Waste Transport in the Aquifer System
bgs	below ground surface
CAS	Chemical Abstracts Service
CDI	chronic daily intake
CERCLA	Comprehensive Environmental Response, Compensation, and Liability Act
COC	contaminant of concern
COPC	chemical of potential concern
DAF	dilution attenuation factor
DOE	U.S. Department of Energy
DQA	data quality assessment
DQO	data quality objective
ED	exposure duration
EE/CA	Engineering Evaluation/Cost Analysis
EF	exposure frequency
ELCR	excess lifetime cancer risk
EPA	U.S. Environmental Protection Agency
EPC	exposure point concentration
FA	fraction absorbed
FFA	Federal Facility Agreement
FS	feasibility study
GI	gastrointestinal
HEAST	Health Effects Assessment Summary Tables
HHRAWG	Human Health Risk Assessment Working Group
HI	hazard index
HQ	hazard quotient
IEUBK	Integrated Exposure Uptake and Biokinetic
IRIS	Integrated Risk Information System
IUR	inhalation unit risk
K_d	adsorption coefficient/distribution coefficient
K_p	permeability coefficient
KDEP	Kentucky Department for Environmental Protection
KYRHB	Kentucky Radiation Health Branch
MARLAP	Multi-Agency Radiological Laboratory Analytical Protocols
MCL	maximum contaminant level
MOC	medium of concern
MQC	minimum quantification concentration
MQO	measurement quality objective
MDC	minimum detectable concentration
MUSLE	Modified Universal Soil Loss Equation
OSWER	Office of Solid Waste and Emergency Response
PA	preliminary assessment
PAH	polycyclic aromatic hydrocarbon
PCB	polychlorinated biphenyl

POC	pathway of concern
PGDP	Paducah Gaseous Diffusion Plant
PRA	probabilistic risk assessment
PRG	preliminary remediation goal
ProUCL	EPA's Upper Confidence Limit Software
RAGS	Risk Assessment Guidance for Superfund
RAO	remedial action objective
RAWG	Risk Assessment Working Group
RCRA	Resource Conservation and Recovery Act
RESRAD	residual radioactivity
RfC	reference concentration
RfD	reference dose
RG	remedial goal
RGA	Regional Gravel Aquifer
RGO	remedial goal option
RI	remedial investigation
RME	reasonable maximum exposure
ROD	record of decision
SADA	Spatial Analysis and Decision Assistance
SESOIL	Seasonal Soil Model
SF	slope factor
SI	site investigation
SMP	Site Management Plan
SQL	sample quantitation limit
SSL	soil screening level
SVOC	semivolatile organic compound
SWMU	solid waste management unit
SWMM	Storm Water Management Model
TCDD	2,3,7,8-tetrachlorodibenzo-p-dioxin
TCE	trichloroethene
TEF	toxicity equivalence factor
TEQ	toxicity equivalents
UCL	upper confidence limit
UCRS	Upper Continental Recharge System
UTL	upper tolerance limit
XRF	X-ray fluorescence

EXECUTIVE SUMMARY

This document describes the methods used to prepare the human health risk assessments and risk evaluations needed to complete remedial activities at the Paducah Gaseous Diffusion Plant (PGDP). This document is not meant to be prescriptive, rather it is meant to provide the framework to complete appropriate risk analyses for projects listed in the Paducah Site Management Plan (DOE 2014) taking into account site-specific conditions at PGDP. The materials and methods presented in this document were developed following agreements reached between the U.S. Department of Energy (DOE) and the regulatory agencies during comment resolution meetings, in the Federal Facility Agreement, and at technical meetings. In this document, the human health risk analyses that will occur during each phase of remedial activities are discussed, analytical techniques are described, and several analytical tools are presented. By providing this material in a single document, consistency of human health risk assessments and evaluations performed for PGDP is ensured, thereby speeding the completion and review of risk assessments and risk evaluations. This document and its appendices, including preliminary remediation goal values, are for use at PGDP and are not applicable to other sites within the Commonwealth of Kentucky. Any endorsement of this document by Commonwealth agencies is limited to its use at PGDP.

This document also discusses some of the methods used to complete dose assessments at PGDP. Dose assessments are conducted to provide information for risk managers and are separate from the risk assessment conducted for decision making. The methods for dose assessment are presented generally, and additional discussion should be held with regulatory agencies prior to initiating any dose assessment project.

This document was prepared by the PGDP Risk Assessment Working Group (RAWG). The RAWG is a multiagency, multidisciplinary group tasked with meeting the following goals:

- Produce tools that can be used to prioritize remedial activities at the PGDP.
- Develop methods to complete risk evaluations for the PGDP.
- Make the results of the risk assessments and evaluations at the PGDP more useful to risk managers.
- Enhance risk communication between the producers of risk assessments and risk evaluations and the users of this information (e.g., risk managers).

Organizations participating in the production of this document and their affiliations are DOE, U.S. Environmental Protection Agency, Commonwealth of Kentucky Division of Waste Management, and Commonwealth of Kentucky Radiation Health Branch.

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1. INTRODUCTION

The purpose of this document is to present the methods and approaches used for screening level, baseline human health, and residual risk assessments and risk evaluations at the Paducah Gaseous Diffusion Plant (PGDP) and provide resources [such as preliminary remediation goals (PRGs) and dose-based concentrations] for completing those assessments. This document is not meant to be prescriptive, rather it is meant to provide the framework to complete appropriate risk analyses for projects listed in the Paducah Site Management Plan (DOE 2014) taking into account site-specific conditions at PGDP. This document is not intended to replace or modify guidance from the U.S. Environmental Protection Agency (EPA), guidance from the Commonwealth of Kentucky, or any of the tripartite agreements. Analyses of risks and hazards presented by environmental contamination at PGDP are integral to the Federal Facility Agreement's (FFA) primary objective of implementing remedies that minimize, control, or eliminate risks to human health and the environment. These analyses begin during the scoping phase [e.g., during scoping meetings and during, for example, the preliminary assessment/site investigation (PA/SI)] when available environmental media and historical information are interpreted and compared with site-specific PRGs and other screening criteria to determine if action may be required at release sites and to plan the timing of that action. These analyses continue during investigation (e.g., the remedial investigation) when historical information, site-specific PRGs, and other screening criteria are used to focus the work plan on the risk-related problems that must be investigated and may need to be addressed during data collection. Subsequently, the results of the risk analyses are used in decision documents to justify why an action is or is not needed at a site.² A more streamlined approach for risk assessments is sometimes used for removal action decision documents. During the production of the decision documents, the risk analyses also are used to develop the risk-based cleanup levels used in subsequent design activities.

Several major decision points occur during the aforementioned process. These decision points often limit the scope of risk analyses performed during investigation and remedy selection, but allow for interim actions to address important environmental concerns and occur several times during the process.

Risk assessors provide information at the decision points and risk managers use that information to make decisions. Risk assessors and managers and their roles are defined as follows (EPA 1989a).

- **Risk Assessor.** An individual, team, or organization that generates site- or media-specific risk assessments for use in site-specific decision making. The assessor relies on existing databases and information [e.g., EPA Integrated Risk Information System (IRIS), health assessment documents, and program-specific toxicity information] and media- or site-specific exposure information in characterizing risk. This group also relies, in part, on regulatory agency risk assessment guidelines and program-specific guidance to address scientific policy issues and scientific uncertainties.
- **Risk Manager.** An individual, team, or organization with responsibility for or authority to take action in response to an identified risk. Risk managers *integrate* the risk characterization information provided by the risk assessor with other considerations specified in applicable statutes to make and justify regulatory decisions. Generally, risk managers include lead and regulatory agency managers and decision makers. Risk managers also play a role in determining the scope of risk assessments.

¹ Reserved.

² There may be scenarios presented pursuant to this document that might not be commensurate with the reasonable foreseeable land use but may serve as a reference point to decision makers.

This document presents the methods to be used to complete the analyses described herein. In addition, this document discusses many of the analytical tools that can be used to complete this process and discusses the sources of the tools. Materials and methods used to complete scoping activities, including the derivation of risk- and dose-based PRGs, the background concentrations of chemicals and radionuclides, and other screening criteria are in Section 2; materials and methods specific to the human health risk assessments, including work plan preparation and baseline human health risk assessment, are in Section 3, "Risk Analyses during the Remedial Investigation"; materials and methods applicable to the feasibility study (FS) risk evaluation, including cleanup level development and consideration of residual risks, are in Section 4. Dose assessments sometimes are provided to risk managers, as well, and also are discussed within these sections. The approach to dose assessments discussed here is based on EPA guidance (EPA 2000a) and is specific to PGDP. The dose-based concentrations are based on Federal Guidance Report 13 (EPA 1999a) and are not appropriate for other activities such as establishment of authorized limits. The exposure parameters used to derive the dose-based concentrations presented are useful inputs when deriving authorized limits.

2. RISK ANALYSES DURING SCOPING ACTIVITIES

Risk analyses during site scoping activities will be performed to do the following:

- Determine if site risks are so great as to require immediate action prior to Remedial Investigation (RI)/FS (i.e., interim action);³
- Determine if site risks are so low as to support a no-further-action decision;
- Prioritize the further investigation of those sites not requiring an interim action or potentially requiring no further action;
- Divide exposure setting into exposure units;⁴ and
- Provide information to be used in subsequent work plan development.

General depictions of the methods that will be followed to complete these analyses are shown in Figure 2.1. Figures 2.2, 2.3, 2.4, and 2.5 present specific issues related to the risk screening process (including issues related to dose).

Generally, analyses completed as part of risk-based site scoping will rely on simple comparisons between site contamination data to PGDP-specific PRGs, including risk-based action and no-action concentrations, dose-based concentrations (if a dose assessment is conducted), background concentrations, and pertinent applicable or relevant and appropriate requirements (ARARs). Table 2.1 shows the significant chemicals of potential concern (COPCs) at PGDP. Significant COPCs are chemicals that have been retained as contaminants of concern (COCs) (sometimes listed as constituents of concern) in prior risk assessments at PGDP. For the purposes of this document, these terms are essentially equivalent. These COPCs therefore are likely to be COPCs for other risk assessments, but the absence of a chemical from the list does not imply that it would not be a COPC at a PGDP site. Risk-based action and no-action concentrations and dose-based concentrations are provided for the significant COPCs and are presented in Tables A.1 through A.11 in Appendix A. Action and no action soil concentrations based on dose limits are derived by following EPA guidance (EPA 2000a) and are used for dose assessments at PGDP.

Table A.1 presents risk-based action concentrations for contaminants in soil and sediment; Table A.2 presents risk-based action levels for contaminants in groundwater; Table A.3 presents risk-based action levels for contaminants in surface water; Table A.4 presents risk-based no-action levels for contaminants in soil and sediment; Table A.5 presents risk-based no-action levels for contaminants in groundwater; Table A.6 presents risk-based no-action levels for contaminants in surface water; Table A.7 presents risk-based no-action levels for contaminants in soil that are protective of groundwater drawn from the Regional Gravel Aquifer (RGA) immediately adjacent to a contaminated area; Table A.8 presents dose-based levels for radionuclide contaminants in soil and sediment; Table A.9 presents dose-based levels for radionuclide contaminants in groundwater; Table A.10 presents dose-based levels for radionuclide contaminants in surface water; and Table A.11 presents dose-based levels for radionuclide contaminants in soil that are

³ The report from this point forward will use references to remedial action documents instead of removal action documents for simplicity. If the approach for removal actions differs in the subsequent discussions, these differences will be noted, as appropriate.

⁴ A default exposure unit of 0.5 acres will be used for sites inside the PGDP industrialized area. For a site outside the industrialized area, the size of the exposure unit will be decided during scoping by agreement among the three parties.

Risk Analyses during Site Scoping at PGDP General Approach

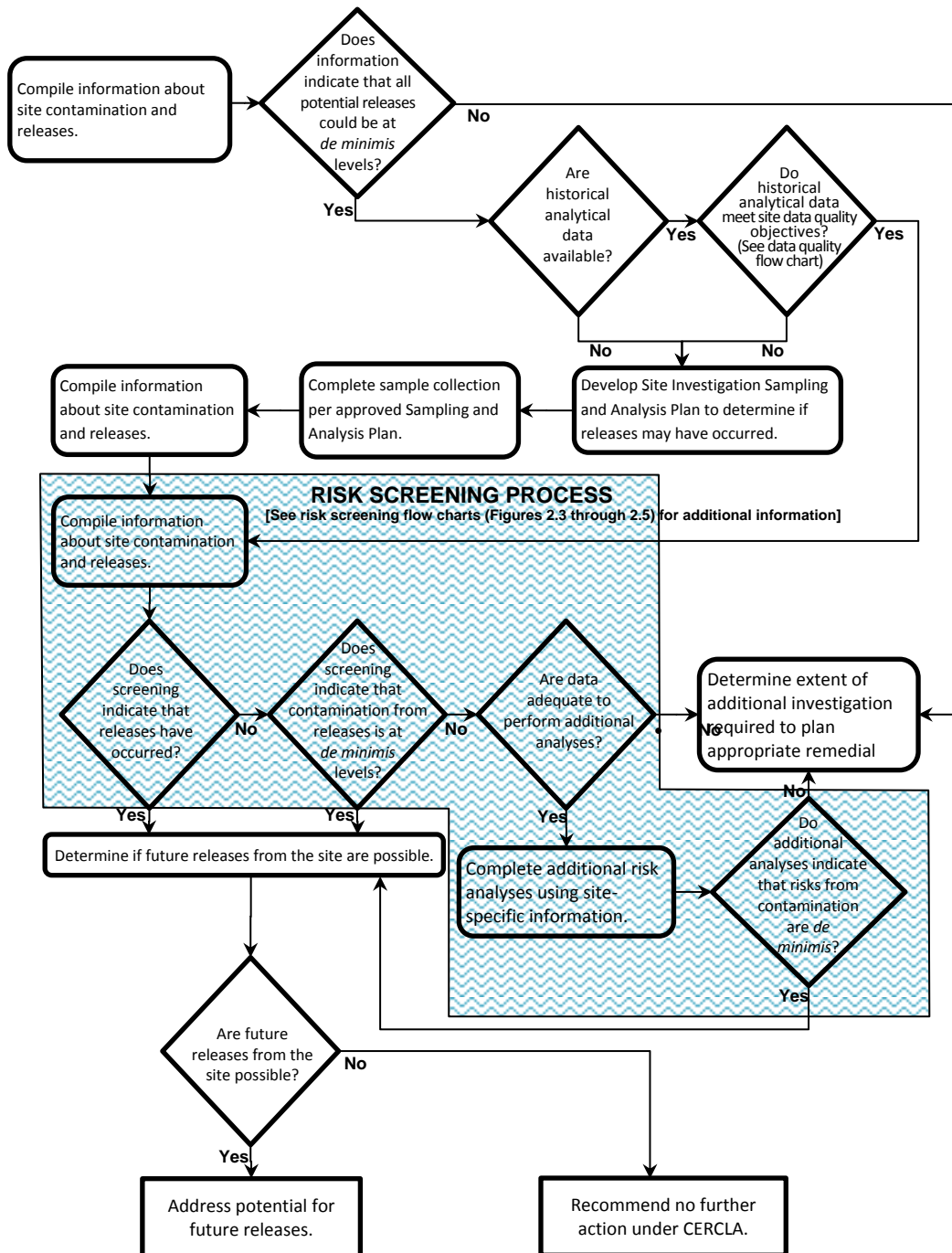
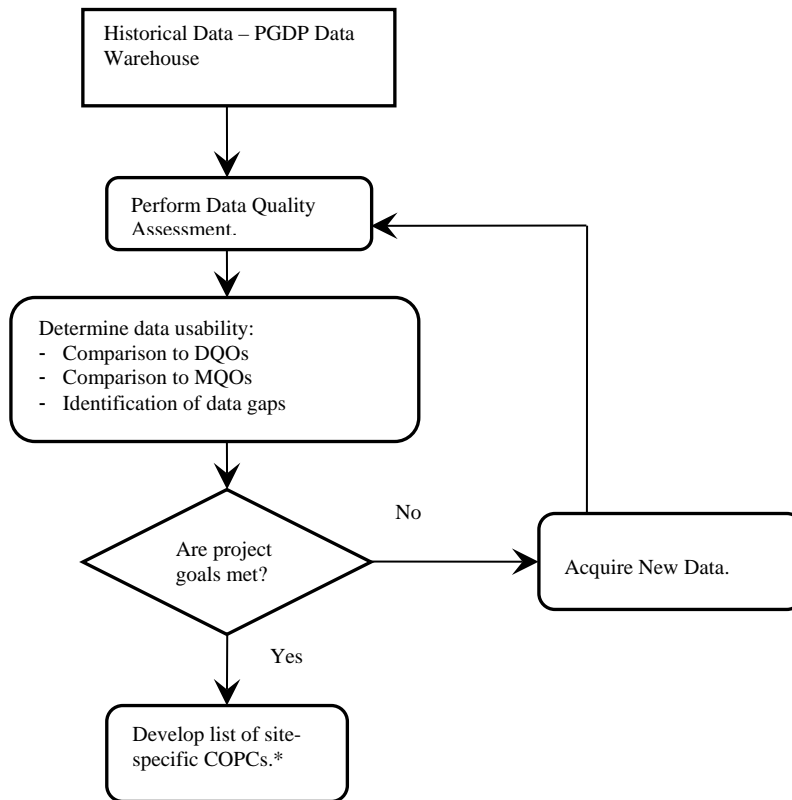


Figure 2.1. General Approach to Risk-Based Site Scoping

Risk-based Site Scoping at PGDP
Data Quality/Data Usability Review



*Identification of site-specific COPCs not currently included in Table 2.1, “Significant Chemicals of Potential Concern at PGDP,” would include the review of additional information (e.g., information identified in the RI process).

Figure 2.2. Data Quality Review to Support Risk-Based Site Scoping

Risk-based Site Scoping at PGDP Part 1 - Human Health Direct Contact Screening

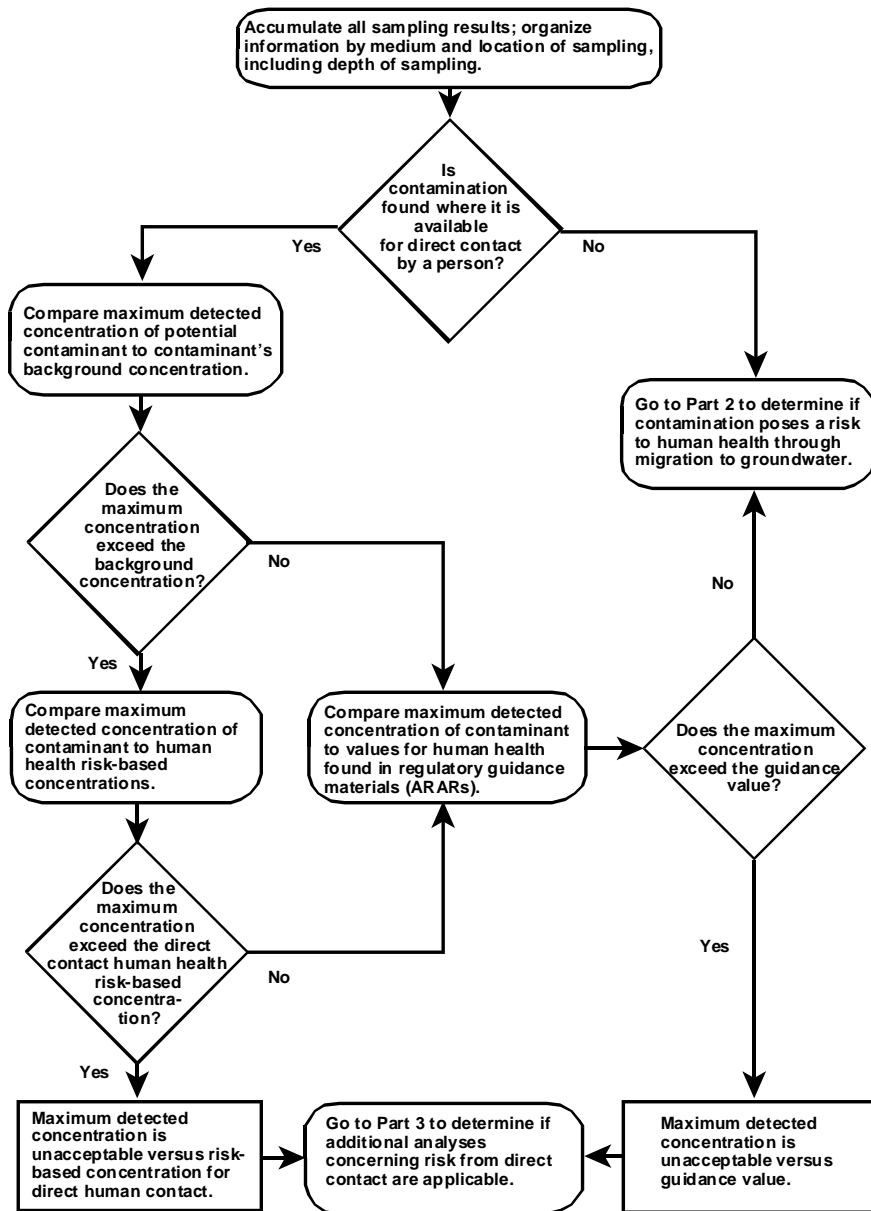


Figure 2.3. Human Health Direct Contact Screening during Risk-Based Site Scoping

**Risk-based Site Scoping at PGDP
Part 2 - Groundwater Protection Screening**

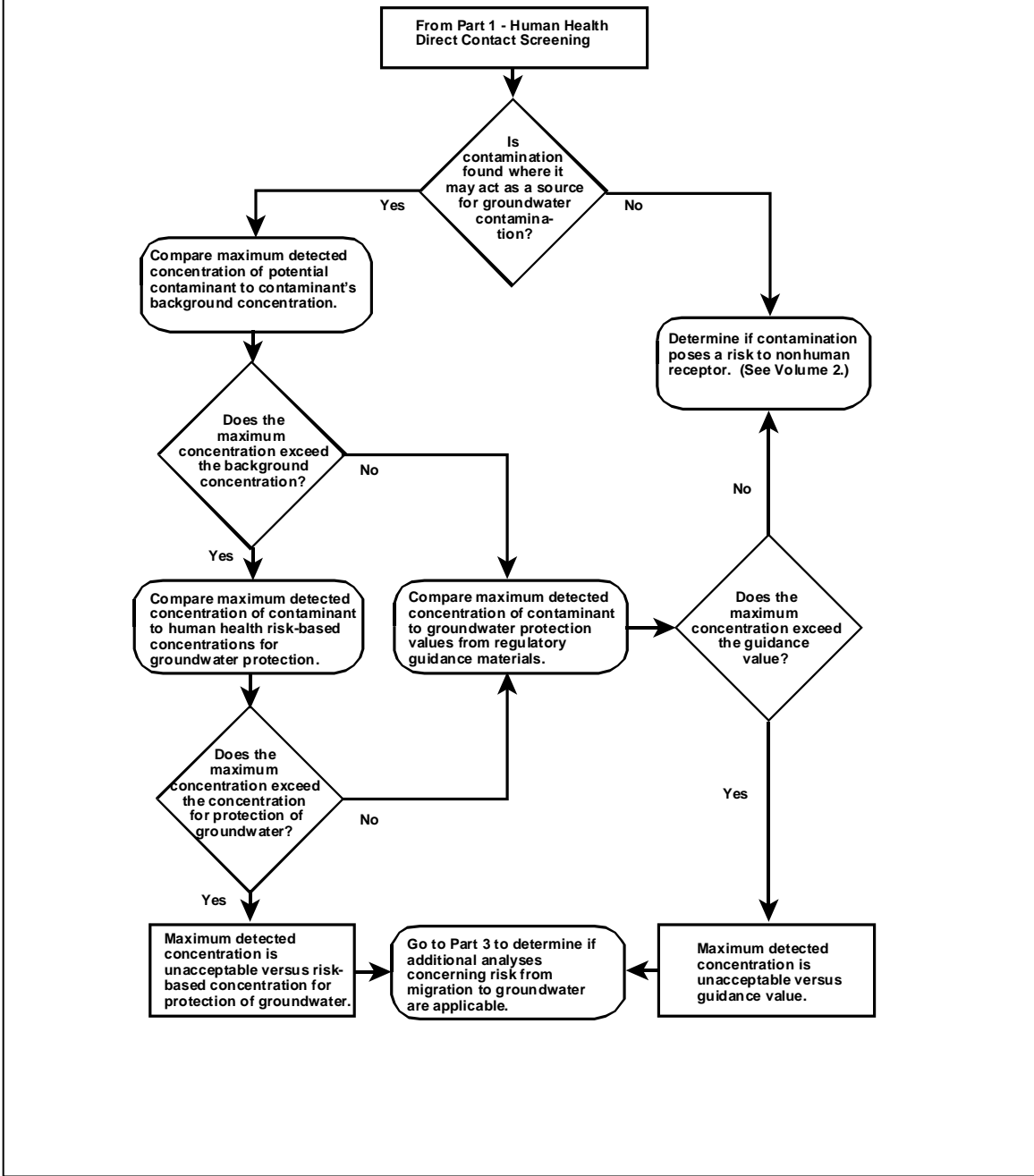


Figure 2.4. Groundwater Protection Screening during Risk-Based Site Scoping

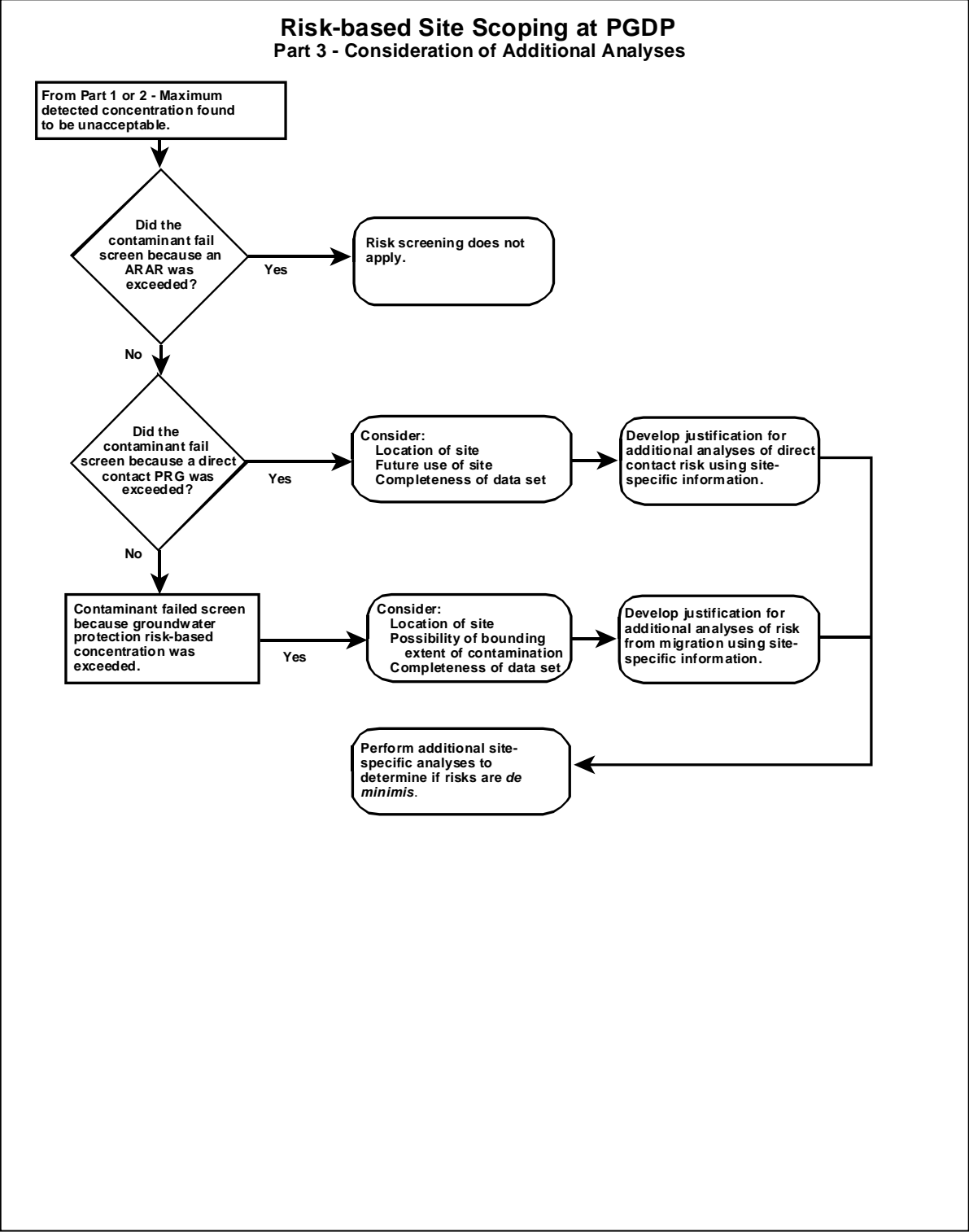


Figure 2.5. Consideration of Additional Analyses during Risk-Based Site Scoping

Table 2.1. Significant Chemicals of Potential Concern at PGDP^{1,2}

Inorganic Chemicals		Organic Compounds		Radionuclides	
Analyte	CAS Number	Analyte	CAS Number	Analyte	CAS Number
Aluminum	7429905	Acenaphthene	83329	Americium-241	14596102
Antimony	7440360	Acenaphthylene	208968	Cesium-137+D	10045973
Arsenic	7440382	Acrylonitrile	107131	Neptunium-237+D	13994202
Barium	7440393	Anthracene	120127	Plutonium-238	13981163
Beryllium	7440417	Benzene	71432	Plutonium-239	15117483
Boron	7440428	Carbazole	86748	Plutonium-240	14119336
Cadmium	7440439	Carbon tetrachloride	56235	Technetium-99	14133767
Chromium III	16065831	Chloroform	67663	Thorium-230	14269637
Chromium VI	18540299	1,1-Dichloroethene	75354	Uranium-234	13966295
Cobalt	7440484	1,2-Dichloroethene (mixed)	540590	Uranium-235+D	15117961
Copper	7440508	<i>trans</i> -1,2-Dichloroethene	156605	Uranium-238+D	7440611
Iron	7439896	<i>cis</i> -1,2-Dichloroethene	156592		
Lead	7439921	Dieldrin	60571		
Manganese	7439965	Ethylbenzene	100414		
Mercury	7439976	Fluoranthene	206440		
Molybdenum	7439987	Fluorene	86737		
Nickel	7440020	Hexachlorobenzene	118741		
Selenium	7782492	Naphthalene	91203		
Silver	7440224	2-Nitroaniline	88744		
Thallium	7440280	N-Nitroso-di-n-propylamine	621647		
Uranium	N/A	Phenanthrene	85018		
Vanadium	7440622	Pyrene	129000		
Zinc	7440666	Tetrachloroethene	127184		
		Trichloroethene	79016		
		Total Dioxins/Furans	1746016		
		2,3,7,8-HpCDD	37871004		
		2,3,7,8-HpCDF	38998753		
		2,3,7,8-HxCDD	34465468		
		2,3,7,8-HxCDF	55684941		
		OCDD	3268879		
		OCDF	39001020		
		2,3,7,8-PeCDD	36088229		
		1,2,3,7,8-PeCDF	57117416		
		2,3,4,7,8-PeCDF	57117314		
		2,3,7,8-TCDD	1746016		
		2,3,7,8-TCDF	5127319		
		Total PAHs	50328		
		Benz(a)anthracene	56553		
		Benzo(a)pyrene	50328		
		Benzo(b)fluoranthene	205992		
		Benzo(k)fluoranthene	207089		
		Chrysene	218019		
		Dibenz(a,h)anthracene	53703		
		Indeno(1,2,3-cd)pyrene	193395		
		Total PCBs	1336363		
		Aroclor 1016	12674112		
		Aroclor 1221	11104282		
		Aroclor 1232	11141165		
		Aroclor 1242	53469219		
		Aroclor 1248	12672296		
		Aroclor 1254	11097691		
		Aroclor 1260	11096825		
		Vinyl chloride	75014		
		Xylenes (Mixture)	1330207		
		p-Xylene	106423		
		m-Xylene	108383		
		o-Xylene	95476		

¹ This list of chemicals, compounds, and radionuclides was compiled from COPCs retained as COCs in baseline risk assessments performed at PGDP between 1990 and 2008 (i.e., DOE 1996a; DOE 1996b; DOE 1999a; DOE 1999b; DOE 2005; and DOE 2008).

² List may be added to during project scoping based on additional information.

CAS = Chemical Abstract Service

protective of groundwater drawn from the RGA immediately adjacent to a contaminated area. Methods used to develop the risk-based and dose-based screening values are presented in Appendix B of this document.

A comparison of analyte concentrations detected in soil and groundwater samples to analyte concentrations detected in background samples will be performed as part of the development of the list of COPCs as shown in Figures 2.3 and 2.4. The values used to represent background are presented in Appendix A. Appendix E also contains a discussion of the derivation of the background values. Only surface soil [0–1 ft below ground surface (bgs)] and subsurface soil (1–16 ft bgs) and groundwater drawn from the RGA and McNairy Formation will be included in comparison with background concentrations because background values are available only for these media at PGDP (DOE 2000). The RGA is the lateral flow system that constitutes the shallow Class II groundwater aquifer beneath PGDP and contiguous lands to the north. The McNairy formation flow system is below the RGA.

Background concentrations for chemicals and radionuclides in soil and RGA and McNairy Formation groundwater to be used during site-scoping activities are presented in Tables A.12 and A.13, respectively. In the background screen for soil and groundwater, the maximum detected concentration of the COPCs will be compared to the values presented in Tables A.12 and A.13. Analytes for which the maximum detected concentrations [or maximum activity for radionuclides with reported values greater than their minimum detectable concentration (MDC)] is less than background will be removed from the data set used in the risk assessment. The background values for soil presented in Table A.12 represent upper tolerance limits (UTLs) of background except as noted in the table footnotes. Additional comparisons of the maximum detected concentration or maximum activity for radionuclides with the range of background values also may be conducted in the uncertainty section of the risk assessment (discussed in Section 3.3.7) to further evaluate if a COPC represents a site contaminant. The maximum detected concentrations or activity for radionuclides for all detected analytes with background values will be included in the prepared summary appendix used for screening against background. Because surface water and sediment are transient media in which concentrations and activities can change rapidly, PGDP does not plan to develop surface water and sediment background. Currently, a comparison of the full range of concentrations and activities in upstream versus downstream samples is to be used to determine if a unit or area is releasing contaminants to the environment. Additionally, as part of the analysis, the data adequacy at both the upgradient location and potentially contaminated site must be considered.

To perform the screening analyses during site scoping, available data must be deemed sufficient to determine the potential contamination at a site. Data used during site scoping will be evaluated using the systematic approach presented in Figure 2.2 to ensure that risk analyses employ data of known quality and that the appropriate quantities and types of data are acquired. This systematic approach also is used to evaluate data during remedial investigation, as discussed in Section 3. Detailed discussions related to data quality/data usability review are provided in Section 3.3.3.1.

In presenting the results of risk-based site scoping analyses, several tables should be prepared using a format that allows for easy identification of those chemicals, compounds, and radioisotopes with the potential to contribute to unacceptable levels of risk. If a dose analysis is conducted, similar tables should be prepared to present the results of the dose-based site scoping analysis. To complete the risk-based screening analyses for site scoping, tables will be prepared for soil and sediment, groundwater, and surface water screening. For soil and sediment, up to four tables will be prepared using the risk-based screening levels. These tables offer comparisons among the following:

- Maximum detected concentrations and action levels,
- Maximum detected concentrations and no-action levels,
- Maximum detected concentrations and levels deemed protective of groundwater, and
- Maximum detected concentrations and established background values for naturally occurring inorganics and radionuclides.

For both groundwater and surface water, two tables will be prepared using the risk-based screening levels. These tables offer comparisons between the following:

- Maximum detected concentrations and action levels and
- Maximum detected concentration and no-action levels.

In addition, summary tables providing the following information will be prepared for each medium:

- Lists of chemicals and radionuclides analyzed for but never detected;
- A presentation of summary statistics, including a comparison of detected analytes with background;
- Lists of sampling stations that contain a contaminant at a concentration greater than the action screening level; and
- Lists of sampling stations that contain a contaminant at a concentration greater than the no-action screening level.

2.1 ANALYSES SUPPORTING ACTION PRIOR TO RI/FS

As discussed in the FFA, interim actions are required at those sites that pose an imminent risk or hazard to human health and the environment. Generally, sites requiring an interim remedial or removal action are those at which contamination with a single or small number of analytes presents a total carcinogenic risk greater than 1×10^{-4} or a systemic toxicity value (i.e., hazard index or HI) greater than one and for which the risk analyses indicate that exposure is occurring under current use patterns. For these sites, the screening risk analyses will be limited to that described here because additional analyses will slow response time; however, to complete later decision documents, estimates of cumulative risk will be developed. [Note: The exact decision point for interim action is a project-specific decision. The values included here are for illustration only. For example, it is possible that a site is a yard that contains source material that might present a principal threat. At such sites, the scoping analyses may not include a risk-based screen. Additionally, note that risks posed to nonhuman receptors (e.g., ecological risk) may call for an interim remedial or removal action even when risks to humans are negligible.] To derive these estimates of cumulative risk, the methods in Equations 1, 2, 3, and 4 will be used. [Methods to derive dose estimates are similar and are not presented. Also, note that for a dose assessment, the benchmarks for dose-based action are 1 mrem/year, 4 mrem/year (for water only), 15 mrem/year, 25 mrem/year, and 100 mrem/year.]⁵

⁵ The radiation dose rates of 1 mrem/year and 15 mrem/year are not DOE or Kentucky standards, and none of these radiation dose rates are EPA standards, including the 15 mrem/year, with the exception of the 4 mrem/year that is the public drinking water standard for beta-emitting radionuclides. The 100 mrem/year dose rate relates to DOE Order 458.1 and the Kentucky public dose limit as established in 902 KAR 100:019, Section 10.

$$\text{Analyte-specific Risk} = \frac{\text{MAX}}{\text{Cancer PRG}} \times \text{Target Risk} \quad \text{[Eq. 1]}$$

where: MAX = Maximum detected concentration in a medium.
Cancer PRG = The medium-specific risk-based no-action screening value for the analyte.
Target Risk = The target risk upon which the risk-based PRG calculation was based (1×10^{-6}).

$$\text{Total Risk} = \sum \text{Analyte-specific Risks} \quad \text{[Eq. 2]}$$

where: Analyte-specific risk is the result from Eq. 1.

$$\text{Analyte-specific Hazard} = \frac{\text{MAX}}{\text{Hazard PRG}} \times \text{Target Hazard} \quad \text{[Eq. 3]}$$

where: MAX = Maximum detected concentration in a medium.
Hazard PRG = The medium-specific risk-based no-action screening value for the analyte.
Target Hazard = The target hazard upon which the risk-based PRG calculation was based (0.1).

$$\text{Total Hazard} = \sum \text{Analyte-specific Hazards} \quad \text{[Eq. 4]}$$

where: Analyte-specific Hazard is the result from Eq. 3.

[Note: When performing these calculations, total risk and hazard estimates will be developed within medium for only the scenario appropriate to the unit's or area's location and use because the reasonably anticipated future land use at a site is significant in defining source material as a principal or low-level threat waste (EPA 1991a). A total risk (or hazard) over all media may be estimated if exposure to contaminants in multiple media may occur. Also, when summarizing this information, the analytes driving the medium-specific total risk and hazard and the major uncertainties in the estimate will be reported, and a total risk or hazard estimate over all media may be reported if this is deemed appropriate.]

The results provided by these analyses may not be sufficient for documentation of final actions, and additional risk assessment and risk evaluation may be needed to meet reporting requirements. Items not provided by these analyses include the following:

- The identification of use scenarios of concern, including consideration of sensitive subpopulations;
- The identification of pathways of concern;
- Consideration of risks due to the transformation, degradation, or migration of contamination (although a comparison of analyte concentrations in soil to screening values protective of groundwater provides this in part); and
- An analysis of uncertainties, including the effect of uncertainties on the resulting risk estimates.

2.2 ANALYSES SUPPORTING NO FURTHER ACTION DECISIONS

No further action can be selected for those sites where it can be demonstrated that no contamination is present that exceeds no action levels (i.e., risks are *de minimis*) or ARARs. (Note; Non-risk issues also must be considered in making this decision. At some sites without unacceptable risk, a no further action decision may not be appropriate because of non-risk concerns.)

In calculating the risk estimate for this decision, the tables discussed earlier and the equations presented earlier will be used. In summarizing this information, the estimated total risk and hazard from all contaminants under the appropriate use will be reported, and the future risk or hazard posed by contaminant transformation, degradation, and migration will be considered qualitatively. In addition, the uncertainties associated with the screening comparison will be discussed, and the effect of these uncertainties on the total risk and hazard estimates for each scenario will be described. Note: As part of this screening analysis, the total risk or hazard over all media will be presented and discussed to ensure that a no further action decision is appropriate.

2.3 ANALYSES USED TO PRIORITIZE FURTHER INVESTIGATIONS

Remedial activities at PGDP are prioritized to ensure that funds allocated to PGDP for remedial actions are directed toward those units or areas that pose the greatest risk to human health and the environment. This prioritization will ensure that these actions provide the maximum benefits in risk reduction. When necessary, risk and hazard estimates for prioritization will be calculated using the tables and equations presented earlier. When summarizing this information, the estimated total risk and hazard from all contaminants under both industrial and residential use will be reported, and the potential future doses and risks posed by contaminant transformation, degradation, and migration will be considered qualitatively. In addition, the uncertainties associated with the screening comparison will be discussed, and the effect of these uncertainties on the total risk and hazard estimates for each receptor group will be estimated qualitatively.

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3. RISK ANALYSES DURING THE REMEDIAL INVESTIGATION

At PGDP, risk analyses occur at three points during the RI of sites: during the preparation of the RI work plan (and some sampling and analysis plans); following implementation of the initial round of work described in the RI work plan (if needed to plan contingency sampling); and during the preparation of the RI report. Analyses occurring at each of these points are discussed in the following sections. (Note that dose assessments are not specifically described in the following. Generally, if a dose assessment is provided, it will be presented in the same format as the risk assessment.)

3.1 ANALYSES DURING WORK PLAN DEVELOPMENT AND IMPLEMENTATION (SCREENING RISK ASSESSMENTS)

As noted in Section 2.4, the screening analyses performed during the site scoping can be used directly in work plan development to reduce the cost of subsequent RI/FS activities. This section discusses the screening analyses that will be performed as part of work plan development and describes the material that will appear in work plans and sampling and analysis plans. (Note: In the following material, “work plan” is used generically for work plans and for those sampling and analysis plans in which risk screening is of use.)

Generally, in work plans, the majority of the risk-related information will appear as part of the initial evaluation. In the work plan’s initial evaluation, the scope, objectives, and methods for the baseline risk assessment will be related; preliminary conceptual site models will be presented; laboratory analytical (or quantitation) limits will be discussed relative to no action screening levels developed specifically for PGDP (i.e., risk-based PRGs in Appendix A); and a preliminary list of COPCs (preliminary COPCs) will be identified. Risk-related information also will appear in the introduction, site characterization summary, and alternatives development description contained in most work plans.

3.1.1 Analyses Appearing in the Introduction of the Integrated RI/FS Work Plan

In the introductory chapter of work plans, the requirements for risk assessments and analyses will be used to help develop the DQOs for the RI. DQOs are qualitative and quantitative criteria used to establish requirements for sample collection and analysis and are based on the needs and intended uses of the data. As a primary user of RI data, the consideration of risk analyses is integral to this process.

Development of DQOs follows a series of steps. The seven steps in the process are shown in a flowchart found in EPA QA/G-4, *Guidance on Systematic Planning Using the Data Quality Objectives Process* (EPA 2006). Similar steps are found in DOE guidance, *Institutionalizing the Data Quality Objectives Process for EM’s Environmental Data Collection Activities* (DOE 1994). The purpose and goal of each step are described in the text in EPA QA/G-4, accompanying the flowchart. EPA QA/G-4 also includes a summary of key elements that also may be of use in developing DQOs for specific investigations. The role of risk assessment within each of these steps is briefly discussed in the remainder of this section.

During Step 1, State the Problem, of the DQO process, risk analyses will be used to identify qualitatively the preliminary COPCs, receptors that may be exposed to contaminants, locations at which exposure may occur, and pathways by which contaminants may reach these locations. This information will be used to develop the conceptual site model against which new data collected as part of the RI can be compared. An example conceptual site model is presented in Figure 3.1.

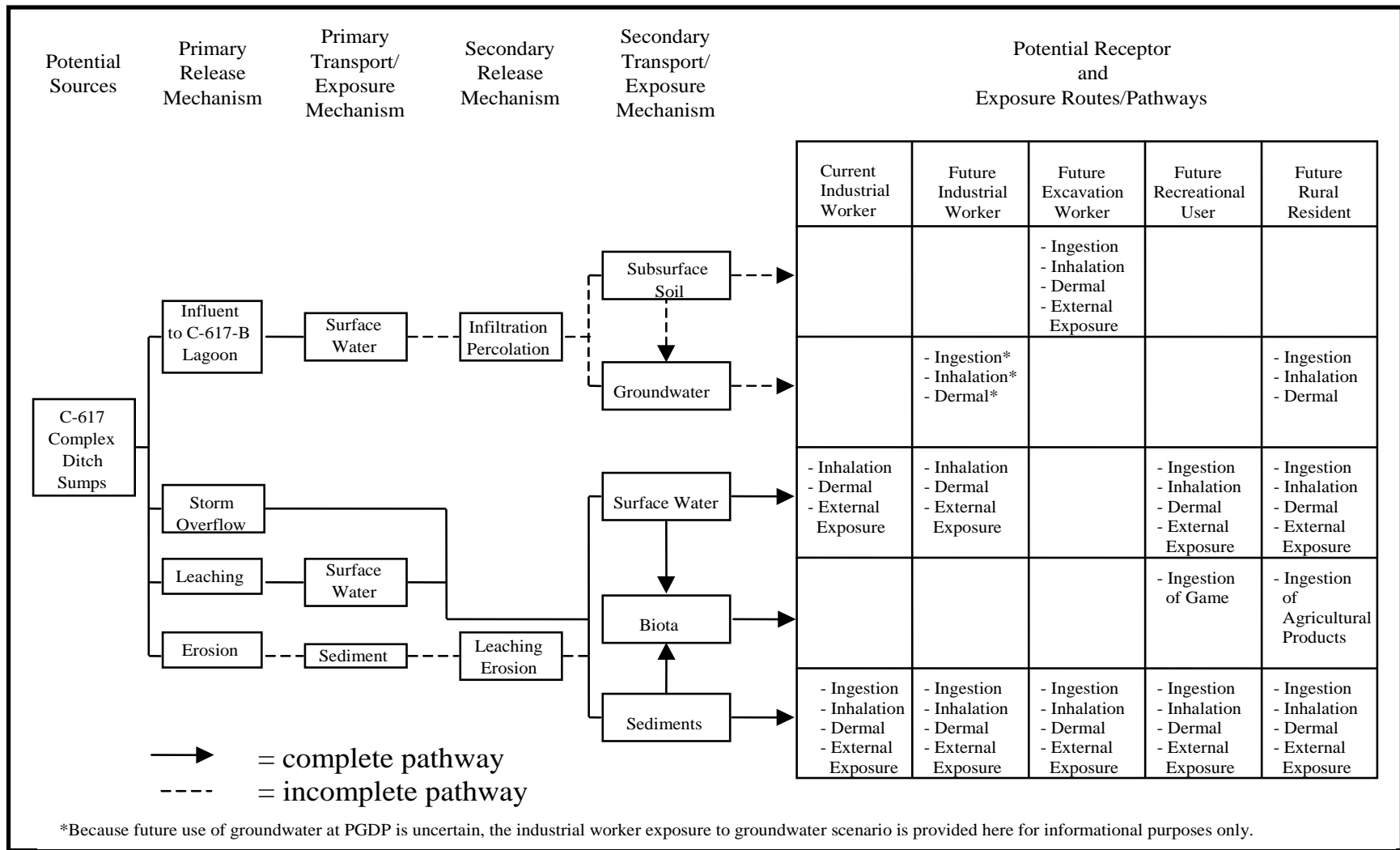


Figure 3.1. Example Risk-Based Conceptual Site Model

Risk analyses also will be used during Step 1 of the DQO process to ensure that the risk management issues are addressed during the investigation. For example, in the approved sampling and analysis plan for SWMU 2 of Waste Area Grouping 22 (DOE 1996a), the problem is stated:

In the past, uranium and multiple COCs were disposed of at SWMU 2. These contaminants have been shown by previous work to be migrating (vertically and horizontally) from the waste cells and show the potential for subsurface migration from the SWMU to the RGA at concentrations or activities that may pose risk to human health and the environment....

Risk analyses will be used during Step 2, Identify the Goals of the Study, of the DQO process to clearly pose questions that must be addressed during the RI. Generally, questions developed during Step 2 of the process will be related to development of contamination concentrations that may remain at or migrate from a site and not pose unacceptable risk; to contaminant migration, and to the activity patterns of present and potential future receptor populations. For example, in the SWMU 2 sampling and analysis plan (DOE 1996a), primary questions related to risk assessment and risk management included the following:

- Will the contaminants migrate (and how) to the RGA at unacceptable concentrations?
- Is there lateral/vertical contaminant movement in the Upper Continental Recharge System (UCRS)?
- What are the chemical characteristics of the waste?

Risk analyses will be used during Step 3, Identify Information Inputs, of the DQO process to establish the preliminary remedial action objectives (RAOs) that must be achieved to mitigate risk to human health and the environment and to provide information useful in determining which alternatives may achieve these objectives. RAOs are criteria used in the FS to aid in the alternative development and selection process. They are site-specific goals that establish the primary objectives and extent of cleanup required by a CERCLA remediation (EPA 1988) and consider COCs, media of concern (MOCs), and potential exposure pathways. The screening levels presented in Section 2 are concentration goals that will make up a portion of the preliminary RAOs for each project. For all investigations at PGDP, the basis of this portion of the human health RAO is to prevent exposure to contaminated media that results in a cumulative (or total) excess lifetime cancer risk (ELCR) greater than 1×10^{-6} or a cumulative (or total) HI greater than or equal to one. This generalized RAO will be enhanced on a project-specific basis as needed (e.g., to include dose concerns).

Risk analyses will be used during Step 4, Define the Boundaries of the Study, of the DQO process to aid in the determination of the spatial and temporal boundaries within which samples must be collected or to which contaminant concentrations must be modeled. Risk analyses will be used to identify spatial boundaries by delimiting the locations both at a SWMU and away from the SWMU at which exposure to contaminants may occur (i.e., exposure points). Risk analyses will be used to identify temporal boundaries by delineating the present and future receptors that may be exposed to contamination and the periods during which these receptors potentially may be present at the exposure points. This information will be used, in turn, to determine the modeling needs for the RI.

Risk analyses will be used during both Steps 3 and 5, Develop the Analytic Approach to the Decision, to set the risk-based limits inherent in these rules and to identify the data required to determine if these limits may be exceeded, consistent with Section XII of the Paducah FFA (EPA 1998a). A primary decision rule that will be included in all work plans for PGDP will note that action must be considered if the risk or hazard posed by contamination at or migrating from a site exceeds allowable limits of an ELCR greater than 1×10^{-6} or HI greater than or equal to one. For example, in the SWMU 2 sampling and analysis plan (DOE 1996a), the leading decision rule (D1) is as follows:

If any of the constituents shown in Table 5.2 are migrating or could migrate (based on RESRAD for uranium and technetium-99 (⁹⁹Tc) and best available 2- or 3-D model for other constituents) from the burial pits, soil matrix, and/or UCRS to the RGA in the future and are found to pose a risk greater than 1×10^{-6} (excess lifetime cancer) or an HI = 1 (noncancer), then an action to control the migration will be evaluated.

Similarly, the following inputs necessary to make this decision are common to all investigations:

- Chemical-specific exposure point concentrations (EPCs) in environmental media, including contaminant concentrations in waste;
- Land-use assumptions (i.e., which scenarios need to be considered);
- Exposure pathways and exposure routes for all current and potential future receptors;
- Exposure units for the investigated area;
- Modeling parameters;
- Risk estimates for each receptor, including sensitive subpopulations, if applicable.

Risk analyses will be used in Step 6, Specify Performance or Acceptance Criteria, by providing the risk-based goals and contaminant concentrations and activities related to these goals that can be used either quantitatively or qualitatively to set decision error limits. As noted previously, consistent with the PGDP FFA, the risk-based goals to be used in all investigations are 1×10^{-6} for ELCR and 1 for HI. For a dose assessment done to provide information for risk managers, the dose-based goal is 1 mrem/year. The concentrations and activities related to these goals are the PRGs presented as the no action levels in Section 2.

Risk analyses will be used in Step 7, Develop the Plan for Obtaining Data, to ensure that the sampling strategy proposed for all investigations meets the minimum requirements needed to achieve answers to the risk-related decision rules. To ensure that this is achieved, all sampling proposed as part of all investigations will be critically reviewed against the needs established under the decision rules for the investigation. Sampling that does not provide information useful to answering risk-related decisions will be justified on another basis.

3.1.2 Analyses Appearing in Prior Characterization Chapter of the Integrated RI/FS Work Plan

In the prior characterization chapter of work plans, results of previous risk evaluations performed for the site under investigation or related to the site will be summarized. Generally, these summaries will consist of results from evaluations performed during the Phases I and II Site Investigations (CH2M HILL 1991 and 1992) or baseline risk assessments and screening analyses performed to support earlier decisions at or near the site, such as prioritization activities.

In presenting the information from previous evaluations, **no attempt will be made to correct any errors or update any values contained in the earlier reports.** All information contained in the earlier report will be presented without change; however, any errors or uncertainties affecting the results will be identified. Additionally, because in earlier baseline risk assessments, results were not summarized in a consistent format, an attempt will be made to present the results taken from these earlier reports in two-way tables. [Note: The format for the two-way table is patterned after the format in Exhibits 8-2 and

8-3 of Risk Assessment Guidance for Superfund (RAGS), Part A, (EPA 1989a) and is consistent with the risk characterization tables found in RAGS, Part D (EPA 1998b).] The exact format for tables presented in RAGS, Part D, is not used for the PGDP risk characterization tables because the Risk Assessment Working Group determined that the tables presented in this Risk Methods Document are adequate to meet the intent of RAGS, Part D. In addition, when summarizing the results of previous assessments, the scenarios, pathways, contaminants, and MOC for each unit or area under investigation will be listed, and major uncertainties affecting the risk assessment results will be noted.

An example of the format for the “two-way table,” adapted from Table 5.78 of Appendix L.1 of the approved *Resource Conservation and Recovery Act Facility Investigation/Remedial Investigation Report for Waste Area Grouping 1 and 7 at Paducah Gaseous Diffusion Plant, Paducah, Kentucky* (DOE 1996b), is shown in Exhibit 3.1. The example table shown in the exhibit will be used to summarize risk assessment results because it allows easy identification of scenarios of concern (i.e., value in column entitled “Total Risk,” COCs (i.e., values in the column entitled “Chemical-Specific Risk”), and pathways of concern (POCs) (i.e., values in the row entitled “Pathway Risk”). In addition, the chemicals and pathways driving total risk can be easily identified, and the risk related to exposure to each environmental medium can be easily derived (i.e., by summing the appropriate pathway totals). Finally, the blank cells in the table and the associated explanation for these blanks show where information was insufficient to allow risks to be characterized.

Exhibit 3.1. Example Two-Way Table for Presentation of Historical Risk Assessment Results

SWMU 136 Excess Lifetime Cancer Risks for Future Rural Resident							
Analyte	Ingestion of Groundwater	Dermal Contact with Groundwater	Ingestion of Soil	Chemical-specific Risk	Total Risk
Trichloroethene	2.30E-05	4.17E-06	8.35E-05	1
Benzo(a)anthracene			8.78E-09	1.35E-06	
Benzo(a)pyrene			1.20E-07	1.83E-05	
.	
.	
.	
Uranium-238			1.53E-09	3.05E-07	
Pathway Risk	2.32E-05	4.23E-06	1.72E-07		
Total Risk							1.10E-04

Note: The reasons for blank cells are discussed as part of the risk assessment/evaluation. Generally, blank cells will result from unavailable or inadequate data.

3.1.3 Analyses Appearing in Initial Evaluation Chapter of the Integrated RI/FS Work Plan

In the initial evaluation chapter of work plans, the methods to be used to complete the baseline risk assessment for the units or areas under investigation will be discussed, and a preliminary evaluation of historical information, including a comparison of concentrations and activities of analytes in environmental samples with health-based standards (i.e., PRGs, ARARs, etc.) and a comparison of analytical limits with background concentrations, will be presented. This information will be used, in turn, to develop the field sampling plan contained in the work plan.

The description of the methods to be used to complete the baseline risk assessments for the units or areas under investigation will follow that presented in Section 3.3 of this document. Generally, this material will delineate clearly the scope and objectives of the baseline risk assessment and briefly describe the activities that will occur during the data evaluation (i.e., identification of COPCs); exposure assessment; toxicity assessment; risk characterization; and remedial goal option (RGO) development stages of the

baseline human health risk assessment. This material also will summarize the results that will be obtained from each stage of the baseline risk assessment. As part of this discussion, conceptual site models for each unit or area under investigation will be presented.

The preliminary evaluation of historical information presented in this chapter of the work plan will summarize the information presented in earlier chapters of the work plan and evaluate this information against the characterization and inventory of wastes, information status of key assessment factors, and release potential from contaminant sources. As part of the characterization and inventory of wastes, comparison tables similar to those discussed in Section 2 will be prepared. Because additional screening criteria may need to be considered, the comparison tables prepared as part of site scoping activities may not be able to be transferred directly to the work plan. An example of the comparison table that will be used in work plans to compare the PGDP screening PRGs to analytical results from soil (and sediment) and groundwater (and surface water) is shown in Exhibit 3.2.

Exhibit 3.2. Presentation of Screening Assessment Results in the RI Work Plan

Analyte	Soil (mg/kg or pCi/g)			Groundwater (µg/L or pCi/L)			
	Maximum ¹	PRG ²	Method Detection Limit ³	Maximum	PRG	MCL ⁴	Method Detection Limit
# 1							
# 2							
.
.
.
# N							

¹ This value will be the maximum detected value for the medium reported in previous investigations. The qualifier codes attached to the value, if any, will be included with the value.

² The risk-based PGDP screening preliminary remediation goal (i.e., PRG) that appears in this table will be the lesser of the cancer- and hazard-based, no action residential use PRGs taken from Appendix A. Additionally, the hazard-based PRG that is included will be that calculated for a child aged 1 to 7.

³ This value will be the project-specific value reported in the Quality Assurance Project Plan of the work plan (or the appropriate chapter of sampling and analysis plans). For radionuclides, this column should have the heading “MDC” or “MDQ” and present MDCs from Multi-Agency Radiological Laboratory Analytical Protocols (MARLAP) guidance.

⁴ The maximum contaminant levels (i.e., MCLs) are drinking water standards and will be taken from the most recent information.

After completing the comparison table for each site, the analytes that previously were detected or are expected to be present and that have detection limits (MDCs for radionuclides) that exceed the PRGs will be reported. The analytes with detection limits exceeding PRGs will be reported because the quantitation limit (or method detection limit for chemicals or MDC for radionuclides) used for samples providing data for risk assessment should be less than those concentrations that may have an impact on human health or the environment. It is important to note that, although this evaluation may show that some quantitation limits exceed their respective screening criteria, this evaluation alone will not be used to establish the analytical quantitation limits for a project. The analytical limits will be established considering this information and factors such as site history and potential actions.

Material in the comparison tables also will be used to compile a list of preliminary COPCs for each unit or area under investigation. An analyte will be placed on this preliminary list if the concentration or activity of the analyte at a unit or area exceeds one or more of the screening criteria. Note: Unless it can be shown that cross-media contamination is not present, the list of preliminary COPCs will be compiled over all media. If it can be demonstrated that cross-media contamination is not likely, then a list of preliminary COPCs will be compiled for each medium to be investigated during the project. These lists will provide risk managers with information that can be used in the initial selection and screening of alternatives. In addition, this list can be used to target the analyte list for the project to ensure that analytical costs are appropriate for the project.

An example of the comparison table that will be used in work plans to compare background values to analytical results for inorganic chemicals and radionuclides in soil and groundwater is shown in Exhibit 3.3. (Note: as discussed earlier, background values are not available for sediment and surface water; therefore, a table comparing analytical results from sediment and surface water to background will not be presented.) This table will be used to justify the analyte list for the project. As with the list of preliminary COPCs, justification of the analyte list is important to ensure that analytical costs are appropriate for the project.

Exhibit 3.3. Presentation of Background Comparison in the RI Work Plan

Analyte	Soil Data for SWMU (mg/kg or pCi/g) ¹			Soil Background Concentration (mg/kg or pCi/g) ²	Groundwater Data for SWMU (µg/L or pCi/L) ³			Groundwater Background Concentration (µg/L or pCi/L) ⁴
	SWMU 1	...	SWMU N		SWMU 1	...	SWMU N	
# 1			
# 2			
.		.				.		
.		.				.		
.		.				.		
# N			

¹ This will be the maximum detected value for soil reported in previous investigations. The qualifier codes attached to the value, if any, will be included with the value.

² The soil background concentration (or activity) will be that presented in Appendix A or updated values.

³ This will be the maximum detected value for groundwater reported in previous investigations. The qualifier codes attached to the value, if any, will be included with the value.

⁴ The groundwater background concentration (or activity) will be that presented in Appendix A or updated values.

3.1.4 Analyses Appearing in Remedial Alternatives Development Chapter of the Integrated RI/FS Work Plan

In the remedial alternatives development chapter of work plans, attention will be paid to the importance of risk reduction in remedial alternatives development and to the method to be used to measure risk reduction during the detailed analysis of remedial alternatives. For example, this chapter will note that remedial alternatives are developed to be protective of human health and the environment and that RAOs will consider COCs, POCs, and MOCs. In addition, this chapter will present the nine criteria used in the detailed analysis of alternatives under CERCLA. Most importantly, this chapter will discuss if a qualitative or quantitative detailed risk analysis of alternatives is anticipated and delineate the data that are required to support this risk analysis. (Determining whether a qualitative or quantitative risk analysis of alternatives is needed is important because additional data may need to be collected during the RI to support a quantitative analysis. Additional discussion concerning qualitative and quantitative risk analysis of alternatives is presented in Section 4.)

3.2 ANALYSES FOLLOWING COMPLETION OF THE INITIAL ROUND OF INVESTIGATION

Many RI work plans will contain a description of contingency sampling that may be used to address the uncertainties in environmental contaminant distribution expected to be encountered during the investigation. If this contingency sampling is to be collected as part of a phased investigation, then analyses may be used to allow the three FFA parties to discuss and agree if contingency soil (or sediment) sampling is necessary. In this case, a formal or informal report may be prepared after the completion of

the initial round of sampling. In this report, results from the initial sampling and relevant historical sampling may be compared to human health screening criteria (i.e., PRGs) for the expected future use of the area and background concentrations of chemicals and radionuclides. To keep this presentation consistent with that used in work plan development, this presentation will use comparison tables similar to those presented earlier. Because the extent of soil (or sediment) contamination needs to be considered, as well as the nature of contamination, tables considering the location of samples (horizontal and vertical), in addition to the tables considering the maximum detected analyte concentrations, will be prepared. A spatial plane view presentation of the data also should be provided.

The format of the comparison table to be used to determine if the nature of contamination in soil may pose an unacceptable risk or hazard is in Exhibit 3.4. In this table, the maximum detected concentration or activity in all soil samples collected at a site is compared to the no action PRG for soil exposure for the expected future land use, the groundwater protection PRG, and the background concentration. This table will be used to refine the list of preliminary COPCs and the analytical list for contingency sampling. In this evaluation, an analyte will become a preliminary COPC if its concentration exceeds any PRG and the background concentration or activity.

Exhibit 3.4. Presentation of Screening Assessment Results to Evaluate Nature of Contamination in Soil after the Initial Round of Sampling

Analyte	Soil (mg/kg or pCi/g)			
	Maximum ¹	PRG ²	Groundwater Protection PRG ³	Background ⁴
# 1				
# 2				
.
.
.
# N				

¹ This value will be the maximum detected value for soil reported in the current and relevant previous investigations. The qualifier codes attached to the value, if any, will be included with the value.

² The PRG will be the lesser of the no action cancer- and no action hazard-based PRGs for exposure to soil for the appropriate future use taken from Appendix A. If residential use PRGs are used, then the no action hazard-based PRG should be that for a child aged 1 to 7.

³ The groundwater protection PRG will be the lesser of the no action cancer- and no action hazard-based PRGs taken from Appendix A. Note: This PRG is protective of groundwater that may be used in the home. A PRG for protection of groundwater used industrially is not relevant to this screening assessment.

⁴ The soil background concentration (or activity) will be that presented in Appendix A or the most recent updated study/report.

The format of the comparison table to be used to determine if the nature of contamination in sediment may pose an unacceptable cancer risk or hazard will be similar to that in Exhibit 3.4; however, for the sediment table, neither the groundwater protection PRG nor the background concentration will appear. The groundwater protection PRG will not be included because migration of contaminants from sediment to groundwater is not expected to be a significant migratory pathway. Background concentrations of chemicals and radionuclides will not be included because these data do not exist for sediment. As with the soil table, the sediment table will be used to refine the list of preliminary COPCs and the analytical list for contingency sampling. In this evaluation, an analyte will become a preliminary COPC if its concentration or activity exceeds any risk-based screening criterion.

The format of the comparison table to be used to evaluate the adequacy of initial sampling in delimiting the extent of contamination in surface soil is in Exhibit 3.5. In this table, the analyte concentrations or activities in surface soil samples collected along migration routes or at the periphery of a site are compared to the no action PRG for soil for the expected future land use and the background concentration or activity. Note that the groundwater protection soil PRG is not used in this comparison because that evaluation is performed as part of the subsurface soil evaluation. Generally, surface sampling will be

deemed adequate if analyte concentrations and activities in samples collected along migration routes do not exceed both the no-action PRGs for soil and background concentrations. In deciding if sampling has adequately determined the extent of contamination, additional factors such as historical information will be considered.

Exhibit 3.5. Presentation of Screening Assessment Results to Evaluate Extent of Contamination in Surface Soil after the Initial Round of Sampling

Analyte	Soil (mg/kg or pCi/g)		
	Maximum ¹	PRG ²	Background ³
# 1			
# 2			
.	.	.	.
.	.	.	.
.	.	.	.
# N			

¹This value will be the maximum detected value for soil reported in a sample collected along migration routes or at the periphery of the unit or area in the current investigation. The qualifier codes attached to the value, if any, will be included with the value.

²The PRG will be the lesser of the no action cancer- and no action hazard-based PRGs for the appropriate future use taken from Appendix A.

³The soil background concentration (or activity) will be that presented in Appendix A or the most recent updated study/report.

The format of the comparison table to be used to evaluate the adequacy of initial sampling in delimiting the extent of contamination in sediment will be similar to that used for soil (Exhibit 3.5); however, the background concentration or activity will not appear in the sediment table because background values for sediment do not exist. The evaluation of this table will be the same as for soil.

The format of the comparison table to be used to evaluate the adequacy of initial sampling in delimiting the extent of contamination in subsurface soil is in Exhibit 3.6. In this table, the analyte concentrations or activities in subsurface soil samples collected at the periphery of the area under investigation will be compared to the groundwater protection PRGs and background concentrations of chemicals and radionuclides. Note: The no action PRGs for soil are not in this table because these criteria are for contact with contaminated soil, and contact with subsurface soil is not expected. Generally, subsurface sampling will be deemed adequate if analyte concentrations and activities in samples collected at the periphery of the unit or area under investigation do not exceed both the groundwater protection PRGs and background concentrations. In deciding if sampling has adequately determined the extent of contamination, additional factors such as historical information will be considered.

Analyses to evaluate groundwater and surface water sampling in determining the nature and extent of contamination in groundwater and surface water will be similar to those for soil. The format of the comparison table to be used to determine if the nature of contamination in groundwater may pose an unacceptable excess cancer risk or systemic toxicity is in Exhibit 3.7. In this table, the maximum detected concentration or activity in all groundwater samples collected at the site will be compared to the no action PRG for residential use of groundwater, the maximum contaminant level (MCL), and the background concentration or activity. This table will be used to refine the list of preliminary COPCs and the analytical list for contingency sampling. In this evaluation, an analyte will become a preliminary COPC if its concentration exceeds any screening criterion and the background concentration or activity. Comparisons to MCLs will not be used to identify COPCs, but will be provided for information only.

Exhibit 3.6. Presentation of Screening Assessment Results to Evaluate Extent of Contamination in Subsurface Soil after the Initial Round of Sampling

Analyte	Soil (mg/kg or pCi/g)		
	Maximum ¹	Groundwater Protection PRG ²	Background ³
# 1			
# 2			
.	.	.	.
.	.	.	.
.	.	.	.
# N			

¹ This value will be the maximum detected value or maximum activity for radionuclides for subsurface soil reported in a sample collected at the periphery of the unit or area in the current investigation. The qualifier codes attached to the value, if any, will be included with the value.

² These values are taken from Appendix A.

³ The soil background concentration (or activity) will be that presented in Appendix A or the most recent updated study/report.

Exhibit 3.7. Presentation of Screening Assessment Results to Evaluate Nature of Contamination in Groundwater after the Initial Round of Sampling

Analyte	Groundwater (µg/l or pCi/l)			
	Maximum ¹	PRG ²	Maximum Contaminant Level ³	Background ⁴
# 1				
# 2				
.
.
.
# N				

¹ This value will be the maximum detected value for groundwater reported in all samples collected around the unit or area during the current and relevant previous investigations. The qualifier codes attached to the value, if any, will be included with the value.

² The PRG will be the lesser of the no action cancer- and no action hazard-based PRGs in Appendix A. Note: The hazard-based PRG should be that for a child aged 1 to 7.

³ The MCL will be taken from Appendix A or the most recent update.

⁴ The groundwater background concentration (or activity) will be that presented in Appendix A or the most recent update.

The table used to determine if contamination in surface water may pose an unacceptable cancer risk or hazard will be similar to that in Exhibit 3.7; however, background concentrations of chemicals and radionuclides will not appear in the surface water table because background data do not exist for surface water. The evaluation of this table will match that for groundwater.

For all investigations except the final RI of the Groundwater Operable Unit, there will be limited evaluation of the extent of existing groundwater contamination during the evaluation of the initial round of sampling. Currently, only the extent of dense nonaqueous-phase liquid contamination (i.e., secondary sources) is addressed during the investigation of the individual units and areas. The method used for the detection of these secondary sources does not rely on risk analysis and will not be discussed here. For the Groundwater Operable Unit investigation, the comparison table used to examine the adequacy of sampling in determining the extent of groundwater contamination will be similar to that in Exhibit 3.7; however, in this evaluation, a table will be prepared for each groundwater sampling location along the suspected periphery of the contaminant plumes. In each of these tables, the maximum detected analyte concentrations and activities will be compared to the no action residential use PRGs, MCLs, and background concentrations. Generally, groundwater sampling will be deemed adequate to determine the extent of contamination if analyte concentrations and activities in samples collected along periphery of the suspected groundwater contaminant plumes do not exceed screening criteria and background

concentrations. In deciding if sampling has adequately determined the extent of contamination, additional factors such as historical information will be considered.

The table to be used to determine the adequacy of sampling in determining the extent of surface water contamination also will be similar to that in Exhibit 3.7. As noted earlier, this table will not contain background concentrations of chemicals and radionuclides because background values are not available for surface water. Generally, surface water sampling will be deemed adequate to determine the extent of contamination if analyte concentrations and activities in samples collected downstream of a unit or area do not exceed screening criteria. In deciding if sampling has adequately determined the extent of contamination, additional factors such as historical information will be considered.

3.3 ANALYSES FOR THE RI REPORT (BASELINE RISK ASSESSMENTS)

Baseline risk assessments will be prepared to support final actions at PGDP. To ensure consistency among assessments and conformity with agreements reached between the U.S. Department of Energy (DOE) and regulatory agencies, all assessments will contain either the material described in succeeding sections or an explanation stating why the material is not presented. Material described herein but not relevant to a particular assessment will be noted in the assessment. The following are specific objectives of the remedial action process to be addressed in this section:

- Delineate the methods PGDP will use in the evaluation, determination, and documentation of baseline risks to human health and the environment at a site; and
- Describe the methods PGDP will use to determine the concentrations and activities of analytes that can remain on-site and still be adequately protective of human health and the environment both on-site and off-site.

In the following sections, the presentation follows the outline to be used in baseline human health risk assessments. Data evaluation methods are discussed in Section 3.3.3, exposure assessment methods are presented in Section 3.3.4, toxicity assessment methods are described in Section 3.3.5, risk characterization methods are delineated in Section 3.3.6, uncertainty in the risk assessment is discussed in Section 3.3.7, and RGO derivation methods are discussed in Section 3.3.8. In addition, the sources used to prepare this material are listed in Section 3.3.1, and general issues are considered in Section 3.3.2.

[Note: The methods for the baseline ecological risk assessment are not considered here. They are described in the companion Ecological Risk Methods Document. Additionally, methods to be used for dose assessment are not presented in detail. The methods for dose assessment generally should follow those used for baseline risk assessments.]

3.3.1 Guidance Documents

The methods discussed in the following sections are consistent with current EPA Region 4 and headquarters risk assessment guidance documents, the Commonwealth of Kentucky Department for Environmental Protection (KDEP) risk assessment guidance, and applicable DOE Orders. In addition, these methods are consistent with agreements reached during meetings among DOE, EPA Region 4, and KDEP risk assessment personnel (DOE 1996c; EPA 1996a; KDEP 1996; and RAWG 2000b, 2000c, 2000d, 2000e, 2000f, 2000g, 2007a, 2007b, 2007c, 2012a, and 2012b) and strategies and methods developed for human health risk assessments for use at other DOE sites located in EPA Region 4 (e.g., K-25, X-10, and Y-12 in Oak Ridge, Tennessee). Some of these methods are different from those used in earlier risk assessments. References for methods and approach should refer to this methods document

and/or the original guidance documents instead of other site-specific project documents to avoid inappropriate references. Many of the documents and other materials used in developing the methods are listed chronologically in the following sections. If newer versions of the listed reference are available, the newer version should be used in place of the specific version listed in the following sections.

3.3.1.1 EPA guidance documents and materials

- *Risk Assessment Guidance for Superfund, Volume I: Human Health Evaluation Manual, Parts A, B, C, D, E, and F* (EPA 1989a, 1991b, 1991c, 1998b, 2004a, and 2009, respectively) (RAGS, Parts A, B, C, D, E, and F, respectively)
- *Exposure Assessment Methods Handbook* (EPA 1989b)
- *Role of the Baseline Risk Assessment in Superfund Remedy Selection Decisions* (EPA 1990a)
- *Guidance for Data Usability in Risk Assessment* (EPA 1990b)
- *Human Health Evaluation Manual, Supplemental Guidance: "Standard Default Exposure Factors"* (EPA 1991d)
- *Dermal Exposure Assessment: Principles and Applications* (EPA 1992a)
- *Risk Assessment Guidance for Superfund, Volume I: Human Health Evaluation Manual, Supplemental Guidance, Dermal Risk Assessment* (EPA 1992b)
- *Supplemental Guidance to RAGS: Calculating the Concentration Term* (EPA 1992c)
- *Guidelines for Exposure Assessment* (EPA 1992d)
- Revisions to Sections 3.3.1 and 3.3.2 of the RAGS, Part B (EPA 1993a)
- *Superfund's Standard Default Exposure Factors for the Central Tendency and Reasonable Maximum Exposure* (EPA 1993b)
- *Guidance Manual for the Integrated Exposure Uptake and Biokinetic (IEUBK) Model for Lead in Children*, EPA/540/R-93/081 (EPA 1994a)
- *OSWER Directive: Revised Interim Soil Lead Guidance for CERCLA Sites and RCRA Corrective Action Facilities, OSWER Dir #9355.4-12*(EPA 1994b)
- *Soil Screening Guidance: Technical Background Document*, EPA/540/R-95/128, Office of Solid Waste and Emergency Response, Washington, DC, July 1996 (EPA 1996b)
- *Exposure Factors Handbook*, EPA 600/P-95/002Fa,b,c (EPA 1997b)
- *Approach for Addressing Dioxin in Soil at CERCLA and RCRA Sites*, OSWER Directive 9200.4-26 (EPA 1998c)
- *Soil Screening Guidance for Radionuclides: User's Guide and Technical Background Document Final Guidance*, OSWER Directive 9355.4-16A and OSWER Directive 9355.4-16 (EPA 2000b)

- *Supplemental Guidance to RAGS: Region 4 Bulletins, Human Health Risk Assessment Bulletins*, EPA Region 4, Website version last updated May 2000 (EPA 2000c)
- *Guidance for Assessing Chemical Contaminant Data for Use in Fish Advisories, Third Edition*, EPA 823-B-00-007 (EPA 2000d)
- *Estimating Dermal and Inhalation Exposure to Volatile Chemicals in Domestic Water* (Schaum et al. 1994)
- Risk Assessment Guidance for Superfund: Volume III-Part A, Process for Conducting Probabilistic Risk Assessment (EPA 2001a)
- *Supplemental Guidance for Developing Soil Screening Levels for Superfund Sites*, Superfund, Office of Solid Waste and Emergency Response, OSWER 9355.4-24 (EPA 2002)
- *Recommendations of the Technical Review Workgroup for Lead for an Approach to Assessing Risk Associated with Adult Exposures to Lead in Soil* (EPA 2003a)
- *Human Health Toxicity Values in Superfund Risk Assessments* (EPA 2003b)
- *Integrated Exposure Uptake Biokinetic Model for Lead in Children, Windows® version* (IEUBKwin v1.1 build 9) (available at www.epa.gov/superfund/lead/products.htm; user's guide is EPA 2004a)
- EPA Regional Screening Level Tables (EPA 2012) at http://www.epa.gov/reg3hwmd/risk/human/rb-concentration_table/Generic_Tables/index.htm
- *Guidance on Systematic Planning Using the Data Quality Objective Process*, EPA QA/G-4 (EPA 2006a)
- *Systematic Planning: A Case Study for Hazardous Waste Site Investigations*, EPA QA/CS-1 (EPA 2006b)
- *National Recommended Water Quality Criteria: 2006* (EPA 2006c)
- 2006 Edition of the Drinking Water Standards and Health Advisories (EPA 2006d)
- Data Quality Assessment: Statistical Methods for Practitioners, EPA QA/G-9S (EPA 2006e)
- *EPA provisional toxicity values support document* available on request from Technical Support Section, EPA Region 4 (EPA-PROV)
- *The 2005 World Health Organization Reevaluation of Human and Mammalian Toxic Equivalency Factors for Dioxins and Dioxin-Like Compounds* (Van den Berg et al. 2006)
- *Brownfields Technology Primer: Vapor Intrusion Considerations for Redevelopment* (EPA 2008)
- *ProUCL Version 4.1.00 Technical Guide (Draft)*. ORD NERL ESC Technical Support Center, Characterization and Monitoring Branch, Las Vegas, NV (EPA/600/R-07/041) (EPA 2010)
- *Human Health Evaluation Manual, Supplemental Guidance: Update of Standard Default Exposure Factors*, OSWER 9200.1-120 (EPA 2014)

3.3.1.2 Commonwealth of Kentucky guidance documents and materials

- *Kentucky Risk Assessment Guidance*, Risk Assessment Branch, Department of Environmental Protection, Commonwealth of Kentucky (KDEP 2002)
- *Kentucky Guidance for Ambient Background Assessment*, Risk Assessment Branch, Department of Environmental Protection, Commonwealth of Kentucky, January 8 (KDEP 2004a)
- *Kentucky Guidance for Groundwater Assessment Screening*, Risk Assessment Branch, Department of Environmental Protection, Commonwealth of Kentucky, January 15 (KDEP 2004b)
- *Trichloroethylene Environmental Levels of Concern*, Risk Assessment Branch, Department of Environmental Protection, Commonwealth of Kentucky, April (KDEP 2004c)

3.3.1.3 Other materials

- Meeting Summary for the Risk Assessment/Risk Evaluation Meeting, February 7, 1996, in Atlanta, February 13, 1996, Conference Call (DOE 1996c)
- *Guidance for Conducting Risk Assessments and Related Risk Activities for the DOE-ORO Environmental Management Program* (Bechtel Jacobs Company LLC 1999)
- Minutes and notes from meetings of the PGDP Human Health Risk Assessment Working Group (RAWG 2000b, 2000c, 2000d, 2000e, 2000f, 2000g, 2007a, 2007b, 2007c, 2012a, 2012b, and Appendix E of this document)

3.3.2 General Methods

The risk methods document generally follows guidance in EPA's RAGS (EPA 1989a) and Kentucky's *Risk Assessment Guidance* (KDEP 2002); however, there are issues for which the two guidance documents differ. In those cases, the Risk Methods Document reconciles these two different approaches.

3.3.2.1 Format for the baseline human health risk assessment

The outline that will be followed when preparing baseline human health risk assessments for PGDP is provided in Appendix C of this document. This outline is consistent with that in RAGS, Part A (EPA 1989a), and in *Kentucky Risk Assessment Guidance* (KDEP 2002) and includes all sections that must be included in a complete baseline human health risk assessment. As such, some portions of the outline may not be applicable to some baseline human health risk assessments of limited scope; however, any baseline human health risk assessment prepared for PGDP will include the major and second level headings in the order presented. Major headings that will appear in all baseline risk assessments are "Results of Previous Studies," "Identification of Chemicals of Potential Concern," "Exposure Assessment," "Toxicity Assessment," "Risk Characterization," "Uncertainty in the Risk Assessment," "Conclusions and Summary," and "Remedial Goal Options Development." In addition, each baseline human health risk assessment will contain introductory material that delineates the scope and objectives of the assessment.

Examples of the format for tables that will be used in the risk assessment are presented in Exhibit 3.8. *List of Chemicals of Potential Concern*; Exhibit 3.9. *Summary of Pathway Analysis in the Exposure Assessment*; Exhibit 3.10. *Presentation of Exposure Point Concentrations*; Exhibit 3.11. *Chemical-Specific Parameters*; Exhibit 3.12. *Daily Intakes (Dose) for Receptor 1*; Exhibit 3.13. *Exposure Route Summary*

for the Current Use Scenario—Systemic Toxicity; Exhibit 3.14. *Driving Contaminants’ Summary for Current Use Scenario—Systemic Toxicity*; Exhibit 3.15. *Summary of Risk Characterization*; Exhibit 3.16. *Summary of Uncertainty Analysis*; and Exhibit 3.17. *Presentation of Remedial Goal Options*. Shorter summary tables for the body of the report will summarize the following information:

- Land use scenarios and media assessed for each source area;
- Scenarios for which human health risk exceeds *de minimis* levels; and
- A table for each source summarizing the COCs and POCs, as well as the contribution of each COC and POC to the total risk and hazard.

3.3.2.2 Presentation of results from previous studies

In all baseline risk assessments prepared for PGDP, the results will be presented from previous risk assessments and other risk evaluations that are relevant to the unit or area being assessed. These results will be included to allow for a comparison between results of earlier work and the results of the current baseline risk assessment. Differences seen will be discussed in the observations section of the current baseline risk assessment.

The format for presenting the results of the earlier risk assessments will follow that which will be used for reporting previous studies in the RI work plan. This is discussed in detail in Section 3.1.2. For risk evaluations, if any, that are not risk assessments, results will be presented verbatim and without interpretation. Relevant results from these studies also may be used in the uncertainty discussion of the current baseline human health risk assessment.

3.3.3 Data Evaluation Methods

The primary purpose of this section of the baseline human health risk assessment will be to develop the list of COPCs used in the assessment. In this section, the data quality/data usability review, procedures to screen data, a summary of the results of the screening, and a final list of COPCs will be presented. Additionally, this section will provide site-specific characterization data used in the exposure assessment. Methods to complete each of these activities are presented in the following.

3.3.3.1 Data quality/data usability review

The overall goal of the data quality/data usability review is to develop a data set of known quality that is representative of the site and is reproducible. Use of this systematic approach is consistent with EPA guidance (EPA 2006f; EPA 2006e). The data quality/data usability review process (Figure 2.2) incorporates the aspects of data quality/data usability [measurement quality objectives (MQOs)] with an evaluation of planned data uses for each project DQOs to make a determination concerning the suitability of historical/current project data for use in risk assessment. The initial steps of data assessment and data validation generally are completed by a subject matter expert before the results are provided to the risk assessor. The data quality assessment (DQA) examines the data set to ensure that the MQOs have been met and that the data are sufficient and representative of the site or source investigated. Figure 3.2 [from the EPA DQA guidance (EPA 2006f)] is provided to illustrate how DQA fits into the data evaluation process.

3.3.3.2 Procedures to screen or evaluate data to determine COPCs

Data screening to develop the list of COPCs will be performed in the following eight steps.

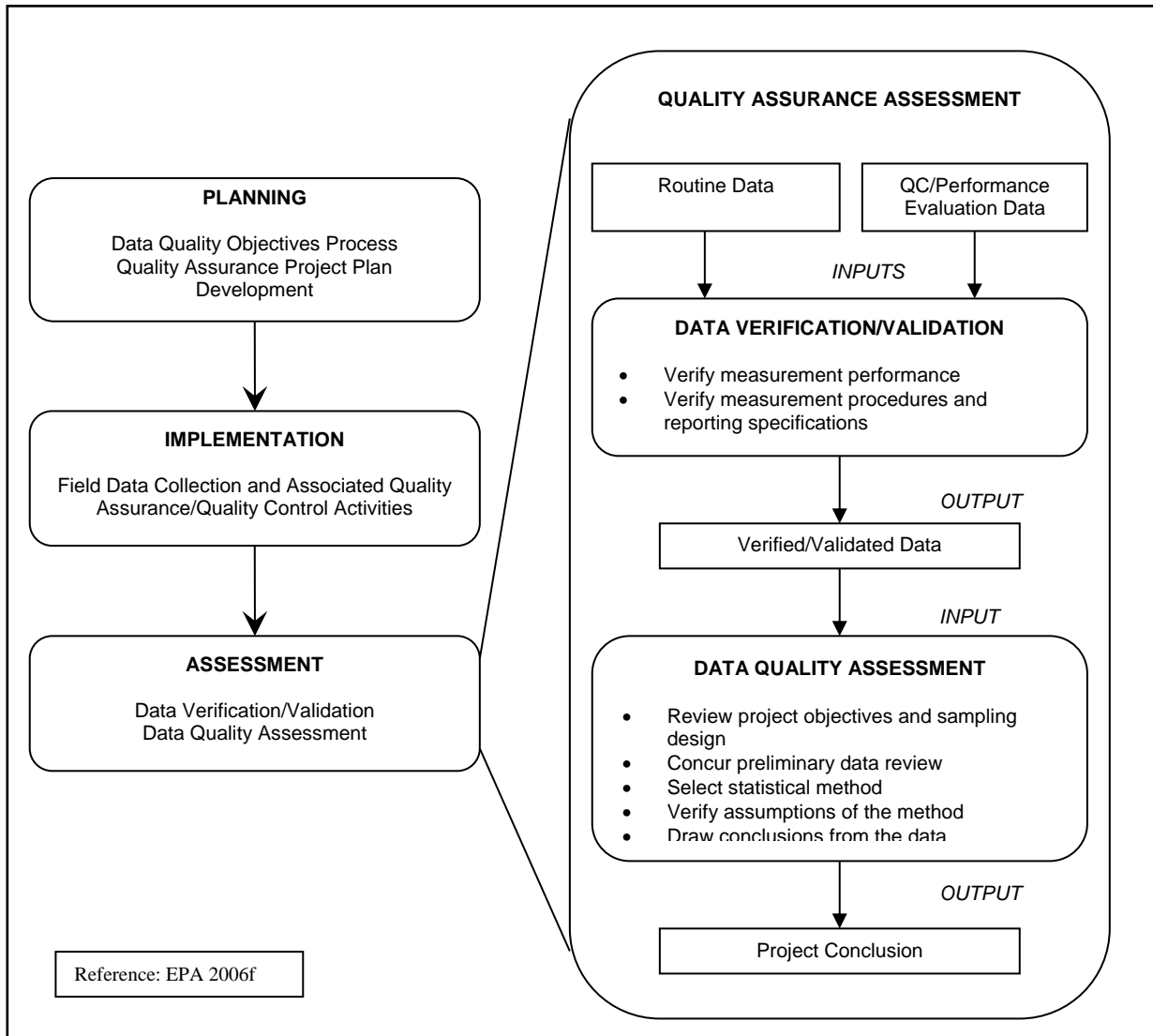


Figure 3.2. Data Life Cycle

- **Step 1: Evaluation of sample design and locations.** Data will be examined to ensure that the samples from which data were derived were collected using sampling methods that are adequate to determine the nature and extent of contamination for the particular unit or area being assessed. Data not from the unit or area under investigation or not useful in determining contaminant migration from the unit or area will not be used quantitatively in the assessment because these data are not representative of the unit or area for which remedial actions are being considered. In particular, when considering groundwater sampling results, only data from samples collected from wells located in contaminant plumes will be used.
- **Step 2: Evaluation of sampling and analytical methods.** Data will be examined to ensure that the sampling methods and analytical methods used in the laboratory are consistent with EPA-approved methods for nonradionuclides. Data for nonradionuclides not from EPA-approved methods will not be used quantitatively in the risk assessment, but may be used qualitatively. Methods for radionuclides will be evaluated during the DQO process to ensure that data quality requirements can be achieved. Also in this step, groundwater and surface water data will be examined, and data from

the analyses of filtered water will be deleted from the data set. Only results from unfiltered samples will be used quantitatively in baseline human risk assessments performed at PGDP. Note: Filtered groundwater and surface water data may be used in the uncertainty section of the assessment when discussing data sources and their effects on risk estimates.

For many sites, survey-type data such as X-ray fluorescence (XRF) data and results from polychlorinated biphenyl (PCB) field test kits are available in addition to the laboratory analytical data. The primary use of such data is for site characterization, but these survey-type data also can play a role in risk-based decision making. Survey-type data assist in determining the distribution of COPCs and can be used to identify which sets of laboratory data should be combined to develop site average contaminant concentrations. Potentially, survey-type data also could be combined with lab data in a risk assessment to determine the average concentrations for contaminants, but this would require demonstrating that the lab and survey-type data possess similar detection limits and analytical uncertainty. In addition, a DQA would need to be completed to show that both types of data sets are comparable and representative of the site conditions. This DQA either could be in the risk assessment or in a report completed prior to or in concert with the risk assessment.

Finally, whenever survey-type data are used for guiding how lab data are handled or are combined with lab data, then the risk assessment would need to have an uncertainty discussion that appropriately identifies (a) how the results of the risk assessment could vary if the survey type data were not used and (b) how the use of the survey data increases or decreases the risk of making an incorrect risk-based decision for a location.

- **Step 3: Evaluation of sample quantitation limits.** See Figure 3.3 for an example of Step 3.

Chemicals. The sample quantitation limits for each analyte and sample will be examined to determine if these limits were below the concentration at which the analyte may pose an unacceptable risk or hazard to human health. If the maximum sample quantitation limit for an analyte (over all samples within a medium) is greater than the concentration that may pose an unacceptable risk or hazard to human health, and the analyte is not detected in any sample, then the data for that analyte will be deemed suspect. Data from these analytes will not be used quantitatively in the risk assessment, but the potential risk or hazard from exposure to media potentially containing these analytes will be examined qualitatively. In developing the qualitative assessment for these data, the maximum quantitation limit for the analyte (in all samples from a medium) will be compared to the appropriate no action residential PRG if historical or process information indicates that the analyte potentially could be present. One-half the maximum quantitation limit for the analyte (in all samples from a medium) will be used in this comparison if historical or process information indicates that the analyte is not expected to be present.

Radionuclides. The analysis for radionuclides will be performed in two steps. In the first step, the MDC/minimum detectable concentration/minimum quantification concentration (MQC) for each analyte and sample will be examined to determine if these limits were below the concentration or activity at which the analyte may pose an unacceptable risk (or dose). If the maximum MDC/MQC for an analyte over all samples within a medium is greater than the concentration or activity that may pose an unacceptable risk (or dose) to human health and the analyte is less than the minimum detectable activity MDC/MQC in any samples, then the data for that analyte will be deemed suspect. The MDCs used for radionuclides should be the MDCs established in the MARLAP Manual (EPA 2004b), which provides guidance for evaluating SQLs for radionuclide data. For radionuclides,

Evaluation of Sample Quantitation Limits

Chemicals:

Consider the following results for Chemicals W, X, Y, and Z. Assume that Chemicals W and Y are site-related contaminants and that Chemicals X and Z are not site-related. Also, let the data qualifier (U) be defined as not detected at the sample quantitation limit (SQL).

Chemical	Sample 1	Sample 2	Sample 3	Sample 4	Screening Value
W	10U	10U	10U	10U	5
X	10U	10U	10U	10U	5
Y	10U	6	10U	10U	5
Z	1U	1U	1U	1U	5

Then, following the rules in Step 3 of the data evaluation process:

- Results for Chemical W are suspect because the maximum SQL overall results (10) is greater than the screening value (5), and Chemical W was not detected in any sample. Because Chemical W is site-related, the qualitative risk analysis of this chemical’s potential effect would use the full SQL.
- Results for Chemical X are suspect because the maximum SQL overall results (10) is greater than the screening value (5), and Chemical X was not detected in any sample. Because Chemical X is not site related, the qualitative risk analysis of this chemical’s potential effect would use one-half the SQL.
- Results for Chemical Y are not suspect even though the maximum SQL exceeds the screening value because Chemical Y was detected in one sample.
- Results for Chemical Z are not suspect because the maximum SQL is less than the screening value.

For radionuclides, SQLs should be evaluated in accordance with the guidance in the Multi-Agency Radiological Laboratory Analytical Protocols (MARLAP) Manual (EPA 2004b).

Note: Other data qualifiers associated with the data must also be considered during data evaluation. Please see Step 4 of the data evaluation process.

**Figure 3.3. Example of Step 3–Evaluation of Sample Quantitation Limits
Laboratory Analytical Data**

all reported values, including negative values,⁶ will be used to derive the EPCs under current conditions.

Survey-type data. When XRF data are used in the derivation of EPCs, all XRF values, including negative values, will be used as reported. Other survey-type data (such as PCB field test kits) should be used in accordance with project-specific review of the data and performance of the method.

- **Step 4: Evaluation of data qualifiers and codes.** Generally, the rules presented in RAGS, Part A, Exhibits 5.4 and 5.5 (EPA 1989a) will be used to evaluate all data qualifiers and codes attached to analytical results for chemicals; however, data with a “B” qualifier (i.e., analyte also found in associated blank) will be examined by analyte to ensure that site-related analytes are not eliminated. For other analytes, the “5 and 10X’s Rule” described in RAGS, Part A, (EPA 1989a) will be considered. In addition, the method used in data validation to examine blank contamination will be evaluated. If data validation qualified sample results as “U” (i.e., analyte not detected) instead of “B”

⁶ Negative results may be reported due to a statistical determination of the counts seen by a detector, minus a background count.

when blank contamination was present and the analyte passed the “5 and 10X’s Rule,” then the data will be reevaluated. Specifically, if chemical data are qualified “B,” and the value is less than that defined by the “5 and 10X’s Rule,” then the data will be assumed to be a nondetect and the reported value will be used to derive the EPC.

- Evaluation of radionuclide data will follow rules agreed upon by the Commonwealth of Kentucky Radiation Health Branch [formerly the Kentucky Radiation Health and Toxic Agents Branch (KYRHTAB)] and DOE (RAWG 2000a through 2000f). The data assessment qualifiers that will appear and their description are as follows:
 - **KYRHTAB-LT:** KYRHTAB has performed an independent data assessment and the results are less than the MDC or detection limit and should not be plotted.
 - **KYRHTAB-50:** KYRHTAB has performed an independent data assessment and the radiation counting uncertainty is greater than 50% of the analytical results.
 - **KYRHTAB-ER:** KYRHTAB has performed an independent data assessment and the data present error problems (i.e., no counting uncertainty or zero counting uncertainty).
 - **KYRHTAB-OK:** KYRHTAB has performed an independent data assessment and the data are acceptable for use.
- **Step 5: Elimination of analytes not detected.** Generally, any chemical not detected in at least one sample from a medium will be deleted from the data set. Any radionuclide for which no analytical results exceed its MARLAP MDC also will be deleted from the project dataset, provided the MDC is an acceptable level for the project.⁷ If a chemical analyte is suspected of being present at very low concentrations (i.e., below the quantitation limit) due to cross-media contamination or is suspected of being present based on historical or process information, the analyte may remain in the data set even though the analyte was not detected. In this case, the concentrations used to determine the representative or EPC for the analyte will be the sample quantitation limits for the analyte in the medium. For classes of analytes such as polycyclic aromatic hydrocarbons (PAHs), PCBs, and dioxins/furans, if one compound is detected at a concentration greater than a screening value and is identified as a COPC, then others in that class will be assumed to be present as well. The method used to analyze these classes of compounds is presented later in this section.
- **Step 6: Examination of toxicity of detected analytes.** The maximum concentrations and activities of analytes remaining in the data set will be compared to no action residential use risk-based PRGs by medium. The PRGs used in this comparison will be the lesser of the lifetime excess cancer-based and child hazard-based no action values found in Appendix A. Those analytes with a maximum detected concentration less than each respective no action risk-based PRG will be eliminated from the data set unless the analyte has a bioaccumulation factor for fish equal to or greater than 100 (DOE 1996d). Note: The uncertainty introduced through the application of this screening procedure will be examined quantitatively in the uncertainty analysis portion of the baseline risk assessment. The derivation of the risk-based PRGs used in this comparison is described in Appendix B of this document.
- **Step 7: Examination of analyte concentrations of essential nutrients detected in site samples.** Analytes not removed from the data set in previous steps will be examined to determine if any are

⁷ These types of decisions (acceptable MDCs) would be a product of the consensus of the FFA parties arrived at during project discussions at the appropriate stage in document development.

essential nutrients. Seven analytes known to be essential nutrients and known to be toxic only at extremely high concentrations will be removed from the data set on the basis of regulatory guidance (EPA 1995). These analytes are calcium, chloride, iodine, magnesium, potassium, sodium, and phosphorus. No other analytes known to be essential nutrients will be deleted from the data set on the basis of this screen. Any uncertainty regarding retention of essential nutrient in the list of COPCs will be discussed in the uncertainty section of the risk assessment.

- **Step 8: Comparison of analyte concentrations detected in soil and groundwater samples to analyte concentrations detected in background.** This comparison is performed as part of the development of the list of COPCs. As a first step, maximum detected concentrations of analytes will be compared to the background concentrations presented in Appendix A. Analytes not detected at a concentration greater than the background concentration will not be retained as COPCs. Analytes detected at concentrations greater than their background concentration may be retained as COPCs, depending upon the outcome of other screening steps. Analytes retained as COPCs, however, may be considered with the full range of background as part of the uncertainty analysis. This analysis, if completed, will be done to determine if the analyte is generally present at concentrations above its background concentration or if the detected concentrations of the analyte above the selected background concentration is consistent with natural enrichment. The impacts on risk characterization of not retaining an analyte on the basis of the background screen will also be considered in the uncertainty analysis.

During the development of the list of COPCs, concentrations of total cancerous PAHs, PCBs, and dioxins/furans (dioxins) will be derived. Total PAHs, total PCBs, and total dioxins will be derived to allow for the correct use of the toxicity screen described in Step 6 and to allow for correct calculation of ELCR from exposure to these organic compounds.

When deriving total PAHs, the toxicity equivalence factors (TEFs) presented in Human Health Risk Assessment Protocol for Hazardous Waste Combustion Facilities (EPA 2005) will be used. These TEFs are presented in Table 3.1. Note that these TEFs will be applied to the concentrations of detected PAHs in each sample and that the total PAH concentration in a sample will be the sum of the products of each PAH and its TEF. For samples in which PAHs are not detected, the value for the minimum detection limit of the PAHs with TEFs will be used in the calculation of the EPC.

When deriving total PCBs [if this analyte (i.e., Total PCBs) is not reported in the data set], the detected concentrations of each PCB within a sample will be summed. For samples in which no PCBs are detected, the value for the minimum detection limit of the PCBs will be used in the calculation of the EPC. If there are detection limits for PCBs exceeding risk-based concentrations, this issue should be discussed in the uncertainty section. Note that there are no TEFs to use when deriving total PCBs from individual Aroclors. If dioxin-like PCBs are detected at a site, they should be added to the total PCBs after weighting with the TEFs for those compounds in Van Den Berg, et al. 2006.

When deriving total dioxin, the TEFs presented in *Federal Register*: May 10, 2007 (Volume 72, Number 90), *Dioxin and Dioxin-like Compounds; Toxic Equivalency Information* will be used. These TEFs are presented in Table 3.1. Note that these TEFs will be applied to both the concentrations of detected dioxins and furans and to one-half the sample quantitation limit of undetected dioxins and furans, when one or more dioxin or furan is detected. The total dioxin concentration in a sample will be the sum of the products of each dioxin/furan and its TEF. For samples in which no dioxin or furan was detected, the minimum detection limit for 2,3,7,8-TCDD will be used as the value for the total dioxin concentration. If there are detection limits for dioxins and furans exceeding risk-based concentrations, this issue should be discussed in the uncertainty section.

Table 3.1. Toxicity Equivalency Factors for PAH Compounds and Dioxins/Furans

PAH Compound ¹	Toxicity Equivalence Factor	Dioxin/Furan Compound ²	Toxicity Equivalence Factor
Benzo(a)pyrene	1.0	2,3,7,8-TCDD	1.0
Benzo(a)anthracene	0.1	1,2,3,7,8-PeCDD	1.0
Benzo(b)fluoranthene	0.1	1,2,3,4,7,8-HxCDD	0.1
Benzo(k)fluoranthene	0.01	1,2,3,6,7,8-HxCDD	0.1
Chrysene	0.001	1,2,3,7,8,9-HxCDD	0.1
Dibenzo(a,h)anthracene	1.0	1,2,3,4,6,7,8-HpCDD	0.01
Indeno(1,2,3-c,d)pyrene	0.1	OCDD	0.0003
All other PAHs	0	2,3,7,8-TCDF	0.1
		1,2,3,7,8-PeCDF	0.03
		2,3,4,7,8-PeCDF	0.3
		1,2,3,4,7,8-HxCDF	0.1
		1,2,3,6,7,8-HxCDF	0.1
		1,2,3,7,8,9-HxCDF	0.1
		2,3,4,6,7,8-HxCDF	0.1
		1,2,3,4,6,7,8-HpCDF	0.01
		1,2,3,4,7,8,9-HpCDF	0.01
		OCDF	0.0003

¹TEFs from EPA 2005

²TEFs from Van Den Berg, et al. 2006

3.3.3.3 Presentation of data evaluation

A summary of the data evaluation will be provided in both narrative and tables. Tables from each step of the data evaluation process may be presented. The detailed data tables, if voluminous, should appear in an appendix to the risk assessment; however, the summary tables described earlier (see Section 3.3.2.1) should appear in the main text of the assessment. At minimum, a table listing the COPCs for the assessment should appear in the main text. An example of the information that should appear in this summary table is in Exhibit 3.8.

Exhibit 3.8. List of Chemicals of Potential Concern

Analyte	Frequency of Detection ¹
Site and Medium²	
Analyte # 1	
Analyte # 2	
.	.
.	.
.	.
Analyte # N	

¹ This value will be the number of samples in which the analyte was detected over the number of samples in which an analysis for the analyte was performed.

² A list of chemicals of potential concern will be presented for each site and medium combination.

3.3.3.4 Site-specific characterization information

Several pieces of site-specific characterization information are relevant to virtually all baseline human health risk assessments performed for PGDP because they explain resource use around PGDP. Because this information is in the form of interviews and letters, it generally is not readily available; therefore, this

information is included in Appendix E of this document to provide a ready source of these materials. Appendix E, presents the following documentation.

- Reference to the Phase I Site Investigation results of surface water and groundwater users survey to determine groundwater use near PGDP (CH2M HILL 1991);
- Summary of agricultural practices in Ballard County, Kentucky;
- Summary of the agricultural practices in McCracken County, Kentucky;
- Area of crop land in Ballard and McCracken County, Kentucky;
- Recreational use of Bayou and Little Bayou Creeks near PGDP;
- Annual harvests of geese, ducks, turkeys, and deer in McCracken and Ballard Counties, Kentucky;
- Reports entitled “Planning Issues for Superfund Site Remediation” and “Quantitative Decision Making in Superfund: A Data Quality Objectives Case Study” from *Hazardous Materials Control* regarding use of exposure units in risk calculations and remedial decisions;
- Kentucky Risk Assessment Guidance, Risk Assessment Branch, Department for Environmental Protection, Commonwealth of Kentucky;
- Kentucky Guidance for Ambient Background Assessment, Risk Assessment Branch, Department for Environmental Protection, Commonwealth of Kentucky, January 8, 2004;
- Kentucky Guidance for Groundwater Assessment Screening, Risk Assessment Branch, Department for Environmental Protection, Commonwealth of Kentucky, January 15, 2004;
- Trichloroethylene Environmental Levels of Concern, Risk Assessment Branch, Department for Environmental Protection, Commonwealth of Kentucky, April 2004;
- Environmental Indicators flowchart submitted to the Hazardous Waste Branch of the Kentucky Division for Waste Management;
- Information used to compile PGDP background;
- The table of parameters for probabilistic risk assessment (PRA) from the Southwest Plume Investigation report. This table provides the parameter values used for the PRA in that report, which should be considered for use in other PRAs. The values in the table do not represent specified default values for use in all PRAs.

3.3.4 Exposure Assessment Methods

The primary purpose of this section of the baseline human health risk assessment will be to report the results of the exposure assessment for each unit or area investigated. In this section, the exposure setting for each unit or area will be characterized, exposure pathways will be identified, exposure will be quantified (i.e., dose or intake calculated), and doses will be presented. Methods to complete each of these steps are discussed in the following sections.

3.3.4.1 Characterize the exposure setting

This section of the exposure assessment or other portions of the document will describe the physical setting of each unit, including meteorology, climate, vegetation, soil type, surface hydrology, groundwater hydrology, and geology. In addition, the surrounding populations will be characterized as needed. Specific note will be given to determining if sensitive subpopulations may be present. In risk assessments in RI reports, the information presented concerning climate, vegetation, soil type, surface hydrology, groundwater hydrology, and geology will be brief, and references will be to material presented in earlier sections of the RI report. (Note: A brief presentation of this material must be included in the baseline risk assessment because the FFA states that the baseline risk assessment is to be written as a stand-alone report.) In baseline risk assessments not in RI reports, the information presented concerning climate, vegetation, soil type, surface hydrology, groundwater hydrology, and geology will be more extensive.

Current and potential future land use and the time frame for future use also will be discussed in this section of the exposure assessment. The most likely future land use will be determined using information in the most recent PGDP Site Management Plan (SMP); however, because future land use over time is uncertain, the use scenarios considered in the baseline risk assessment will not be governed by that information alone. Use scenarios that will be considered in all baseline risk assessments under future conditions are rural residential, recreational, industrial, outdoor worker, and excavation. Appropriate use scenarios may be evaluated during project scoping.

Finally, this section of the baseline human health risk assessment will integrate the preceding information and declare the unit or area under investigation either as a source or integrator unit and identify exposure points. Definitions used to determine whether the area or unit is a source or integrator are as follows:

- **Source unit.** Those units or areas that may release contaminants to other units or areas.
- **Integrator unit.** Those units or areas that accumulate contaminants from source units or areas.

Generally, application of these definitions to units and areas to be investigated at PGDP shows that all areas on-site where contamination exists (e.g., the soil and other material at burial grounds, spill areas, and landfills) are source areas. Integrator units identified using these definitions are air, groundwater (e.g., RGA), and surface water (e.g., Bayou and Little Bayou Creek watersheds and the Ohio River).

Also in this section of the exposure assessment, exposure points will be evaluated. For source units, the exposure points that will be evaluated under current conditions are at the unit or area (“hot spots” may be evaluated separately) and at points downgradient to which contamination may migrate. Downgradient points that will be evaluated for risk communication purposes include at the PGDP industrialized area [i.e., the area corresponding to the industrial land use delineated in the SMP (DOE 2014)] (if applicable), at the DOE property boundary (if applicable), and at Little Bayou Creek (if applicable). Note that for units or areas outside the industrialized area at PGDP, exposure at the industrialized area will not be considered because it is not necessary for remedial decisions. For integrator units, exposure points that will be considered are those within the contaminated area (e.g., above the contaminated groundwater plume or along the contaminated ditch) and at areas downgradient. Generally, exposure points that consider migration from a source will consider the time of exposure. For example, for exposure to groundwater both at a source and at the facility boundary, risk or hazard from exposure to measured concentrations under current conditions and future conditions will be determined. In addition, risk or hazard from exposure to expected future concentrations or activities will be modeled to determine the risk or hazard that may occur under potential future conditions as contaminants migrate from the source to the underlying aquifer. Exposure to contaminants in or migrating to the surface water integrator unit will be handled similarly. The mechanism that will be

<p>Industrialized Area Area corresponding to the industrial land use delineated in the Site Management Plan.</p>

used to determine the extent of modeling that will be used in a baseline human health risk assessment is discussed later.

3.3.4.2 Identification of exposure pathways

This section of the exposure assessment will delineate the pathways through which the receptors may be exposed under both current and future conditions. For current receptors, these pathways and their parameters should be based on realistic exposures; for future receptors, these pathways and their parameters should be based on reasonable maximum exposure (RME) values. The goal of this material will be to provide a complete depiction of all exposure pathways for current and future uses. To achieve this goal, this section will present conceptual site models and supporting text. Also, in this section, each pathway will be described in terms of source, exposure route, exposure point, and receptor. This format will be followed because all four must be present for a complete pathway to exist. Note: Potential pathways not containing all four items will be described as being incomplete, and text justifying their omission from the assessment will be provided. Potential pathways that will be considered in all assessments are described herein.

Exposure assessments in baseline human health risk assessments completed in the past indicate that at least 24 exposure pathways should be considered as potential pathways in all assessments. These pathways are listed. (Note: Additional pathways, such as contact with buried waste, may be reasonable for some units or areas; these pathways are not included.)

- Ingestion of groundwater as a drinking water source
- Inhalation of volatile constituents emitted from groundwater during household use
- Dermal contact with groundwater while showering
- External exposure to ionizing radiation emitted by constituents in groundwater while showering
- Inhalation of volatile constituents emitted from groundwater during irrigation
- Incidental ingestion of soil
- Dermal contact with soil
- Inhalation of particulates emitted from soil
- Inhalation of volatile constituents emitted from soil
- External exposure to ionizing radiation emitted by constituents in soil
- Incidental ingestion of surface water while swimming or wading in creeks or natural or man-made ponds
- Dermal contact with surface water while swimming or wading in creeks or natural or man-made ponds
- External exposure to ionizing radiation emitted by constituents in surface water while swimming or wading in creeks or natural or man-made ponds
- Incidental ingestion of sediment while swimming or wading in creeks or natural or man-made ponds

- Dermal contact with sediment while swimming or wading in creeks or natural or man-made ponds
- External exposure to ionizing radiation emitted by constituents in sediment while swimming or wading in creeks or natural or man-made ponds
- Consumption of fish taken from creeks or natural or man-made ponds
- Consumption of vegetables and produce raised in contaminated soil
- Consumption of irrigated vegetables
- Consumption of beef from animals contaminated by consuming vegetation (pasture and concentrates) irrigated with contaminated water or grown on contaminated soil, by drinking contaminated water, or ingesting contaminated soil
- Consumption of dairy products (i.e., milk) from animals contaminated by consuming vegetation (pasture and concentrates) irrigated with contaminated water or grown on contaminated soil, by drinking contaminated water, or ingesting contaminated soil
- Consumption of pork from animals contaminated by consuming vegetation (concentrates) irrigated with contaminated water or grown on contaminated soil or by drinking contaminated water
- Consumption of poultry products from animals drinking contaminated water
- Consumption of game (i.e., deer, rabbits, and quail) contaminated by consuming contaminated vegetation or soil and ingesting water.

While these pathways have been found to be reasonable in past assessments, not all may be reasonable, or complete, for future assessments; therefore, the decision as to which pathways to quantify will be made on a project-specific basis. In any case, the rationale for the inclusion or exclusion of any of the pathways listed herein will be included in the exposure assessment.

It is important to note that the pathways relating to livestock consumption are not reasonable for most source units. This is because most source units are too small to support livestock in addition to a homestead and garden. Generally, a source unit will be required to be larger than two acres to be considered for livestock production. (This requirement assumes that a minimum of two acres is required for a home and associated garden.) Note: Under this definition, all integrator unit assessments will contain an assessment of risk from consumption of livestock because the area they cover is greater than two acres. In assessments where livestock consumption is included, the range size for each beef or cow will be two acres per head (Morrison 1959).

Using the characterization information and pathway analysis, a conceptual site model will be developed for each unit or area. The format that will be used for the conceptual site models is that in Figure 3.1. Note: When presenting the conceptual site models for multiple units or areas in a single baseline human health risk assessment, the units or areas may be grouped to reduce the number of figures that need to be presented.

3.3.4.4 Quantification of exposure

To quantify exposure or dose, both the EPC and the exposure factors are required. Here, the EPC can be defined as the concentration or activity of the COPC in the environmental medium ingested, inhaled, contacted, or consumed, and the exposure factor can be defined as the product of the exposure parameters describing the degree of exposure to the environmental medium in terms of duration or frequency of exposure and mass of the receptor.

EPCs under current conditions of all COPCs for which environmental samples were taken will be determined using the following procedure.

- (1) If results from fewer than ten samples are available, then the EPC will be the maximum detected concentration.
- (2) If results from ten or more samples are available, then the most recent version of EPA's ProUCL software version of EPA's ProUCL software will be used to determine the EPC. The value selected as the EPC will be the value recommended by ProUCL, noted as the "Potential UCL to Use." EPA's ProUCL software⁸ incorporates a number of different distributional tests that may be used to perform the distributional tests and calculates the most appropriate EPC (EPA 2010).

Options to determine the ten or more samples may include use of grid values. It is recommended that a geostatistical approach utilizing Spatial Analysis and Decision Assistance (SADA) or similar software be used to estimate values for empty grids. SADA is available at <http://www.tiem.utk.edu/~sada/index.shtml>. Alternately, an average value may be used. An example is shown in the text box [from Soils Operable Unit RI Report (DOE 2012)]. These options should be discussed and agreed to in the planning phases of projects.

In determining the UCL when the medium is soil, data will be segregated into depth intervals relevant to receptors. For all scenarios except the outdoor worker, data from samples collected from 0 to 1 ft bgs will

From Soils Operable Unit RI Report (DOE 2012):

The representative sampling design for the SWMUs was gridding. In some instances (such as SWMUs/AOCs not grid sampled in summer 2010), when a grid was applied to the SWMUs/AOCs, a grid lacking a sample result resulted. In order to fill a grid lacking a sample result, the average of the grids within the exposure unit with sampling results was used. Attachment D2 presents an uncertainty evaluation in determining EPC values using these averages against EPC values calculated without using the averages or the maximum value, as applicable. An example for determining the EPC through averaging is illustrated below.

If the SWMU/exposure unit combination had less than 10 grids, the maximum grid result was used as the EPC. If the SWMU/exposure unit combination had 10 or more grids, the grid values were used to determine the EPC. Grid values were determined following guidance in the work plan. Basically, the maximum detected result from within the grid applies to the grid. If not detected, the minimum detection limit applies to the grid.

If a grid had no result (detect or nondetect) for the COPC, an average of the results for the grids with results was used.

⁸ Software is available at www.epa.gov/osp/hstl/tsc/software.htm.

be used to estimate the EPC.⁹ For excavation activities performed by the outdoor worker, data collected from 0 to 10 ft bgs will be used to estimate the EPC, unless information indicates that results from samples collected at deeper depths (i.e., 0–16 ft bgs in areas where infrastructure is found) should be included in the derivation of the EPC.

In determining the UCL when the medium is groundwater, data from samples from each potable aquifer (i.e., RGA and McNairy Formation) will be used; however, data will be summarized within and not over aquifers. Note: For the groundwater integrator investigations (e.g., that for the Groundwater Operable Unit), the representative concentration for groundwater may be the average concentration of the samples taken from wells within the contaminant plume if data are sufficient. In addition, as with soil, the wells used in each calculation may be grouped so that risk or hazard at differing contaminant concentrations and in various areas may be estimated. Decisions concerning the method that will be used to estimate the concentration of COPCs for the groundwater integrator unit will be made on a case-by-case basis and will be justified in the baseline risk assessment.

Risks from water drawn from the UCRS will not be presented in the main body of the risk assessment because this water source is not considered to be an aquifer due to low yield. However, risks from ingestion of water from this source will be considered at least qualitatively in the uncertainty section of the risk assessment.

Finally, for some samples, duplicate or split-sample analyses may be available. When calculating the representative concentration, the maximum value reported in the duplicate or split-sample analysis will be used. Duplicate and split-sample results will not be averaged when calculating the representative concentration in baseline risk assessments performed for PGDP.

The EPCs and activities used for future conditions will depend on the time frame for which risk or hazard is being quantified. At minimum, for all assessments for PGDP, risk and hazard to potential future users, will be quantified using the current EPCs and activities. In addition, for those sites and areas where future concentrations or activities may increase, modeled concentrations will be used. To determine if modeling is needed, the maximum soil concentrations and activities at the source (over all depths) for each analyte will be compared to the appropriate groundwater protection PRG (PRGs appear in Appendix A). If the maximum soil concentration exceeds the groundwater protection PRG, then future concentrations in groundwater and surface water (if appropriate) will be modeled. Models to be used to determine future concentrations and activities at the source and in groundwater will be based on the modeling matrix presented in Table 3.2. Tier 1 values are existing sets of screening levels used for the initial screening of a site. Tier 2 values also are used for scoping, but account for more specific estimates of model parameters than the default Tier 1 values. Tiers 3 and 4 are models used with primarily site-specific values for site decision making.

⁹ A single set of exposure equations and parameters is provided for the outdoor worker scenario in Appendix D. The exposure parameters provided in Appendix D should be used without changes when assessing potential risk from exposure to surface soils in locations outside the industrialized area at the Paducah site. When assessing potential risk from exposure to both surface and subsurface soil in locations outside the industrialized areas, all exposure parameters, except exposure duration (ED) and exposure frequency (EF), should be used without changes to assess an outdoor worker. ED and EF for exposure by the outdoor worker to surface and subsurface soil in locations outside the industrialized area should be established considering guidance in the Exposure Factors Handbook (EPA 1997b) or similar sources (e.g., site-specific information) and should be documented. Similarly, when assessing potential risk from exposure either to surface soil or subsurface soil in locations inside the industrialized area, all exposure parameters, except ED and EF, should be used without changes to assess both an outdoor worker and a construction/excavation worker. As above, ED and EF for exposure by the outdoor worker or a construction/excavation worker to surface soil or subsurface soil or both in locations inside the industrialized area should be established considering guidance in the Exposure Factors Handbook (EPA 1997b) or similar sources and should be documented.

Table 3.2. Modeling Matrix for Groundwater, Surface Water, and Biota

	Values for Soil to Protect Groundwater	Model	Point of Exposure	Notes
INVESTIGATION DOCUMENTS	Tier 1 (Used for scoping)	SSLs and/or RESRAD	At source unit	Value to be used for initial scoping, use DAF of 1 for SSLs unless site-specific values are available. Groundwater Protection value based on residential use and targets of 1E-6, 0.1, and 1 for risk, hazard, and dose, respectively. If site-specific DAF values are used, then need to justify these values. The depth of water needs to be considered in the calculation.
		Vapor intrusion model	At source unit	Initial vapor intrusion model will use default values.
	Tier 2 (Used for scoping)	SESOIL and/or RESRAD	At source unit	Includes source delimitation. Recognize SESOIL limitations when modeling inorganic COPCs-refine K_{ds} .
DECISION DOCUMENTS	Tier 3 (Enhanced modeling used in decision documents if needed)	SESOIL and RESRAD suite of codes (including RESRAD-OFFSITE) with AT123D	At source unit and at Downgradient points (Industrialized area, DOE property boundary, creek, river)	Uses source delimitation and refined K_{ds} from above. Use values from this effort to set initial cleanup levels. On the Terrace (southern portion of PGDP), different points of exposure will apply.
	Tier 4 (Enhanced modeling used in decision and design documents if needed)	Source modeling and MODFLOW/MT3D/RT3D	At source unit and at Downgradient points appropriate to the selected remedy	To be used to refine cleanup levels (if needed). May be especially important to set monitoring goals. On the Terrace (southern portion of PGDP), different points of exposure will apply.

Table 3.2. Modeling Matrix for Groundwater, Surface Water, and Biota (Continued)

	Values for Soil and Sediment to Protect Surface Water	Model	Point of Exposure	Notes
INVESTIGATION DOCUMENTS	Tier 1 (Used for scoping)	SSLs and/or RESRAD	At source unit	Value to be used for initial scoping by Project Team. Use DAF of 1 for SSLs. Groundwater Protection value based on recreational use and targets of 1E-6, 0.1, and 1 for risk, hazard, and dose, respectively. If site-specific DAF values are used, then need to justify these values.
	Tier 2 (Used for scoping)	MUSLE	At source unit	Includes source delimitation. Value to be used during follow-up meetings by Project Team.
DECISION DOCUMENTS	Tier 3 (Enhanced modeling used in decision documents if needed)	SWMM	At source unit and at Downgradient points (Industrialized area, creek)	Uses source delimitation from above. Initial cleanup level calculations.
	Tier 4 (Enhanced modeling used in decision and design documents if needed)	Enhanced SWMM	At source unit and at Downgradient points appropriate to the selected remedy (Industrialized area, creek)	To be used to refine cleanup levels (if needed). May be especially important to set monitoring goals.

Table 3.2. Modeling Matrix for Groundwater, Surface Water, and Biota (Continued)

	Values for Soil and Sediment to Protect Biota	Model	Point of Exposure	Notes
INVESTIGATION DOCUMENTS	Tier 1	NONE	NONE	The RAWG determined that development of screening values based on biota modeling would not be appropriate; therefore, these values do not exist.
	Tier 2 (Used in Baseline Risk Assessments)	Those contained in current Methods Document, Appendix D	At source unit	Includes source delimitation.
DECISION DOCUMENTS	Tier 3 (Enhanced modeling used in Decision Documents if needed)	Those contained in current Methods Document, Appendix D for biota and transport models presented earlier for receiving media.	At source unit and at Downgradient points (Industrialized area, creek)	Uses source delimitation from above. Initial cleanup level calculations.
	Tier 4 (Enhanced modeling used in Decision and Design Documents if needed)	Those contained in current Methods Document, Appendix D for biota and transport models presented earlier for receiving media.	At source unit and at Downgradient points (Industrialized area, creek)	To be used to refine cleanup levels (if needed). May be especially important to set monitoring goals.

AT123D = Analytical Transient 1-, 2-, 3-Dimensional Simulation of Waste Transport in the Aquifer System

COPC = chemical of potential concern

DAF = dilution/attenuation factor

HHRAWG = Human Health Risk Assessment Working Group

K_d = adsorption coefficient/distribution coefficient

PGDP = Paducah Gaseous Diffusion Plant

RESRAD = Residual Radiation

SESOIL = Seasonal Soil Model

SSL = Soil Screening Level

SWMM = Storm Water Management Model

MODFLOW/MT3D/RT3D = three-dimensional finite-difference groundwater model

MUSLE = Modified Universal Soil Loss Equation

Because all models contain significant uncertainty, the baseline risk assessment's analysis of off-site migration also will include risks calculated using current contaminant concentrations at source units in addition to modeled values. This analysis will be included in the uncertainty section of all baseline risk assessments that contain modeling.

In baseline risk assessments for the integrator units, analyte degradation, attenuation, and transformation will be considered in addition to migration when calculating future concentrations, if possible. The analysis of these factors will rely upon the analysis presented in earlier sections of the remedial investigation report.

The equations to be used to combine the EPCs and exposure factors to estimate dose will follow the general format presented in RAGS, Part A (EPA 1989a). This general equation is shown in Equation 5. Specific equations are presented in Appendix D of this document. In this appendix, references are presented for each exposure parameter (e.g., CR, BW) included in the equation. Generally, these parameters were taken from guidance documents (e.g., EPA 1989a; KDEP 2002) unless site-specific values are available. (Equations used to derive radionuclide dose are similar to those presented in Appendix D.)

$$\text{Intake} = C \times \frac{\text{CR} \times \text{EFD}}{\text{BW}} \times \frac{1}{\text{AT}} \quad \text{Eq. 5}$$

where: Intake = The dose (mg/(kg × day))

C = The average concentration contacted over the exposure period. See Eqs. 6 and 7 and associated discussion.

CR = The contact rate or amount of contaminated medium contacted per unit time or event.

EFD = The exposure frequency and duration describing how long and how often exposure occurs.

BW = The average body weight of the receptor over the term of exposure.

AT = The averaging time or period over which exposure is averaged.

In the material in Appendix D, equations that can be used to calculate the concentrations of COPCs in selected biota (e.g., vegetables, fish, game, and livestock) also are presented. Generally, for baseline human health risk assessments for source units inside the industrialized area at PGDP, concentrations of COPCs in biota will be estimated using these equations because biota sampling cannot be performed. (These biota are not present.)

For assessments for source units outside the industrialized area and for integrator unit baseline risk assessments, results from biota sampling may be available. In cases where this information is available, the EPC will be calculated using the methods presented earlier in this section. In cases where this information is not available, the equations presented in Appendix D will be used to estimate the concentrations in biota. (Note: Because concentrations in biota can differ markedly with time of sampling, tissue sampled, species sampled, age of animal, and other factors, the use of analytical results from biota sampling in the risk assessment also may give results that are very uncertain; therefore, the uncertainty in the results calculated using biota analytical results also will be considered completely.)

3.3.4.5 Presentation of the results of the exposure assessment

Several figures and tables will be used to report the results of the exposure assessment in baseline human health risk assessments performed for PGDP. As noted earlier, conceptual site models for each unit, group of units, or area under investigation will be presented, and tables presenting exposure and risk information will be prepared. In addition, this section also will present a summary of the decisions made concerning the selection of pathways to be quantified for each unit or area under investigation; the representative (i.e., exposure point) concentration of COPCs in each medium, including biota; any chemical-specific

values used in the calculations; and the daily intakes resulting from the application of the exposure equations.

The material appearing in this summary will be taken from the larger tables presented in the appendix to the risk assessment. Formats to present this summary information are in Exhibits 3.9–3.12.

Exhibit 3.9. Summary of Pathway Analysis in the Exposure Assessment

Potentially Exposed Population	Exposure route, medium, and exposure point ¹	Pathway selected? (yes/no)	Reason for pathway selection or dismissal ²
Time period ³			
Population 1 ⁴			
	Pathway 1		
	Pathway 2		
	.	.	.
	.	.	.
	.	.	.
	Pathway N		

¹ Each of the pathways presented in this section will be included.

² A short statement drawn from the discussion in the text will be provided for the decision.

³ Summary tables will be prepared for both the current or future time period. If multiple future time periods are assessed, a summary table will be included for each.

⁴ The populations will be rural residential, recreational, industrial, and excavator. Only populations relevant to the time period will be included.

Exhibit 3.10. Presentation of Exposure Point Concentrations¹

Chemical of Potential Concern ²	Medium 1 ³	Medium 2	...	Medium N
Unit or Area 1 ⁴				
Analyte 1			...	
Analyte 2			...	
.
.
.
Analyte N			...	

¹ A table will be made for each time period if models are used to estimate future representative concentrations.

² All chemicals of potential concern across all media will be presented for each unit or area.

³ All media will be listed. The order will be groundwater, soil, sediment, surface water, and biota if possible. More than one EPC may be derived for a media if different depths are used for exposures under different scenarios.

⁴ Each unit or area will be presented separately, but only one table will be used if possible.

Exhibit 3.11. Chemical-Specific Parameters

Chemical of Potential Concern ¹	Parameter 1 ²	Parameter 2	...	Parameter N
Analyte 1			...	
Analyte 2			...	
.
.
.
Analyte N			...	

¹ All chemicals of potential concern over all units or areas investigated will be presented. A separate list will not be presented for each unit unless unit-specific, chemical-specific parameters are used in the assessment.

² All chemical-specific parameters will be listed so that the calculations in the assessment can be duplicated by reviewers or users.

Exhibit 3.12. Daily Intakes (Dose) for Receptor 1¹

Chemical of Potential Concern ²	Pathway 1 ³	Pathway 2	...	Pathway N
Unit of Area 1 ⁴				
Analyte 1			...	
Analyte 2			...	
.
.
.
Analyte N			...	

¹ A separate table will be made for each receptor. If use patterns are assumed to differ between time periods, separate tables for each time period will also be provided.

² COPCs across all media will be listed for each unit or area.

³ Each pathway included in the assessment will be listed. The order followed will be groundwater pathways, soil pathways, surface water pathways, sediment pathways, and biota pathways, if possible.

⁴ A separate presentation will be made for each unit or area; however, only one table will be used if possible.

3.3.4.6 Probabilistic Risk Assessment

Initially, all baseline risk assessments will be conducted as deterministic (point estimate) risk assessments. COPCs with high variability and uncertainty in exposure concentrations or for which individual exposure parameters greatly influence the risk or hazard estimate may be considered for PRAs. These assessments evaluate the variability and uncertainty in risk estimates, and are used to determine the likelihood of exceeding a risk level of concern. PRAs will be conducted following the guidance in *RAGS Volume III-Part A* (EPA 2001a). Scoping is an extremely important component of a PRA to determine which parameters should vary and develop appropriate ranges of values for those parameters. Ranges of values for variables in the risk equations that were used in a previous PRA for the Southwest Plume are provided in Appendix E of this document. The values for variables listed in Appendix E are appropriate as a starting point for other PRAs, but should be reviewed to ensure they are applicable to a specific project and modified if necessary. Documents using PRA also will need to include additional sections providing explanation of how the PRA was conducted, the interpretation of the results, and the appropriate application of the results to decision making to ensure that the PRA and its results are understandable to both the regulatory agencies and the public.

3.3.5 Toxicity Assessment Methods

The primary purpose of this section of the baseline human health risk assessment will be to report the toxic effects of the COPCs on exposed populations. In addition, this section will briefly describe the methods used by EPA and in the toxicity assessment, to develop toxicity parameters, delineate the sources used to acquire the toxicity parameters, and present tables summarizing the toxicity information used in the risk assessment. In closing, this section will summarize the amount of toxicity information available on the COPCs in the risk assessment and discuss general toxicity assessment uncertainties. Requirements for each of these activities are discussed below.

3.3.5.1 Toxicity summaries

A toxicity summary for each COPC will be presented in the toxicity assessment. Each summary will contain a short description of the toxic effects of the chemical and the source of the toxicity values. Included in each description will be information on the effects associated with exposure to the chemical; the concentrations at which adverse effects are expected to occur in humans; a brief description of the database used to derive each toxicity value, including the particular study from which the toxicity value used in risk characterization was derived; and the approval status of any toxicity values. Each toxicity

summary will conclude with a listing of the toxicity values used in the risk assessment for administered and absorbed dose routes of exposure.

3.3.5.2 Sources of toxicity information

The sources that will be used in developing toxicity information for risk assessments performed for PGDP are listed below. These will be examined in the order presented.

- Tier 1 sources: *IRIS* (EPA 2007)
- Tier 2 sources: EPA Provisional Peer Reviewed Toxicity Values
- Tier 3 sources:
 - Health Effects Assessment Summary Tables (HEAST) (EPA 1997c, 2001b)
 - Other sources identified in OSWER Directive 9285.7-53
 - Agency for Toxic Substances and Disease Registry toxicological profiles

When compiling toxicity information, provisional and withdrawn values and toxicity values withdrawn from *IRIS* or *HEAST* will be included, and provisional values will be clearly identified. If toxicity information is not available from the sources listed above, surrogate chemicals with toxicity values may be identified through consideration of chemical structure and characteristics. Selection of surrogate chemicals requires consultation with and approval from EPA and KDEP.

Note: Toxicity values will not be developed for PGDP risk assessments without consultation with the regulatory agencies.

Three additional issues will be addressed when reporting the sources of toxicity information. These are the use of toxicity values for chronic versus subchronic effects, the calculation of toxicity values for absorbed versus administered dose, and the use of oral administered dose toxicity values for the inhalation exposure route. Each of these is discussed herein.

Generally, all risk assessments performed for PGDP will only use toxicity values for chronic exposure when characterizing risk. Although *RAGS, Part A*, (EPA 1989a) states that toxicity values for subchronic exposure should be used for exposure durations less than seven years in length, these will not be used because they are not available for many chemicals (in which case the chronic value should be used). The receptor groups that are affected by this decision are the child rural resident, the recreational user, and the outdoor worker. In no case will toxicity values based on subchronic exposure be used for child or teen receptors. For outdoor workers, toxicity values based in subchronic exposure may be used if the information provided by their use is beneficial in remedial action decision making.

To properly characterize risk from absorbed dose (e.g., dose from dermal absorption across the skin), it is necessary to have toxicity values that are based on absorbed dose. Generally, all toxicity values in *IRIS* and *HEAST* are based on administered dose and cannot be used directly with the chronic daily absorbed doses calculated using the exposure equations in Appendix D. To convert administered dose toxicity values to absorbed dose toxicity values, the guidance provided in *Risk Assessment Guidance for Superfund, Volume I: Human Health Evaluation Manual. Supplemental Guidance, Dermal Risk Assessment, Interim Guidance* (EPA 1992b) will be used. The method delineated in this guidance is depicted in Eqs. 6 and 7. Equation 6 shows that the administered dose toxicity value for cancer effects (administered dose slope factor) is converted to an absorbed dose toxicity value (absorbed dose slope factor) by dividing by the chemical-specific gastrointestinal absorption efficiency of the respective chemical or compound. Equation 7 shows that the administered dose toxicity value for systemic toxicity [administered reference dose (RfD)] are converted to an absorbed dose toxicity value (absorbed RfD) by multiplying by the chemical-specific gastrointestinal absorption efficiency of the respective chemical or compound.

As stated in RAGS Part E (EPA 2004):

For those organic chemicals that do not appear on the table, the recommendation is to assume a 100% ABS_{GI} value, based on review of literature, indicating that organic chemicals are generally well absorbed (>50%) across the GI tract. Absorption data for inorganics are also provided in Exhibit 4-1 [see text box], indicating a wide range of absorption values for inorganics. Despite the wide range of absorption values for inorganics, the recommendation is to assume a 100% ABS_{GI} value for inorganics that do not appear in this table. This assumption may contribute to an underestimation of risk for those inorganics that are actually poorly absorbed. The extent of this underestimation is inversely proportional to the actual GI absorption.

$$Absorbed\ SF = \frac{Administered\ SF}{GI\ Efficiency} \quad \text{Eq. 6}$$

where: Absorbed SF = The absorbed dose slope factor for cancer effects
 Administered SF = The administered dose slope factor for cancer effects
 GI Efficiency = The chemical-specific gastrointestinal absorption efficiency

$$Absorbed\ RfD = Administered\ RfD \times GI\ Efficiency \quad \text{Eq. 7}$$

where: Absorbed RfD = The absorbed reference dose for systemic toxicity
 Administered RfD = The administered reference dose for systemic toxicity
 GI Efficiency = The chemical-specific gastrointestinal absorption efficiency

The dermal dose derived with this methodology provides an estimate of the contribution of the dermal pathway to the systemic dose. Dermal exposure for baseline risk assessments will follow the *Risk Assessment Guidance for Superfund Volume I: Human Health Evaluation Manual* (Part E, Supplemental Guidance for Dermal Risk Assessment) (EPA 2004c). The EPA guidance provides specific values for eleven compounds or groups of compounds in Exhibit 3-4 of the dermal guidance. For an alternative evaluation, providing more restrictive values for the dermal-soil pathway, the default values of 25% dermal absorption for volatile organic compounds (VOCs), 10% dermal absorption from soil for all semivolatile organic compounds (SVOCs) without specific absorption values specified in RAGS, Part E: and 5% dermal absorption from soil for all inorganic compounds without specific absorption specified in RAGS, Part E, may (based on project-specific goals) be applied to a quantitative risk assessment. This approach is consistent with guidance from KDEP. KDEP-specific values for dermal absorption are provided in Appendix B. See text box for additional information. For the dermal-water pathway, absorption should be calculated using the methods described in RAGS, Part E. For inorganic chemicals, the K_p (permeability coefficient) parameter has been identified as one of the major parameters contributing to uncertainty in the assessment of

In RAGS Part E 2004, Exhibit 4-1, the following GI absorption efficiencies are listed that are below the 5% dermal absorption KDEP has recommended as a default value for inorganics. For these constituents, the dermal absorption value should be modified from 5% to mimic the GI absorption efficiencies, as follows:

Beryllium	0.007 = 0.7%
Chromium III	0.013 = 1.3%
Chromium VI	0.025 = 2.5%
Manganese	0.04 = 4%
Nickel	0.04 = 4%
Silver	0.04 = 4%
Vanadium	0.026 = 2.6%

This is in addition to the chemical-specific dermal absorption fractions listed in RAGS Part E Exhibit 3-4, including:

Arsenic	0.03 = 3%
Cadmium	0.001 = 0.1%

dermal exposures to contaminants in aqueous media. The EPA guidance recommends the use of predicted K_p values. For chemicals that fall outside the Effective Prediction Domain for determining K_p , a fraction-absorbed (FA) term should be applied. This Risk Methods Document recommends the EPA default exposure values for all variables for the dermal-water and dermal-soil pathways. These include the residential scenario for water exposure and residential and industrial for soil exposure. For dermal-water exposures, the entire skin surface area is assumed to be available for exposure when bathing and swimming occurs, but the surface area available for a wading scenario includes the portions of the body specified in Appendix D for the dermal equations. Default values for the soil adherence factor (AF) also are provided with the equations in Appendix D. The guidance does not include a method for assessing dermal absorption of chemicals in the vapor phase, with the assumption that inhalation will be the major exposure route for vapors.

3.3.5.3 Tables summarizing the toxicity information

To facilitate review of the toxicity assessment, summary tables of toxicity information will be prepared following the examples in the previous sections of this guidance document. Additional tables may be prepared for the main body of the risk assessment, if needed to clarify the toxicity assessment process.

3.3.5.4 Summary of toxicity information available on the COPCs

This section of the toxicity assessment will provide a listing of the chemical classes and the number of chemicals within each class that have toxicity information ordered by medium within the unit or area under investigation. This summary will be presented to illustrate the total amount of toxicity information available to characterize risk in the following section.

3.3.6 Risk Characterization Methods

The primary purpose of this section of the baseline human health risk assessment will be to integrate the dose information developed in the exposure assessment with the effects information presented in the toxicity assessment to characterize the risk and hazard posed by environmental contamination at PGDP. In this section, the methods used to integrate the information to characterize risk and hazard and the tables and narrative summarizing the risk characterization for each exposure unit under each current and potential future use scenario will be presented. This section will conclude with a listing of use scenarios of concern for each location and a listing of COCs, POCs, and MOCs for each use scenario of concern.

3.3.6.1 Methods used to integrate dose and toxicity

In all baseline human health risk assessments performed for PGDP, the methods outlined in RAGS, Part A, will be used to integrate dose and toxicity information and characterize risk. To characterize risk from inhaled contaminants, the methods outlined in RAGS, Part F will be followed (EPA 2009). The following presents the equations that will be used for these calculations and describes the result of each equation. Note: In this presentation, the calculations for systemic toxicity (i.e., hazard) and cancer risk are presented separately because they differ slightly. Also, note that the values for systemic toxicity are estimates of whether the daily doses from each COPC, from each exposure pathway, and over all pathways and COPCs exceed that which may result in toxic effects in the receptor. However, the values for cancer risk are estimates of the excess cancer incidence that may result from exposure to each COPC, from each exposure pathway, and over all pathways.

Equations 8, 9, and 10 will be used to characterize the potential for systemic toxicity in all baseline human health risk assessments performed for PGDP. The result of Eq. 8 is a numeric estimate of the potential for systemic toxicity posed by a single chemical within a single pathway of exposure. The result

of Eq. 9 is a numeric estimate of the potential for systemic toxicity posed by all chemicals reaching a receptor through a single pathway. The result of Eq. 10 is a numeric estimate of the potential for systemic toxicity posed to a receptor by exposure to all chemicals over all pathways. (This last value is often called an estimate of “total noncarcinogenic risk.”)

$$HQ_i = \frac{CDI_i}{RfD_i} \quad \text{Eq. 8}$$

where: HQ_i = The hazard quotient, an estimate of the systemic toxicity posed by a single chemical
 CDI_i = The estimate of chronic daily intake (or absorbed dose for some exposure routes) from the exposure assessment (calculated from the chemical intake equations in Appendix D)
 RfD_i = The chronic reference dose for administered or absorbed dose, as appropriate

$$HQ = \frac{EC (\mu\text{g}/\text{m}^3)}{[RfC (\text{mg}/\text{m}^3) \times 1000 (\mu\text{g}/\text{mg})]} \quad \text{Eq. 8a}$$

where: HQ_i = The hazard quotient, an estimate of the systemic toxicity posed by a single chemical for inhalation
 EC_i = The exposure concentration for chronic exposure (calculated from the equations in Appendix D)
 RfC_i = The reference concentration for chronic inhalation exposure

$$HI_p = \sum_{i=1}^n HQ_i \quad \text{Eq. 9}$$

where: HI_p = The pathway hazard index, an estimate of the systemic toxicity posed by all chemicals within a single pathway
 HQ_i = The individual chemical hazard quotients for chemicals reaching the receptor through a single pathway (from Eq. 8 or Eq. 8a)

$$HI_{total} = \sum_{p=1}^n HI_p \quad \text{Eq. 10}$$

where: HI_{total} = The total hazard index, an estimate of the systemic toxicity posed by all chemicals over all pathways
 HI_p = The pathway hazard indices from Eq. 9

Equations 11, 12, and 13 will be used to characterize the potential excess lifetime cancer incidence (i.e., ELCR) in all baseline human health risk assessments performed for PGDP. The result of Eq. 11 is an estimate of the increased cancer incidence (i.e., a probability) to a receptor that results from exposure to a single chemical (or radionuclide) within a single pathway. The result of Eq. 12 is an estimate of the increased cancer incidence (i.e., probability) that results from exposure to all chemicals (or radionuclides) reaching a receptor through a single pathway. The result of Eq. 13 is an estimate of the increased cancer incidence (i.e., probability) that results from exposure to all chemicals (or radionuclides) reaching a receptor over all pathways. (This last value is often called an estimate of “total carcinogenic risk.”)

$$ELCR_i = CDI_i \times SF_i \quad \text{Eq. 11}$$

where: $ELCR_i$ = The chemical-specific excess cancer incidence
 CDI_i = The estimate of chronic daily intake (or absorbed dose) from the exposure assessment (calculated from the chemical intake equations in Appendix D)
 SF_i = The slope factor for administered or absorbed dose, as appropriate

$$ELCR = EC (\mu\text{g}/\text{m}^3) \times IUR (\mu\text{g}/\text{m}^3)^{-1} \quad \text{Eq. 11a}$$

where: $ELCR_i$ = The chemical-specific excess cancer incidence
 EC_i = The exposure concentration for chronic exposure (calculated from the equations in Appendix D)
 IUR_i = The unit risk for chronic inhalation exposure

$$ELCR_p = \sum_{i=1}^n ELCR_i \quad \text{Eq. 12}$$

where: $ELCR_p$ = The pathway-specific excess cancer incidence
 $ELCR_i$ = The chemical-specific excess cancer incidence from Eq. 11 or Eq. 11a

$$ELCR_{total} = \sum_{p=1}^n ELCR_p \quad \text{Eq. 13}$$

where: $ELCR_{total}$ = The total excess cancer incidence posed by all chemicals over all pathways
 $ELCR_p$ = The pathway-specific excess cancer incidence from Eq. 12

3.3.6.2 Presentation of risk characterization

In the baseline human health risk assessment, risk will be characterized for each exposure unit under each current and potential future use scenario. The results of the characterization will be presented in both tables and as narrative. The tables that will be used for each time, exposure unit, and receptor combination will be consistent with the two-way table presented in RAGS, Part D (EPA 1998b). At this time, scenarios are evaluated independently. Scenarios may be combined if it is determined that it is appropriate to do so to represent cumulative risk on a site-specific basis. The exact format presented in RAGS Part D is not used for the PGDP risk characterization tables because the FFA team discussed table presentation and agreed that the tables presented in this guidance document are adequate to meet the intent of RAGS, Part D. The narrative that explains this table, which may include summary tables, will present the exposure unit; the receptor, HI_{total} (from Equation 10) or $ELCR_{total}$ (from Equation 13); the primary pathways contributing to HI_{total} or $ELCR_{total}$ (i.e., “driving pathways”); and the primary chemicals contributing to HI_{total} or $ELCR_{total}$ (i.e., “driving chemicals”). An example of a narrative description of risk taken from DOE 1996e is presented below.

Exhibit 3.13 summarizes the HIs for exposure routes for the current industrial worker over all locations. As shown in this exhibit, the total scenario HI (i.e., Location Total in Exhibit 3.13) is greater than 1 for Sectors 5, 6, and 9. For each location, the driving exposure route is dermal contact with soil, which accounts for more than 95% of the total HI. Also, for each location, the inhalation exposure route contributes insignificantly to the location total HI.

Exhibit 3.14 summarizes the contaminants contributing more than 1% of the total systemic toxicity for the current industrial worker over all locations for those locations where the total systemic toxicity for the location exceeds 1. As shown in this exhibit, in each case, metals are the primary driving contaminants; however, PCBs and PAHs are minor contributors for Sector 6.

In the tables prepared for risk characterization, all COPCs will be listed, even those that do not have a value. Also, these tables will present the total chemical-specific hazard (or risk) over all pathways, the total pathway-specific hazard (or risk) over all chemicals, the total hazard or risk over all pathways and

chemicals, and the total risk and hazard over all media within the exposure unit (consistent with the Conceptual Site Model).

Exhibit 3.13. Exposure Route Summary for the Current Use Scenario—Systemic Toxicity^a

Scenario and Location	Exposure Routes for Soil			Location Total
	Incidental Ingestion	Dermal Contact	Inhalation of Vapors/Particles	
<i>Current industrial worker</i>				
Sector 1	NA	NA	NA	NV
% of Total	NV	NV	NV	
Sector 2	< 0.1	0.4	NV	0.4
% of Total	1%	99%	NV	
Sector 3	< 0.1	0.3	< 0.1	0.3
% of Total	2%	98%	< 1%	
Sector 4	< 0.1	1.0	< 0.1	1.0
% of Total	1%	99%	< 1%	
Sector 5	< 0.1	1.7	< 0.1	1.8
% of Total	2%	98%	< 1%	
Sector 6	< 0.1	1.2	< 0.1	1.2
% of Total	5%	95%	< 1%	
Sector 8	< 0.1	1.0	< 0.1	1.0
% of Total	< 1%	99%	< 1%	
Sector 9	< 0.1	1.3	NV	1.3
% of Total	1%	99%	NV	

NA indicates that the scenario is not applicable for this location.

NV indicates that a value is not available.

Current convention is to use one significant digit for presentation of hazard indices. Two significant digits are used here when the hazard index is greater than 1 to enable the reader to match the numbers reported in the exhibit with those in its associated risk characterization table. Additionally, use of two significant digits, when the exposure route's value is greater than 1, allows the reader to sum the route values and check the location total.

Exhibit 3.14. Driving Contaminants Summary for Current Use Scenario—Systemic Toxicity

Scenario and Location	Driving Contaminants Over All Exposure Routes	Location Total
<i>Current industrial worker</i>		
Sector 1	HI < 1	NV
Sector 2	HI < 1	0.4
Sector 3	HI < 1	0.3
Sector 4	HI < 1	1.0
Sector 5	iron (47%); chromium (26%); antimony (22%); uranium (3%)	1.8
Sector 6	chromium (22%); antimony (22%); arsenic (20%); PCB (13%); aluminum (13%); pyrene (2%); fluoranthene (1%)	1.2
Sector 8	HI < 1	1.0
Sector 9	antimony (58%); aluminum (23%); chromium (17%); uranium (2%)	1.3

NA indicates that the scenario is not applicable for this location.

NV indicates that a value is not available.

HI < 1 indicates that total scenario hazard index is less than 1; therefore, analytes are not listed.

3.3.6.3 Risk characterization for lead

Risk characterization for lead is a special case. Although it is known that exposure to lead can result in systemic toxic effects and possibly cancer, the approved toxicity values required to estimate potential for systemic toxicity and carcinogenesis are not available. The risk characterization for lead will consist of a comparison of the maximum detected concentration from the site/source to the no action screening levels from EPA and the Commonwealth of Kentucky. The no action screening levels are 400 mg/kg in soil and sediment for the residential and recreational scenarios, 800 mg/kg in soil and sediment for the industrial, and outdoor worker scenarios) and 15 µg/L in groundwater and surface water for all scenarios

(residential, recreational, industrial, and outdoor worker). Sites with lead concentrations exceeding these levels will undergo additional analysis for risk using the results of EPA's IEUBK (EPA 2004a) for evaluating residential and recreational exposures of children and the results of the EPA Adult Lead Model (ALM) (EPA 2003a) for evaluating industrial and outdoor worker exposures. The parameters for use in each of these models are presented in Appendix B. Screening values for lead appear in the tables presented in Appendix A.

3.3.6.4 Selection of use scenarios, pathways, contaminants, and MOC

Use scenarios, pathways, contaminants, and MOC will be identified for each unit or area under investigation. If any unit or area is divided into exposure units during the exposure assessment, use scenarios, pathways, contaminants, and MOC will be identified for each exposure unit.

In identifying use scenarios, pathways, contaminants, and MOC, specific rules will be followed as discussed below.

- **Identification of use scenarios of concern.** To determine use scenarios of concern on the basis of risk, risk characterization results for total systemic toxicity (HI_{total}) and total risk ($ELCR_{total}$) will be compared to benchmarks of $HI = 1.0$ and $ELCR = 1 \times 10^{-6}$. Use scenarios with HI_{total} or $ELCR_{total}$ exceeding either of these benchmarks will be deemed use scenarios of concern. Note: The results in the example narrative provided in Section 3.3.6.2 indicate the teen recreational use scenario is a use scenario of concern for SWMU 8a ($HI_{total} = 71.5$). This value would be found in the lower right hand corner of a two-way table consistent with RAGS, Part D (EPA 1998b).
- **Identification of POCs.** To determine POCs, risk characterization results for pathway hazard (HI_p) and risk ($ELCR_p$) over all chemicals *within a use scenario of concern* will be compared to benchmarks of $HI = 0.1$ and $ELCR = 1 \times 10^{-6}$. Pathways within a use scenario of concern exceeding either of these benchmarks will be deemed POCs for the use scenario of concern. Note: The results in the example narrative provided in Section 3.3.6.2 indicate that the POCs for the teen recreational user are dermal contact with surface water ($HI_p = 2.0$), dermal contact with leachate ($HI_p = 0.6$), ingestion of fish ($HI_p = 60.5$), ingestion of sediment ($HI_p = 0.1$), dermal contact with sediment ($HI_p = 8.2$), and ingestion of venison ($HI_p = 0.2$). These values would be found along the bottom margin of a two-way table consistent with RAGS, Part D (EPA 1998b).
- **Identification of COCs.** To determine COCs, risk characterization results for chemical hazard (HQ_i) and risk ($ELCR_i$) over all pathways *within a use scenario of concern* will be compared to benchmarks of $HQ = 0.1$ and $ELCR = 1 \times 10^{-6}$. Chemicals of potential concern within a use scenario of concern exceeding either of these benchmarks will be deemed COCs for the use scenario of concern. [Note: For dioxins and furans, PAHs, and PCBs, the total risk over all congeners (for dioxins and furans) or compounds (for PAHs and PCBs) will be used when determining if these are COCs.] The results in the example narrative provided in Section 3.3.6.2 indicate that the COCs for the teen recreational user are aluminum ($HQ_i = 0.2$), antimony ($HQ_i = 6.1$), arsenic ($HQ_i = 0.2$), cadmium ($HQ_i = 0.6$), iron ($HQ_i = 9.4$), manganese ($HQ_i = 48.4$), strontium ($HQ_i = 0.1$), vanadium ($HQ_i = 4.7$), and zinc ($HQ_i = 1.7$). These values would be found along the right margin of a two-way table.
- **Identification of Priority COCs.** To determine priority COCs (i.e., those COCs contributing most to cumulative HI and ELCR), risk characterization results for chemical hazard (HQ_i) and risk ($ELCR_i$) over all pathways *within a use scenario of concern* will be compared to benchmarks of $HQ = 1$ and $ELCR = 1 \times 10^{-4}$. Chemicals of concern exceeding either of these benchmarks will be deemed priority COCs for the use scenario of concern. [Note: For dioxins and furans, PAHs, and PCBs, the total risk over

all congeners (for dioxins and furans) or compounds (for PAHs and PCBs) will be used when determining if these chemicals are priority COCs.]

- **Identification of MOCs.** To determine MOCs, the POCs are reviewed, and those media in these pathways are deemed to be MOC. This is equivalent to screening the total risk and hazard posed by COPCs in the various media against benchmarks of HI = 0.1 and ELCR = 1×10^{-6} . For the results presented in the example narrative in Section 3.3.6.2, the MOCs are surface water, leachate, fish, sediment, and venison.
- **Identification of scenarios of concern, POCs, COCs, and MOCs in Dose Assessment.** If a dose assessment is conducted to provide additional information to risk managers, a scenario of concern will be one that has a total dose exceeding the PGDP *de minimis* dose of 1 mrem/year. A COC will be one that has a contaminant-specific dose exceeding 1 mrem/year. A POC will be an exposure route that has a route-specific dose exceeding 1 mrem/year. An MOC will be those media appearing in any POC.

3.3.6.5 Consideration of COPCs for which risk cannot be estimated

For some COPCs, information is insufficient for risk characterization. Generally, risk cannot be characterized for these chemicals because toxicity values are not available. When this occurs in risk assessments performed for PGDP, these COPCs will be deemed COCs during risk characterization, and they will be reported along with the COCs chosen by the rules outlined above.

3.3.6.6 Summary of risk characterization

To provide a summary of risk characterization for each unit or area under investigation, a table will be prepared and included as a summary of risk characterization in all baseline human health risk assessments. This table will follow the format shown in Exhibit 3.15 and list the risk and hazard posed within each use scenario of concern, the percent contribution of each POC to HI_{total} and ELCR_{total}, and the percent contribution of each COC to HI_{total} and ELCR_{total}. A similar table will be prepared to summarize the results of the dose assessment if a dose assessment is conducted for the site.

Exhibit 3.15. Summary of Risk Characterization

Use Scenario ¹	Total ELCR ²	COCs ³	% Total ELCR ⁴	POCs ⁵	% Total ELCR ⁶	Total HI ⁷	COCs	% Total HI ⁸	POCs	% Total HI ⁹
# 1										
# 2										
.
.
.
# N										

¹ All use scenarios will be listed.

² These values will be those found at the lower right of each unit's two-way table for the scenario of interest.

³ These constituents will be the COCs selected applying the rules listed earlier.

⁴ This value will be calculated by dividing the chemical-specific ELCR (ELCR_i) by the total ELCR (ELCR_{total}).

⁵ These pathways will be the POCs selected applying the rules listed earlier.

⁶ This value will be calculated by dividing the pathway-specific ELCR (ELCR_p) by the total ELCR (ELCR_{total}).

⁷ These values will be those found at the lower right of each unit's two-way table for the scenario of interest.

⁸ This value will be calculated by dividing the chemical-specific hazard quotient (HQ_i) by the total HI (HI_{total}).

⁹ This value will be calculated by dividing the pathway-specific HI (HI_p) by the total HI (HI_{total}).

3.3.7 Consideration of Uncertainty in the Risk Assessment

Uncertainties are associated with each of the steps of the baseline risk assessment. Following a general discussion of uncertainties in risk assessment, this section presents the uncertainties that will be addressed in baseline human health risk assessments prepared for PGDP and provides a format for summarizing this information (when a qualitative uncertainty analysis or sensitivity analysis is performed).

The potential effect of the uncertainties on the final risk characterization must be considered when interpreting the results of the risk characterization because the uncertainties directly affect the final risk estimates. Types of uncertainties that must be considered can be divided into four broad categories. These are uncertainties associated with data and data evaluation (i.e., identification of COPCs); exposure assessment; toxicity assessment; and risk characterization. Specific uncertainties under each of these broad categories that will be addressed in baseline human health risk assessments completed for PGDP are listed in the following material.

The exact method that will be used to present the uncertainty analysis in all baseline risk assessments cannot be included here. This is due, in large part, to the fact that the rigor of the uncertainty analysis will depend on the unit or area under investigation, the decisions that must be made for the unit or area, and the uncertainties affecting the risk estimates. At minimum, all baseline risk assessments will contain a qualitative uncertainty analysis that will include a quantitative sensitivity analysis of salient uncertainties. In the qualitative uncertainty analysis, the magnitude of the uncertainty on the risk characterization will be categorized as small, moderate, or large. Uncertainties categorized as small will be those that should not cause the risk estimates to vary by more than one order of magnitude; uncertainties categorized as moderate will be those that may cause the risk estimates to vary by between one and two orders of magnitude; and, uncertainties categorized as large will be those that may cause the risk estimates to vary by more than two orders of magnitude.

In the qualitative uncertainty analysis, a note will be made that the uncertainties listed and evaluated are neither independent nor mutually exclusive. It also will be noted that the total effect of all uncertainties upon the risk estimates is not the sum of the estimated effects of each uncertainty evaluated.

3.3.7.1 Uncertainties in data, data evaluation, and identification of COPCs

- Retention of common laboratory contaminants in the list of COPC
- Retention of infrequently detected analytes (i.e., detected in less than 10% of the samples analyzed) in the list of COPCs
- Lack of consideration in temporal patterns when selecting COPCs
- Spatial distribution and number of sampling locations (representativeness)
- Quantitation limits for some analytes exceeding their respective human health risk-based screening criteria (i.e., PRGs)
- Use of historical data in addition to data collected as part of the RI field investigation
- Removal of analytes from the list of COPCs on the basis of a comparison to background concentrations

- Removal of analytes from the list of COPCs on the basis of comparison to concentrations found in associated blanks
- Removal of analytes from the list of COPCs on the basis of a toxicity screen
- Characterization of EPCs for environmental media under current conditions
- Consideration of temporal changes in analyte concentrations and activities
- Use of results from analyses of unfiltered groundwater samples versus filtered groundwater samples
- Use of results from analyses of unfiltered surface water samples versus filtered surface water samples
- Uncertainties in exposure assessment
- Incorporation of biota fate and transport modeling into risk and hazard estimates (if this type of modeling were performed)
- Uncertainties in modeled concentrations, including the consideration of solubility as defined by differences between contaminant concentrations in filtered and unfiltered water samples
- Use of reasonable maximum exposure parameters versus average parameters for all exposure routes and associated pathways
- General issues in the development of conceptual site models
- Consideration of livestock scenarios
- Summation of risk and hazard across units or areas under investigation
- Use of default values from KDEP 2002 when estimating dermal absorbed dose (especially from soil and sediment)
- Difference in gamma walkover survey results and associated laboratory analyses

3.3.7.2 Uncertainties in toxicity assessment

- Use of provisional or withdrawn toxicity values
- Difference in risk estimates for TCE based on use of Kentucky DEP oral slope factor and EPA TCE oral slope factor (currently CalEPA value)
- Extrapolation of oral administered dose toxicity values to inhalation dose toxicity values
- Derivation of absorbed dose toxicity values from oral administered dose toxicity values
- Lack of toxicity information, toxicity values, or both for some COPCs
- Use of chronic exposure toxicity values for exposures that are subchronic

3.3.7.3 Uncertainties in risk characterization

- Combination of chemical-specific risk and hazard estimates (ELCR_i and HQ_i, respectively) to derive pathway-specific and use scenario risk and hazard estimates (ELCR_p and ELCR_{total} and HI_p and HI_{total}, respectively) (i.e., effect of chemical mixtures)
- Combination of risk estimates from chemical and radioisotope exposure
- Summing cancer risks across pathways and across target organs

(Note: Uncertainties regarding the risk characterization are discussed in the accompanying text box.)

3.3.7.4 Summary of qualitative uncertainty analysis

Because uncertainties in the baseline risk assessment must be addressed when screening potential remedial actions, developing revised preliminary remedial goals from RGOs and selecting the final action, the effect of all uncertainties on the risk and hazard estimates will be summarized in a single table. Note: Exhibit 3.16, is most useful when summarizing a qualitative uncertainty analysis; other formats may be used for a quantitative uncertainty analysis.

In addition to the summary table, a narrative (i.e., an Observations section) discussing the joint effects of the various uncertainties on the risk characterization results will be prepared. The overall goal of the narrative will be to focus the list of COCs to those COCs that contribute significantly to the risk and for which the risk estimate or the revised risk estimate in the uncertainty analysis is believed to reasonably reflect the risks posed to receptors under the most likely future use. This narrative in the Observations section will discuss how uncertainties affect the identification of COCs and evaluate scenarios that reflect the most likely future exposure. It also will describe how the inclusion of certain pathways (dermal, food ingestion, etc.) may lead to an overestimate of risks and summarizes which contaminants and/or pathways exceed *de minimis* levels. The narrative will address each of the COCs individually.

Uncertainty in Combining Chemical-Specific Risk and Hazard Estimates and Pathway-Specific Risk and Hazard Estimates

One uncertainty in the risk characterization guidance contained in this document is the method used to combine HQs and chemical-specific ELCRs across pathways and to combine pathway HIs and ELCRs to calculate total HI and ELCR. The method to be used to calculate pathway HIs and ELCRs follows EPA protocols (EPA 1989a). This method calls for the simple addition of HQs and chemical-specific ELCRs to calculate pathway HIs and ELCRs, respectively, and assumes that all effects between chemicals are additive. As explained in EPA 1989a, this assumption is made because information concerning the effects of chemical mixtures is lacking.

The following limitations of this approach for systemic toxicity effects are reported by EPA:

- Little is known about the effects of chemical mixtures; although additivity is assumed, the interaction of multiple chemicals could possibly be synergistic or antagonistic.
- The RfDs and RfCs do not have equal accuracy or precision and are not based on the same severity of effects.
- Dose additivity is most properly applied to compounds that induce the same effect by the same mechanism of action. While the approach recommended by EPA is a useful screening-level approach, the cumulative systemic toxicity could be overestimated for chemicals that act by different mechanisms and/or on different target organs.

The following limitations of this approach for chemical carcinogenesis are reported by EPA:

- Cancer risks (i.e., ELCRs) are based on slope factors that represent an upper 95th percentile estimate of potency; the upper 95th percentiles of probability distributions are not strictly additive. Summing these risks can result in an overly conservative estimate of lifetime ELCR.
- Cancer risks may not be additive. By analogy to systemic toxicity effects, the endpoints may differ, and mechanisms of effect may vary.
- Not all slope factors contain the same weight-of-evidence for human carcinogenicity. EPA recognizes this by placing weight-of-evidence classifications on all slope factors. Those contaminants with a weight-of-evidence classification of A should probably receive more attention in the selection of a remedial design than contaminants with a B or C classification. Similarly, a contaminant with a B classification should probably receive greater attention than one with a C classification. The simple combination of ELCRs does not take this hierarchy into account.

Uncertainty in Combining Risk Estimated for Chemical Exposure to Those for Risk Estimated for Radioisotope Exposure

Uncertainty associated with adding risks from chemical exposure to those from exposure to radionuclides arises from two sources. First, the slope factors used to characterize the risk from chemicals are derived differently from the slope factors used to characterize risk from radionuclides. This difference results in estimates of chemical exposure risks that may be considered to be upper-bound risk estimates and estimates of radionuclide exposure risks that may be considered to be central tendency (i.e., "best") estimates; therefore, combining chemical exposure and radionuclide exposure risk estimates to estimate total risk for a land use scenario may place too much emphasis on chemical exposure risk. Second, the mechanism by which chemicals may cause cancer varies from the mechanism by which radionuclides may cause cancer. This difference in mechanism of action inflates the uncertainties that assume cancer risks are additive.

Exhibit 3.16. Summary of Uncertainty Analysis

Description of Uncertainty	Estimated Effect ¹		
	Small	Moderate	Large
Uncertainties related to data, data evaluation, and identification of chemicals of potential concern ²			
Data uncertainty 1			
Data uncertainty 2			
.	.	.	.
.	.	.	.
.	.	.	.
Data uncertainty n			

¹ Definitions of effects are as follows:

- Small—Uncertainty should not cause the risk or hazard estimate to vary by more than one order of magnitude;
- Moderate—Uncertainty may cause the risk or hazard estimate to vary by between one and two orders of magnitude; and
- Large—Uncertainty may cause the risk or hazard estimate to vary by more than two orders of magnitude.

² A similar heading will appear for each of the major portions of the baseline human health risk assessment. The other headings are “Uncertainties related to exposure assessment,” “Uncertainties related to toxicity assessment,” and “Uncertainties related to risk characterization.”

3.3.8 Remedial Goal Option Derivation Methods

This section of the baseline human health risk assessment will delineate the methods used to derive and present RGOs. It is important to note that RGOs are not cleanup levels, but are site-specific, risk- or dose-based criteria that may be used to guide the development of revised PRGs in the FS and cleanup levels in the Record of Decision (ROD) by risk managers. Cleanup levels are developed as part of the risk analysis in the ROD (EPA 2006b).

3.3.8.1 Calculation of remedial goal options

Guidance in EPA (2000b) directs that multiple RGOs must be calculated for each COC identified in a baseline human health risk assessment. To do this, the goals are calculated by rearranging the exposure equations quantified in the risk assessment so that they solve for a concentration or activity in a medium that results in a specific “target risk,” “target hazard,” or “target dose.” Target risks that will be used to derive RGOs at PGDP are 1×10^{-4} , 1×10^{-5} , and 1×10^{-6} . Target hazards that will be used to derive RGOs are 3, 1, and 0.1. Target doses for all media but groundwater are 1, 15, 25, and 100 mrem/year. For groundwater, the dose targets are 1, 4 (for beta and photon emitters), 15, 25, and 100 mrem/year. As noted above, an RGO must be developed for each COC. Because the selection of a COC is medium- and use scenario-specific, RGOs will be developed for each COC identified for each use scenario of concern at a unit or area. Also, because RGOs must be medium-specific, exposure routes that integrate contaminant contributions from more than one medium (e.g., consumption of vegetables) will be segregated so that each medium contributing to the exposure route is evaluated separately. This segregation will be done by assuming that the concentration or activity of contaminants in the medium not under evaluation is zero.

In addition to calling for the development of RGOs, EPA (2000b) provides two methods that may be used to develop these values. The first involves rearranging and combining all the exposure equations utilized to determine risk or hazard and using the rearranged equation to calculate the RGO. The second simply uses ratios of concentrations or activities and level of risk, hazard, or dose to derive the RGO. Although the first method is of greater utility because the rearranged equation can be used to directly solve for RGOs, its use involves rearranging a large complex equation in which the chance for error abounds, especially if the estimated contaminant concentrations at the exposure point rely on fate and transport modeling. Similarly, although the second method is simpler mathematically, it can result in an incorrect solution if risk, hazard, or dose determined for COCs at the source in the baseline human health risk

assessment is not linearly and directly related to the concentration or activity of the COCs at the exposure point. Fortunately, the concentration or activity in each of the exposure equations that will be used in baseline human health risk assessments at PGDP (see Appendix D) is linearly and directly related to the resulting risk, hazard, or dose; therefore, the second method will be used in risk assessments at PGDP and is presented in Eqs. 14 and 15. Note: If additional exposure equations beyond those in Appendix D are used in an assessment performed for PGDP, these equations will be checked to ensure that the concentration or activity of COCs is directly and linearly related to risk or hazard.

$$\frac{\text{Conc}_{\text{observed}}}{\text{ELCR}_{\text{derived}}} = \frac{\text{RGO}}{\text{Target ELCR}} \quad \text{Eq. 14}$$

where: $\text{Conc}_{\text{observed}}$ = The representative EPC for the COC
 $\text{ELCR}_{\text{derived}}$ = The chemical-specific ELCR of a COC due to exposure to a single medium across all exposure routes
 RGO = The remedial goal option
 Target Risk = Either 1×10^{-4} , 1×10^{-5} , or 1×10^{-6}

$$\frac{\text{Conc}_{\text{observed}}}{\text{HI}_{\text{derived}}} = \frac{\text{RGO}}{\text{Target HI}} \quad \text{Eq. 15}$$

where: $\text{Conc}_{\text{observed}}$ = The representative EPC for the COC
 HI = The chemical-specific HI of a COC from exposure to a single medium across all exposure routes
 RGO = The remedial goal option
 Target Hazard = Either 3, 1, or 0.1

As noted, dose-based RGOs will be calculated using similar methods. The targets to be used for all media except groundwater are 1, 15, 25, and 100 mrem/year. For groundwater, the dose targets are 1, 4, 15, 25, and 100 mrem/year.

3.3.8.2 Presentation of remedial goal options

As noted, RGOs must be calculated for each COC within each MOC for each use scenario of concern identified in the baseline human health risk assessment; therefore, many RGOs will be developed in most risk assessments considering multiple units or areas. To simplify the consideration of the RGOs by users of the risk assessment, the format in Exhibit 3.17 will be used to present the RGOs in all baseline human health risk assessments prepared for PGDP. Note: Using this format will result in the preparation of a single table containing all COCs within each MOC for each use scenario of concern; therefore, this table or relevant portions of it can be used directly in the FS.

3.3.8.3 Revising exposure parameters and calculations in the uncertainty section

As part of the uncertainty analysis for the risk assessment, risk may be recalculated with default exposure factors replaced using exposure parameters consistent with site-specific values. The decision to recalculate risks using these alternative exposure parameters would be a product of the consensus of the FFA parties arrived at during project discussions at the appropriate stage in document development. For example, the exposure duration of 25 years for the outdoor worker may be replaced with a shorter duration of 1 to 5 years that is more likely to reflect the potential exposures at the site. The shorter exposure duration and possibly a revised exposure frequency combined with the other default parameters for the outdoor worker scenario also may be used to produce an excavation worker scenario. Also, risk

from dermal exposure to soil/sediment could be evaluated quantitatively to determine the impact of the use of default dermal absorption (ABS) values on the risk characterization. These revised calculations may be considered in the development of revised PRGs and cleanup levels to be used in the preparation of remedy selection documents.

Exhibit 3.17. Presentation of Remedial Goal Options¹

Chemical of Concern	Rep. Conc. ²	Regulatory Value ³	ELCR at Conc. ⁴	HI at Conc. ⁵	RGO at HI=0.1	RGO at HI=1	RGO at HI=3	RGO at ELCR= 1×10^{-6}	RGO at ELCR= 1×10^{-5}	RGO at ELCR= 1×10^{-4}	Units
Scenario and medium ⁶											
# 1 ⁷											
# 2											
.
.
.
# N											

¹ A separate table will be made for each unit or area under investigation.

² This value will be the representative concentration used in the calculation of risk or hazard in the baseline human health risk assessment.

³ Regulatory values (taken from ARARs) may not be available for some media.

⁴ This value will be the chemical-specific, medium-specific ELCR presented in the baseline human health risk assessment for the scenario of concern.

⁵ This value will be the chemical-specific, medium-specific ELCR presented in the baseline human health risk assessment for the scenario of concern.

⁶ Each MOC within a scenario of concern will be presented. The current use scenario RGOs will be presented first followed by the options for the most likely future use. The options for the least likely future use will appear last. Also, for the ground and surface water RGO tables, the appropriate MCLs will be listed.

⁷ All COCs should be listed, including those that could not be evaluated quantitatively.

A separate table following a similar format will be prepared for dose-based RGOs.

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4. RISK ANALYSES IN THE PREPARATION OF REMEDY SELECTION DOCUMENTS

As noted in RAGS, Part C, (EPA 1991c) and in *A Guide to Preparing Superfund Proposed Plans, Records of Decision, and Other Remedy Selection Documents* (EPA 1999b), risk analyses are an integral part of the remedy selection documents (e.g., FS, Proposed Plan, and ROD). The role of risk evaluations in these documents is discussed in this section. Risk evaluations that appear in other documents, including SI documents and Engineering Evaluations/Cost Analyses (EE/CAs), should be equivalent in data quality and content to risk assessments in the documents described in this section. Risk assessments in SI and EE/CA documents may vary from those described in the following section depending on how that risk assessment is used in decision-making for the specific project. A more streamlined approach for risk assessments is sometimes used for removal action decision documents.

Risk evaluations begin in the development and screening stage of the FS, extend through the detailed analysis of alternatives in the FS, and are reported in varying level of detail in the Proposed Plan and ROD. The primary goal of risk analyses here is to provide risk managers with the information needed to choose among specific remedial alternatives and to verify that a cleanup level was achieved. Generally, if a piece of risk information is not needed to choose among alternatives or to verify cleanup, it does not need to be generated; however, it should be noted that it is not uncommon for additional risk analyses to occur after the completion and signing of a ROD (e.g., during the design and implementation of the chosen remedy and after the implementation is complete). Generally, additional analyses occur because additional information relevant to the chosen remedy is required. Because the need for and form of these analyses is determined on a project-specific basis, the analyses that may occur after the completion of the FS are not discussed in detail here. The information provided in Sections 2 and 3 should be used to guide any additional work to ensure technical adequacy.

4.1 RISK ANALYSES DURING THE FEASIBILITY STUDY

Risk analyses impact four significant portions of the FS. These are the reporting of baseline or screening risk assessment results (including any dose assessment), the evaluation of the risk analyses to determine the need for remedial action, the identification and screening of technologies and alternatives, and the detailed analysis of alternatives. These areas are discussed in Sections 4.1.1, 4.1.2, 4.1.3, and 4.1.4, respectively.

4.1.1 Presentation of Risk Assessment Results in the Feasibility Study

Section 7, Summary and Conclusions, of the baseline human health risk assessment can be copied directly to the FS report. Additionally, following guidance in EPA 1999b, the tables consistent with RAGS, Part D, or relevant parts of them can be inserted directly into the FS. The material placed in the FS will contain a summary of the methods used to identify the COPCs and to complete the exposure assessment, toxicity assessment, and risk characterization, including the identification of significant uncertainties affecting the risk results. In addition, the risk characterization summary tables (Exhibit 3.15) and the relevant portions of the RGO summary tables (Exhibit 3.17) can be transported directly to the FS report. Electronic copies of this material will be made available to the authors of the FS report to simplify the reporting of this information and ensure consistency between the RI and FS reports.

As noted in RAGS Part C (EPA 1991c), the primary use of the baseline risk assessment from the RI is to assess what the relative effectiveness of each remedy would be in reducing the baseline risk. For some FS reports, recalculation of risk or dose estimates may be required to differentiate between remedial

alternatives; these changes to the baseline risk assessment should be conducted within the scope of Chapter 2 of RAGS Part C (EPA 1991c). The level of risk evaluation to be conducted in the FS should be determined and agreed to by the three FFA parties during scoping for the FS. Situations where risk estimates may need to be recalculated for the FS report include the following:

- The time between the completion of the RI report and the preparation of the FS report is such that additional information not considered in the RI report becomes available (e.g., additional samples or updated toxicity values).
- It is determined that the remedial technologies will produce new contaminants that were not present at the site under baseline conditions.
- The decision to include in the FS more advanced modeling from the matrix in Table 3.2 (including probabilistic risk assessment) in the FS than was used in the RI in order to provide refined estimates of risk necessary for determining the long-term or short-term effectiveness of remedial options or the differences in residual risk between remedial options.

Revised PRGs consistent with the alternatives will be derived in the FS. These revised PRGs will utilize the site-specific information in the RI report and the risk assessment in their calculation.

If additional risk assessment computations are required in the FS, then these computations will follow the methods outlined in Section 3. Most importantly, the exposure equations presented in Appendix D will be used for all risk computations that appear in the FS report, and the methods presented in Section 3.3.8 for RGO development will be followed.

In all FS reports, the summary of the risk assessment results will be followed by an evaluation of these results. This evaluation will consider the risk estimates, their basis, and the uncertainties deemed relevant to selection of a remedy. This evaluation will provide the focus for RAO development later in the FS report. The information that follows identifies typical decisions made when determining the need for remedial action in the FS report.

4.1.2 Modifications to Baseline Human Health Risk Assessment Parameters that Could Appear in the Feasibility Study

The evaluation of risks in the FS report focuses on those issues that are important in making decisions about whether remedial action is necessary and choosing between the proposed remedial alternatives; therefore, only a few parameters related to long-term risks should be conducted in the FS.

Uncertainties in the risk assessment can affect the values generated for risk and hazard, which affect the importance of the magnitude of differences in the residual risk and hazard associated with different remedial options. The uncertainty section of the baseline human health risk assessment will identify whether an uncertainty is small, moderate, or large for the investigation. If the uncertainty is small, it probably will not be necessary to reevaluate the risk assessment results. If, however, the uncertainty is moderate to large, then the FS will evaluate the uncertainty in more detail and may recalculate risk values as determined by agreement of the three parties.

Calculation of short-term risks during the detailed analysis of remedial alternatives (see Section 4.1.3) may require significant recalculation of risks from the baseline risk assessment to account for differences between the exposures to current workers and off-site residents and the default values used for the baseline risk assessment in the RI. For example, current industrial workers and current off-site residents do not consume groundwater from the facility for drinking. In addition, current industrial workers have

lower dermal exposure and shorter duration of exposure than is assumed for future industrial workers under a default exposure scenario. Outdoor workers also will have lower exposures than the default parameters due to the use of personal protective equipment and engineering controls. These differences need to be accounted for in the evaluation of short-term risks in the FS.

4.1.2.1 Land use considerations for determining appropriate response actions to protect future potential receptors

Land use is an important consideration when determining appropriate response actions based on potential future receptors. Uncertainties associated with future land use are largely due to the inability to predict if existing controls will be in place in the future. There may be scenarios presented pursuant to this document that may not be commensurate with the reasonable foreseeable land use but may serve as a reference point to decision makers. Consequently, the results of the baseline human health risk assessment will not be modified when determining potential risks to future receptors. The alternatives developed in the FS report will have to ensure protection of potential future receptors. Protection may be accomplished through continuation of existing controls in some instances. Consequently, potential future scenarios will be evaluated in the FS report to supply decision makers with the information needed to choose appropriate remedial actions. The information that follows provides examples of scenarios that may be evaluated for future receptors in the FS report.

Site-specific exposures for current industrial workers and the inability to predict potential future exposures have been discussed earlier. For a future industrial worker, the risks to a default industrial worker as determined in the baseline human health risk assessment will be used when estimating risks to determine the need for action. This evaluation includes potential risks as a result of contact with contaminated RGA groundwater, which also is a possibility in the future. Additional evaluations that will be included for the future industrial worker may include an evaluation of the continuation of existing institutional controls (i.e., controls and procedures that limit access and an alternative water source); continuation of controls and procedures (i.e., continuation of current industrial scenario) assuming contact with contaminated RGA groundwater (i.e., no separate water source); and default exposure (i.e., no controls or procedures) without contact with contaminated RGA groundwater (i.e., assuming a separate water supply).

Future recreational users and residential users inside the DOE property boundary (including area within the restricted access area, but not the surrounding West Kentucky Wildlife Management Area) will be assessed in the FS report based on the results of the baseline human health risk assessment. The risk manager will assume that no controls would be in place to restrict a future on-site recreational user or resident from contact with surface contamination.

Modeling during the baseline human health risk assessment typically involves a large degree of uncertainty. For this reason, modeling parameters may be reevaluated during the preparation of the FS report, as discussed in the modeling matrix presented in Table 3.2, if needed to reduce uncertainty and aid in choosing between the proposed remedial alternatives. For the same reason, the FS may consider use of probabilistic models for risk assessment in place of the deterministic models used during the RI if these additional analyses are deemed necessary through scoping agreements by the three parties.

4.1.2.2 Identification of use scenarios, pathways, contaminants, and MOC for decision making purposes

Following evaluation of the results and uncertainties in the baseline human health risk assessment and finalization of risk management decisions, a list of use scenarios, pathways, contaminants, and MOC for decision making purposes will be developed.

In the FS report, each item of concern will be identified based on the guidance presented in Section 3.3.6.4.

4.1.3 Risk Analyses during the Identification and Screening of Technologies and Alternatives

During the identification and screening stage of the FS, a range of remedial alternatives is identified, and each alternative is evaluated with respect to effectiveness, implementability, and cost (EPA 1991c). As part of the evaluation of effectiveness, human health risks to the community (e.g., short- and long-term health risks from releases during remediation and after remediation, respectively) and remediation workers (i.e., short-term health risks during remedial activities) will be considered. At PGDP, this evaluation will be performed qualitatively to be consistent with guidance in RAGS, Part C.

4.1.4 Risk Analyses during the Detailed Analysis of Alternatives

The overall objective of the detailed analysis of alternatives is to obtain and present the information needed by risk managers to select a remedial alternative for a site (EPA 1991c). Risk analysis affects three of the selection criteria against which alternatives are evaluated: long-term effectiveness, short-term effectiveness, and overall protection of human health and the environment.

Generally, the human health risk analyses performed during the FS follow the same procedures as the baseline human health risk assessment. Unlike the baseline human health risk assessment, where the goal is to estimate the risk posed by environmental contamination, the goal of the FS risk analyses is to determine to what extent the various remedial alternatives reduce risk, so that unacceptable levels of risk are not posed by residual environmental contamination.

Consistent with RAGS, Part C, (EPA 1991c), at PGDP the risk analyses performed during the detailed analysis of alternatives may be either qualitative or quantitative. In most cases, a qualitative analysis will be sufficient as indicated in RAGS, Part C; however, a quantitative analysis may be required in some cases. The decision of whether a qualitative or quantitative analysis of alternatives is needed will be made using the guidance in RAGS, Part C. In this guidance, EPA notes that the type of analysis that is required depends on (1) whether the relative short-term or long-term effectiveness is an important consideration in selecting the alternative and (2) the “perceived risk” associated with the alternative. In RAGS, Part C, EPA defines “perceived risk” as that leading to the belief by site engineers, risk assessors, and neighboring communities, including workers, that an alternative either may not be adequately protective or lead to increased risk. Specific parameters that will be taken into account at PGDP when examining “perceived risk” and determining if a quantitative analysis is required include the following (adapted from RAGS, Part C):

- Proximity of populations to the unit or area;
- Presence of highly or acutely toxic chemicals;
- Technologies with high release potential, either planned or unplanned;
- High uncertainties in the nature of releases;
- Multiple contaminants or exposure routes or both affecting the same receptor;
- Releases from neighboring units or areas, including uncontrolled releases from units or areas not yet addressed;

- Releases that occur over a long period; and
- Level of community concern.

4.1.4.1 Qualitative risk evaluations

As noted herein, a qualitative analysis will be sufficient for most units or areas. In this type of analysis, the risk evaluation will qualitatively evaluate each alternative against the RAOs defined during the FS. In all cases, the qualitative analysis will evaluate whether the alternative can reduce exposure to probable and potential receptor populations to acceptable levels. In many evaluations, this will involve qualitatively determining if an alternative is effective in reducing contaminant concentrations at a unit or area to the cleanup level (i.e., the RGO or revised PRG consistent with the alternative being evaluated).¹⁰ In other cases, this will involve determining if an alternative is effective in changing activity patterns of receptors so that the rate of contact by receptors to the contaminated materials is reduced, resulting in a lowered exposure. Finally, the qualitative risk evaluation in the detailed analysis of alternatives for PGDP will examine the potential for an alternative to produce new contaminants that were not at a unit or area during the RI.

In developing the risk evaluation portion of the qualitative detailed analysis of alternatives, several sources of information will be used. These sources are listed below [adapted from RAGS, Part C, (EPA 1991c)] and include information from the baseline or screening risk assessment (as modified during the risk management to determine the need for action), treatability studies, and results at other sites. Material from the risk assessment includes the following:

- The exposure setting, including exposed populations and future land use;
- The exposure pathways, including sources of contamination, COCs, fate and transport of chemicals (i.e., migration, degradation, and transformation), and exposure points;
- General exposure considerations, including rate of contact, exposure frequency, and exposure duration;
- Exposure concentrations, including temporal effects;
- Estimates of chemical intake and uptake;
- Toxicity information, including uncertainty in toxicity values; and
- Methods used to quantify risks from exposure to media containing multiple chemicals and radionuclides.

¹⁰ “Preliminary remediation goals...may be revised...based on the consideration of appropriate factors including, but not limited to: exposure factors, uncertainty factors, and technical factors. Included under exposure factors are: cumulative effect of multiple contaminants, the potential for human exposure from other pathways at the site, population sensitivities, potential impacts on environmental receptors, and cross-media impacts of alternatives. Factors related to uncertainty may include: the reliability of alternatives, the weight of scientific evidence concerning exposures and individual and cumulative health effects, and the reliability of exposure data. Technical factors may include: detection/quantification limits for contaminants, technical limitations to remediation, the ability to monitor and control movement of contaminants, and background levels of contaminants. The final selection of the appropriate risk level is made when the remedy is selected based on the balancing of criteria...” [taken from the National Contingency Plan Preamble: Exposure, Technical, and Uncertainty Factors (55 Fed. Reg. 8717, March 8, 1990)]. Also, see RAGS Volume 1, Part B, Section 2.3 and 2.8 (EPA 1993a) and OSWER Directive 9355.0-30, “Role of the Baseline Risk Assessment in Superfund Remedy Selection Decisions” (EPA 1990a).

Material found in treatability studies that will be used in the qualitative risk evaluation includes the following:

- Effectiveness at reducing potential for exposure, either through reduction in contaminant concentrations and activities or through making the medium containing the contaminant unavailable for contact;
- Potential for short-term emissions; and
- Potential for production of new contaminants.

Materials found when examining results from other sites that will be used in the qualitative risk evaluation include the following:

- Actual contaminant reductions achieved;
- Conditions in which the technology was not effective; and
- Actual release rates of current or new contaminants.

4.1.4.2 Quantitative risk evaluations

Methods for quantitative risk evaluations during the detailed analysis of alternatives have not yet been developed for PGDP. These will be included when they become available. It is anticipated that these methods will follow, in large part, the guidance and requirements for quantitative risk evaluations during the detailed analysis of alternatives in RAGS, Part C (EPA 1991c) and the more detailed guidance presented in Section 3 of this report.

4.2 RISK ANALYSES AFTER THE FEASIBILITY STUDY

After the FS is completed, a remedy is proposed in the Proposed Plan and documented in the ROD. Following this, the remedy is designed and implemented and, depending on the remedy, the site either is deleted or is placed within the group for which five-year reviews are required. This section discusses the risk evaluation activities that will occur during and after the preparation of the Proposed Plan. These risk evaluation activities should be consistent with EPA guidance in the *Guide to Preparing Superfund Proposed Plans, Records of Decision, and other Remedy Selection Decision Documents* (EPA 1999b). Some of the material presented here was taken from RAGS, Part C (EPA 1991c).

4.2.1 Risk Evaluation for the Proposed Remedial Action Plan

Generally, no new risk evaluations will take place during the preparation of the Proposed Plan. The material presented in the Proposed Plan should be taken entirely from the supporting FS. This includes a summary of site risks, the site COCs, and, if applicable, the revised PRGs for the selected alternative or a description of the basis for them (i.e., risk or dose target). Consistent with EPA 1999b, the material presented in the “Summary of Site Risks” section of the Proposed Plan primarily will be presented as narrative and limited to approximately three paragraphs. Key information from the baseline risk assessment (or its equivalent screening assessment from scoping activities) that will be presented includes all the following:

- Major COCs in each medium
- Land- and groundwater-use assumptions

- Potentially exposed populations under current and future use scenarios
- Major pathways and routes of exposure
- Summary of risk characterization

The risk section of the Proposed Plan also will contain a text box of standard language from the Proposed Plan/ROD guidance (EPA 1999b). This standard language will contain a definition of risk assessment and the meaning of the results from a risk assessment.

The risk section of the Proposed Plan will conclude with language similar to the following text taken from EPA 1999b.

It is the lead agency's current judgment that the Preferred Alternative identified in this Proposed Plan, or one of the other active measures considered in the Proposed Plan, is necessary to protect public health or welfare or the environment from actual or threatened releases of pollutants or contaminants from this site. These pollutants or contaminants may present an imminent and substantial endangerment to public health or welfare.

If new information becomes available during the public comment period, then additional analysis of the alternatives, or possibly the baseline risks, may be needed. (Note: These analyses will encompass all alternatives and be performed qualitatively to the extent possible.)

4.2.2 Risk Evaluation for the ROD

The primary risk evaluation-related activities that will occur during the ROD will be to document the results of the risk assessment and the risk evaluation portions of the comparison of alternatives performed in the FS and to document the derivation of the chemical-specific cleanup levels. Consistent with EPA guidance in both *Guide to Preparing Superfund Proposed Plans, Records of Decision, and other Remedy Selection Decision Documents* (EPA 1999b) and RAGS, Part C (EPA 1991c), the appropriate risk assessment materials will be discussed in relation to three of the nine CERCLA alternative analysis criteria: long-term effectiveness, short-term effectiveness, and overall protection of human health and the environment. The discussion of overall protection of human health and the environment will consider, to the extent possible, any residual risks that may remain after the alternative is implemented. Specific information to be presented includes the following:

- Chemical-specific cleanup levels to be attained at the conclusion of the response action;
- Corresponding chemical-specific risk levels;
- Areas of attainment for cleanup levels for groundwater being addressed; and
- Lead agency's basis for the cleanup levels (e.g., risk calculation, ARARs, background, etc.).

To the extent possible, the "Summary of Site Risks" section of the ROD will be presented following the outline contained in EPA 1999b; therefore, this material will include the following:

- A statement of basis for taking action and
- A brief summary of the relevant portions of the risk assessment.

Additionally, this section will focus on the risk drivers as defined in the FS and the exposure scenarios and pathways driving the need for action. The conceptual site model (which should be presented in the *Summary of Site Characteristics* section of the ROD) will be used to support the presentation of site risks.

The standard language to be used for the statement of basis for action will be similar to that which also appears in the Proposed Plan. For the ROD, this statement will appear at the beginning of the site risk summary instead of at the end.

In most cases, the tabular information that appears in the ROD will be drawn directly from EPA 1999b; however, additional tables or tables of a slightly different format may be used to explain the risk assessment results, as needed. Note that the primary purpose for including the detailed risk characterization tables in an appendix of the baseline risk assessment is to streamline the preparation of these tables for the FS and ROD.

4.2.3 Risk Analyses for Residual Risks

As noted in RAGS, Part C, (EPA 1991c) analyses to examine residual risks may be required for some locations after implementation of a remedy. Additionally, as discussed in the SMP (DOE 2014), after completion of all investigations and remedial actions at PGDP, the FFA requires that PGDP determine the residual risks remaining at the facility. In addition, the five-year review of some sites may require additional residual risk analyses. These residual risk analyses should be conducted consistent with guidance on the five-year review process from both EPA (EPA 2001c; EPA 2003c) and DOE (DOE 2002). The methods to be used to complete the analyses of residual risks at most units will be qualitative. If quantitative, these analyses will be consistent with the methods in either Section 2 or that in Section 3 of this document. Additionally, any quantitative analyses will be consistent with Section 3.3.4 of RAGS, Part C (EPA 1991c). Generally, these analyses will determine the risks remaining after remediation due to contamination remaining at or migrating from multiple sources. In these analyses, the measured concentrations and activities of contaminants remaining at the various source units and in the integrator unit will be used. The cleanup levels in the ROD for the various source units and areas in the integrator units will not be used in these analyses.

Other issues that will be considered when evaluating residual risk will be the following:

- Concentrations and activities of new analytes formed as a result of remedial activities or because of natural processes;
- Changes in land use or proposed future use since the completion of the baseline risk assessment;
- Updated toxicity values; and
- Reduction of migration because of engineered controls and expected future performance of these controls.

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APPENDIX A

SCREENING LEVELS
(Current as of November 2013)

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SCREENING LEVELS

This appendix presents lists of values that can be used during screening and baseline human health risk assessments at the Paducah Gaseous Diffusion Plant (PGDP). These values include risk- and dose-based values for soil, sediment, groundwater, and surface water; background values for soil and groundwater; and regulatory values. All information is current as of the production dates listed in this document, and all values were calculated using the best available information. Methods used to derive the risk- and dose-based values are presented in Appendix B. The screening values presented in this appendix were developed specifically for PGDP and may not be applicable to sites outside that facility. Values are provided in these tables for significant chemicals of potential concern (COPCs) for PGDP. Values for other chemicals can be obtained using the electronic Preliminary Remediation Goal (PRG) calculator.

Please consider the following notes before using the values presented in this appendix.

- (1) Action values are the lesser of a hazard-based value calculated using a target hazard index (HI) of 3 and a cancer-based value calculated using a target excess lifetime cancer risk (ELCR) of 1E-04.
- (2) HI values are calculated separately for each receptor. Cancer risks for receptors within a scenario are combined to give one lifetime cancer risk value. For the residential scenario, the cancer risk reflects the adult and child combined. For the recreational scenario, the cancer risk reflects the combined risk to adult, child, and teen.
- (3) Action values and no action values are calculated using only direct exposure pathways. Please see Appendix B for a listing of exposure parameters included in the PRG calculations. Because the action values are not calculated using PGDP default exposure parameters, these values should be used as benchmarks only. Cumulative risk calculations should not be based upon these values.
- (4) No action values are the lesser of a hazard-based value calculated using a target HI of 0.1 and a cancer-based value calculated using a target ELCR of 1E-06. If more than five COPCs are identified for the site, it also may be appropriate to generate no action levels based on 1×10^{-7} risk to account for additivity of risk. These values were calculated using the exposure parameters listed with the exposure equations in Appendix D. These parameters also are listed in Appendix B. Because the no action values are consistent with the PGDP default exposure parameters, these values can be used to derive cumulative risk estimates in addition to their use as benchmarks.
- (5) Background values for soil and groundwater presented in this appendix are provisional. These values are subject to change.
- (6) Soil screening levels for chemicals for protection of groundwater were derived using information presented in the U.S. Environmental Protection Agency (EPA) Soil Screening Level (SSL) Web site. The SSL values based upon a dilution attenuation factor of 1 should be considered to be “no action values.” “Action” SSLs have not been selected to date for the PGDP.
- (7) Regulatory values are for planning purposes only. A qualified regulatory specialist should be consulted before using these values for other purposes.
- (8) The outdoor worker scenario replaces the scenario listed in the 2001 version as “excavation worker” and uses the same exposure parameters. Based on consensus of the work group, the outdoor worker can be modified by reducing the exposure duration from 25 years to a value between one and five years to generate site-specific values for exposures during excavation.

(9) Chemical-specific notes for risk-based and dose-based screening values:

- a) General—Several screening values for soil/sediment (especially those on the action level tables) are listed with a value of 100,000 mg/kg. This value was assigned to the chemical because the screening value derived for the contaminant exceeded the upper limit value deemed reasonable by the PGDP Risk Assessment Working Group; therefore, the screening value was reduced to an upper limit value (100,000 mg/kg). If the chemical's environmental concentration exceeds the upper limit value, then additional risk evaluations for the chemical should be performed before accepting the results of a simple comparison.
- b) Chromium—The screening value for Chromium VI presented in these tables should only be used if the comparison is to a Chromium VI result. For a 'Total Chromium' result, the screening value listed for 'Total Chromium' should be used. The cancer-based screening value for Total Chromium was derived using the cancer slope factor for Chromium VI reported in the EPA Integrated Risk Information System database. Please see the toxicity value tables for additional information regarding this value.
- c) Lead—The screening values for lead were selected by the PGDP Risk Assessment Working Group. These values were not derived using the methods presented in Appendix B and are not included in the electronic PRG calculator. No action levels are 400 mg/kg for soil/sediment for the resident and the recreator scenarios and 800 mg/kg for the industrial worker and outdoor worker scenarios. These values represent the current screening values provided by the Kentucky Department for Environmental Protection. Action levels for soil/sediment are set preliminarily equivalent to the no action levels. Sites at which the 400 mg/kg concentration in soil is exceeded should be evaluated using site specific Integrated Exposure Uptake Biokinetic (IEUBK) modeling for a level resulting in a child exceeding a target blood level of 2.5 µg/dl and a target blood level of 10 µg/dl and Adult Lead Model (ALM) modeling for an adult exceeding the same target blood lead levels. Parameters for use in the IEUBK model are provided in Table B.6 of Appendix B. Parameters for the ALM model should be developed for each site. No action and action levels for groundwater and for surface water are unchanged from those agreed to by the PGDP Risk Assessment Working Group in the 2001 version of this document.
- d) Reserved.
- e) Carcinogenic polycyclic aromatic hydrocarbons (cPAHs)—(These organic compounds include benz(a)anthracene, benzo(a)pyrene, benzo(b)fluoranthene, benzo(k)fluoranthene, chrysene, dibenz(a,h)anthracene, and indeno(1,2,3-cd)pyrene.) The PGDP Risk Assessment Working Group has determined that these compounds should be evaluated as a group using the PAH (Total) screening values. Please see Section 3.3.3.2, Step 8, of the main text of the methods document for guidance on deriving total PAH concentration from results for individual compounds.
- f) Polychlorinated biphenyls (PCB)—These organic compounds include those listed as Aroclors in the screening tables.) The PGDP Risk Assessment Working Group has determined that the cancer effects of these organic compound mixtures should be evaluated as a group using the PCB (Total) screening values. (The screening value associated with the highest risk value is to be used.) Please see Section 3.3.3.2, Step 8, of the main text of the methods document for guidance on deriving total PCB concentration from results for individual mixtures.

- g) Dioxins/Furans—(These organic compounds include the following chlorinated dioxins and furans: 2,3,7,8-TCDD; 1,2,3,7,8-PeCDD; 2,3,4,7,8-PeCDD; 2,3,5,7,8-PeCDD; 2,3,6,7,8-PeCDD; 1,2,3,4,7,8-HxCDD; 1,2,3,5,7,8-HxCDD; 1,2,3,6,7,8-HxCDD; 2,3,4,5,7,8-HxCDD; 2,3,4,6,7,8-HxCDD; 2,3,5,6,7,8-HxCDD; 1,2,3,4,5,7,8-HpCDD; 1,2,3,4,6,7,8-HpCDD; 2,3,4,5,6,7,8-HpCDD; OCDD; 2,3,7,8-TCDF; 1,2,3,7,8-PeCDF; 2,3,4,7,8-PeCDF; 1,2,3,4,7,8-HxCDF; 1,2,3,5,7,8-HxCDF; 1,2,3,6,7,8-HxCDF; 2,3,4,5,7,8-HxCDF; 2,3,4,6,7,8-HxCDF; 2,3,5,6,7,8-HxCDF; 1,2,3,4,5,7,8-HpCDF; 1,2,3,4,6,7,8-HpCDF; 2,3,4,5,6,7,8-HpCDF; and OCDF.) The PGDP Risk Assessment Working Group has determined that these organic compounds should be evaluated as a group using the Dioxins/Furans (Total) screening values. Please see Section 3.3.3.2, Step 8, of the main text of the methods document for guidance on deriving the total dioxin/furan concentration from results for individual compounds.
- h) Radionuclides—For cesium-137, neptunium-237, uranium-235, and uranium-238, only screening values derived considering the contribution from short-lived decay products should be used. These screening values are listed with a “+D” in the following tables.

Radionuclides—Dose levels are (1) 1 mrem/year (from NRCRP Report No. 116, Section 17, Negligible Individual Dose and ANSI/HPS standard N13.12); (2) 15 mrem/year (from Establishment of Cleanup Levels for CERCLA Sites with Radioactive Contamination” OSWER No. 9200.4-18, August 22, 1997); (3) 25 mrem/year (derived from the public dose limit of 100 mrem/year limit in DOE Order 458.1 and considering ALARA principles); and (4) 100 mrem/year. A value of 4 mrem/year is used for groundwater (from <http://www.epa.gov/safewater/contaminants/index.html>).

Due to the nature of Appendix A, not all acronyms are defined within the text. An acronym list is provided on page A-9.

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ACRONYMS

ALARA	as low as reasonably achievable
AL	action level
ALM	Adult Lead Model
ANSI	American National Standards Institute
BaP	benzo(a)pyrene
CERCLA	Comprehensive Environmental Response, Compensation, and Liability Act
CLP	Contract Laboratory Program
COPC	chemical of potential concern
cPAHs	carcinogenic polycyclic aromatic hydrocarbons
DAF	dilution attenuation factor
DOE	U.S. Department of Energy
ELCR	excess lifetime cancer risk
EPA	U.S. Environmental Protection Agency
GW	groundwater
HI	hazard index
HPS	Health Physics Society
IEUBK	Integrated Exposure Uptake Biokinetic
KAR	Kentucky Administrative Record
K_d	chemical-specific distribution coefficient
MCL	maximum contaminant level
MCLG	maximum contaminant level goal
n/a	not available
NAL	no action level
NRCRP	Nuclear Regulatory Commission Report
OSWER	Office of Solid Waste and Emergency Response
PAH	polycyclic aromatic hydrocarbon
PCB	polychlorinated biphenyl
PGDP	Paducah Gaseous Diffusion Plant
PRG	Preliminary Remediation Goal
RAIS	Risk Assessment Information System
RCRA	Resource Conservation and Recovery Act
RESRAD	residual radioactivity
RGA	Regional Gravel Aquifer
SSL	Soil Screening Level
UTL	upper tolerance limit
WQC	water quality criteria

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Table A.1. Soil/Sediment Action Levels for Significant COPCs at PGDP

(Values calculated on 11/5/2013 and are based on best available information.)

CAS	Analyte	Units	Outdoor Worker			Industrial Worker			Adult Recreational User		
			Hazard	Cancer	Action	Hazard	Cancer	Action	Hazard	Cancer	Action
7429905	Aluminum	mg/kg	1.00E+05	-	1.00E+05	1.00E+05	-	1.00E+05	1.00E+05	-	1.00E+05
7440360	Antimony (metallic)	mg/kg	3.45E+02	-	3.45E+02	2.45E+03	-	2.45E+03	2.95E+03	-	2.95E+03
7440382	Arsenic, Inorganic	mg/kg	2.00E+02	4.15E+01	4.15E+01	1.83E+03	3.81E+02	3.81E+02	8.16E+02	3.53E+02	3.53E+02
7440393	Barium	mg/kg	1.00E+05	-	1.00E+05	1.00E+05	-	1.00E+05	1.00E+05	-	1.00E+05
7440417	Beryllium and compounds	mg/kg	1.72E+03	1.00E+05	1.72E+03	1.19E+04	1.00E+05	1.19E+04	1.47E+04	1.00E+05	1.47E+04
7440428	Boron And Borates Only	mg/kg	1.00E+05	-	1.00E+05	1.00E+05	-	1.00E+05	1.00E+05	-	1.00E+05
7440439	Cadmium (Diet)	mg/kg	6.18E+02	1.00E+05	6.18E+02	5.94E+03	1.00E+05	5.94E+03	2.25E+03	1.00E+05	2.25E+03
16065831	Chromium (Total) ^a	mg/kg	1.00E+05	2.68E+04	2.68E+04	1.00E+05	1.98E+04	1.98E+04	1.00E+05	1.59E+05	1.00E+05
18540299	Chromium(III), Insoluble Salts	mg/kg	1.00E+05	-	1.00E+05	1.00E+05	-	1.00E+05	1.00E+05	-	1.00E+05
7440473	Chromium(VI)	mg/kg	2.59E+03	1.60E+02	1.60E+02	1.82E+04	1.08E+03	1.08E+03	2.21E+04	2.82E+03	2.82E+03
7440484	Cobalt	mg/kg	2.59E+02	1.00E+05	2.59E+02	1.81E+03	1.00E+05	1.81E+03	2.21E+03	1.00E+05	2.21E+03
7440508	Copper	mg/kg	3.45E+04	-	3.45E+04	1.00E+05	-	1.00E+05	1.00E+05	-	1.00E+05
7439896	Iron	mg/kg	1.00E+05	-	1.00E+05	1.00E+05	-	1.00E+05	1.00E+05	-	1.00E+05
7439921	Lead	mg/kg	-	-	8.00E+02	-	-	8.00E+02	-	-	4.00E+02
7439965	Manganese	mg/kg	2.04E+04	-	2.04E+04	1.00E+05	-	1.00E+05	1.00E+05	-	1.00E+05
7439976	Mercury, Inorganic Salts	mg/kg	2.59E+02	-	2.59E+02	1.84E+03	-	1.84E+03	2.21E+03	-	2.21E+03
7439987	Molybdenum	mg/kg	4.32E+03	-	4.32E+03	3.06E+04	-	3.06E+04	3.69E+04	-	3.69E+04
7440020	Nickel Soluble Salts	mg/kg	1.71E+04	1.00E+05	1.71E+04	1.00E+05	1.00E+05	1.00E+05	1.00E+05	1.00E+05	1.00E+05
7782492	Selenium	mg/kg	4.32E+03	-	4.32E+03	3.06E+04	-	3.06E+04	3.69E+04	-	3.69E+04
7440224	Silver	mg/kg	4.32E+03	-	4.32E+03	3.06E+04	-	3.06E+04	3.69E+04	-	3.69E+04
7791120	Thallium (Soluble Salts)	mg/kg	8.64E+00	-	8.64E+00	6.12E+01	-	6.12E+01	7.38E+01	-	7.38E+01
238	Uranium (Soluble Salts)	mg/kg	2.58E+03	-	2.58E+03	1.79E+04	-	1.79E+04	2.21E+04	-	2.21E+04
7440622	Vanadium and Compounds	mg/kg	4.35E+03	-	4.35E+03	3.03E+04	-	3.03E+04	3.72E+04	-	3.72E+04
7440666	Zinc and Compounds	mg/kg	1.00E+05	-	1.00E+05	1.00E+05	-	1.00E+05	1.00E+05	-	1.00E+05
83329	Acenaphthene	mg/kg	2.28E+04	-	2.28E+04	1.00E+05	-	1.00E+05	5.25E+04	-	5.25E+04
208968	Acenaphthylene	mg/kg	-	-	-	-	-	-	-	-	-
107131	Acrylonitrile	mg/kg	2.91E+02	8.57E+01	8.57E+01	2.17E+02	1.31E+02	1.31E+02	1.00E+05	8.24E+02	8.24E+02
120127	Anthracene	mg/kg	1.00E+05	-	1.00E+05	1.00E+05	-	1.00E+05	1.00E+05	-	1.00E+05
71432	Benzene	mg/kg	1.28E+03	5.21E+02	5.21E+02	1.41E+03	5.66E+02	5.66E+02	2.95E+04	4.05E+03	4.05E+03
86748	Carbazole	mg/kg	-	2.04E+03	2.04E+03	-	2.86E+04	2.86E+04	-	1.07E+04	1.07E+04
56235	Carbon Tetrachloride	mg/kg	1.56E+03	3.21E+02	3.21E+02	1.95E+03	3.16E+02	3.16E+02	2.95E+04	2.33E+03	2.33E+03
67663	Chloroform	mg/kg	3.12E+03	1.89E+02	1.89E+02	3.42E+03	1.50E+02	1.50E+02	7.38E+04	1.18E+03	1.18E+03
75354	Dichloroethylene, 1,1-	mg/kg	4.02E+03	-	4.02E+03	3.24E+03	-	3.24E+03	1.00E+05	-	1.00E+05
540590	Dichloroethylene, 1,2- (Mixed Isomers)	mg/kg	7.77E+03	-	7.77E+03	5.52E+04	-	5.52E+04	6.63E+04	-	6.63E+04
156592	Dichloroethylene, 1,2-cis-	mg/kg	1.73E+03	-	1.73E+03	1.23E+04	-	1.23E+04	1.47E+04	-	1.47E+04
156605	Dichloroethylene, 1,2-trans-	mg/kg	2.47E+03	-	2.47E+03	2.09E+03	-	2.09E+03	1.00E+05	-	1.00E+05
60571	Dieldrin	mg/kg	2.18E+01	2.54E+00	2.54E+00	3.06E+02	3.58E+01	3.58E+01	5.49E+01	1.34E+01	1.34E+01

Hazard-based value calculated using target HI of 3.

Cancer-based value calculated using target ELCR of 1E-04.

Action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.1. Soil/Sediment Action Levels for Significant COPCs at PGDP (Continued)

(Values calculated on 11/5/2013 and are based on best available information.)

CAS	Analyte	Units	Outdoor Worker			Industrial Worker			Adult Recreational User		
			Hazard	Cancer	Action	Hazard	Cancer	Action	Hazard	Cancer	Action
1746016	Dioxins/Furans, Total (as TCDD) ^b	mg/kg	4.68E-04	4.79E-04	4.68E-04	4.29E-03	4.40E-03	4.29E-03	1.91E-03	4.07E-03	1.91E-03
37871004	~HpCDD, 2,3,7,8-	mg/kg	4.68E-02	4.79E-02	4.68E-02	4.29E-01	4.40E-01	4.29E-01	1.91E-01	4.07E-01	1.91E-01
38998753	~HpCDF, 2,3,7,8-	mg/kg	3.06E-02	3.13E-02	3.06E-02	4.29E-01	4.40E-01	4.29E-01	7.71E-02	1.65E-01	7.71E-02
34465468	~HxCDD, 2,3,7,8-	mg/kg	4.68E-03	4.79E-03	4.68E-03	4.29E-02	4.40E-02	4.29E-02	1.91E-02	4.07E-02	1.91E-02
55684941	~HxCDF, 2,3,7,8-	mg/kg	3.06E-03	3.13E-03	3.06E-03	4.29E-02	4.40E-02	4.29E-02	7.71E-03	1.65E-02	7.71E-03
3268879	~OCDD	mg/kg	1.56E+00	1.60E+00	1.56E+00	1.43E+01	1.47E+01	1.43E+01	6.36E+00	1.36E+01	6.36E+00
39001020	~OCDF	mg/kg	1.02E+00	1.04E+00	1.02E+00	1.43E+01	1.47E+01	1.43E+01	2.57E+00	5.48E+00	2.57E+00
36088229	~PeCDD, 2,3,7,8-	mg/kg	4.68E-04	4.79E-04	4.68E-04	4.29E-03	4.40E-03	4.29E-03	1.91E-03	4.07E-03	1.91E-03
57117416	~PeCDF, 1,2,3,7,8-	mg/kg	1.02E-02	1.04E-02	1.02E-02	1.43E-01	1.47E-01	1.43E-01	2.57E-02	5.48E-02	2.57E-02
57117314	~PeCDF, 2,3,4,7,8-	mg/kg	1.02E-03	1.04E-03	1.02E-03	1.43E-02	1.47E-02	1.43E-02	2.57E-03	5.48E-03	2.57E-03
1746016	~TCDD, 2,3,7,8-	mg/kg	4.68E-04	4.79E-04	4.68E-04	4.29E-03	4.40E-03	4.29E-03	1.91E-03	4.07E-03	1.91E-03
51207319	~TCDF, 2,3,7,8-	mg/kg	3.06E-03	3.13E-03	3.06E-03	4.29E-02	4.40E-02	4.29E-02	7.71E-03	1.65E-02	7.71E-03
100414	Ethylbenzene	mg/kg	4.80E+04	2.61E+03	2.61E+03	7.08E+04	2.83E+03	2.83E+03	1.00E+05	2.02E+04	2.02E+04
206440	Fluoranthene	mg/kg	1.52E+04	-	1.52E+04	1.00E+05	-	1.00E+05	3.51E+04	-	3.51E+04
86737	Fluorene	mg/kg	1.52E+04	-	1.52E+04	1.00E+05	-	1.00E+05	3.51E+04	-	3.51E+04
118741	Hexachlorobenzene	mg/kg	3.48E+02	2.54E+01	2.54E+01	4.92E+03	3.58E+02	3.58E+02	8.79E+02	1.34E+02	1.34E+02
91203	Naphthalene	mg/kg	1.97E+03	2.43E+03	1.97E+03	1.94E+03	1.80E+03	1.80E+03	1.75E+04	1.44E+04	1.44E+04
88744	Nitroaniline, 2-	mg/kg	4.35E+03	-	4.35E+03	5.73E+04	-	5.73E+04	1.10E+04	-	1.10E+04
621647	Nitroso-di-N-propylamine, N-	mg/kg	-	5.82E+00	5.82E+00	-	8.18E+01	8.18E+01	-	3.06E+01	3.06E+01
85018	Phenanthrene	mg/kg	-	-	-	-	-	-	-	-	-
1336363	Polychlorinated Biphenyls, Total ^c	mg/kg	-	1.70E+01	1.70E+01	-	2.86E+02	2.86E+02	-	7.98E+01	7.98E+01
12674112	~Aroclor 1016	mg/kg	2.55E+01	4.85E+02	2.55E+01	4.29E+02	8.18E+03	4.29E+02	5.76E+01	2.28E+03	5.76E+01
11104282	~Aroclor 1221	mg/kg	-	1.60E+01	1.60E+01	-	1.17E+02	1.17E+02	-	7.59E+01	7.59E+01
11141165	~Aroclor 1232	mg/kg	-	1.60E+01	1.60E+01	-	1.17E+02	1.17E+02	-	7.59E+01	7.59E+01
53469219	~Aroclor 1242	mg/kg	-	1.70E+01	1.70E+01	-	2.86E+02	2.86E+02	-	7.98E+01	7.98E+01
12672296	~Aroclor 1248	mg/kg	-	1.70E+01	1.70E+01	-	2.86E+02	2.86E+02	-	7.98E+01	7.98E+01
11097691	~Aroclor 1254	mg/kg	7.29E+00	1.70E+01	7.29E+00	1.23E+02	2.86E+02	1.23E+02	1.64E+01	7.98E+01	1.64E+01
11096825	~Aroclor 1260	mg/kg	-	1.70E+01	1.70E+01	-	2.86E+02	2.86E+02	-	7.98E+01	7.98E+01
50328	Polycyclic aromatic hydrocarbons, Total Carcinogenic ^d	mg/kg	-	4.86E+00	4.86E+00	-	7.84E+01	7.84E+01	-	2.33E+01	2.33E+01
56553	~Benz[a]anthracene	mg/kg	-	4.86E+01	4.86E+01	-	7.84E+02	7.84E+02	-	2.33E+02	2.33E+02
50328	~Benzo[a]pyrene	mg/kg	-	4.86E+00	4.86E+00	-	7.84E+01	7.84E+01	-	2.33E+01	2.33E+01
205992	~Benzo[b]fluoranthene	mg/kg	-	4.86E+01	4.86E+01	-	7.84E+02	7.84E+02	-	2.33E+02	2.33E+02
207089	~Benzo[k]fluoranthene	mg/kg	-	4.86E+02	4.86E+02	-	7.84E+03	7.84E+03	-	2.33E+03	2.33E+03
218019	~Chrysene	mg/kg	-	4.86E+03	4.86E+03	-	7.84E+04	7.84E+04	-	2.33E+04	2.33E+04
53703	~Dibenz[a,h]anthracene	mg/kg	-	4.86E+00	4.86E+00	-	7.84E+01	7.84E+01	-	2.33E+01	2.33E+01
193395	~Indeno[1,2,3-cd]pyrene	mg/kg	-	4.86E+01	4.86E+01	-	7.84E+02	7.84E+02	-	2.33E+02	2.33E+02

Hazard-based value calculated using target HI of 3.

Cancer-based value calculated using target ELCR of 1E-04.

Action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.1. Soil/Sediment Action Levels for Significant COPCs at PGDP (Continued)

(Values calculated on 11/5/2013 and are based on best available information.)

CAS	Analyte	Units	Outdoor Worker			Industrial Worker			Adult Recreational User		
			Hazard	Cancer	Action	Hazard	Cancer	Action	Hazard	Cancer	Action
129000	Pyrene	mg/kg	1.14E+04	-	1.14E+04	1.00E+05	-	1.00E+05	2.63E+04	-	2.63E+04
127184	Tetrachloroethylene	mg/kg	1.33E+03	1.13E+04	1.33E+03	1.28E+03	1.14E+04	1.28E+03	4.41E+04	8.38E+04	4.41E+04
79016	Trichloroethylene	mg/kg	7.05E+01	6.21E+02	7.05E+01	6.12E+01	6.73E+02	6.12E+01	3.69E+03	4.82E+03	3.69E+03
75014	Vinyl Chloride	mg/kg	1.07E+03	8.68E+01	8.68E+01	1.26E+03	2.11E+02	2.11E+02	2.21E+04	2.31E+01	2.31E+01
1330207	Xylene, m-	mg/kg	9.87E+03	-	9.87E+03	7.68E+03	-	7.68E+03	1.00E+05	-	1.00E+05
106423	Xylene, Mixture	mg/kg	1.04E+04	-	1.04E+04	8.19E+03	-	8.19E+03	1.00E+05	-	1.00E+05
108383	Xylene, o-	mg/kg	1.15E+04	-	1.15E+04	9.06E+03	-	9.06E+03	1.00E+05	-	1.00E+05
95476	Xylene, p-	mg/kg	1.01E+04	-	1.01E+04	7.86E+03	-	7.86E+03	1.00E+05	-	1.00E+05
14596102	Am-241	pCi/g	-	3.33E+02	3.33E+02	-	1.79E+03	1.79E+03	-	2.28E+03	2.28E+03
10045973	Cs-137+D	pCi/g	-	1.37E+01	1.37E+01	-	5.08E+01	5.08E+01	-	8.48E+01	8.48E+01
13994202	Np-237+D	pCi/g	-	3.22E+01	3.22E+01	-	1.21E+02	1.21E+02	-	1.88E+02	1.88E+02
13981163	Pu-238	pCi/g	-	4.23E+02	4.23E+02	-	2.97E+03	2.97E+03	-	3.30E+03	3.30E+03
15117483	Pu-239	pCi/g	-	3.70E+02	3.70E+02	-	2.60E+03	2.60E+03	-	2.89E+03	2.89E+03
14119336	Pu-240	pCi/g	-	3.71E+02	3.71E+02	-	2.62E+03	2.62E+03	-	2.89E+03	2.89E+03
14133767	Tc-99	pCi/g	-	3.09E+04	3.09E+04	-	1.00E+05	1.00E+05	-	9.92E+04	9.92E+04
14269637	Th-230	pCi/g	-	5.70E+02	5.70E+02	-	3.95E+03	3.95E+03	-	3.88E+03	3.88E+03
13966295	U-234	pCi/g	-	8.72E+02	8.72E+02	-	6.11E+03	6.11E+03	-	5.02E+03	5.02E+03
15117961	U-235+D	pCi/g	-	4.84E+01	4.84E+01	-	1.84E+02	1.84E+02	-	2.83E+02	2.83E+02
7440611	U-238+D	pCi/g	-	1.81E+02	1.81E+02	-	7.48E+02	7.48E+02	-	1.01E+03	1.01E+03

Hazard-based value calculated using target HI of 3.

Cancer-based value calculated using target ELCR of 1E-04.

Action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.1. Soil/Sediment Action Levels for Significant COPCs at PGDP (Continued)

(Values calculated on 11/5/2013 and are based on best available information.)

Analyte	Child Recreational User			Teen Recreational User			Resident	Adult Resident		Child Resident	
	Hazard	Cancer	Action	Hazard	Cancer	Action	Cancer ^e	Hazard	Action	Hazard	Action
Aluminum	1.00E+05	-	1.00E+05	1.00E+05	-	1.00E+05	-	1.00E+05	1.00E+05	1.00E+05	1.00E+05
Antimony (metallic)	2.35E+02	-	2.35E+02	1.34E+03	-	1.34E+03	-	8.76E+02	8.76E+02	9.39E+01	9.39E+01
Arsenic, Inorganic	1.24E+02	1.07E+02	1.07E+02	3.09E+02	1.34E+02	1.34E+02	2.36E+01	2.42E+02	2.36E+01	4.95E+01	2.36E+01
Barium	1.00E+05	-	1.00E+05	1.00E+05	-	1.00E+05	-	1.00E+05	1.00E+05	4.59E+04	4.59E+04
Beryllium and compounds	1.17E+03	1.00E+05	1.17E+03	6.72E+03	1.00E+05	6.72E+03	1.00E+05	4.17E+03	4.17E+03	4.68E+02	4.68E+02
Boron And Borates Only	1.00E+05	-	1.00E+05	1.00E+05	-	1.00E+05	-	1.00E+05	1.00E+05	4.68E+04	4.68E+04
Cadmium (Diet)	3.75E+02	1.00E+05	3.75E+02	8.40E+02	1.00E+05	8.40E+02	1.00E+05	6.57E+02	6.57E+02	1.50E+02	1.50E+02
Chromium (Total) ^a	1.00E+05	4.43E+04	4.43E+04	1.00E+05	4.43E+04	4.43E+04	1.55E+03	1.00E+05	1.55E+03	1.00E+05	1.55E+03
Chromium(III), Insoluble Salts	1.00E+05	-	1.00E+05	1.00E+05	-	1.00E+05	-	1.00E+05	1.00E+05	1.00E+05	1.00E+05
Chromium(VI)	1.76E+03	8.54E+01	8.54E+01	1.01E+04	4.85E+02	4.85E+02	2.93E+01	6.48E+03	2.93E+01	7.02E+02	2.93E+01
Cobalt	1.76E+02	1.00E+05	1.76E+02	1.01E+03	1.00E+05	1.01E+03	3.68E+04	6.42E+02	6.42E+02	7.02E+01	7.02E+01
Copper	2.35E+04	-	2.35E+04	1.00E+05	-	1.00E+05	-	8.76E+04	8.76E+04	9.39E+03	9.39E+03
Iron	1.00E+05	-	1.00E+05	1.00E+05	-	1.00E+05	-	1.00E+05	1.00E+05	1.00E+05	1.00E+05
Lead	-	-	4.00E+02	-	-	4.00E+02	-	-	4.00E+02	-	4.00E+02
Manganese	1.40E+04	-	1.40E+04	1.00E+05	-	1.00E+05	-	4.20E+04	4.20E+04	5.49E+03	5.49E+03
Mercury, Inorganic Salts	1.76E+02	-	1.76E+02	1.01E+03	-	1.01E+03	-	6.57E+02	6.57E+02	7.05E+01	7.05E+01
Molybdenum	2.93E+03	-	2.93E+03	1.68E+04	-	1.68E+04	-	1.10E+04	1.10E+04	1.17E+03	1.17E+03
Nickel Soluble Salts	1.17E+04	1.00E+05	1.17E+04	6.72E+04	1.00E+05	6.72E+04	1.00E+05	3.93E+04	3.93E+04	4.65E+03	4.65E+03
Selenium	2.93E+03	-	2.93E+03	1.68E+04	-	1.68E+04	-	1.10E+04	1.10E+04	1.17E+03	1.17E+03
Silver	2.93E+03	-	2.93E+03	1.68E+04	-	1.68E+04	-	1.10E+04	1.10E+04	1.17E+03	1.17E+03
Thallium (Soluble Salts)	5.88E+00	-	5.88E+00	3.36E+01	-	3.36E+01	-	2.19E+01	2.19E+01	2.35E+00	2.35E+00
Uranium (Soluble Salts)	1.76E+03	-	1.76E+03	1.01E+04	-	1.01E+04	-	6.33E+03	6.33E+03	7.02E+02	7.02E+02
Vanadium and Compounds	2.96E+03	-	2.96E+03	1.70E+04	-	1.70E+04	-	1.08E+04	1.08E+04	1.18E+03	1.18E+03
Zinc and Compounds	1.00E+05	-	1.00E+05	1.00E+05	-	1.00E+05	-	1.00E+05	1.00E+05	7.05E+04	7.05E+04
Acenaphthene	1.25E+04	-	1.25E+04	1.88E+04	-	1.88E+04	-	1.56E+04	1.56E+04	4.98E+03	4.98E+03
Acenaphthylene	-	-	-	-	-	-	-	-	-	-	-
Acrylonitrile	6.06E+02	3.41E+02	3.41E+02	1.00E+05	5.12E+02	5.12E+02	2.37E+01	5.16E+01	2.37E+01	5.16E+01	2.37E+01
Anthracene	6.24E+04	-	6.24E+04	9.39E+04	-	9.39E+04	-	7.80E+04	7.80E+04	2.50E+04	2.50E+04
Benzene	1.52E+03	2.62E+03	1.52E+03	1.34E+04	2.74E+03	2.74E+03	1.08E+02	3.42E+02	1.08E+02	2.59E+02	1.08E+02
Carbazole	-	4.75E+03	4.75E+03	-	3.85E+03	3.85E+03	8.66E+02	-	8.66E+02	-	8.66E+02
Carbon Tetrachloride	1.69E+03	1.78E+03	1.69E+03	1.34E+04	1.62E+03	1.62E+03	6.09E+01	4.77E+02	6.09E+01	3.27E+02	6.09E+01
Chloroform	3.75E+03	1.44E+03	1.44E+03	3.36E+04	8.61E+02	8.61E+02	2.95E+01	8.31E+02	2.95E+01	6.33E+02	2.95E+01
Dichloroethylene, 1,1-	7.08E+03	-	7.08E+03	1.00E+05	-	1.00E+05	-	7.74E+02	7.74E+02	7.29E+02	7.29E+02
Dichloroethylene, 1,2- (Mixed Isomers)	5.28E+03	-	5.28E+03	3.03E+04	-	3.03E+04	-	1.97E+04	1.97E+04	2.11E+03	2.11E+03
Dichloroethylene, 1,2-cis-	1.17E+03	-	1.17E+03	6.72E+03	-	6.72E+03	-	4.38E+03	4.38E+03	4.68E+02	4.68E+02
Dichloroethylene, 1,2-trans-	4.02E+03	-	4.02E+03	6.72E+04	-	6.72E+04	-	5.01E+02	5.01E+02	4.59E+02	4.59E+02
Dieldrin	1.22E+01	5.94E+00	5.94E+00	1.98E+01	4.81E+00	4.81E+00	1.08E+00	1.64E+01	1.08E+00	4.89E+00	1.08E+00

Hazard-based value calculated using target HI of 3.

Cancer-based value calculated using target ELCR of 1E-04.

Action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.1. Soil/Sediment Action Levels for Significant COPCs at PGDP (Continued)

(Values calculated on 11/5/2013 and are based on best available information.)

Analyte	Child Recreational User			Teen Recreational User			Resident	Adult Resident		Child Resident	
	Hazard	Cancer	Action	Hazard	Cancer	Action	Cancer ^e	Hazard	Action	Hazard	Action
Dioxins/Furans, Total (as TCDD) ^b	2.89E-04	1.24E-03	2.89E-04	7.23E-04	1.55E-03	7.23E-04	2.72E-04	5.67E-04	2.72E-04	1.16E-04	1.16E-04
~HpCDD, 2,3,7,8-	2.89E-02	1.24E-01	2.89E-02	7.23E-02	1.55E-01	7.23E-02	2.72E-02	5.67E-02	2.72E-02	1.16E-02	1.16E-02
~HpCDF, 2,3,7,8-	1.71E-02	7.31E-02	1.71E-02	2.77E-02	5.92E-02	2.77E-02	1.33E-02	2.29E-02	1.33E-02	6.84E-03	6.84E-03
~HxCDD, 2,3,7,8-	2.89E-03	1.24E-02	2.89E-03	7.23E-03	1.55E-02	7.23E-03	2.72E-03	5.67E-03	2.72E-03	1.16E-03	1.16E-03
~HxCDF, 2,3,7,8-	1.71E-03	7.31E-03	1.71E-03	2.77E-03	5.92E-03	2.77E-03	1.33E-03	2.29E-03	1.33E-03	6.84E-04	6.84E-04
~OCDD	9.63E-01	4.12E+00	9.63E-01	2.42E+00	5.16E+00	2.42E+00	9.06E-01	1.89E+00	9.06E-01	3.87E-01	3.87E-01
~OCDF	5.70E-01	2.44E+00	5.70E-01	9.24E-01	1.97E+00	9.24E-01	4.44E-01	7.62E-01	4.44E-01	2.28E-01	2.28E-01
~PeCDD, 2,3,7,8-	2.89E-04	1.24E-03	2.89E-04	7.23E-04	1.55E-03	7.23E-04	2.72E-04	5.67E-04	2.72E-04	1.16E-04	1.16E-04
~PeCDF, 1,2,3,7,8-	5.70E-03	2.44E-02	5.70E-03	9.24E-03	1.97E-02	9.24E-03	4.44E-03	7.62E-03	4.44E-03	2.28E-03	2.28E-03
~PeCDF, 2,3,4,7,8-	5.70E-04	2.44E-03	5.70E-04	9.24E-04	1.97E-03	9.24E-04	4.44E-04	7.62E-04	4.44E-04	2.28E-04	2.28E-04
~TCDD, 2,3,7,8-	2.89E-04	1.24E-03	2.89E-04	7.23E-04	1.55E-03	7.23E-04	2.72E-04	5.67E-04	2.72E-04	1.16E-04	1.16E-04
~TCDF, 2,3,7,8-	1.71E-03	7.31E-03	1.71E-03	2.77E-03	5.92E-03	2.77E-03	1.33E-03	2.29E-03	1.33E-03	6.84E-04	6.84E-04
Ethylbenzene	4.68E+04	1.31E+04	1.31E+04	1.00E+05	1.37E+04	1.37E+04	5.39E+02	1.76E+04	5.39E+02	1.05E+04	5.39E+02
Fluoranthene	8.31E+03	-	8.31E+03	1.25E+04	-	1.25E+04	-	1.04E+04	1.04E+04	3.33E+03	3.33E+03
Fluorene	8.31E+03	-	8.31E+03	1.25E+04	-	1.25E+04	-	1.04E+04	1.04E+04	3.33E+03	3.33E+03
Hexachlorobenzene	1.96E+02	5.94E+01	5.94E+01	3.18E+02	4.81E+01	4.81E+01	1.08E+01	2.62E+02	1.08E+01	7.83E+01	1.08E+01
Naphthalene	2.39E+03	2.14E+04	2.39E+03	6.27E+03	1.07E+04	6.27E+03	3.57E+02	4.29E+02	3.57E+02	3.66E+02	3.57E+02
Nitroaniline, 2-	2.44E+03	-	2.44E+03	3.96E+03	-	3.96E+03	-	3.21E+03	3.21E+03	9.72E+02	9.72E+02
Nitroso-di-N-propylamine, N-	-	1.36E+01	1.36E+01	-	1.10E+01	1.10E+01	2.47E+00	-	2.47E+00	-	2.47E+00
Phenanthrene	-	-	-	-	-	-	-	-	-	-	-
Polychlorinated Biphenyls, Total ^c	-	3.85E+01	3.85E+01	-	2.84E+01	2.84E+01	6.70E+00	-	6.70E+00	-	6.70E+00
~Aroclor 1016	1.39E+01	1.10E+03	1.39E+01	2.05E+01	8.12E+02	2.05E+01	1.91E+02	1.71E+01	1.71E+01	5.55E+00	5.55E+00
~Aroclor 1221	-	3.79E+01	3.79E+01	-	2.78E+01	2.78E+01	5.72E+00	-	5.72E+00	-	5.72E+00
~Aroclor 1232	-	3.79E+01	3.79E+01	-	2.78E+01	2.78E+01	5.72E+00	-	5.72E+00	-	5.72E+00
~Aroclor 1242	-	3.85E+01	3.85E+01	-	2.84E+01	2.84E+01	6.70E+00	-	6.70E+00	-	6.70E+00
~Aroclor 1248	-	3.85E+01	3.85E+01	-	2.84E+01	2.84E+01	6.70E+00	-	6.70E+00	-	6.70E+00
~Aroclor 1254	3.96E+00	3.85E+01	3.96E+00	5.85E+00	2.84E+01	5.85E+00	6.70E+00	4.89E+00	4.89E+00	1.58E+00	1.58E+00
~Aroclor 1260	-	3.85E+01	3.85E+01	-	2.84E+01	2.84E+01	6.70E+00	-	6.70E+00	-	6.70E+00
Polycyclic aromatic hydrocarbons, Total Carcinogenic ^d	-	2.08E+00	2.08E+00	-	3.12E+00	3.12E+00	5.77E-01	-	5.77E-01	-	5.77E-01
~Benz[a]anthracene	-	2.08E+01	2.08E+01	-	3.12E+01	3.12E+01	5.77E+00	-	5.77E+00	-	5.77E+00
~Benzo[a]pyrene	-	2.08E+00	2.08E+00	-	3.12E+00	3.12E+00	5.77E-01	-	5.77E-01	-	5.77E-01
~Benzo[b]fluoranthene	-	2.08E+01	2.08E+01	-	3.12E+01	3.12E+01	5.77E+00	-	5.77E+00	-	5.77E+00
~Benzo[k]fluoranthene	-	2.08E+02	2.08E+02	-	3.12E+02	3.12E+02	5.77E+01	-	5.77E+01	-	5.77E+01
~Chrysene	-	2.08E+03	2.08E+03	-	3.12E+03	3.12E+03	5.77E+02	-	5.77E+02	-	5.77E+02
~Dibenz[a,h]anthracene	-	2.08E+00	2.08E+00	-	3.12E+00	3.12E+00	5.77E-01	-	5.77E-01	-	5.77E-01
~Indeno[1,2,3-cd]pyrene	-	2.08E+01	2.08E+01	-	3.12E+01	3.12E+01	5.77E+00	-	5.77E+00	-	5.77E+00

Hazard-based value calculated using target HI of 3.

Cancer-based value calculated using target ELCR of 1E-04.

Action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.1. Soil/Sediment Action Levels for Significant COPCs at PGDP (Continued)

(Values calculated on 11/5/2013 and are based on best available information.)

Analyte	Child Recreational User			Teen Recreational User			Resident	Adult Resident		Child Resident	
	Hazard	Cancer	Action	Hazard	Cancer	Action	Cancer ^e	Hazard	Action	Hazard	Action
Pyrene	6.24E+03	-	6.24E+03	9.39E+03	-	9.39E+03	-	7.80E+03	7.80E+03	2.50E+03	2.50E+03
Tetrachloroethylene	1.83E+03	6.15E+04	1.83E+03	2.02E+04	5.78E+04	2.02E+04	2.19E+03	3.09E+02	3.09E+02	2.58E+02	2.58E+02
Trichloroethylene	1.11E+02	5.86E+02	1.11E+02	1.68E+03	1.22E+03	1.22E+03	4.75E+01	1.47E+01	1.47E+01	1.32E+01	1.32E+01
Vinyl Chloride	1.21E+03	7.03E+00	7.03E+00	1.01E+04	2.31E+01	2.31E+01	5.97E+00	3.06E+02	5.97E+00	2.21E+02	5.97E+00
Xylene, m-	1.86E+04	-	1.86E+04	1.00E+05	-	1.00E+05	-	1.84E+03	1.84E+03	1.77E+03	1.77E+03
Xylene, Mixture	1.96E+04	-	1.96E+04	1.00E+05	-	1.00E+05	-	1.95E+03	1.95E+03	1.88E+03	1.88E+03
Xylene, o-	2.13E+04	-	2.13E+04	1.00E+05	-	1.00E+05	-	2.16E+03	2.16E+03	2.08E+03	2.08E+03
Xylene, p-	1.89E+04	-	1.89E+04	1.00E+05	-	1.00E+05	-	1.87E+03	1.87E+03	1.81E+03	1.81E+03
Am-241	-	2.12E+03	2.12E+03	-	1.70E+03	1.70E+03	2.41E+02	-	2.41E+02	-	2.41E+02
Cs-137+D	-	1.26E+02	1.26E+02	-	6.30E+01	6.30E+01	1.00E+01	-	1.00E+01	-	1.00E+01
Np-237+D	-	2.70E+02	2.70E+02	-	1.40E+02	1.40E+02	2.21E+01	-	2.21E+01	-	2.21E+01
Pu-238	-	2.45E+03	2.45E+03	-	2.45E+03	2.45E+03	3.26E+02	-	3.26E+02	-	3.26E+02
Pu-239	-	2.15E+03	2.15E+03	-	2.14E+03	2.14E+03	2.85E+02	-	2.85E+02	-	2.85E+02
Pu-240	-	2.15E+03	2.15E+03	-	2.14E+03	2.14E+03	2.85E+02	-	2.85E+02	-	2.85E+02
Tc-99	-	7.56E+04	7.56E+04	-	7.37E+04	7.37E+04	9.91E+03	-	9.91E+03	-	9.91E+03
Th-230	-	2.91E+03	2.91E+03	-	2.88E+03	2.88E+03	3.84E+02	-	3.84E+02	-	3.84E+02
U-234	-	3.75E+03	3.75E+03	-	3.73E+03	3.73E+03	4.97E+02	-	4.97E+02	-	4.97E+02
U-235+D	-	3.98E+02	3.98E+02	-	2.10E+02	2.10E+02	3.31E+01	-	3.31E+01	-	3.31E+01
U-238+D	-	1.18E+03	1.18E+03	-	7.48E+02	7.48E+02	1.13E+02	-	1.13E+02	-	1.13E+02

NOTE: The action level for HI is 3 because the range of values for HI (based on RGO tables) are 0.1, 1, and 3. Please see Figure 1.1 of the Risk Methods Document.

^a Chromium (Total) utilizes Chromium III hazard values and Chromium VI inhalation risk values (to be conservative in terms of protecting human health).

^b Total dioxins/furans uses values for 2,3,7,8-TCDD, see screening note 9g in the Appendix A introduction, 'Screening Levels' on pages A-3 through A-5.

^c Total PCBs ALs are based on the high risk toxicity values.

^d Total carcinogenic PAHs uses values for benzo(a)pyrene, see screening note 9e in the Appendix A introduction, 'Screening Levels' on pages A-3 through A-5.

^e For the resident, ELCRs (i.e. cancer ALs) were calculated using the child/adult age-adjusted lifetime scenario.

Table A.2. Groundwater Action Levels for Significant COPCs at PGDP
(Values calculated on 11/5/2013 and are based on best available information.)

CAS	Analyte	Units	Resident	Adult Resident		Child Resident	
			Cancer ^c	Hazard	Action	Hazard	Action
7429905	Aluminum	mg/L	-	1.09E+02	1.09E+02	3.12E+01	3.12E+01
7440360	Antimony (metallic)	mg/L	-	4.23E-02	4.23E-02	1.25E-02	1.25E-02
7440382	Arsenic, Inorganic	mg/L	3.78E-03	3.27E-02	3.78E-03	9.39E-03	3.78E-03
7440393	Barium	mg/L	-	2.04E+01	2.04E+01	6.18E+00	6.18E+00
7440417	Beryllium and compounds	mg/L	-	1.25E-01	1.25E-01	5.58E-02	5.58E-02
7440428	Boron And Borates Only	mg/L	-	2.18E+01	2.18E+01	6.24E+00	6.24E+00
7440439	Cadmium (Water)	mg/L	-	4.95E-02	4.95E-02	1.54E-02	1.54E-02
16065831	Chromium (Total) ^a	mg/L	-	1.17E+02	1.17E+02	4.41E+01	4.41E+01
18540299	Chromium(III), Insoluble Salts	mg/L	-	1.17E+02	1.17E+02	4.41E+01	4.41E+01
7440473	Chromium(VI)	mg/L	3.00E-03	2.32E-01	3.00E-03	8.79E-02	3.00E-03
7440484	Cobalt	mg/L	-	3.27E-02	3.27E-02	9.39E-03	9.39E-03
7440508	Copper	mg/L	-	4.35E+00	4.35E+00	1.25E+00	1.25E+00
7439896	Iron	mg/L	-	7.62E+01	7.62E+01	2.19E+01	2.19E+01
7439921	Lead	mg/L	-	-	3.00E-02	-	3.00E-02
7439965	Manganese	mg/L	-	2.33E+00	2.33E+00	7.35E-01	7.35E-01
7439976	Mercury, Inorganic Salts	mg/L	-	3.06E-02	3.06E-02	9.27E-03	9.27E-03
7439987	Molybdenum	mg/L	-	5.46E-01	5.46E-01	1.56E-01	1.56E-01
7440020	Nickel Soluble Salts	mg/L	-	2.13E+00	2.13E+00	6.24E-01	6.24E-01
7782492	Selenium	mg/L	-	5.46E-01	5.46E-01	1.56E-01	1.56E-01
7440224	Silver	mg/L	-	5.07E-01	5.07E-01	1.55E-01	1.55E-01
7791120	Thallium (Soluble Salts)	mg/L	-	1.09E-03	1.09E-03	3.12E-04	3.12E-04
238	Uranium (Soluble Salts)	mg/L	-	3.27E-01	3.27E-01	9.39E-02	9.39E-02
7440622	Vanadium and Compounds	mg/L	-	4.59E-01	4.59E-01	1.53E-01	1.53E-01
7440666	Zinc and Compounds	mg/L	-	3.27E+01	3.27E+01	9.39E+00	9.39E+00
83329	Acenaphthene	mg/L	-	2.71E+00	2.71E+00	1.34E+00	1.34E+00
208968	Acenaphthylene	mg/L	-	-	-	-	-
107131	Acrylonitrile	mg/L	4.25E-03	1.25E-02	4.25E-03	1.24E-02	4.25E-03
120127	Anthracene	mg/L	-	8.76E+00	8.76E+00	5.28E+00	5.28E+00
71432	Benzene	mg/L	3.79E-02	1.26E-01	3.79E-02	7.32E-02	3.79E-02
86748	Carbazole	mg/L	1.98E-01	-	1.98E-01	-	1.98E-01
56235	Carbon Tetrachloride	mg/L	3.82E-02	2.22E-01	3.82E-02	9.81E-02	3.82E-02
67663	Chloroform	mg/L	1.89E-02	3.81E-01	1.89E-02	2.03E-01	1.89E-02
75354	Dichloroethylene, 1,1-	mg/L	-	9.93E-01	9.93E-01	6.84E-01	6.84E-01
540590	Dichloroethylene, 1,2- (Mixed Isomers)	mg/L	-	8.76E-01	8.76E-01	2.72E-01	2.72E-01
156592	Dichloroethylene, 1,2-cis-	mg/L	-	1.94E-01	1.94E-01	6.03E-02	6.03E-02
156605	Dichloroethylene, 1,2-trans-	mg/L	-	3.15E-01	3.15E-01	2.32E-01	2.32E-01
60571	Dieldrin	mg/L	1.93E-04	1.91E-03	1.93E-04	1.02E-03	1.93E-04
1746016	Dioxins/Furans, Total (as TCDD) ^b	mg/L	4.37E-08	7.68E-08	4.37E-08	2.19E-08	2.19E-08
37871004	~HpCDD, 2,3,7,8-	mg/L	4.37E-06	7.68E-06	4.37E-06	2.19E-06	2.19E-06
38998753	~HpCDF, 2,3,7,8-	mg/L	4.37E-06	7.68E-06	4.37E-06	2.19E-06	2.19E-06
34465468	~HxCDD, 2,3,7,8-	mg/L	4.37E-07	7.68E-07	4.37E-07	2.19E-07	2.19E-07
55684941	~HxCDF, 2,3,7,8-	mg/L	4.37E-07	7.68E-07	4.37E-07	2.19E-07	2.19E-07
3268879	~OCDD	mg/L	1.46E-04	2.56E-04	1.46E-04	7.29E-05	7.29E-05
39001020	~OCDF	mg/L	1.46E-04	2.56E-04	1.46E-04	7.29E-05	7.29E-05

Hazard-based value calculated using target HI of 3.

Cancer-based value calculated using target ELCR of 1E-04.

Action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.2. Groundwater Action Levels for Significant COPCs at PGDP (Continued)
 (Values calculated on 11/5/2013 and are based on best available information.)

CAS	Analyte	Units	Resident	Adult Resident		Child Resident	
			Cancer ^e	Hazard	Action	Hazard	Action
36088229	~PeCDD, 2,3,7,8-	mg/L	4.37E-08	7.68E-08	4.37E-08	2.19E-08	2.19E-08
57117416	~PeCDF, 1,2,3,7,8-	mg/L	1.46E-06	2.56E-06	1.46E-06	7.29E-07	7.29E-07
57117314	~PeCDF, 2,3,4,7,8-	mg/L	1.46E-07	2.56E-07	1.46E-07	7.29E-08	7.29E-08
1746016	~TCDD, 2,3,7,8-	mg/L	4.37E-08	7.68E-08	4.37E-08	2.19E-08	2.19E-08
51207319	~TCDF, 2,3,7,8-	mg/L	4.37E-07	7.68E-07	4.37E-07	2.19E-07	2.19E-07
100414	Ethylbenzene	mg/L	1.32E-01	3.27E+00	1.32E-01	1.88E+00	1.32E-01
206440	Fluoranthene	mg/L	-	4.38E+00	4.38E+00	1.25E+00	1.25E+00
86737	Fluorene	mg/L	-	1.47E+00	1.47E+00	8.04E-01	8.04E-01
118741	Hexachlorobenzene	mg/L	3.55E-03	8.76E-02	3.55E-03	2.50E-02	3.55E-03
91203	Naphthalene	mg/L	1.43E-02	1.85E-02	1.43E-02	1.81E-02	1.43E-02
88744	Nitroaniline, 2-	mg/L	-	1.03E+00	1.03E+00	3.06E-01	3.06E-01
621647	Nitroso-di-N-propylamine, N-	mg/L	7.99E-04	-	7.99E-04	-	7.99E-04
85018	Phenanthrene	mg/L	-	-	-	-	-
1336363	Polychlorinated Biphenyls, Total ^c	mg/L	1.42E-02	-	1.42E-02	-	1.42E-02
12674112	~Aroclor 1016	mg/L	8.11E-02	7.68E-03	7.68E-03	2.19E-03	2.19E-03
11104282	~Aroclor 1221	mg/L	4.82E-04	-	4.82E-04	-	4.82E-04
11141165	~Aroclor 1232	mg/L	4.82E-04	-	4.82E-04	-	4.82E-04
53469219	~Aroclor 1242	mg/L	2.84E-03	-	2.84E-03	-	2.84E-03
12672296	~Aroclor 1248	mg/L	2.84E-03	-	2.84E-03	-	2.84E-03
11097691	~Aroclor 1254	mg/L	2.84E-03	2.19E-03	2.19E-03	6.27E-04	6.27E-04
11096825	~Aroclor 1260	mg/L	2.84E-03	-	2.84E-03	-	2.84E-03
50328	Polycyclic aromatic hydrocarbons, Total Carcinogenic ^d	mg/L	2.24E-04	-	2.24E-04	-	2.24E-04
56553	~Benz[a]anthracene	mg/L	2.24E-03	-	2.24E-03	-	2.24E-03
50328	~Benzo[a]pyrene	mg/L	2.24E-04	-	2.24E-04	-	2.24E-04
205992	~Benzo[b]fluoranthene	mg/L	2.24E-03	-	2.24E-03	-	2.24E-03
207089	~Benzo[k]fluoranthene	mg/L	2.24E-02	-	2.24E-02	-	2.24E-02
218019	~Chrysene	mg/L	2.24E-01	-	2.24E-01	-	2.24E-01
53703	~Dibenz[a,h]anthracene	mg/L	2.24E-04	-	2.24E-04	-	2.24E-04
193395	~Indeno[1,2,3-cd]pyrene	mg/L	2.24E-03	-	2.24E-03	-	2.24E-03
129000	Pyrene	mg/L	-	5.91E-01	5.91E-01	4.11E-01	4.11E-01
127184	Tetrachloroethylene	mg/L	9.97E-01	1.56E-01	1.56E-01	9.78E-02	9.78E-02
79016	Trichloroethylene	mg/L	1.95E-02	9.87E-03	9.87E-03	6.81E-03	6.81E-03
75014	Vinyl Chloride	mg/L	1.11E-03	2.05E-01	1.11E-03	8.01E-02	1.11E-03
1330207	Xylene, m-	mg/L	-	5.97E-01	5.97E-01	5.61E-01	5.61E-01
106423	Xylene, Mixture	mg/L	-	5.97E-01	5.97E-01	5.61E-01	5.61E-01
108383	Xylene, o-	mg/L	-	6.00E-01	6.00E-01	5.61E-01	5.61E-01
95476	Xylene, p-	mg/L	-	5.97E-01	5.97E-01	5.61E-01	5.61E-01
14596102	Am-241	pCi/L	4.82E+01	-	4.82E+01	-	4.82E+01
10045973	Cs-137+D	pCi/L	1.65E+02	-	1.65E+02	-	1.65E+02
13994202	Np-237+D	pCi/L	7.44E+01	-	7.44E+01	-	7.44E+01
13981163	Pu-238	pCi/L	3.83E+01	-	3.83E+01	-	3.83E+01
15117483	Pu-239	pCi/L	3.71E+01	-	3.71E+01	-	3.71E+01
14119336	Pu-240	pCi/L	3.71E+01	-	3.71E+01	-	3.71E+01
14133767	Tc-99	pCi/L	1.82E+03	-	1.82E+03	-	1.82E+03
14269637	Th-230	pCi/L	5.51E+01	-	5.51E+01	-	5.51E+01

Hazard-based value calculated using target HI of 3.

Cancer-based value calculated using target ELCR of 1E-04.

Action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.2. Groundwater Action Levels for Significant COPCs at PGDP (Continued)
 (Values calculated on 11/5/2013 and are based on best available information.)

CAS	Analyte	Units	Resident	Adult Resident		Child Resident	
			Cancer ^e	Hazard	Action	Hazard	Action
13966295	U-234	pCi/L	7.09E+01	-	7.09E+01	-	7.09E+01
15117961	U-235+D	pCi/L	6.98E+01	-	6.98E+01	-	6.98E+01
7440611	U-238+D	pCi/L	5.75E+01	-	5.75E+01	-	5.75E+01

NOTE: The action level for HI is 3 because the range of values for HI (based on RGO tables) are 0.1, 1, and 3.

Please see Figure 1.1 of the Risk Methods Document.

^a Chromium (Total) utilizes Chromium III hazard values and Chromium VI inhalation risk values (to be conservative in terms of protecting human health).

^b Total dioxins/furans uses values for 2,3,7,8-TCDD, see screening note 9g in the Appendix A introduction, 'Screening Levels' on pages A-3 through A-5.

^c Total PCBs ALs are based on the high risk toxicity values.

^d Total carcinogenic PAHs uses values for benzo(a)pyrene, see screening note 9e in the Appendix A introduction, 'Screening Levels' on pages A-3 through A-5.

^e For the resident, ELCRs (i.e. cancer ALs) were calculated using the child/adult age-adjusted lifetime scenario (i.e., lifetime exposure).

Hazard-based value calculated using target HI of 3.

Cancer-based value calculated using target ELCR of 1E-04.

Action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.3. Surface Water Action Levels for Significant COPCs at PGDP

(Values calculated on 11/6/2013 and are based on best available information.)

CAS	Analyte	Units	Outdoor Worker Wading ^c			Industrial Worker Wading ^c		
			Hazard	Cancer	Action	Hazard	Cancer	Action
7429905	Aluminum	mg/L	1.02E+05	-	1.02E+05	2.51E+04	-	2.51E+04
7440360	Antimony (metallic)	mg/L	6.12E+00	-	6.12E+00	1.51E+00	-	1.51E+00
7440382	Arsenic, Inorganic	mg/L	3.06E+01	6.34E+00	6.34E+00	7.53E+00	1.56E+00	1.56E+00
7440393	Barium	mg/L	1.43E+03	-	1.43E+03	3.51E+02	-	3.51E+02
7440417	Beryllium and compounds	mg/L	1.43E+00	-	1.43E+00	3.51E-01	-	3.51E-01
7440428	Boron And Borates Only	mg/L	2.04E+04	-	2.04E+04	5.01E+03	-	5.01E+03
7440439	Cadmium (Water)	mg/L	2.55E+00	-	2.55E+00	6.27E-01	-	6.27E-01
16065831	Chromium (Total) ^d	mg/L	1.99E+03	-	1.99E+03	4.89E+02	-	4.89E+02
18540299	Chromium(III), Insoluble Salts	mg/L	1.99E+03	-	1.99E+03	4.89E+02	-	4.89E+02
7440473	Chromium(VI)	mg/L	3.81E+00	1.27E-01	1.27E-01	9.42E-01	3.11E-02	3.11E-02
7440484	Cobalt	mg/L	7.65E+01	-	7.65E+01	1.88E+01	-	1.88E+01
7440508	Copper	mg/L	4.08E+03	-	4.08E+03	1.01E+03	-	1.01E+03
7439896	Iron	mg/L	7.14E+04	-	7.14E+04	1.76E+04	-	1.76E+04
7439921	Lead	mg/L	-	-	3.00E-02	-	-	3.00E-02
7439965	Manganese	mg/L	9.78E+01	-	9.78E+01	2.41E+01	-	2.41E+01
7439976	Mercury, Inorganic Salts	mg/L	2.14E+00	-	2.14E+00	5.28E-01	-	5.28E-01
7439987	Molybdenum	mg/L	5.10E+02	-	5.10E+02	1.25E+02	-	1.25E+02
7440020	Nickel Soluble Salts	mg/L	4.08E+02	-	4.08E+02	1.01E+02	-	1.01E+02
7782492	Selenium	mg/L	5.10E+02	-	5.10E+02	1.25E+02	-	1.25E+02
7440224	Silver	mg/L	3.39E+01	-	3.39E+01	8.37E+00	-	8.37E+00
7791120	Thallium (Soluble Salts)	mg/L	1.02E+00	-	1.02E+00	2.51E-01	-	2.51E-01
238	Uranium (Soluble Salts)	mg/L	3.06E+02	-	3.06E+02	7.53E+01	-	7.53E+01
7440622	Vanadium and Compounds	mg/L	1.34E+01	-	1.34E+01	3.30E+00	-	3.30E+00
7440666	Zinc and Compounds	mg/L	5.10E+04	-	5.10E+04	1.25E+04	-	1.25E+04
83329	Acenaphthene	mg/L	2.94E+01	-	2.94E+01	1.15E+01	-	1.15E+01
208968	Acenaphthylene	mg/L	2.94E+01	-	-	1.15E+01	-	-
107131	Acrylonitrile	mg/L	2.48E+03	1.07E+01	1.07E+01	7.47E+02	3.23E+00	3.23E+00
120127	Anthracene	mg/L	7.62E+01	-	7.62E+01	3.03E+01	-	3.03E+01
71432	Benzene	mg/L	1.76E+01	7.46E+00	7.46E+00	5.70E+00	2.41E+00	2.41E+00
86748	Carbazole	mg/L	-	3.37E+00	3.37E+00	-	1.31E+00	1.31E+00
56235	Carbon Tetrachloride	mg/L	1.04E+01	3.45E+00	3.45E+00	3.93E+00	1.31E+00	1.31E+00
67663	Chloroform	mg/L	7.71E+01	2.32E+01	2.32E+01	2.70E+01	8.13E+00	8.13E+00
75354	Dichloroethylene, 1,1-	mg/L	2.53E+02	-	2.53E+02	8.55E+01	-	8.55E+01
540590	Dichloroethylene, 1,2- (Mixed Isomers)	mg/L	4.83E+01	-	4.83E+01	1.64E+01	-	1.64E+01
156592	Dichloroethylene, 1,2-cis-	mg/L	1.07E+01	-	1.07E+01	3.63E+00	-	3.63E+00
156605	Dichloroethylene, 1,2-trans-	mg/L	1.07E+02	-	1.07E+02	3.63E+01	-	3.63E+01
60571	Dieldrin	mg/L	1.87E-02	2.18E-03	2.18E-03	7.41E-03	8.66E-04	8.66E-04

Hazard-based value calculated using target HI of 3.

Cancer-based value calculated using target ELCR of 1E-04.

Action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.3. Surface Water Action Levels for Significant COPCs at PGDP (Continued)
(Values calculated on 11/6/2013 and are based on best available information.)

CAS	Analyte	Units	Outdoor Worker Wading ^c			Industrial Worker Wading ^c		
			Hazard	Cancer	Action	Hazard	Cancer	Action
1746016	Dioxins/Furans, Total (as TCDD) ^b	mg/L	-	-	-	-	-	-
37871004	~HpCDD, 2,3,7,8-	mg/L	-	-	-	-	-	-
38998753	~HpCDF, 2,3,7,8-	mg/L	-	-	-	-	-	-
34465468	~HxCDD, 2,3,7,8-	mg/L	-	-	-	-	-	-
55684941	~HxCDF, 2,3,7,8-	mg/L	-	-	-	-	-	-
3268879	~OCDD	mg/L	-	-	-	-	-	-
39001020	~OCDF	mg/L	-	-	-	-	-	-
36088229	~PeCDD, 2,3,7,8-	mg/L	-	-	-	-	-	-
57117416	~PeCDF, 1,2,3,7,8-	mg/L	-	-	-	-	-	-
57117314	~PeCDF, 2,3,4,7,8-	mg/L	-	-	-	-	-	-
1746016	~TCDD, 2,3,7,8-	mg/L	-	-	-	-	-	-
51207319	~TCDF, 2,3,7,8-	mg/L	-	-	-	-	-	-
100414	Ethylbenzene	mg/L	1.14E+02	9.64E+00	9.64E+00	4.20E+01	3.55E+00	3.55E+00
206440	Fluoranthene	mg/L	-	-	-	-	-	-
86737	Fluorene	mg/L	1.42E+01	-	1.42E+01	5.49E+00	-	5.49E+00
118741	Hexachlorobenzene	mg/L	-	-	-	-	-	-
91203	Naphthalene	mg/L	2.14E+01	-	2.14E+01	8.04E+00	-	8.04E+00
88744	Nitroaniline, 2-	mg/L	1.05E+02	-	1.05E+02	3.84E+01	-	3.84E+01
621647	Nitroso-di-N-propylamine, N-	mg/L	-	2.81E-01	2.81E-01	-	1.01E-01	1.01E-01
85018	Phenanthrene	mg/L	2.94E+01	-	-	-	-	-
1336363	Polychlorinated Biphenyls, Total ^c	mg/L	-	-	-	-	-	-
12674112	~Aroclor 1016	mg/L	-	-	-	-	-	-
11104282	~Aroclor 1221	mg/L	-	9.36E-03	9.36E-03	-	3.72E-03	3.72E-03
11141165	~Aroclor 1232	mg/L	-	9.36E-03	9.36E-03	-	3.72E-03	3.72E-03
53469219	~Aroclor 1242	mg/L	-	-	-	-	-	-
12672296	~Aroclor 1248	mg/L	-	-	-	-	-	-
11097691	~Aroclor 1254	mg/L	-	-	-	-	-	-
11096825	~Aroclor 1260	mg/L	-	-	-	-	-	-
50328	Polycyclic aromatic hydrocarbons, Total Carcinogenic ^d	mg/L	-	-	-	-	-	-
56553	~Benz[a]anthracene	mg/L	-	-	-	-	-	-
50328	~Benzo[a]pyrene	mg/L	-	-	-	-	-	-
205992	~Benzo[b]fluoranthene	mg/L	-	-	-	-	-	-
207089	~Benzo[k]fluoranthene	mg/L	-	-	-	-	-	-
218019	~Chrysene	mg/L	-	-	-	-	-	-
53703	~Dibenz[a,h]anthracene	mg/L	-	-	-	-	-	-
193395	~Indeno[1,2,3-cd]pyrene	mg/L	-	-	-	-	-	-

Hazard-based value calculated using target HI of 3.

Cancer-based value calculated using target ELCR of 1E-04.

Action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.3. Surface Water Action Levels for Significant COPCs at PGDP (Continued)
 (Values calculated on 11/6/2013 and are based on best available information.)

CAS	Analyte	Units	Outdoor Worker Wading ^c			Industrial Worker Wading ^c		
			Hazard	Cancer	Action	Hazard	Cancer	Action
129000	Pyrene	mg/L	4.62E+00	-	4.62E+00	1.83E+00	-	1.83E+00
127184	Tetrachloroethylene	mg/L	7.02E+00	5.20E+01	7.02E+00	2.72E+00	2.02E+01	2.72E+00
79016	Trichloroethylene	mg/L	2.10E+00	4.54E+00	2.10E+00	7.65E-01	1.65E+00	7.65E-01
75014	Vinyl Chloride	mg/L	2.50E+01	2.08E-02	2.08E-02	7.74E+00	6.57E-02	6.57E-02
1330207	Xylene, m-	mg/L	2.10E+02	-	2.10E+02	7.80E+01	-	7.80E+01
106423	Xylene, Mixture	mg/L	2.24E+02	-	2.24E+02	8.28E+01	-	8.28E+01
108383	Xylene, o-	mg/L	2.38E+02	-	2.38E+02	8.73E+01	-	8.73E+01
95476	Xylene, p-	mg/L	2.27E+02	-	2.27E+02	8.37E+01	-	8.37E+01
14596102	Am-241	pCi/L	-	-	-	-	-	-
10045973	Cs-137+D	pCi/L	-	-	-	-	-	-
13994202	Np-237+D	pCi/L	-	-	-	-	-	-
13981163	Pu-238	pCi/L	-	-	-	-	-	-
15117483	Pu-239	pCi/L	-	-	-	-	-	-
14119336	Pu-240	pCi/L	-	-	-	-	-	-
14133767	Tc-99	pCi/L	-	-	-	-	-	-
14269637	Th-230	pCi/L	-	-	-	-	-	-
13966295	U-234	pCi/L	-	-	-	-	-	-
15117961	U-235+D	pCi/L	-	-	-	-	-	-
7440611	U-238+D	pCi/L	-	-	-	-	-	-

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Hazard-based value calculated using target HI of 3.

Cancer-based value calculated using target ELCR of 1E-04.

Action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.3. Surface Water Action Levels for Significant COPCs at PGDP (Continued)

(Values calculated on 11/6/2013 and are based on best available information.)

Analyte	Units	Adult Recreational User Swimming			Child Recreational User Swimming			Teen Recreational User Swimming		
		Hazard	Cancer	Action	Hazard	Cancer	Action	Hazard	Cancer	Action
Aluminum	mg/L	2.51E+04	-	2.51E+04	2.48E+03	-	2.48E+03	2.51E+04	-	2.51E+04
Antimony (metallic)	mg/L	4.02E+00	-	4.02E+00	6.03E-01	-	6.03E-01	4.02E+00	-	4.02E+00
Arsenic, Inorganic	mg/L	7.50E+00	1.25E+00	1.25E+00	7.44E-01	6.44E-01	6.44E-01	7.50E+00	8.27E-01	8.27E-01
Barium	mg/L	1.11E+03	-	1.11E+03	1.97E+02	-	1.97E+02	1.11E+03	-	1.11E+03
Beryllium and compounds	mg/L	1.30E+00	-	1.30E+00	2.87E-01	-	2.87E-01	1.30E+00	-	1.30E+00
Boron And Borates Only	mg/L	5.01E+03	-	5.01E+03	4.98E+02	-	4.98E+02	5.01E+03	-	5.01E+03
Cadmium (Diet)	mg/L	2.08E+00	-	2.08E+00	3.90E-01	-	3.90E-01	2.08E+00	-	2.08E+00
Chromium (Total) ^a	mg/L	1.78E+03	-	1.78E+03	3.84E+02	-	3.84E+02	1.78E+03	-	1.78E+03
Chromium(III), Insoluble Salts	mg/L	1.78E+03	-	1.78E+03	3.84E+02	-	3.84E+02	1.78E+03	-	1.78E+03
Chromium(VI)	mg/L	3.42E+00	1.70E-01	1.70E-01	7.38E-01	3.59E-02	3.59E-02	3.42E+00	5.35E-02	5.35E-02
Cobalt	mg/L	8.94E+00	-	8.94E+00	8.01E-01	-	8.01E-01	8.94E+00	-	8.94E+00
Copper	mg/L	1.00E+03	-	1.00E+03	9.93E+01	-	9.93E+01	1.00E+03	-	1.00E+03
Iron	mg/L	1.75E+04	-	1.75E+04	1.74E+03	-	1.74E+03	1.75E+04	-	1.75E+04
Lead	mg/L	-	-	3.00E-02	-	-	3.00E-02	-	-	3.00E-02
Manganese	mg/L	8.19E+01	-	8.19E+01	1.59E+01	-	1.59E+01	8.19E+01	-	8.19E+01
Mercury, Inorganic Salts	mg/L	1.67E+00	-	1.67E+00	2.95E-01	-	2.95E-01	1.67E+00	-	1.67E+00
Molybdenum	mg/L	1.25E+02	-	1.25E+02	1.24E+01	-	1.24E+01	1.25E+02	-	1.25E+02
Nickel Soluble Salts	mg/L	2.43E+02	-	2.43E+02	3.39E+01	-	3.39E+01	2.43E+02	-	2.43E+02
Selenium	mg/L	1.25E+02	-	1.25E+02	1.24E+01	-	1.24E+01	1.25E+02	-	1.25E+02
Silver	mg/L	2.66E+01	-	2.66E+01	4.77E+00	-	4.77E+00	2.66E+01	-	2.66E+01
Thallium (Soluble Salts)	mg/L	2.51E-01	-	2.51E-01	2.48E-02	-	2.48E-02	2.51E-01	-	2.51E-01
Uranium (Soluble Salts)	mg/L	7.50E+01	-	7.50E+01	7.44E+00	-	7.44E+00	7.50E+01	-	7.50E+01
Vanadium and Compounds	mg/L	1.16E+01	-	1.16E+01	2.36E+00	-	2.36E+00	1.16E+01	-	1.16E+01
Zinc and Compounds	mg/L	8.40E+03	-	8.40E+03	7.80E+02	-	7.80E+02	8.40E+03	-	8.40E+03
Acenaphthene	mg/L	2.69E+01	-	2.69E+01	9.36E+00	-	9.36E+00	2.69E+01	-	2.69E+01
Acenaphthylene	mg/L	2.69E+01	-	-	9.36E+00	-	-	2.69E+01	-	-
Acrylonitrile	mg/L	8.58E+02	3.17E+00	3.17E+00	9.57E+01	1.72E+00	1.72E+00	8.58E+02	2.14E+00	2.14E+00
Anthracene	mg/L	7.02E+01	-	7.02E+01	2.52E+01	-	2.52E+01	7.02E+01	-	7.02E+01
Benzene	mg/L	1.46E+01	6.25E+00	6.25E+00	3.42E+00	6.03E+00	3.42E+00	1.46E+01	5.06E+00	5.06E+00
Carbazole	mg/L	-	3.81E+00	3.81E+00	-	4.33E+00	4.33E+00	-	3.21E+00	3.21E+00
Carbon Tetrachloride	mg/L	8.97E+00	3.56E+00	3.56E+00	2.61E+00	3.62E+00	2.61E+00	8.97E+00	2.92E+00	2.92E+00
Chloroform	mg/L	5.91E+01	1.88E+01	1.88E+01	1.27E+01	1.60E+01	1.27E+01	5.91E+01	1.47E+01	1.47E+01
Dichloroethylene, 1,1-	mg/L	2.06E+02	-	2.06E+02	4.83E+01	-	4.83E+01	2.06E+02	-	2.06E+02
Dichloroethylene, 1,2- (Mixed Isomers)	mg/L	3.93E+01	-	3.93E+01	9.06E+00	-	9.06E+00	3.93E+01	-	3.93E+01
Dichloroethylene, 1,2-cis-	mg/L	8.70E+00	-	8.70E+00	2.01E+00	-	2.01E+00	8.70E+00	-	8.70E+00
Dichloroethylene, 1,2-trans-	mg/L	8.70E+01	-	8.70E+01	2.01E+01	-	2.01E+01	8.70E+01	-	8.70E+01
Dieldrin	mg/L	1.72E-02	2.55E-03	2.55E-03	6.12E-03	2.97E-03	2.97E-03	1.72E-02	2.16E-03	2.16E-03

Hazard-based value calculated using target HI of 3.

Cancer-based value calculated using target ELCR of 1E-04.

Action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.3. Surface Water Action Levels for Significant COPCs at PGDP (Continued)

(Values calculated on 11/6/2013 and are based on best available information.)

Analyte	Units	Adult Recreational User Swimming			Child Recreational User Swimming			Teen Recreational User Swimming		
		Hazard	Cancer	Action	Hazard	Cancer	Action	Hazard	Cancer	Action
Dioxins/Furans, Total (as TCDD) ^b	mg/L	2.39E-05	1.96E-05	1.96E-05	1.97E-06	8.40E-06	1.97E-06	2.39E-05	1.20E-05	1.20E-05
~HpCDD, 2,3,7,8-	mg/L	2.39E-03	1.96E-03	1.96E-03	1.97E-04	8.40E-04	1.97E-04	2.39E-03	1.20E-03	1.20E-03
~HpCDF, 2,3,7,8-	mg/L	2.39E-03	1.96E-03	1.96E-03	1.97E-04	8.40E-04	1.97E-04	2.39E-03	1.20E-03	1.20E-03
~HxCDD, 2,3,7,8-	mg/L	2.39E-04	1.96E-04	1.96E-04	1.97E-05	8.40E-05	1.97E-05	2.39E-04	1.20E-04	1.20E-04
~HxCDF, 2,3,7,8-	mg/L	2.39E-04	1.96E-04	1.96E-04	1.97E-05	8.40E-05	1.97E-05	2.39E-04	1.20E-04	1.20E-04
~OCDD	mg/L	7.95E-02	6.53E-02	6.53E-02	6.54E-03	2.80E-02	6.54E-03	7.95E-02	4.01E-02	4.01E-02
~OCDF	mg/L	7.95E-02	6.53E-02	6.53E-02	6.54E-03	2.80E-02	6.54E-03	7.95E-02	4.01E-02	4.01E-02
~PeCDD, 2,3,7,8-	mg/L	2.39E-05	1.96E-05	1.96E-05	1.97E-06	8.40E-06	1.97E-06	2.39E-05	1.20E-05	1.20E-05
~PeCDF, 1,2,3,7,8-	mg/L	7.95E-04	6.53E-04	6.53E-04	6.54E-05	2.80E-04	6.54E-05	7.95E-04	4.01E-04	4.01E-04
~PeCDF, 2,3,4,7,8-	mg/L	7.95E-05	6.53E-05	6.53E-05	6.54E-06	2.80E-05	6.54E-06	7.95E-05	4.01E-05	4.01E-05
~TCDD, 2,3,7,8-	mg/L	2.39E-05	1.96E-05	1.96E-05	1.97E-06	8.40E-06	1.97E-06	2.39E-05	1.20E-05	1.20E-05
~TCDF, 2,3,7,8-	mg/L	2.39E-04	1.96E-04	1.96E-04	1.97E-05	8.40E-05	1.97E-05	2.39E-04	1.20E-04	1.20E-04
Ethylbenzene	mg/L	1.02E+02	1.02E+01	1.02E+01	3.21E+01	1.13E+01	1.13E+01	1.02E+02	8.52E+00	8.52E+00
Fluoranthene	mg/L	1.36E+03	-	1.36E+03	1.12E+02	-	1.12E+02	1.36E+03	-	1.36E+03
Fluorene	mg/L	1.30E+01	-	1.30E+01	4.53E+00	-	4.53E+00	1.30E+01	-	1.30E+01
Hexachlorobenzene	mg/L	2.72E+01	1.59E+00	1.59E+00	2.25E+00	6.82E-01	6.82E-01	2.72E+01	9.78E-01	9.78E-01
Naphthalene	mg/L	1.93E+01	-	1.93E+01	6.18E+00	-	6.18E+00	1.93E+01	-	1.93E+01
Nitroaniline, 2-	mg/L	7.56E+01	-	7.56E+01	1.52E+01	-	1.52E+01	7.56E+01	-	7.56E+01
Nitroso-di-N-propylamine, N-	mg/L	-	1.65E-01	1.65E-01	-	1.09E-01	1.09E-01	-	1.19E-01	1.19E-01
Phenanthrene	mg/L	-	-	-	-	-	-	-	-	-
Polychlorinated Biphenyls, Total ^c	mg/L	-	6.37E+00	6.37E+00	-	2.73E+00	2.73E+00	-	3.91E+00	3.91E+00
~Aroclor 1016	mg/L	2.39E+00	3.64E+01	2.39E+00	1.97E-01	1.56E+01	1.97E-01	2.39E+00	2.24E+01	2.39E+00
~Aroclor 1221	mg/L	-	1.10E-02	1.10E-02	-	1.30E-02	1.30E-02	-	9.36E-03	9.36E-03
~Aroclor 1232	mg/L	-	1.10E-02	1.10E-02	-	1.30E-02	1.30E-02	-	9.36E-03	9.36E-03
~Aroclor 1242	mg/L	-	1.27E+00	1.27E+00	-	5.46E-01	5.46E-01	-	7.82E-01	7.82E-01
~Aroclor 1248	mg/L	-	1.27E+00	1.27E+00	-	5.46E-01	5.46E-01	-	7.82E-01	7.82E-01
~Aroclor 1254	mg/L	6.81E-01	1.27E+00	6.81E-01	5.61E-02	5.46E-01	5.61E-02	6.81E-01	7.82E-01	6.81E-01
~Aroclor 1260	mg/L	-	1.27E+00	1.27E+00	-	5.46E-01	5.46E-01	-	7.82E-01	7.82E-01
Polycyclic aromatic hydrocarbons, Total Carcinogenic ^d	mg/L	-	3.49E-01	3.49E-01	-	2.80E-02	2.80E-02	-	8.04E-02	8.04E-02
~Benz[a]anthracene	mg/L	-	3.49E+00	3.49E+00	-	2.80E-01	2.80E-01	-	8.04E-01	8.04E-01
~Benzo[a]pyrene	mg/L	-	3.49E-01	3.49E-01	-	2.80E-02	2.80E-02	-	8.04E-02	8.04E-02
~Benzo[b]fluoranthene	mg/L	-	3.49E+00	3.49E+00	-	2.80E-01	2.80E-01	-	8.04E-01	8.04E-01
~Benzo[k]fluoranthene	mg/L	-	3.49E+01	3.49E+01	-	2.80E+00	2.80E+00	-	8.04E+00	8.04E+00
~Chrysene	mg/L	-	3.49E+02	3.49E+02	-	2.80E+01	2.80E+01	-	8.04E+01	8.04E+01
~Dibenz[a,h]anthracene	mg/L	-	3.49E-01	3.49E-01	-	2.80E-02	2.80E-02	-	8.04E-02	8.04E-02
~Indeno[1,2,3-cd]pyrene	mg/L	-	3.49E+00	3.49E+00	-	2.80E-01	2.80E-01	-	8.04E-01	8.04E-01

Hazard-based value calculated using target HI of 3.

Cancer-based value calculated using target ELCR of 1E-04.

Action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.3. Surface Water Action Levels for Significant COPCs at PGDP (Continued)
 (Values calculated on 11/6/2013 and are based on best available information.)

Analyte	Units	Adult Recreational User Swimming			Child Recreational User Swimming			Teen Recreational User Swimming		
		Hazard	Cancer	Action	Hazard	Cancer	Action	Hazard	Cancer	Action
Pyrene	mg/L	4.26E+00	-	4.26E+00	1.55E+00	-	1.55E+00	4.26E+00	-	4.26E+00
Tetrachloroethylene	mg/L	6.30E+00	5.76E+01	6.30E+00	2.06E+00	6.35E+01	2.06E+00	6.30E+00	4.81E+01	6.30E+00
Trichloroethylene	mg/L	1.75E+00	7.96E+00	1.75E+00	4.47E-01	1.42E+00	4.47E-01	1.75E+00	2.41E+00	1.75E+00
Vinyl Chloride	mg/L	1.89E+01	-	1.89E+01	3.72E+00	7.00E-03	7.00E-03	1.89E+01	-	1.89E+01
Xylene, m-	mg/L	1.90E+02	-	1.90E+02	6.00E+01	-	6.00E+01	1.90E+02	-	1.90E+02
Xylene, Mixture	mg/L	2.02E+02	-	2.02E+02	6.33E+01	-	6.33E+01	2.02E+02	-	2.02E+02
Xylene, o-	mg/L	2.14E+02	-	2.14E+02	6.63E+01	-	6.63E+01	2.14E+02	-	2.14E+02
Xylene, p-	mg/L	2.05E+02	-	2.05E+02	6.39E+01	-	6.39E+01	2.05E+02	-	2.05E+02
Am-241	pCi/L	-	1.37E+04	1.37E+04	-	2.74E+04	2.74E+04	-	1.37E+04	1.37E+04
Cs-137+D	pCi/L	-	4.69E+04	4.69E+04	-	9.37E+04	9.37E+04	-	4.69E+04	4.69E+04
Np-237+D	pCi/L	-	2.11E+04	2.11E+04	-	4.23E+04	4.23E+04	-	2.11E+04	2.11E+04
Pu-238	pCi/L	-	1.09E+04	1.09E+04	-	2.17E+04	2.17E+04	-	1.09E+04	1.09E+04
Pu-239	pCi/L	-	1.06E+04	1.06E+04	-	2.11E+04	2.11E+04	-	1.06E+04	1.06E+04
Pu-240	pCi/L	-	1.06E+04	1.06E+04	-	2.11E+04	2.11E+04	-	1.06E+04	1.06E+04
Tc-99	pCi/L	-	5.18E+05	5.18E+05	-	1.04E+06	1.04E+06	-	5.18E+05	5.18E+05
Th-230	pCi/L	-	1.57E+04	1.57E+04	-	3.13E+04	3.13E+04	-	1.57E+04	1.57E+04
U-234	pCi/L	-	2.01E+04	2.01E+04	-	4.03E+04	4.03E+04	-	2.01E+04	2.01E+04
U-235+D	pCi/L	-	1.98E+04	1.98E+04	-	3.97E+04	3.97E+04	-	1.98E+04	1.98E+04
U-238+D	pCi/L	-	1.64E+04	1.64E+04	-	3.27E+04	3.27E+04	-	1.64E+04	1.64E+04

Hazard-based value calculated using target HI of 3.

Cancer-based value calculated using target ELCR of 1E-04.

Action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.3. Surface Water Action Levels for Significant COPCs at PGDP (Continued)

(Values calculated on 11/6/2013 and are based on best available information.)

Analyte	Units	Adult Recreational User			Child Recreational User			Teen Recreational User		
		Wading ^e			Wading ^e			Wading ^e		
		Hazard	Cancer	Action	Hazard	Cancer	Action	Hazard	Cancer	Action
Aluminum	mg/L	9.45E+04	-	9.45E+04	1.37E+04	-	1.37E+04	9.45E+04	-	9.45E+04
Antimony (metallic)	mg/L	5.67E+00	-	5.67E+00	8.19E-01	-	8.19E-01	5.67E+00	-	5.67E+00
Arsenic, Inorganic	mg/L	2.84E+01	6.93E+00	6.93E+00	4.11E+00	3.55E+00	3.55E+00	2.84E+01	2.24E+00	2.24E+00
Barium	mg/L	1.33E+03	-	1.33E+03	1.91E+02	-	1.91E+02	1.33E+03	-	1.33E+03
Beryllium and compounds	mg/L	1.33E+00	-	1.33E+00	1.91E-01	-	1.91E-01	1.33E+00	-	1.33E+00
Boron And Borates Only	mg/L	1.89E+04	-	1.89E+04	2.74E+03	-	2.74E+03	1.89E+04	-	1.89E+04
Cadmium (Diet)	mg/L	2.37E+00	-	2.37E+00	3.42E-01	-	3.42E-01	2.37E+00	-	2.37E+00
Chromium (Total) ^a	mg/L	1.85E+03	-	1.85E+03	2.67E+02	-	2.67E+02	1.85E+03	-	1.85E+03
Chromium(III), Insoluble Salts	mg/L	1.85E+03	-	1.85E+03	2.67E+02	-	2.67E+02	1.85E+03	-	1.85E+03
Chromium(VI)	mg/L	3.54E+00	2.60E-01	2.60E-01	5.13E-01	2.49E-02	2.49E-02	3.54E+00	3.14E-02	3.14E-02
Cobalt	mg/L	7.11E+01	-	7.11E+01	1.03E+01	-	1.03E+01	7.11E+01	-	7.11E+01
Copper	mg/L	3.78E+03	-	3.78E+03	5.46E+02	-	5.46E+02	3.78E+03	-	3.78E+03
Iron	mg/L	6.63E+04	-	6.63E+04	9.57E+03	-	9.57E+03	6.63E+04	-	6.63E+04
Lead	mg/L	-	-	3.00E-02	-	-	3.00E-02	-	-	3.00E-02
Manganese	mg/L	9.09E+01	-	9.09E+01	1.31E+01	-	1.31E+01	9.09E+01	-	9.09E+01
Mercury, Inorganic Salts	mg/L	1.99E+00	-	1.99E+00	2.87E-01	-	2.87E-01	1.99E+00	-	1.99E+00
Molybdenum	mg/L	4.74E+02	-	4.74E+02	6.84E+01	-	6.84E+01	4.74E+02	-	4.74E+02
Nickel Soluble Salts	mg/L	3.78E+02	-	3.78E+02	5.46E+01	-	5.46E+01	3.78E+02	-	3.78E+02
Selenium	mg/L	4.74E+02	-	4.74E+02	6.84E+01	-	6.84E+01	4.74E+02	-	4.74E+02
Silver	mg/L	3.15E+01	-	3.15E+01	4.56E+00	-	4.56E+00	3.15E+01	-	3.15E+01
Thallium (Soluble Salts)	mg/L	9.45E-01	-	9.45E-01	1.37E-01	-	1.37E-01	9.45E-01	-	9.45E-01
Uranium (Soluble Salts)	mg/L	2.84E+02	-	2.84E+02	4.11E+01	-	4.11E+01	2.84E+02	-	2.84E+02
Vanadium and Compounds	mg/L	1.24E+01	-	1.24E+01	1.79E+00	-	1.79E+00	1.24E+01	-	1.24E+01
Zinc and Compounds	mg/L	4.74E+04	-	4.74E+04	6.84E+03	-	6.84E+03	4.74E+04	-	4.74E+04
Acenaphthene	mg/L	2.72E+01	-	2.72E+01	6.27E+00	-	6.27E+00	2.72E+01	-	2.72E+01
Acenaphthylene	mg/L	2.72E+01	-	9.08E+01	6.27E+00	-	2.09E+01	2.72E+01	-	9.08E+01
Acrylonitrile	mg/L	2.31E+03	1.43E+01	1.43E+01	4.08E+02	7.33E+00	7.33E+00	2.31E+03	4.62E+00	4.62E+00
Anthracene	mg/L	7.08E+01	-	7.08E+01	1.65E+01	-	1.65E+01	7.08E+01	-	7.08E+01
Benzene	mg/L	1.63E+01	1.07E+01	1.07E+01	3.09E+00	5.48E+00	3.09E+00	1.63E+01	3.45E+00	3.45E+00
Carbazole	mg/L	-	5.82E+00	5.82E+00	-	2.98E+00	2.98E+00	-	1.88E+00	1.88E+00
Carbon Tetrachloride	mg/L	9.60E+00	5.84E+00	5.84E+00	2.15E+00	2.99E+00	2.15E+00	9.60E+00	1.88E+00	1.88E+00
Chloroform	mg/L	7.17E+01	3.61E+01	3.61E+01	1.47E+01	1.85E+01	1.47E+01	7.17E+01	1.16E+01	1.16E+01
Dichloroethylene, 1,1-	mg/L	2.35E+02	-	2.35E+02	4.68E+01	-	4.68E+01	2.35E+02	-	2.35E+02
Dichloroethylene, 1,2- (Mixed Isomers)	mg/L	4.50E+01	-	4.50E+01	8.91E+00	-	8.91E+00	4.50E+01	-	4.50E+01
Dichloroethylene, 1,2-cis-	mg/L	9.99E+00	-	9.99E+00	1.98E+00	-	1.98E+00	9.99E+00	-	9.99E+00
Dichloroethylene, 1,2-trans-	mg/L	9.99E+01	-	9.99E+01	1.98E+01	-	1.98E+01	9.99E+01	-	9.99E+01
Dieldrin	mg/L	1.74E-02	3.85E-03	3.85E-03	4.05E-03	1.97E-03	1.97E-03	1.74E-02	1.24E-03	1.24E-03

Hazard-based value calculated using target HI of 3.

Cancer-based value calculated using target ELCR of 1E-04.

Action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.3. Surface Water Action Levels for Significant COPCs at PGDP (Continued)

(Values calculated on 11/6/2013 and are based on best available information.)

Analyte	Units	Adult Recreational User Wading ^e			Child Recreational User Wading ^e			Teen Recreational User Wading ^e		
		Hazard	Cancer	Action	Hazard	Cancer	Action	Hazard	Cancer	Action
Dioxins/Furans, Total (as TCDD) ^b	mg/L	-	-	-	-	-	-	-	-	-
~HpCDD, 2,3,7,8-	mg/L	-	-	-	-	-	-	-	-	-
~HpCDF, 2,3,7,8-	mg/L	-	-	-	-	-	-	-	-	-
~HxCDD, 2,3,7,8-	mg/L	-	-	-	-	-	-	-	-	-
~HxCDF, 2,3,7,8-	mg/L	-	-	-	-	-	-	-	-	-
~OCDD	mg/L	-	-	-	-	-	-	-	-	-
~OCDF	mg/L	-	-	-	-	-	-	-	-	-
~PeCDD, 2,3,7,8-	mg/L	-	-	-	-	-	-	-	-	-
~PeCDF, 1,2,3,7,8-	mg/L	-	-	-	-	-	-	-	-	-
~PeCDF, 2,3,4,7,8-	mg/L	-	-	-	-	-	-	-	-	-
~TCDD, 2,3,7,8-	mg/L	-	-	-	-	-	-	-	-	-
~TCDF, 2,3,7,8-	mg/L	-	-	-	-	-	-	-	-	-
Ethylbenzene	mg/L	1.05E+02	1.58E+01	1.58E+01	2.28E+01	8.07E+00	8.07E+00	1.05E+02	5.09E+00	5.09E+00
Fluoranthene	mg/L	-	-	-	-	-	-	-	-	-
Fluorene	mg/L	1.31E+01	-	1.31E+01	2.99E+00	-	2.99E+00	1.31E+01	-	1.31E+01
Hexachlorobenzene	mg/L	-	-	-	-	-	-	-	-	-
Naphthalene	mg/L	1.98E+01	-	1.98E+01	4.38E+00	-	4.38E+00	1.98E+01	-	1.98E+01
Nitroaniline, 2-	mg/L	9.72E+01	-	9.72E+01	2.09E+01	-	2.09E+01	9.72E+01	-	9.72E+01
Nitroso-di-N-propylamine, N-	mg/L	-	4.47E-01	4.47E-01	-	2.28E-01	2.28E-01	-	1.44E-01	1.44E-01
Phenanthrene	mg/L	-	-	-	-	-	-	-	-	-
Polychlorinated Biphenyls, Total ^c	mg/L	-	-	-	-	-	-	-	-	-
~Aroclor 1016	mg/L	-	-	-	-	-	-	-	-	-
~Aroclor 1221	mg/L	-	1.65E-02	1.65E-02	-	8.44E-03	8.44E-03	-	5.32E-03	5.32E-03
~Aroclor 1232	mg/L	-	1.65E-02	1.65E-02	-	8.44E-03	8.44E-03	-	5.32E-03	5.32E-03
~Aroclor 1242	mg/L	-	-	-	-	-	-	-	-	-
~Aroclor 1248	mg/L	-	-	-	-	-	-	-	-	-
~Aroclor 1254	mg/L	-	-	-	-	-	-	-	-	-
~Aroclor 1260	mg/L	-	-	-	-	-	-	-	-	-
Polycyclic aromatic hydrocarbons, Total Carcinogenic ^d	mg/L	-	-	-	-	-	-	-	-	-
~Benz[a]anthracene	mg/L	-	-	-	-	-	-	-	-	-
~Benzo[a]pyrene	mg/L	-	-	-	-	-	-	-	-	-
~Benzo[b]fluoranthene	mg/L	-	-	-	-	-	-	-	-	-
~Benzo[k]fluoranthene	mg/L	-	-	-	-	-	-	-	-	-
~Chrysene	mg/L	-	-	-	-	-	-	-	-	-
~Dibenz[a,h]anthracene	mg/L	-	-	-	-	-	-	-	-	-
~Indeno[1,2,3-cd]pyrene	mg/L	-	-	-	-	-	-	-	-	-

Hazard-based value calculated using target HI of 3.

Cancer-based value calculated using target ELCR of 1E-04.

Action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.3. Surface Water Action Levels for Significant COPCs at PGDP (Continued)

(Values calculated on 11/6/2013 and are based on best available information.)

Analyte	Units	Adult Recreational User Wading ^e			Child Recreational User Wading ^e			Teen Recreational User Wading ^e		
		Hazard	Cancer	Action	Hazard	Cancer	Action	Hazard	Cancer	Action
Pyrene	mg/L	4.29E+00	-	4.29E+00	9.96E-01	-	9.96E-01	4.29E+00	-	4.29E+00
Tetrachloroethylene	mg/L	6.51E+00	8.96E+01	6.51E+00	1.49E+00	4.58E+01	1.49E+00	6.51E+00	2.89E+01	6.51E+00
Trichloroethylene	mg/L	1.95E+00	1.38E+01	1.95E+00	4.17E-01	1.32E+00	4.17E-01	1.95E+00	1.67E+00	1.67E+00
Vinyl Chloride	mg/L	2.33E+01	-	2.33E+01	4.23E+00	2.42E-02	2.42E-02	2.33E+01	-	2.33E+01
Xylene, m-	mg/L	1.95E+02	-	1.95E+02	4.26E+01	-	4.26E+01	1.95E+02	-	1.95E+02
Xylene, Mixture	mg/L	2.08E+02	-	2.08E+02	4.50E+01	-	4.50E+01	2.08E+02	-	2.08E+02
Xylene, o-	mg/L	2.21E+02	-	2.21E+02	4.77E+01	-	4.77E+01	2.21E+02	-	2.21E+02
Xylene, p-	mg/L	2.11E+02	-	2.11E+02	4.56E+01	-	4.56E+01	2.11E+02	-	2.11E+02
Am-241	pCi/L	-	-	-	-	-	-	-	-	-
Cs-137+D	pCi/L	-	-	-	-	-	-	-	-	-
Np-237+D	pCi/L	-	-	-	-	-	-	-	-	-
Pu-238	pCi/L	-	-	-	-	-	-	-	-	-
Pu-239	pCi/L	-	-	-	-	-	-	-	-	-
Pu-240	pCi/L	-	-	-	-	-	-	-	-	-
Tc-99	pCi/L	-	-	-	-	-	-	-	-	-
Th-230	pCi/L	-	-	-	-	-	-	-	-	-
U-234	pCi/L	-	-	-	-	-	-	-	-	-
U-235+D	pCi/L	-	-	-	-	-	-	-	-	-
U-238+D	pCi/L	-	-	-	-	-	-	-	-	-

NOTE: The action level for HI is 3 because the range of values for HI (based on RGO tables) are 0.1, 1, and 3. Please see Figure 1.1 of the Risk Methods Document.

^a Chromium (Total) utilizes Chromium III hazard values and Chromium VI inhalation risk values (to be conservative in terms of protecting human health).

^b Total dioxins/furans uses values for 2,3,7,8-TCDD, see screening note 9g in the Appendix A introduction, 'Screening Levels' on pages A-3 through A-5.

^c Total PCBs ALs are based on the high risk toxicity values.

^d Total carcinogenic PAHs uses values for benzo(a)pyrene, see screening note 9e in the Appendix A introduction, 'Screening Levels' on pages A-3 through A-5.

^e Wading scenario considers dermal contact only.

Hazard-based value calculated using target HI of 3.

Cancer-based value calculated using target ELCR of 1E-04.

Action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.4. Soil/Sediment No Action Levels for Significant COPCs at PGDP

(Values calculated on 11/5/2013 and are based on best available information.)

CAS	Analyte	Units	Outdoor Worker			Industrial Worker			Adult Recreational User		
			Hazard	Cancer	No Action	Hazard	Cancer	No Action	Hazard	Cancer	No Action
7429905	Aluminum	mg/kg	2.86E+04	-	2.86E+04	1.00E+05	-	1.00E+05	1.00E+05	-	1.00E+05
7440360	Antimony (metallic)	mg/kg	1.15E+01	-	1.15E+01	8.18E+01	-	8.18E+01	9.83E+01	-	9.83E+01
7440382	Arsenic, Inorganic	mg/kg	6.67E+00	4.15E-01	4.15E-01	6.09E+01	3.81E+00	3.81E+00	2.72E+01	3.53E+00	3.53E+00
7440393	Barium	mg/kg	5.67E+03	-	5.67E+03	3.59E+04	-	3.59E+04	4.91E+04	-	4.91E+04
7440417	Beryllium and compounds	mg/kg	5.73E+01	9.39E+03	5.73E+01	3.95E+02	6.95E+03	3.95E+02	4.91E+02	5.57E+04	4.91E+02
7440428	Boron And Borates Only	mg/kg	5.75E+03	-	5.75E+03	4.07E+04	-	4.07E+04	4.91E+04	-	4.91E+04
7440439	Cadmium (Diet)	mg/kg	2.06E+01	1.25E+04	2.06E+01	1.98E+02	9.26E+03	1.98E+02	7.49E+01	7.42E+04	7.49E+01
16065831	Chromium (Total) ^a	mg/kg	4.32E+04	2.68E+02	2.68E+02	1.00E+05	1.98E+02	1.98E+02	1.00E+05	1.59E+03	1.59E+03
18540299	Chromium(III), Insoluble Salts	mg/kg	4.32E+04	-	4.32E+04	1.00E+05	-	1.00E+05	1.00E+05	-	1.00E+05
7440473	Chromium(VI)	mg/kg	8.62E+01	1.60E+00	1.60E+00	6.07E+02	1.08E+01	1.08E+01	7.37E+02	2.82E+01	2.82E+01
7440484	Cobalt	mg/kg	8.62E+00	2.50E+03	8.62E+00	6.03E+01	1.85E+03	6.03E+01	7.37E+01	1.48E+04	7.37E+01
7440508	Copper	mg/kg	1.15E+03	-	1.15E+03	8.18E+03	-	8.18E+03	9.83E+03	-	9.83E+03
7439896	Iron	mg/kg	2.01E+04	-	2.01E+04	1.00E+05	-	1.00E+05	1.00E+05	-	1.00E+05
7439921	Lead	mg/kg	-	-	8.00E+02	-	-	8.00E+02	-	-	4.00E+02
7439965	Manganese	mg/kg	6.79E+02	-	6.79E+02	4.21E+03	-	4.21E+03	5.90E+03	-	5.90E+03
7439976	Mercury, Inorganic Salts	mg/kg	8.63E+00	-	8.63E+00	6.13E+01	-	6.13E+01	7.37E+01	-	7.37E+01
7439987	Molybdenum	mg/kg	1.44E+02	-	1.44E+02	1.02E+03	-	1.02E+03	1.23E+03	-	1.23E+03
7440020	Nickel Soluble Salts	mg/kg	5.71E+02	8.66E+04	5.71E+02	3.80E+03	6.41E+04	3.80E+03	4.91E+03	1.00E+05	4.91E+03
7782492	Selenium	mg/kg	1.44E+02	-	1.44E+02	1.02E+03	-	1.02E+03	1.23E+03	-	1.23E+03
7440224	Silver	mg/kg	1.44E+02	-	1.44E+02	1.02E+03	-	1.02E+03	1.23E+03	-	1.23E+03
7791120	Thallium (Soluble Salts)	mg/kg	2.88E-01	-	2.88E-01	2.04E+00	-	2.04E+00	2.46E+00	-	2.46E+00
238	Uranium (Soluble Salts)	mg/kg	8.61E+01	-	8.61E+01	5.98E+02	-	5.98E+02	7.37E+02	-	7.37E+02
7440622	Vanadium and Compounds	mg/kg	1.45E+02	-	1.45E+02	1.01E+03	-	1.01E+03	1.24E+03	-	1.24E+03
7440666	Zinc and Compounds	mg/kg	8.63E+03	-	8.63E+03	6.13E+04	-	6.13E+04	7.37E+04	-	7.37E+04
83329	Acenaphthene	mg/kg	7.60E+02	-	7.60E+02	1.23E+04	-	1.23E+04	1.75E+03	-	1.75E+03
208968	Acenaphthylene	mg/kg	-	-	-	-	-	-	-	-	-
107131	Acrylonitrile	mg/kg	9.71E+00	8.57E-01	8.57E-01	7.24E+00	1.31E+00	1.31E+00	9.83E+03	8.24E+00	8.24E+00
120127	Anthracene	mg/kg	3.80E+03	-	3.80E+03	6.13E+04	-	6.13E+04	8.76E+03	-	8.76E+03
71432	Benzene	mg/kg	4.26E+01	5.21E+00	5.21E+00	4.71E+01	5.66E+00	5.66E+00	9.83E+02	4.05E+01	4.05E+01
86748	Carbazole	mg/kg	-	2.04E+01	2.04E+01	-	2.86E+02	2.86E+02	-	1.07E+02	1.07E+02
56235	Carbon Tetrachloride	mg/kg	5.21E+01	3.21E+00	3.21E+00	6.49E+01	3.16E+00	3.16E+00	9.83E+02	2.33E+01	2.33E+01
67663	Chloroform	mg/kg	1.04E+02	1.89E+00	1.89E+00	1.14E+02	1.50E+00	1.50E+00	2.46E+03	1.18E+01	1.18E+01
75354	Dichloroethylene, 1,1-	mg/kg	1.34E+02	-	1.34E+02	1.08E+02	-	1.08E+02	1.23E+04	-	1.23E+04
540590	Dichloroethylene, 1,2- (Mixed Isomers)	mg/kg	2.59E+02	-	2.59E+02	1.84E+03	-	1.84E+03	2.21E+03	-	2.21E+03
156592	Dichloroethylene, 1,2-cis-	mg/kg	5.75E+01	-	5.75E+01	4.09E+02	-	4.09E+02	4.91E+02	-	4.91E+02
156605	Dichloroethylene, 1,2-trans-	mg/kg	8.23E+01	-	8.23E+01	6.98E+01	-	6.98E+01	4.91E+03	-	4.91E+03
60571	Dieldrin	mg/kg	7.27E-01	2.54E-02	2.54E-02	1.02E+01	3.58E-01	3.58E-01	1.83E+00	1.34E-01	1.34E-01

Hazard-based value calculated using target HI of 0.1.

Cancer-based value calculated using target ELCR of 1E-06.

No action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.4. Soil/Sediment No Action Levels for Significant COPCs at PGDP (Continued)

(Values calculated on 11/5/2013 and are based on best available information.)

CAS	Analyte	Units	Outdoor Worker			Industrial Worker			Adult Recreational User		
			Hazard	Cancer	No Action	Hazard	Cancer	No Action	Hazard	Cancer	No Action
1746016	Dioxins/Furans, Total (as TCDD) ^b	mg/kg	1.56E-05	4.79E-06	4.79E-06	1.43E-04	4.40E-05	4.40E-05	6.35E-05	4.07E-05	4.07E-05
37871004	~HpCDD, 2,3,7,8-	mg/kg	1.56E-03	4.79E-04	4.79E-04	1.43E-02	4.40E-03	4.40E-03	6.35E-03	4.07E-03	4.07E-03
38998753	~HpCDF, 2,3,7,8-	mg/kg	1.02E-03	3.13E-04	3.13E-04	1.43E-02	4.40E-03	4.40E-03	2.57E-03	1.65E-03	1.65E-03
34465468	~HxCDD, 2,3,7,8-	mg/kg	1.56E-04	4.79E-05	4.79E-05	1.43E-03	4.40E-04	4.40E-04	6.35E-04	4.07E-04	4.07E-04
55684941	~HxCDF, 2,3,7,8-	mg/kg	1.02E-04	3.13E-05	3.13E-05	1.43E-03	4.40E-04	4.40E-04	2.57E-04	1.65E-04	1.65E-04
3268879	~OCDD	mg/kg	5.19E-02	1.60E-02	1.60E-02	4.77E-01	1.47E-01	1.47E-01	2.12E-01	1.36E-01	1.36E-01
39001020	~OCDF	mg/kg	3.39E-02	1.04E-02	1.04E-02	4.77E-01	1.47E-01	1.47E-01	8.56E-02	5.48E-02	5.48E-02
36088229	~PeCDD, 2,3,7,8-	mg/kg	1.56E-05	4.79E-06	4.79E-06	1.43E-04	4.40E-05	4.40E-05	6.35E-05	4.07E-05	4.07E-05
57117416	~PeCDF, 1,2,3,7,8-	mg/kg	3.39E-04	1.04E-04	1.04E-04	4.77E-03	1.47E-03	1.47E-03	8.56E-04	5.48E-04	5.48E-04
57117314	~PeCDF, 2,3,4,7,8-	mg/kg	3.39E-05	1.04E-05	1.04E-05	4.77E-04	1.47E-04	1.47E-04	8.56E-05	5.48E-05	5.48E-05
1746016	~TCDD, 2,3,7,8-	mg/kg	1.56E-05	4.79E-06	4.79E-06	1.43E-04	4.40E-05	4.40E-05	6.35E-05	4.07E-05	4.07E-05
51207319	~TCDF, 2,3,7,8-	mg/kg	1.02E-04	3.13E-05	3.13E-05	1.43E-03	4.40E-04	4.40E-04	2.57E-04	1.65E-04	1.65E-04
100414	Ethylbenzene	mg/kg	1.60E+03	2.61E+01	2.61E+01	2.36E+03	2.83E+01	2.83E+01	2.46E+04	2.02E+02	2.02E+02
206440	Fluoranthene	mg/kg	5.06E+02	-	5.06E+02	8.18E+03	-	8.18E+03	1.17E+03	-	1.17E+03
86737	Fluorene	mg/kg	5.06E+02	-	5.06E+02	8.18E+03	-	8.18E+03	1.17E+03	-	1.17E+03
118741	Hexachlorobenzene	mg/kg	1.16E+01	2.54E-01	2.54E-01	1.64E+02	3.58E+00	3.58E+00	2.93E+01	1.34E+00	1.34E+00
91203	Naphthalene	mg/kg	6.56E+01	2.43E+01	2.43E+01	6.45E+01	1.80E+01	1.80E+01	5.84E+02	1.44E+02	1.44E+02
88744	Nitroaniline, 2-	mg/kg	1.45E+02	-	1.45E+02	1.91E+03	-	1.91E+03	3.67E+02	-	3.67E+02
621647	Nitroso-di-N-propylamine, N-	mg/kg	-	5.82E-02	5.82E-02	-	8.18E-01	8.18E-01	-	3.06E-01	3.06E-01
85018	Phenanthrene	mg/kg	-	-	-	-	-	-	-	-	-
1336363	Polychlorinated Biphenyls, Total ^c	mg/kg	-	1.70E-01	1.70E-01	-	2.86E+00	2.86E+00	-	7.98E-01	7.98E-01
12674112	~Aroclor 1016	mg/kg	8.50E-01	4.85E+00	8.50E-01	1.43E+01	8.18E+01	1.43E+01	1.92E+00	2.28E+01	1.92E+00
11104282	~Aroclor 1221	mg/kg	-	1.60E-01	1.60E-01	-	1.17E+00	1.17E+00	-	7.59E-01	7.59E-01
11141165	~Aroclor 1232	mg/kg	-	1.60E-01	1.60E-01	-	1.17E+00	1.17E+00	-	7.59E-01	7.59E-01
53469219	~Aroclor 1242	mg/kg	-	1.70E-01	1.70E-01	-	2.86E+00	2.86E+00	-	7.98E-01	7.98E-01
12672296	~Aroclor 1248	mg/kg	-	1.70E-01	1.70E-01	-	2.86E+00	2.86E+00	-	7.98E-01	7.98E-01
11097691	~Aroclor 1254	mg/kg	2.43E-01	1.70E-01	1.70E-01	4.09E+00	2.86E+00	2.86E+00	5.47E-01	7.98E-01	5.47E-01
11096825	~Aroclor 1260	mg/kg	-	1.70E-01	1.70E-01	-	2.86E+00	2.86E+00	-	7.98E-01	7.98E-01
50328	Polycyclic aromatic hydrocarbons, Total Carcinogenic ^d	mg/kg	-	4.86E-02	4.86E-02	-	7.84E-01	7.84E-01	-	2.33E-01	2.33E-01
56553	~Benz[a]anthracene	mg/kg	-	4.86E-01	4.86E-01	-	7.84E+00	7.84E+00	-	2.33E+00	2.33E+00
50328	~Benzo[a]pyrene	mg/kg	-	4.86E-02	4.86E-02	-	7.84E-01	7.84E-01	-	2.33E-01	2.33E-01
205992	~Benzo[b]fluoranthene	mg/kg	-	4.86E-01	4.86E-01	-	7.84E+00	7.84E+00	-	2.33E+00	2.33E+00
207089	~Benzo[k]fluoranthene	mg/kg	-	4.86E+00	4.86E+00	-	7.84E+01	7.84E+01	-	2.33E+01	2.33E+01
218019	~Chrysene	mg/kg	-	4.86E+01	4.86E+01	-	7.84E+02	7.84E+02	-	2.33E+02	2.33E+02
53703	~Dibenz[a,h]anthracene	mg/kg	-	4.86E-02	4.86E-02	-	7.84E-01	7.84E-01	-	2.33E-01	2.33E-01
193395	~Indeno[1,2,3-cd]pyrene	mg/kg	-	4.86E-01	4.86E-01	-	7.84E+00	7.84E+00	-	2.33E+00	2.33E+00

Hazard-based value calculated using target HI of 0.1.

Cancer-based value calculated using target ELCR of 1E-06.

No action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.4. Soil/Sediment No Action Levels for Significant COPCs at PGDP (Continued)

(Values calculated on 11/5/2013 and are based on best available information.)

CAS	Analyte	Units	Outdoor Worker			Industrial Worker			Adult Recreational User		
			Hazard	Cancer	No Action	Hazard	Cancer	No Action	Hazard	Cancer	No Action
129000	Pyrene	mg/kg	3.80E+02	-	3.80E+02	6.13E+03	-	6.13E+03	8.76E+02	-	8.76E+02
127184	Tetrachloroethylene	mg/kg	4.44E+01	1.13E+02	4.44E+01	4.27E+01	1.14E+02	4.27E+01	1.47E+03	8.38E+02	8.38E+02
79016	Trichloroethylene	mg/kg	2.35E+00	6.21E+00	2.35E+00	2.04E+00	6.73E+00	2.04E+00	1.23E+02	4.82E+01	4.82E+01
75014	Vinyl Chloride	mg/kg	3.57E+01	8.68E-01	8.68E-01	4.20E+01	2.11E+00	2.11E+00	7.37E+02	2.31E-01	2.31E-01
1330207	Xylene, m-	mg/kg	3.29E+02	-	3.29E+02	2.56E+02	-	2.56E+02	4.91E+04	-	4.91E+04
106423	Xylene, Mixture	mg/kg	3.48E+02	-	3.48E+02	2.73E+02	-	2.73E+02	4.91E+04	-	4.91E+04
108383	Xylene, o-	mg/kg	3.84E+02	-	3.84E+02	3.02E+02	-	3.02E+02	4.91E+04	-	4.91E+04
95476	Xylene, p-	mg/kg	3.35E+02	-	3.35E+02	2.62E+02	-	2.62E+02	4.91E+04	-	4.91E+04
14596102	Am-241	pCi/g	-	3.33E+00	3.33E+00	-	1.79E+01	1.79E+01	-	2.28E+01	2.28E+01
10045973	Cs-137+D	pCi/g	-	1.37E-01	1.37E-01	-	5.08E-01	5.08E-01	-	8.48E-01	8.48E-01
13994202	Np-237+D	pCi/g	-	3.22E-01	3.22E-01	-	1.21E+00	1.21E+00	-	1.88E+00	1.88E+00
13981163	Pu-238	pCi/g	-	4.23E+00	4.23E+00	-	2.97E+01	2.97E+01	-	3.30E+01	3.30E+01
15117483	Pu-239	pCi/g	-	3.70E+00	3.70E+00	-	2.60E+01	2.60E+01	-	2.89E+01	2.89E+01
14119336	Pu-240	pCi/g	-	3.71E+00	3.71E+00	-	2.62E+01	2.62E+01	-	2.89E+01	2.89E+01
14133767	Tc-99	pCi/g	-	3.09E+02	3.09E+02	-	2.02E+03	2.02E+03	-	9.92E+02	9.92E+02
14269637	Th-230	pCi/g	-	5.70E+00	5.70E+00	-	3.95E+01	3.95E+01	-	3.88E+01	3.88E+01
13966295	U-234	pCi/g	-	8.72E+00	8.72E+00	-	6.11E+01	6.11E+01	-	5.02E+01	5.02E+01
15117961	U-235+D	pCi/g	-	4.84E-01	4.84E-01	-	1.84E+00	1.84E+00	-	2.83E+00	2.83E+00
7440611	U-238+D	pCi/g	-	1.81E+00	1.81E+00	-	7.48E+00	7.48E+00	-	1.01E+01	1.01E+01

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Hazard-based value calculated using target HI of 0.1.

Cancer-based value calculated using target ELCR of 1E-06.

No action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.4. Soil/Sediment No Action Levels for Significant COPCs at PGDP (Continued)

(Values calculated on 11/5/2013 and are based on best available information.)

Analyte	Child Recreational User			Teen Recreational User			Resident	Adult Resident		Child Resident	
	Hazard	Cancer	No Action	Hazard	Cancer	No Action	Cancer ^e	Hazard	No Action	Hazard	No Action
Aluminum	1.95E+04	-	1.95E+04	1.00E+05	-	1.00E+05	-	6.62E+04	6.62E+04	7.74E+03	7.74E+03
Antimony (metallic)	7.82E+00	-	7.82E+00	4.48E+01	-	4.48E+01	-	2.92E+01	2.92E+01	3.13E+00	3.13E+00
Arsenic, Inorganic	4.13E+00	1.07E+00	1.07E+00	1.03E+01	1.34E+00	1.34E+00	2.36E-01	8.05E+00	2.36E-01	1.65E+00	2.36E-01
Barium	3.89E+03	-	3.89E+03	2.24E+04	-	2.24E+04	-	1.21E+04	1.21E+04	1.53E+03	1.53E+03
Beryllium and compounds	3.91E+01	8.27E+04	3.91E+01	2.24E+02	4.13E+04	2.24E+02	1.38E+03	1.39E+02	1.39E+02	1.56E+01	1.56E+01
Boron And Borates Only	3.91E+03	-	3.91E+03	2.24E+04	-	2.24E+04	-	1.45E+04	1.45E+04	1.56E+03	1.56E+03
Cadmium (Diet)	1.25E+01	1.00E+05	1.25E+01	2.80E+01	5.51E+04	2.80E+01	1.84E+03	2.19E+01	2.19E+01	5.00E+00	5.00E+00
Chromium (Total) ^a	2.93E+04	4.43E+02	4.43E+02	1.00E+05	4.43E+02	4.43E+02	1.55E+01	1.00E+05	1.55E+01	1.17E+04	1.55E+01
Chromium(III), Insoluble Salts	2.93E+04	-	2.93E+04	1.00E+05	-	1.00E+05	-	1.00E+05	1.00E+05	1.17E+04	1.17E+04
Chromium(VI)	5.86E+01	8.54E-01	8.54E-01	3.36E+02	4.85E+00	4.85E+00	2.93E-01	2.16E+02	2.93E-01	2.34E+01	2.93E-01
Cobalt	5.86E+00	2.21E+04	5.86E+00	3.36E+01	1.10E+04	3.36E+01	3.68E+02	2.14E+01	2.14E+01	2.34E+00	2.34E+00
Copper	7.82E+02	-	7.82E+02	4.48E+03	-	4.48E+03	-	2.92E+03	2.92E+03	3.13E+02	3.13E+02
Iron	1.37E+04	-	1.37E+04	7.85E+04	-	7.85E+04	-	5.11E+04	5.11E+04	5.48E+03	5.48E+03
Lead	-	-	4.00E+02	-	-	4.00E+02	-	-	4.00E+02	-	4.00E+02
Manganese	4.67E+02	-	4.67E+02	2.69E+03	-	2.69E+03	-	1.40E+03	1.40E+03	1.83E+02	1.83E+02
Mercury, Inorganic Salts	5.87E+00	-	5.87E+00	3.36E+01	-	3.36E+01	-	2.19E+01	2.19E+01	2.35E+00	2.35E+00
Molybdenum	9.78E+01	-	9.78E+01	5.61E+02	-	5.61E+02	-	3.65E+02	3.65E+02	3.91E+01	3.91E+01
Nickel Soluble Salts	3.90E+02	1.00E+05	3.90E+02	2.24E+03	1.00E+05	2.24E+03	1.27E+04	1.31E+03	1.31E+03	1.55E+02	1.55E+02
Selenium	9.78E+01	-	9.78E+01	5.61E+02	-	5.61E+02	-	3.65E+02	3.65E+02	3.91E+01	3.91E+01
Silver	9.78E+01	-	9.78E+01	5.61E+02	-	5.61E+02	-	3.65E+02	3.65E+02	3.91E+01	3.91E+01
Thallium (Soluble Salts)	1.96E-01	-	1.96E-01	1.12E+00	-	1.12E+00	-	7.30E-01	7.30E-01	7.82E-02	7.82E-02
Uranium (Soluble Salts)	5.86E+01	-	5.86E+01	3.36E+02	-	3.36E+02	-	2.11E+02	2.11E+02	2.34E+01	2.34E+01
Vanadium and Compounds	9.85E+01	-	9.85E+01	5.65E+02	-	5.65E+02	-	3.59E+02	3.59E+02	3.93E+01	3.93E+01
Zinc and Compounds	5.87E+03	-	5.87E+03	3.36E+04	-	3.36E+04	-	2.19E+04	2.19E+04	2.35E+03	2.35E+03
Acenaphthene	4.16E+02	-	4.16E+02	6.26E+02	-	6.26E+02	-	5.21E+02	5.21E+02	1.66E+02	1.66E+02
Acenaphthylene	-	-	-	-	-	-	-	-	-	-	-
Acrylonitrile	2.02E+01	3.41E+00	3.41E+00	4.48E+03	5.12E+00	5.12E+00	2.37E-01	1.72E+00	2.37E-01	1.72E+00	2.37E-01
Anthracene	2.08E+03	-	2.08E+03	3.13E+03	-	3.13E+03	-	2.60E+03	2.60E+03	8.32E+02	8.32E+02
Benzene	5.05E+01	2.62E+01	2.62E+01	4.48E+02	2.74E+01	2.74E+01	1.08E+00	1.14E+01	1.08E+00	8.62E+00	1.08E+00
Carbazole	-	4.75E+01	4.75E+01	-	3.85E+01	3.85E+01	8.66E+00	-	8.66E+00	-	8.66E+00
Carbon Tetrachloride	5.63E+01	1.78E+01	1.78E+01	4.48E+02	1.62E+01	1.62E+01	6.09E-01	1.59E+01	6.09E-01	1.09E+01	6.09E-01
Chloroform	1.25E+02	1.44E+01	1.44E+01	1.12E+03	8.61E+00	8.61E+00	2.95E-01	2.77E+01	2.95E-01	2.11E+01	2.95E-01
Dichloroethylene, 1,1-	2.36E+02	-	2.36E+02	5.61E+03	-	5.61E+03	-	2.58E+01	2.58E+01	2.43E+01	2.43E+01
Dichloroethylene, 1,2- (Mixed Isomers)	1.76E+02	-	1.76E+02	1.01E+03	-	1.01E+03	-	6.57E+02	6.57E+02	7.04E+01	7.04E+01
Dichloroethylene, 1,2-cis-	3.91E+01	-	3.91E+01	2.24E+02	-	2.24E+02	-	1.46E+02	1.46E+02	1.56E+01	1.56E+01
Dichloroethylene, 1,2-trans-	1.34E+02	-	1.34E+02	2.24E+03	-	2.24E+03	-	1.67E+01	1.67E+01	1.53E+01	1.53E+01
Dieldrin	4.07E-01	5.94E-02	5.94E-02	6.59E-01	4.81E-02	4.81E-02	1.08E-02	5.45E-01	1.08E-02	1.63E-01	1.08E-02

Hazard-based value calculated using target HI of 0.1.

Cancer-based value calculated using target ELCR of 1E-06.

No action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.4. Soil/Sediment No Action Levels for Significant COPCs at PGDP (Continued)

(Values calculated on 11/5/2013 and are based on best available information.)

Analyte	Child Recreational User			Teen Recreational User			Resident	Adult Resident		Child Resident	
	Hazard	Cancer	No Action	Hazard	Cancer	No Action	Cancer ^e	Hazard	No Action	Hazard	No Action
Dioxins/Furans, Total (as TCDD) ^b	9.64E-06	1.24E-05	9.64E-06	2.41E-05	1.55E-05	1.55E-05	2.72E-06	1.89E-05	2.72E-06	3.86E-06	2.72E-06
~HpCDD, 2,3,7,8-	9.64E-04	1.24E-03	9.64E-04	2.41E-03	1.55E-03	1.55E-03	2.72E-04	1.89E-03	2.72E-04	3.86E-04	2.72E-04
~HpCDF, 2,3,7,8-	5.70E-04	7.31E-04	5.70E-04	9.23E-04	5.92E-04	5.92E-04	1.33E-04	7.63E-04	1.33E-04	2.28E-04	1.33E-04
~HxCDD, 2,3,7,8-	9.64E-05	1.24E-04	9.64E-05	2.41E-04	1.55E-04	1.55E-04	2.72E-05	1.89E-04	2.72E-05	3.86E-05	2.72E-05
~HxCDF, 2,3,7,8-	5.70E-05	7.31E-05	5.70E-05	9.23E-05	5.92E-05	5.92E-05	1.33E-05	7.63E-05	1.33E-05	2.28E-05	1.33E-05
~OCDD	3.21E-02	4.12E-02	3.21E-02	8.05E-02	5.16E-02	5.16E-02	9.06E-03	6.29E-02	9.06E-03	1.29E-02	9.06E-03
~OCDF	1.90E-02	2.44E-02	1.90E-02	3.08E-02	1.97E-02	1.97E-02	4.44E-03	2.54E-02	4.44E-03	7.60E-03	4.44E-03
~PeCDD, 2,3,7,8-	9.64E-06	1.24E-05	9.64E-06	2.41E-05	1.55E-05	1.55E-05	2.72E-06	1.89E-05	2.72E-06	3.86E-06	2.72E-06
~PeCDF, 1,2,3,7,8-	1.90E-04	2.44E-04	1.90E-04	3.08E-04	1.97E-04	1.97E-04	4.44E-05	2.54E-04	4.44E-05	7.60E-05	4.44E-05
~PeCDF, 2,3,4,7,8-	1.90E-05	2.44E-05	1.90E-05	3.08E-05	1.97E-05	1.97E-05	4.44E-06	2.54E-05	4.44E-06	7.60E-06	4.44E-06
~TCDD, 2,3,7,8-	9.64E-06	1.24E-05	9.64E-06	2.41E-05	1.55E-05	1.55E-05	2.72E-06	1.89E-05	2.72E-06	3.86E-06	2.72E-06
~TCDF, 2,3,7,8-	5.70E-05	7.31E-05	5.70E-05	9.23E-05	5.92E-05	5.92E-05	1.33E-05	7.63E-05	1.33E-05	2.28E-05	1.33E-05
Ethylbenzene	1.56E+03	1.31E+02	1.31E+02	1.12E+04	1.37E+02	1.37E+02	5.39E+00	5.85E+02	5.39E+00	3.51E+02	5.39E+00
Fluoranthene	2.77E+02	-	2.77E+02	4.17E+02	-	4.17E+02	-	3.47E+02	3.47E+02	1.11E+02	1.11E+02
Fluorene	2.77E+02	-	2.77E+02	4.17E+02	-	4.17E+02	-	3.47E+02	3.47E+02	1.11E+02	1.11E+02
Hexachlorobenzene	6.52E+00	5.94E-01	5.94E-01	1.06E+01	4.81E-01	4.81E-01	1.08E-01	8.72E+00	1.08E-01	2.61E+00	1.08E-01
Naphthalene	7.97E+01	2.14E+02	7.97E+01	2.09E+02	1.07E+02	1.07E+02	3.57E+00	1.43E+01	3.57E+00	1.22E+01	3.57E+00
Nitroaniline, 2-	8.14E+01	-	8.14E+01	1.32E+02	-	1.32E+02	-	1.07E+02	1.07E+02	3.24E+01	3.24E+01
Nitroso-di-N-propylamine, N-	-	1.36E-01	1.36E-01	-	1.10E-01	1.10E-01	2.47E-02	-	2.47E-02	-	2.47E-02
Phenanthrene	-	-	-	-	-	-	-	-	-	-	-
Polychlorinated Biphenyls, Total ^c	-	3.85E-01	3.85E-01	-	2.84E-01	2.84E-01	6.70E-02	-	6.70E-02	-	6.70E-02
~Aroclor 1016	4.62E-01	1.10E+01	4.62E-01	6.82E-01	8.12E+00	6.82E-01	1.91E+00	5.69E-01	5.69E-01	1.85E-01	1.85E-01
~Aroclor 1221	-	3.79E-01	3.79E-01	-	2.78E-01	2.78E-01	5.72E-02	-	5.72E-02	-	5.72E-02
~Aroclor 1232	-	3.79E-01	3.79E-01	-	2.78E-01	2.78E-01	5.72E-02	-	5.72E-02	-	5.72E-02
~Aroclor 1242	-	3.85E-01	3.85E-01	-	2.84E-01	2.84E-01	6.70E-02	-	6.70E-02	-	6.70E-02
~Aroclor 1248	-	3.85E-01	3.85E-01	-	2.84E-01	2.84E-01	6.70E-02	-	6.70E-02	-	6.70E-02
~Aroclor 1254	1.32E-01	3.85E-01	1.32E-01	1.95E-01	2.84E-01	1.95E-01	6.70E-02	1.63E-01	6.70E-02	5.28E-02	5.28E-02
~Aroclor 1260	-	3.85E-01	3.85E-01	-	2.84E-01	2.84E-01	6.70E-02	-	6.70E-02	-	6.70E-02
Polycyclic aromatic hydrocarbons, Total Carcinogenic ^d	-	2.08E-02	2.08E-02	-	3.12E-02	3.12E-02	5.77E-03	-	5.77E-03	-	5.77E-03
~Benz[a]anthracene	-	2.08E-01	2.08E-01	-	3.12E-01	3.12E-01	5.77E-02	-	5.77E-02	-	5.77E-02
~Benzo[a]pyrene	-	2.08E-02	2.08E-02	-	3.12E-02	3.12E-02	5.77E-03	-	5.77E-03	-	5.77E-03
~Benzo[b]fluoranthene	-	2.08E-01	2.08E-01	-	3.12E-01	3.12E-01	5.77E-02	-	5.77E-02	-	5.77E-02
~Benzo[k]fluoranthene	-	2.08E+00	2.08E+00	-	3.12E+00	3.12E+00	5.77E-01	-	5.77E-01	-	5.77E-01
~Chrysene	-	2.08E+01	2.08E+01	-	3.12E+01	3.12E+01	5.77E+00	-	5.77E+00	-	5.77E+00
~Dibenz[a,h]anthracene	-	2.08E-02	2.08E-02	-	3.12E-02	3.12E-02	5.77E-03	-	5.77E-03	-	5.77E-03
~Indeno[1,2,3-cd]pyrene	-	2.08E-01	2.08E-01	-	3.12E-01	3.12E-01	5.77E-02	-	5.77E-02	-	5.77E-02

Hazard-based value calculated using target HI of 0.1.

Cancer-based value calculated using target ELCR of 1E-06.

No action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.4. Soil/Sediment No Action Levels for Significant COPCs at PGDP (Continued)

(Values calculated on 11/5/2013 and are based on best available information.)

Analyte	Child Recreational User			Teen Recreational User			Resident	Adult Resident		Child Resident	
	Hazard	Cancer	No Action	Hazard	Cancer	No Action	Cancer ^e	Hazard	No Action	Hazard	No Action
Pyrene	2.08E+02	-	2.08E+02	3.13E+02	-	3.13E+02	-	2.60E+02	2.60E+02	8.32E+01	8.32E+01
Tetrachloroethylene	6.09E+01	6.15E+02	6.09E+01	6.73E+02	5.78E+02	5.78E+02	2.19E+01	1.03E+01	1.03E+01	8.61E+00	8.61E+00
Trichloroethylene	3.70E+00	5.86E+00	3.70E+00	5.61E+01	1.22E+01	1.22E+01	4.75E-01	4.89E-01	4.75E-01	4.40E-01	4.40E-01
Vinyl Chloride	4.03E+01	7.03E-02	7.03E-02	3.36E+02	2.31E-01	2.31E-01	5.97E-02	1.02E+01	5.97E-02	7.36E+00	5.97E-02
Xylene, m-	6.20E+02	-	6.20E+02	2.24E+04	-	2.24E+04	-	6.12E+01	6.12E+01	5.91E+01	5.91E+01
Xylene, Mixture	6.53E+02	-	6.53E+02	2.24E+04	-	2.24E+04	-	6.51E+01	6.51E+01	6.27E+01	6.27E+01
Xylene, o-	7.11E+02	-	7.11E+02	2.24E+04	-	2.24E+04	-	7.21E+01	7.21E+01	6.93E+01	6.93E+01
Xylene, p-	6.31E+02	-	6.31E+02	2.24E+04	-	2.24E+04	-	6.24E+01	6.24E+01	6.03E+01	6.03E+01
Am-241	-	2.12E+01	2.12E+01	-	1.70E+01	1.70E+01	2.41E+00	-	2.41E+00	-	2.41E+00
Cs-137+D	-	1.26E+00	1.26E+00	-	6.30E-01	6.30E-01	1.00E-01	-	1.00E-01	-	1.00E-01
Np-237+D	-	2.70E+00	2.70E+00	-	1.40E+00	1.40E+00	2.21E-01	-	2.21E-01	-	2.21E-01
Pu-238	-	2.45E+01	2.45E+01	-	2.45E+01	2.45E+01	3.26E+00	-	3.26E+00	-	3.26E+00
Pu-239	-	2.15E+01	2.15E+01	-	2.14E+01	2.14E+01	2.85E+00	-	2.85E+00	-	2.85E+00
Pu-240	-	2.15E+01	2.15E+01	-	2.14E+01	2.14E+01	2.85E+00	-	2.85E+00	-	2.85E+00
Tc-99	-	7.56E+02	7.56E+02	-	7.37E+02	7.37E+02	9.91E+01	-	9.91E+01	-	9.91E+01
Th-230	-	2.91E+01	2.91E+01	-	2.88E+01	2.88E+01	3.84E+00	-	3.84E+00	-	3.84E+00
U-234	-	3.75E+01	3.75E+01	-	3.73E+01	3.73E+01	4.97E+00	-	4.97E+00	-	4.97E+00
U-235+D	-	3.98E+00	3.98E+00	-	2.10E+00	2.10E+00	3.31E-01	-	3.31E-01	-	3.31E-01
U-238+D	-	1.18E+01	1.18E+01	-	7.48E+00	7.48E+00	1.13E+00	-	1.13E+00	-	1.13E+00

^a Chromium (Total) utilizes Chromium III hazard values and Chromium VI inhalation risk values (to be conservative in terms of protecting human health).

^b Total dioxins/furans uses values for 2,3,7,8-TCDD, see screening note 9g in the Appendix A introduction, 'Screening Levels' on pages A-3 through A-5.

^c Total PCBs NALs are based on the high risk toxicity values.

^d Total carcinogenic PAHs uses values for benzo(a)pyrene, see screening note 9e in the Appendix A introduction, 'Screening Levels' on pages A-3 through A-5.

^e For the resident, ELCRs (i.e. cancer NALs) were calculated using the child/adult age-adjusted lifetime scenario.

Hazard-based value calculated using target HI of 0.1.

Cancer-based value calculated using target ELCR of 1E-06.

No action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.5. Groundwater No Action Levels for Significant COPCs at PGDP

(Values calculated on 11/5/2013 and are based on best available information.)

CAS	Analyte	Units	Resident	Adult Resident		Child Resident	
			Cancer ^e	Hazard	No Action	Hazard	No Action
7429905	Aluminum	mg/L	-	3.63E+00	3.63E+00	1.04E+00	1.04E+00
7440360	Antimony (metallic)	mg/L	-	1.41E-03	1.41E-03	4.15E-04	4.15E-04
7440382	Arsenic, Inorganic	mg/L	3.78E-05	1.09E-03	3.78E-05	3.13E-04	3.78E-05
7440393	Barium	mg/L	-	6.79E-01	6.79E-01	2.06E-01	2.06E-01
7440417	Beryllium and compounds	mg/L	-	4.18E-03	4.18E-03	1.86E-03	1.86E-03
7440428	Boron And Borates Only	mg/L	-	7.26E-01	7.26E-01	2.08E-01	2.08E-01
7440439	Cadmium (Water)	mg/L	-	1.65E-03	1.65E-03	5.13E-04	5.13E-04
16065831	Chromium (Total) ^a	mg/L	-	3.91E+00	3.91E+00	1.47E+00	1.47E+00
18540299	Chromium(III), Insoluble Salts	mg/L	-	3.91E+00	3.91E+00	1.47E+00	1.47E+00
7440473	Chromium(VI)	mg/L	3.00E-05	7.72E-03	3.00E-05	2.93E-03	3.00E-05
7440484	Cobalt	mg/L	-	1.09E-03	1.09E-03	3.13E-04	3.13E-04
7440508	Copper	mg/L	-	1.45E-01	1.45E-01	4.17E-02	4.17E-02
7439896	Iron	mg/L	-	2.54E+00	2.54E+00	7.29E-01	7.29E-01
7439921	Lead	mg/L	-	-	1.50E-02	-	1.50E-02
7439965	Manganese	mg/L	-	7.75E-02	7.75E-02	2.45E-02	2.45E-02
7439976	Mercury, Inorganic Salts	mg/L	-	1.02E-03	1.02E-03	3.09E-04	3.09E-04
7439987	Molybdenum	mg/L	-	1.82E-02	1.82E-02	5.21E-03	5.21E-03
7440020	Nickel Soluble Salts	mg/L	-	7.11E-02	7.11E-02	2.08E-02	2.08E-02
7782492	Selenium	mg/L	-	1.82E-02	1.82E-02	5.21E-03	5.21E-03
7440224	Silver	mg/L	-	1.69E-02	1.69E-02	5.15E-03	5.15E-03
7791120	Thallium (Soluble Salts)	mg/L	-	3.63E-05	3.63E-05	1.04E-05	1.04E-05
238	Uranium (Soluble Salts)	mg/L	-	1.09E-02	1.09E-02	3.13E-03	3.13E-03
7440622	Vanadium and Compounds	mg/L	-	1.53E-02	1.53E-02	5.09E-03	5.09E-03
7440666	Zinc and Compounds	mg/L	-	1.09E+00	1.09E+00	3.13E-01	3.13E-01
83329	Acenaphthene	mg/L	-	9.02E-02	9.02E-02	4.46E-02	4.46E-02
208968	Acenaphthylene	mg/L	-	-	-	-	-
107131	Acrylonitrile	mg/L	4.25E-05	4.16E-04	4.25E-05	4.13E-04	4.25E-05
120127	Anthracene	mg/L	-	2.92E-01	2.92E-01	1.76E-01	1.76E-01
71432	Benzene	mg/L	3.79E-04	4.19E-03	3.79E-04	2.44E-03	3.79E-04
86748	Carbazole	mg/L	1.98E-03	-	1.98E-03	-	1.98E-03
56235	Carbon Tetrachloride	mg/L	3.82E-04	7.41E-03	3.82E-04	3.27E-03	3.82E-04
67663	Chloroform	mg/L	1.89E-04	1.27E-02	1.89E-04	6.78E-03	1.89E-04
75354	Dichloroethylene, 1,1-	mg/L	-	3.31E-02	3.31E-02	2.28E-02	2.28E-02
540590	Dichloroethylene, 1,2- (Mixed Isomers)	mg/L	-	2.92E-02	2.92E-02	9.06E-03	9.06E-03
156592	Dichloroethylene, 1,2-cis-	mg/L	-	6.48E-03	6.48E-03	2.01E-03	2.01E-03
156605	Dichloroethylene, 1,2-trans-	mg/L	-	1.05E-02	1.05E-02	7.72E-03	7.72E-03
60571	Dieldrin	mg/L	1.93E-06	6.36E-05	1.93E-06	3.41E-05	1.93E-06

Hazard-based value calculated using target HI of 0.1.

Cancer-based value calculated using target ELCR of 1E-06.

No action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.5. Groundwater No Action Levels for Significant COPCs at PGDP (Continued)

(Values calculated on 11/5/2013 and are based on best available information.)

CAS	Analyte	Units	Resident	Adult Resident		Child Resident	
			Cancer ^e	Hazard	No Action	Hazard	No Action
1746016	Dioxins/Furans, Total (as TCDD) ^b	mg/L	4.37E-10	2.56E-09	4.37E-10	7.30E-10	4.37E-10
37871004	~HpCDD, 2,3,7,8-	mg/L	4.37E-08	2.56E-07	4.37E-08	7.30E-08	4.37E-08
38998753	~HpCDF, 2,3,7,8-	mg/L	4.37E-08	2.56E-07	4.37E-08	7.30E-08	4.37E-08
34465468	~HxCDD, 2,3,7,8-	mg/L	4.37E-09	2.56E-08	4.37E-09	7.30E-09	4.37E-09
55684941	~HxCDF, 2,3,7,8-	mg/L	4.37E-09	2.56E-08	4.37E-09	7.30E-09	4.37E-09
3268879	~OCDD	mg/L	1.46E-06	8.52E-06	1.46E-06	2.43E-06	1.46E-06
39001020	~OCDF	mg/L	1.46E-06	8.52E-06	1.46E-06	2.43E-06	1.46E-06
36088229	~PeCDD, 2,3,7,8-	mg/L	4.37E-10	2.56E-09	4.37E-10	7.30E-10	4.37E-10
57117416	~PeCDF, 1,2,3,7,8-	mg/L	1.46E-08	8.52E-08	1.46E-08	2.43E-08	1.46E-08
57117314	~PeCDF, 2,3,4,7,8-	mg/L	1.46E-09	8.52E-09	1.46E-09	2.43E-09	1.46E-09
1746016	~TCDD, 2,3,7,8-	mg/L	4.37E-10	2.56E-09	4.37E-10	7.30E-10	4.37E-10
51207319	~TCDF, 2,3,7,8-	mg/L	4.37E-09	2.56E-08	4.37E-09	7.30E-09	4.37E-09
100414	Ethylbenzene	mg/L	1.32E-03	1.09E-01	1.32E-03	6.25E-02	1.32E-03
206440	Fluoranthene	mg/L	-	1.46E-01	1.46E-01	4.17E-02	4.17E-02
86737	Fluorene	mg/L	-	4.91E-02	4.91E-02	2.68E-02	2.68E-02
118741	Hexachlorobenzene	mg/L	3.55E-05	2.92E-03	3.55E-05	8.34E-04	3.55E-05
91203	Naphthalene	mg/L	1.43E-04	6.17E-04	1.43E-04	6.04E-04	1.43E-04
88744	Nitroaniline, 2-	mg/L	-	3.42E-02	3.42E-02	1.02E-02	1.02E-02
621647	Nitroso-di-N-propylamine, N-	mg/L	7.99E-06	-	7.99E-06	-	7.99E-06
85018	Phenanthrene	mg/L	-	-	-	-	-
1336363	Polychlorinated Biphenyls, Total ^c	mg/L	1.42E-04	-	1.42E-04	-	1.42E-04
12674112	~Aroclor 1016	mg/L	8.11E-04	2.56E-04	2.56E-04	7.30E-05	7.30E-05
11104282	~Aroclor 1221	mg/L	4.82E-06	-	4.82E-06	-	4.82E-06
11141165	~Aroclor 1232	mg/L	4.82E-06	-	4.82E-06	-	4.82E-06
53469219	~Aroclor 1242	mg/L	2.84E-05	-	2.84E-05	-	2.84E-05
12672296	~Aroclor 1248	mg/L	2.84E-05	-	2.84E-05	-	2.84E-05
11097691	~Aroclor 1254	mg/L	2.84E-05	7.30E-05	2.84E-05	2.09E-05	2.09E-05
11096825	~Aroclor 1260	mg/L	2.84E-05	-	2.84E-05	-	2.84E-05
50328	Polycyclic aromatic hydrocarbons, Total Carcinogenic ^d	mg/L	2.24E-06	-	2.24E-06	-	2.24E-06
56553	~Benz[a]anthracene	mg/L	2.24E-05	-	2.24E-05	-	2.24E-05
50328	~Benzo[a]pyrene	mg/L	2.24E-06	-	2.24E-06	-	2.24E-06
205992	~Benzo[b]fluoranthene	mg/L	2.24E-05	-	2.24E-05	-	2.24E-05
207089	~Benzo[k]fluoranthene	mg/L	2.24E-04	-	2.24E-04	-	2.24E-04
218019	~Chrysene	mg/L	2.24E-03	-	2.24E-03	-	2.24E-03
53703	~Dibenz[a,h]anthracene	mg/L	2.24E-06	-	2.24E-06	-	2.24E-06
193395	~Indeno[1,2,3-cd]pyrene	mg/L	2.24E-05	-	2.24E-05	-	2.24E-05

Hazard-based value calculated using target HI of 0.1.

Cancer-based value calculated using target ELCR of 1E-06.

No action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.5. Groundwater No Action Levels for Significant COPCs at PGDP (Continued)

(Values calculated on 11/5/2013 and are based on best available information.)

CAS	Analyte	Units	Resident	Adult Resident		Child Resident	
			Cancer ^e	Hazard	No Action	Hazard	No Action
129000	Pyrene	mg/L	-	1.97E-02	1.97E-02	1.37E-02	1.37E-02
127184	Tetrachloroethylene	mg/L	9.97E-03	5.19E-03	5.19E-03	3.26E-03	3.26E-03
79016	Trichloroethylene	mg/L	1.95E-04	3.29E-04	1.95E-04	2.27E-04	1.95E-04
75014	Vinyl Chloride	mg/L	1.11E-05	6.83E-03	1.11E-05	2.67E-03	1.11E-05
1330207	Xylene, m-	mg/L	-	1.99E-02	1.99E-02	1.87E-02	1.87E-02
106423	Xylene, Mixture	mg/L	-	1.99E-02	1.99E-02	1.87E-02	1.87E-02
108383	Xylene, o-	mg/L	-	2.00E-02	2.00E-02	1.87E-02	1.87E-02
95476	Xylene, p-	mg/L	-	1.99E-02	1.99E-02	1.87E-02	1.87E-02
14596102	Am-241	pCi/L	4.82E-01	-	4.82E-01	-	4.82E-01
10045973	Cs-137+D	pCi/L	1.65E+00	-	1.65E+00	-	1.65E+00
13994202	Np-237+D	pCi/L	7.44E-01	-	7.44E-01	-	7.44E-01
13981163	Pu-238	pCi/L	3.83E-01	-	3.83E-01	-	3.83E-01
15117483	Pu-239	pCi/L	3.71E-01	-	3.71E-01	-	3.71E-01
14119336	Pu-240	pCi/L	3.71E-01	-	3.71E-01	-	3.71E-01
14133767	Tc-99	pCi/L	1.82E+01	-	1.82E+01	-	1.82E+01
14269637	Th-230	pCi/L	5.51E-01	-	5.51E-01	-	5.51E-01
13966295	U-234	pCi/L	7.09E-01	-	7.09E-01	-	7.09E-01
15117961	U-235+D	pCi/L	6.98E-01	-	6.98E-01	-	6.98E-01
7440611	U-238+D	pCi/L	5.75E-01	-	5.75E-01	-	5.75E-01

^a Chromium (Total) utilizes Chromium III hazard values and Chromium VI inhalation risk values (to be conservative in terms of protecting human health).

^b Total dioxins/furans uses values for 2,3,7,8-TCDD, see screening note 9g in the Appendix A introduction, 'Screening Levels' on pages A-3 through A-5.

^c Total PCBs NALs are based on the high risk toxicity values.

^d Total carcinogenic PAHs uses values for benzo(a)pyrene, see screening note 9e in the Appendix A introduction, 'Screening Levels' on pages A-3 through A-5.

^e For the resident, ELCRs (i.e. cancer NALs) were calculated using the child/adult age-adjusted lifetime scenario (i.e., lifetime exposure)

Hazard-based value calculated using target HI of 0.1.

Cancer-based value calculated using target ELCR of 1E-06.

No action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.6. Surface Water No Action Levels for Significant COPCs at PGDP
(Values calculated on 11/6/2013 and are based on best available information.)

CAS	Analyte	Units	Outdoor Worker Wading ^c			Industrial Worker Wading ^c		
			Hazard	Cancer	No Action	Hazard	Cancer	No Action
7429905	Aluminum	mg/L	3.40E+03	-	3.40E+03	8.36E+02	-	8.36E+02
7440360	Antimony (metallic)	mg/L	2.04E-01	-	2.04E-01	5.02E-02	-	5.02E-02
7440382	Arsenic, Inorganic	mg/L	1.02E+00	6.34E-02	6.34E-02	2.51E-01	1.56E-02	1.56E-02
7440393	Barium	mg/L	4.76E+01	-	4.76E+01	1.17E+01	-	1.17E+01
7440417	Beryllium and compounds	mg/L	4.76E-02	-	4.76E-02	1.17E-02	-	1.17E-02
7440428	Boron And Borates Only	mg/L	6.80E+02	-	6.80E+02	1.67E+02	-	1.67E+02
7440439	Cadmium (Water)	mg/L	8.49E-02	-	8.49E-02	2.09E-02	-	2.09E-02
16065831	Chromium (Total) ^a	mg/L	6.63E+01	-	6.63E+01	1.63E+01	-	1.63E+01
18540299	Chromium(III), Insoluble Salts	mg/L	6.63E+01	-	6.63E+01	1.63E+01	-	1.63E+01
7440473	Chromium(VI)	mg/L	1.27E-01	1.27E-03	1.27E-03	3.14E-02	3.11E-04	3.11E-04
7440484	Cobalt	mg/L	2.55E+00	-	2.55E+00	6.27E-01	-	6.27E-01
7440508	Copper	mg/L	1.36E+02	-	1.36E+02	3.35E+01	-	3.35E+01
7439896	Iron	mg/L	2.38E+03	-	2.38E+03	5.85E+02	-	5.85E+02
7439921	Lead	mg/L	-	-	1.50E-02	-	-	1.50E-02
7439965	Manganese	mg/L	3.26E+00	-	3.26E+00	8.03E-01	-	8.03E-01
7439976	Mercury, Inorganic Salts	mg/L	7.13E-02	-	7.13E-02	1.76E-02	-	1.76E-02
7439987	Molybdenum	mg/L	1.70E+01	-	1.70E+01	4.18E+00	-	4.18E+00
7440020	Nickel Soluble Salts	mg/L	1.36E+01	-	1.36E+01	3.35E+00	-	3.35E+00
7782492	Selenium	mg/L	1.70E+01	-	1.70E+01	4.18E+00	-	4.18E+00
7440224	Silver	mg/L	1.13E+00	-	1.13E+00	2.79E-01	-	2.79E-01
7791120	Thallium (Soluble Salts)	mg/L	3.40E-02	-	3.40E-02	8.36E-03	-	8.36E-03
238	Uranium (Soluble Salts)	mg/L	1.02E+01	-	1.02E+01	2.51E+00	-	2.51E+00
7440622	Vanadium and Compounds	mg/L	4.45E-01	-	4.45E-01	1.10E-01	-	1.10E-01
7440666	Zinc and Compounds	mg/L	1.70E+03	-	1.70E+03	4.18E+02	-	4.18E+02
83329	Acenaphthene	mg/L	9.79E-01	-	9.79E-01	3.83E-01	-	3.83E-01
208968	Acenaphthylene ^f	mg/L	9.79E-01	-	9.79E-01	3.83E-01	-	3.83E-01
107131	Acrylonitrile	mg/L	8.28E+01	1.07E-01	1.07E-01	2.49E+01	3.23E-02	3.23E-02
120127	Anthracene	mg/L	2.54E+00	-	2.54E+00	1.01E+00	-	1.01E+00
71432	Benzene	mg/L	5.86E-01	7.46E-02	7.46E-02	1.90E-01	2.41E-02	2.41E-02
86748	Carbazole	mg/L	-	3.37E-02	3.37E-02	-	1.31E-02	1.31E-02
56235	Carbon Tetrachloride	mg/L	3.45E-01	3.45E-02	3.45E-02	1.31E-01	1.31E-02	1.31E-02
67663	Chloroform	mg/L	2.57E+00	2.32E-01	2.32E-01	9.00E-01	8.13E-02	8.13E-02
75354	Dichloroethylene, 1,1-	mg/L	8.42E+00	-	8.42E+00	2.85E+00	-	2.85E+00
540590	Dichloroethylene, 1,2- (Mixed Isomers)	mg/L	1.61E+00	-	1.61E+00	5.46E-01	-	5.46E-01
156592	Dichloroethylene, 1,2-cis-	mg/L	3.58E-01	-	3.58E-01	1.21E-01	-	1.21E-01
156605	Dichloroethylene, 1,2-trans-	mg/L	3.58E+00	-	3.58E+00	1.21E+00	-	1.21E+00
60571	Diieldrin	mg/L	6.23E-04	2.18E-05	2.18E-05	2.47E-04	8.66E-06	8.66E-06

Hazard-based value calculated using target HI of 0.1.

Cancer-based value calculated using target ELCR of 1E-06.

No action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.6. Surface Water No Action Levels for Significant COPCs at PGDP (Continued)
(Values calculated on 11/6/2013 and are based on best available information.)

CAS	Analyte	Units	Outdoor Worker Wading ^c			Industrial Worker Wading ^c		
			Hazard	Cancer	No Action	Hazard	Cancer	No Action
1746016	Dioxins/Furans, Total (as TCDD) ^b	mg/L	-	-	-	-	-	-
37871004	~HpCDD, 2,3,7,8-	mg/L	-	-	-	-	-	-
38998753	~HpCDF, 2,3,7,8-	mg/L	-	-	-	-	-	-
34465468	~HxCDD, 2,3,7,8-	mg/L	-	-	-	-	-	-
55684941	~HxCDF, 2,3,7,8-	mg/L	-	-	-	-	-	-
3268879	~OCDD	mg/L	-	-	-	-	-	-
39001020	~OCDF	mg/L	-	-	-	-	-	-
36088229	~PeCDD, 2,3,7,8-	mg/L	-	-	-	-	-	-
57117416	~PeCDF, 1,2,3,7,8-	mg/L	-	-	-	-	-	-
57117314	~PeCDF, 2,3,4,7,8-	mg/L	-	-	-	-	-	-
1746016	~TCDD, 2,3,7,8-	mg/L	-	-	-	-	-	-
51207319	~TCDF, 2,3,7,8-	mg/L	-	-	-	-	-	-
100414	Ethylbenzene	mg/L	3.79E+00	9.64E-02	9.64E-02	1.40E+00	3.55E-02	3.55E-02
206440	Fluoranthene	mg/L	-	-	-	-	-	-
86737	Fluorene	mg/L	4.72E-01	-	4.72E-01	1.83E-01	-	1.83E-01
118741	Hexachlorobenzene	mg/L	-	-	-	-	-	-
91203	Naphthalene	mg/L	7.12E-01	-	7.12E-01	2.68E-01	-	2.68E-01
88744	Nitroaniline, 2-	mg/L	3.49E+00	-	3.49E+00	1.28E+00	-	1.28E+00
621647	Nitroso-di-N-propylamine, N-	mg/L	-	2.81E-03	2.81E-03	-	1.01E-03	1.01E-03
85018	Phenanthrene ^f	mg/L	9.79E-01	-	9.79E-01	-	-	-
1336363	Polychlorinated Biphenyls, Total ^c	mg/L	-	-	-	-	-	-
12674112	~Aroclor 1016	mg/L	-	-	-	-	-	-
11104282	~Aroclor 1221	mg/L	-	9.36E-05	9.36E-05	-	3.72E-05	3.72E-05
11141165	~Aroclor 1232	mg/L	-	9.36E-05	9.36E-05	-	3.72E-05	3.72E-05
53469219	~Aroclor 1242	mg/L	-	-	-	-	-	-
12672296	~Aroclor 1248	mg/L	-	-	-	-	-	-
11097691	~Aroclor 1254	mg/L	-	-	-	-	-	-
11096825	~Aroclor 1260	mg/L	-	-	-	-	-	-
50328	Polycyclic aromatic hydrocarbons, Total Carcinogenic ^d	mg/L	-	-	-	-	-	-
56553	~Benz[a]anthracene	mg/L	-	-	-	-	-	-
50328	~Benzo[a]pyrene	mg/L	-	-	-	-	-	-
205992	~Benzo[b]fluoranthene	mg/L	-	-	-	-	-	-
207089	~Benzo[k]fluoranthene	mg/L	-	-	-	-	-	-
218019	~Chrysene	mg/L	-	-	-	-	-	-
53703	~Dibenz[a,h]anthracene	mg/L	-	-	-	-	-	-
193395	~Indeno[1,2,3-cd]pyrene	mg/L	-	-	-	-	-	-

Hazard-based value calculated using target HI of 0.1.

Cancer-based value calculated using target ELCR of 1E-06.

No action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.6. Surface Water No Action Levels for Significant COPCs at PGDP (Continued)
 (Values calculated on 11/6/2013 and are based on best available information.)

CAS	Analyte	Units	Outdoor Worker Wading ^c			Industrial Worker Wading ^c		
			Hazard	Cancer	No Action	Hazard	Cancer	No Action
129000	Pyrene	mg/L	1.54E-01	-	1.54E-01	6.10E-02	-	6.10E-02
127184	Tetrachloroethylene	mg/L	2.34E-01	5.20E-01	2.34E-01	9.07E-02	2.02E-01	9.07E-02
79016	Trichloroethylene	mg/L	7.00E-02	4.54E-02	4.54E-02	2.55E-02	1.65E-02	1.65E-02
75014	Vinyl Chloride	mg/L	8.34E-01	2.08E-04	2.08E-04	2.58E-01	6.57E-04	6.57E-04
1330207	Xylene, m-	mg/L	7.01E+00	-	7.01E+00	2.60E+00	-	2.60E+00
106423	Xylene, Mixture	mg/L	7.46E+00	-	7.46E+00	2.76E+00	-	2.76E+00
108383	Xylene, o-	mg/L	7.93E+00	-	7.93E+00	2.91E+00	-	2.91E+00
95476	Xylene, p-	mg/L	7.57E+00	-	7.57E+00	2.79E+00	-	2.79E+00
14596102	Am-241	pCi/L	-	-	-	-	-	-
10045973	Cs-137+D	pCi/L	-	-	-	-	-	-
13994202	Np-237+D	pCi/L	-	-	-	-	-	-
13981163	Pu-238	pCi/L	-	-	-	-	-	-
15117483	Pu-239	pCi/L	-	-	-	-	-	-
14119336	Pu-240	pCi/L	-	-	-	-	-	-
14133767	Tc-99	pCi/L	-	-	-	-	-	-
14269637	Th-230	pCi/L	-	-	-	-	-	-
13966295	U-234	pCi/L	-	-	-	-	-	-
15117961	U-235+D	pCi/L	-	-	-	-	-	-
7440611	U-238+D	pCi/L	-	-	-	-	-	-

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Hazard-based value calculated using target HI of 0.1.

Cancer-based value calculated using target ELCR of 1E-06.

No action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.6. Surface Water No Action Levels for Significant COPCs at PGDP (Continued)

(Values calculated on 11/6/2013 and are based on best available information.)

Analyte	Units	Adult Recreational User Swimming			Child Recreational User Swimming			Teen Recreational User Swimming		
		Hazard	Cancer	No Action	Hazard	Cancer	No Action	Hazard	Cancer	No Action
Aluminum	mg/L	8.35E+02	-	8.35E+02	8.28E+01	-	8.28E+01	8.35E+02	-	8.35E+02
Antimony (metallic)	mg/L	1.34E-01	-	1.34E-01	2.01E-02	-	2.01E-02	1.34E-01	-	1.34E-01
Arsenic, Inorganic	mg/L	2.50E-01	1.25E-02	1.25E-02	2.48E-02	6.44E-03	6.44E-03	2.50E-01	8.27E-03	8.27E-03
Barium	mg/L	3.70E+01	-	3.70E+01	6.55E+00	-	6.55E+00	3.70E+01	-	3.70E+01
Beryllium and compounds	mg/L	4.33E-02	-	4.33E-02	9.56E-03	-	9.56E-03	4.33E-02	-	4.33E-02
Boron And Borates Only	mg/L	1.67E+02	-	1.67E+02	1.66E+01	-	1.66E+01	1.67E+02	-	1.67E+02
Cadmium (Diet)	mg/L	6.92E-02	-	6.92E-02	1.30E-02	-	1.30E-02	6.92E-02	-	6.92E-02
Chromium (Total) ^a	mg/L	5.94E+01	-	5.94E+01	1.28E+01	-	1.28E+01	5.94E+01	-	5.94E+01
Chromium(III), Insoluble Salts	mg/L	5.94E+01	-	5.94E+01	1.28E+01	-	1.28E+01	5.94E+01	-	5.94E+01
Chromium(VI)	mg/L	1.14E-01	1.70E-03	1.70E-03	2.46E-02	3.59E-04	3.59E-04	1.14E-01	5.35E-04	5.35E-04
Cobalt	mg/L	2.98E-01	-	2.98E-01	2.67E-02	-	2.67E-02	2.98E-01	-	2.98E-01
Copper	mg/L	3.34E+01	-	3.34E+01	3.31E+00	-	3.31E+00	3.34E+01	-	3.34E+01
Iron	mg/L	5.84E+02	-	5.84E+02	5.80E+01	-	5.80E+01	5.84E+02	-	5.84E+02
Lead	mg/L	-	-	1.50E-02	-	-	1.50E-02	-	-	1.50E-02
Manganese	mg/L	2.73E+00	-	2.73E+00	5.29E-01	-	5.29E-01	2.73E+00	-	2.73E+00
Mercury, Inorganic Salts	mg/L	5.55E-02	-	5.55E-02	9.83E-03	-	9.83E-03	5.55E-02	-	5.55E-02
Molybdenum	mg/L	4.17E+00	-	4.17E+00	4.14E-01	-	4.14E-01	4.17E+00	-	4.17E+00
Nickel Soluble Salts	mg/L	8.11E+00	-	8.11E+00	1.13E+00	-	1.13E+00	8.11E+00	-	8.11E+00
Selenium	mg/L	4.17E+00	-	4.17E+00	4.14E-01	-	4.14E-01	4.17E+00	-	4.17E+00
Silver	mg/L	8.87E-01	-	8.87E-01	1.59E-01	-	1.59E-01	8.87E-01	-	8.87E-01
Thallium (Soluble Salts)	mg/L	8.35E-03	-	8.35E-03	8.28E-04	-	8.28E-04	8.35E-03	-	8.35E-03
Uranium (Soluble Salts)	mg/L	2.50E+00	-	2.50E+00	2.48E-01	-	2.48E-01	2.50E+00	-	2.50E+00
Vanadium and Compounds	mg/L	3.86E-01	-	3.86E-01	7.86E-02	-	7.86E-02	3.86E-01	-	3.86E-01
Zinc and Compounds	mg/L	2.80E+02	-	2.80E+02	2.60E+01	-	2.60E+01	2.80E+02	-	2.80E+02
Acenaphthene	mg/L	8.97E-01	-	8.97E-01	3.12E-01	-	3.12E-01	8.97E-01	-	8.97E-01
Acenaphthylene ^f	mg/L	8.97E-01	-	8.97E-01	3.12E-01	-	3.12E-01	8.97E-01	-	8.97E-01
Acrylonitrile	mg/L	2.86E+01	3.17E-02	3.17E-02	3.19E+00	1.72E-02	1.72E-02	2.86E+01	2.14E-02	2.14E-02
Anthracene	mg/L	2.34E+00	-	2.34E+00	8.41E-01	-	8.41E-01	2.34E+00	-	2.34E+00
Benzene	mg/L	4.86E-01	6.25E-02	6.25E-02	1.14E-01	6.03E-02	6.03E-02	4.86E-01	5.06E-02	5.06E-02
Carbazole	mg/L	-	3.81E-02	3.81E-02	-	4.33E-02	4.33E-02	-	3.21E-02	3.21E-02
Carbon Tetrachloride	mg/L	2.99E-01	3.56E-02	3.56E-02	8.69E-02	3.62E-02	3.62E-02	2.99E-01	2.92E-02	2.92E-02
Chloroform	mg/L	1.97E+00	1.88E-01	1.88E-01	4.24E-01	1.60E-01	1.60E-01	1.97E+00	1.47E-01	1.47E-01
Dichloroethylene, 1,1-	mg/L	6.87E+00	-	6.87E+00	1.61E+00	-	1.61E+00	6.87E+00	-	6.87E+00
Dichloroethylene, 1,2- (Mixed Isomers)	mg/L	1.31E+00	-	1.31E+00	3.02E-01	-	3.02E-01	1.31E+00	-	1.31E+00
Dichloroethylene, 1,2-cis-	mg/L	2.90E-01	-	2.90E-01	6.70E-02	-	6.70E-02	2.90E-01	-	2.90E-01
Dichloroethylene, 1,2-trans-	mg/L	2.90E+00	-	2.90E+00	6.70E-01	-	6.70E-01	2.90E+00	-	2.90E+00
Dieldrin	mg/L	5.73E-04	2.55E-05	2.55E-05	2.04E-04	2.97E-05	2.97E-05	5.73E-04	2.16E-05	2.16E-05

Hazard-based value calculated using target HI of 0.1.

Cancer-based value calculated using target ELCR of 1E-06.

No action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.6. Surface Water No Action Levels for Significant COPCs at PGDP (Continued)

(Values calculated on 11/6/2013 and are based on best available information.)

Analyte	Units	Adult Recreational User Swimming			Child Recreational User Swimming			Teen Recreational User Swimming		
		Hazard	Cancer	No Action	Hazard	Cancer	No Action	Hazard	Cancer	No Action
Dioxins/Furans, Total (as TCDD) ^b	mg/L	7.95E-07	1.96E-07	1.96E-07	6.55E-08	8.40E-08	6.55E-08	7.95E-07	1.20E-07	1.20E-07
~HpCDD, 2,3,7,8-	mg/L	7.95E-05	1.96E-05	1.96E-05	6.55E-06	8.40E-06	6.55E-06	7.95E-05	1.20E-05	1.20E-05
~HpCDF, 2,3,7,8-	mg/L	7.95E-05	1.96E-05	1.96E-05	6.55E-06	8.40E-06	6.55E-06	7.95E-05	1.20E-05	1.20E-05
~HxCDD, 2,3,7,8-	mg/L	7.95E-06	1.96E-06	1.96E-06	6.55E-07	8.40E-07	6.55E-07	7.95E-06	1.20E-06	1.20E-06
~HxCDF, 2,3,7,8-	mg/L	7.95E-06	1.96E-06	1.96E-06	6.55E-07	8.40E-07	6.55E-07	7.95E-06	1.20E-06	1.20E-06
~OCDD	mg/L	2.65E-03	6.53E-04	6.53E-04	2.18E-04	2.80E-04	2.18E-04	2.65E-03	4.01E-04	4.01E-04
~OCDF	mg/L	2.65E-03	6.53E-04	6.53E-04	2.18E-04	2.80E-04	2.18E-04	2.65E-03	4.01E-04	4.01E-04
~PeCDD, 2,3,7,8-	mg/L	7.95E-07	1.96E-07	1.96E-07	6.55E-08	8.40E-08	6.55E-08	7.95E-07	1.20E-07	1.20E-07
~PeCDF, 1,2,3,7,8-	mg/L	2.65E-05	6.53E-06	6.53E-06	2.18E-06	2.80E-06	2.18E-06	2.65E-05	4.01E-06	4.01E-06
~PeCDF, 2,3,4,7,8-	mg/L	2.65E-06	6.53E-07	6.53E-07	2.18E-07	2.80E-07	2.18E-07	2.65E-06	4.01E-07	4.01E-07
~TCDD, 2,3,7,8-	mg/L	7.95E-07	1.96E-07	1.96E-07	6.55E-08	8.40E-08	6.55E-08	7.95E-07	1.20E-07	1.20E-07
~TCDF, 2,3,7,8-	mg/L	7.95E-06	1.96E-06	1.96E-06	6.55E-07	8.40E-07	6.55E-07	7.95E-06	1.20E-06	1.20E-06
Ethylbenzene	mg/L	3.41E+00	1.02E-01	1.02E-01	1.07E+00	1.13E-01	1.13E-01	3.41E+00	8.52E-02	8.52E-02
Fluoranthene	mg/L	4.54E+01	-	4.54E+01	3.74E+00	-	3.74E+00	4.54E+01	-	4.54E+01
Fluorene	mg/L	4.34E-01	-	4.34E-01	1.51E-01	-	1.51E-01	4.34E-01	-	4.34E-01
Hexachlorobenzene	mg/L	9.08E-01	1.59E-02	1.59E-02	7.49E-02	6.82E-03	6.82E-03	9.08E-01	9.78E-03	9.78E-03
Naphthalene	mg/L	6.42E-01	-	6.42E-01	2.06E-01	-	2.06E-01	6.42E-01	-	6.42E-01
Nitroaniline, 2-	mg/L	2.52E+00	-	2.52E+00	5.05E-01	-	5.05E-01	2.52E+00	-	2.52E+00
Nitroso-di-N-propylamine, N-	mg/L	-	1.65E-03	1.65E-03	-	1.09E-03	1.09E-03	-	1.19E-03	1.19E-03
Phenanthrene ^f	mg/L	-	-	0.00E+00	-	-	0.00E+00	-	-	0.00E+00
Polychlorinated Biphenyls, Total ^c	mg/L	-	6.37E-02	6.37E-02	-	2.73E-02	2.73E-02	-	3.91E-02	3.91E-02
~Aroclor 1016	mg/L	7.95E-02	3.64E-01	7.95E-02	6.55E-03	1.56E-01	6.55E-03	7.95E-02	2.24E-01	7.95E-02
~Aroclor 1221	mg/L	-	1.10E-04	1.10E-04	-	1.30E-04	1.30E-04	-	9.36E-05	9.36E-05
~Aroclor 1232	mg/L	-	1.10E-04	1.10E-04	-	1.30E-04	1.30E-04	-	9.36E-05	9.36E-05
~Aroclor 1242	mg/L	-	1.27E-02	1.27E-02	-	5.46E-03	5.46E-03	-	7.82E-03	7.82E-03
~Aroclor 1248	mg/L	-	1.27E-02	1.27E-02	-	5.46E-03	5.46E-03	-	7.82E-03	7.82E-03
~Aroclor 1254	mg/L	2.27E-02	1.27E-02	1.27E-02	1.87E-03	5.46E-03	1.87E-03	2.27E-02	7.82E-03	7.82E-03
~Aroclor 1260	mg/L	-	1.27E-02	1.27E-02	-	5.46E-03	5.46E-03	-	7.82E-03	7.82E-03
Polycyclic aromatic hydrocarbons, Total Carcinogenic ^d	mg/L	-	3.49E-03	3.49E-03	-	2.80E-04	2.80E-04	-	8.04E-04	8.04E-04
~Benz[a]anthracene	mg/L	-	3.49E-02	3.49E-02	-	2.80E-03	2.80E-03	-	8.04E-03	8.04E-03
~Benzo[a]pyrene	mg/L	-	3.49E-03	3.49E-03	-	2.80E-04	2.80E-04	-	8.04E-04	8.04E-04
~Benzo[b]fluoranthene	mg/L	-	3.49E-02	3.49E-02	-	2.80E-03	2.80E-03	-	8.04E-03	8.04E-03
~Benzo[k]fluoranthene	mg/L	-	3.49E-01	3.49E-01	-	2.80E-02	2.80E-02	-	8.04E-02	8.04E-02
~Chrysene	mg/L	-	3.49E+00	3.49E+00	-	2.80E-01	2.80E-01	-	8.04E-01	8.04E-01
~Dibenz[a,h]anthracene	mg/L	-	3.49E-03	3.49E-03	-	2.80E-04	2.80E-04	-	8.04E-04	8.04E-04
~Indeno[1,2,3-cd]pyrene	mg/L	-	3.49E-02	3.49E-02	-	2.80E-03	2.80E-03	-	8.04E-03	8.04E-03

Hazard-based value calculated using target HI of 0.1.

Cancer-based value calculated using target ELCR of 1E-06.

No action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.6. Surface Water No Action Levels for Significant COPCs at PGDP (Continued)

(Values calculated on 11/6/2013 and are based on best available information.)

Analyte	Units	Adult Recreational User Swimming			Child Recreational User Swimming			Teen Recreational User Swimming		
		Hazard	Cancer	No Action	Hazard	Cancer	No Action	Hazard	Cancer	No Action
Pyrene	mg/L	1.42E-01	-	1.42E-01	5.15E-02	-	5.15E-02	1.42E-01	-	1.42E-01
Tetrachloroethylene	mg/L	2.10E-01	5.76E-01	2.10E-01	6.86E-02	6.35E-01	6.86E-02	2.10E-01	4.81E-01	2.10E-01
Trichloroethylene	mg/L	5.83E-02	7.96E-02	5.83E-02	1.49E-02	1.42E-02	1.42E-02	5.83E-02	2.41E-02	2.41E-02
Vinyl Chloride	mg/L	6.31E-01	-	6.31E-01	1.24E-01	7.00E-05	7.00E-05	6.31E-01	-	6.31E-01
Xylene, m-	mg/L	6.33E+00	-	6.33E+00	2.00E+00	-	2.00E+00	6.33E+00	-	6.33E+00
Xylene, Mixture	mg/L	6.72E+00	-	6.72E+00	2.11E+00	-	2.11E+00	6.72E+00	-	6.72E+00
Xylene, o-	mg/L	7.13E+00	-	7.13E+00	2.21E+00	-	2.21E+00	7.13E+00	-	7.13E+00
Xylene, p-	mg/L	6.82E+00	-	6.82E+00	2.13E+00	-	2.13E+00	6.82E+00	-	6.82E+00
Am-241	pCi/L	-	1.37E+02	1.37E+02	-	2.74E+02	2.74E+02	-	1.37E+02	1.37E+02
Cs-137+D	pCi/L	-	4.69E+02	4.69E+02	-	9.37E+02	9.37E+02	-	4.69E+02	4.69E+02
Np-237+D	pCi/L	-	2.11E+02	2.11E+02	-	4.23E+02	4.23E+02	-	2.11E+02	2.11E+02
Pu-238	pCi/L	-	1.09E+02	1.09E+02	-	2.17E+02	2.17E+02	-	1.09E+02	1.09E+02
Pu-239	pCi/L	-	1.06E+02	1.06E+02	-	2.11E+02	2.11E+02	-	1.06E+02	1.06E+02
Pu-240	pCi/L	-	1.06E+02	1.06E+02	-	2.11E+02	2.11E+02	-	1.06E+02	1.06E+02
Tc-99	pCi/L	-	5.18E+03	5.18E+03	-	1.04E+04	1.04E+04	-	5.18E+03	5.18E+03
Th-230	pCi/L	-	1.57E+02	1.57E+02	-	3.13E+02	3.13E+02	-	1.57E+02	1.57E+02
U-234	pCi/L	-	2.01E+02	2.01E+02	-	4.03E+02	4.03E+02	-	2.01E+02	2.01E+02
U-235+D	pCi/L	-	1.98E+02	1.98E+02	-	3.97E+02	3.97E+02	-	1.98E+02	1.98E+02
U-238+D	pCi/L	-	1.64E+02	1.64E+02	-	3.27E+02	3.27E+02	-	1.64E+02	1.64E+02

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Hazard-based value calculated using target HI of 0.1.

Cancer-based value calculated using target ELCR of 1E-06.

No action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.6. Surface Water No Action Levels for Significant COPCs at PGDP (Continued)

(Values calculated on 11/6/2013 and are based on best available information.)

Analyte	Units	Adult Recreational User Wading ^c			Child Recreational User Wading ^c			Teen Recreational User Wading ^c		
		Hazard	Cancer	No Action	Hazard	Cancer	No Action	Hazard	Cancer	No Action
Aluminum	mg/L	3.15E+03	-	3.15E+03	4.56E+02	-	4.56E+02	3.15E+03	-	3.15E+03
Antimony (metallic)	mg/L	1.89E-01	-	1.89E-01	2.73E-02	-	2.73E-02	1.89E-01	-	1.89E-01
Arsenic, Inorganic	mg/L	9.46E-01	6.93E-02	6.93E-02	1.37E-01	3.55E-02	3.55E-02	9.46E-01	2.24E-02	2.24E-02
Barium	mg/L	4.42E+01	-	4.42E+01	6.38E+00	-	6.38E+00	4.42E+01	-	4.42E+01
Beryllium and compounds	mg/L	4.42E-02	-	4.42E-02	6.38E-03	-	6.38E-03	4.42E-02	-	4.42E-02
Boron And Borates Only	mg/L	6.31E+02	-	6.31E+02	9.12E+01	-	9.12E+01	6.31E+02	-	6.31E+02
Cadmium (Diet)	mg/L	7.89E-02	-	7.89E-02	1.14E-02	-	1.14E-02	7.89E-02	-	7.89E-02
Chromium (Total) ^a	mg/L	6.15E+01	-	6.15E+01	8.89E+00	-	8.89E+00	6.15E+01	-	6.15E+01
Chromium(III), Insoluble Salts	mg/L	6.15E+01	-	6.15E+01	8.89E+00	-	8.89E+00	6.15E+01	-	6.15E+01
Chromium(VI)	mg/L	1.18E-01	2.60E-03	2.60E-03	1.71E-02	2.49E-04	2.49E-04	1.18E-01	3.14E-04	3.14E-04
Cobalt	mg/L	2.37E+00	-	2.37E+00	3.42E-01	-	3.42E-01	2.37E+00	-	2.37E+00
Copper	mg/L	1.26E+02	-	1.26E+02	1.82E+01	-	1.82E+01	1.26E+02	-	1.26E+02
Iron	mg/L	2.21E+03	-	2.21E+03	3.19E+02	-	3.19E+02	2.21E+03	-	2.21E+03
Lead	mg/L	-	-	1.50E-02	-	-	1.50E-02	-	-	1.50E-02
Manganese	mg/L	3.03E+00	-	3.03E+00	4.38E-01	-	4.38E-01	3.03E+00	-	3.03E+00
Mercury, Inorganic Salts	mg/L	6.62E-02	-	6.62E-02	9.57E-03	-	9.57E-03	6.62E-02	-	6.62E-02
Molybdenum	mg/L	1.58E+01	-	1.58E+01	2.28E+00	-	2.28E+00	1.58E+01	-	1.58E+01
Nickel Soluble Salts	mg/L	1.26E+01	-	1.26E+01	1.82E+00	-	1.82E+00	1.26E+01	-	1.26E+01
Selenium	mg/L	1.58E+01	-	1.58E+01	2.28E+00	-	2.28E+00	1.58E+01	-	1.58E+01
Silver	mg/L	1.05E+00	-	1.05E+00	1.52E-01	-	1.52E-01	1.05E+00	-	1.05E+00
Thallium (Soluble Salts)	mg/L	3.15E-02	-	3.15E-02	4.56E-03	-	4.56E-03	3.15E-02	-	3.15E-02
Uranium (Soluble Salts)	mg/L	9.46E+00	-	9.46E+00	1.37E+00	-	1.37E+00	9.46E+00	-	9.46E+00
Vanadium and Compounds	mg/L	4.13E-01	-	4.13E-01	5.97E-02	-	5.97E-02	4.13E-01	-	4.13E-01
Zinc and Compounds	mg/L	1.58E+03	-	1.58E+03	2.28E+02	-	2.28E+02	1.58E+03	-	1.58E+03
Acenaphthene	mg/L	9.08E-01	-	9.08E-01	2.09E-01	-	2.09E-01	9.08E-01	-	9.08E-01
Acenaphthylene ⁱ	mg/L	9.08E-01	-	9.08E-01	2.09E-01	-	2.09E-01	9.08E-01	-	9.08E-01
Acrylonitrile	mg/L	7.69E+01	1.43E-01	1.43E-01	1.36E+01	7.33E-02	7.33E-02	7.69E+01	4.62E-02	4.62E-02
Anthracene	mg/L	2.36E+00	-	2.36E+00	5.49E-01	-	5.49E-01	2.36E+00	-	2.36E+00
Benzene	mg/L	5.44E-01	1.07E-01	1.07E-01	1.03E-01	5.48E-02	5.48E-02	5.44E-01	3.45E-02	3.45E-02
Carbazole	mg/L	-	5.82E-02	5.82E-02	-	2.98E-02	2.98E-02	-	1.88E-02	1.88E-02
Carbon Tetrachloride	mg/L	3.20E-01	5.84E-02	5.84E-02	7.16E-02	2.99E-02	2.99E-02	3.20E-01	1.88E-02	1.88E-02
Chloroform	mg/L	2.39E+00	3.61E-01	3.61E-01	4.91E-01	1.85E-01	1.85E-01	2.39E+00	1.16E-01	1.16E-01
Dichloroethylene, 1,1-	mg/L	7.82E+00	-	7.82E+00	1.56E+00	-	1.56E+00	7.82E+00	-	7.82E+00
Dichloroethylene, 1,2- (Mixed Isomers)	mg/L	1.50E+00	-	1.50E+00	2.97E-01	-	2.97E-01	1.50E+00	-	1.50E+00
Dichloroethylene, 1,2-cis-	mg/L	3.33E-01	-	3.33E-01	6.61E-02	-	6.61E-02	3.33E-01	-	3.33E-01
Dichloroethylene, 1,2-trans-	mg/L	3.33E+00	-	3.33E+00	6.61E-01	-	6.61E-01	3.33E+00	-	3.33E+00
Dieldrin	mg/L	5.79E-04	3.85E-05	3.85E-05	1.35E-04	1.97E-05	1.97E-05	5.79E-04	1.24E-05	1.24E-05

Hazard-based value calculated using target HI of 0.1.

Cancer-based value calculated using target ELCR of 1E-06.

No action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.6. Surface Water No Action Levels for Significant COPCs at PGDP (Continued)

(Values calculated on 11/6/2013 and are based on best available information.)

Analyte	Units	Adult Recreational User Wading ^c			Child Recreational User Wading ^c			Teen Recreational User Wading ^c		
		Hazard	Cancer	No Action	Hazard	Cancer	No Action	Hazard	Cancer	No Action
Dioxins/Furans, Total (as TCDD) ^b	mg/L	-	-	-	-	-	-	-	-	-
~HpCDD, 2,3,7,8-	mg/L	-	-	-	-	-	-	-	-	-
~HpCDF, 2,3,7,8-	mg/L	-	-	-	-	-	-	-	-	-
~HxCDD, 2,3,7,8-	mg/L	-	-	-	-	-	-	-	-	-
~HxCDF, 2,3,7,8-	mg/L	-	-	-	-	-	-	-	-	-
~OCDD	mg/L	-	-	-	-	-	-	-	-	-
~OCDF	mg/L	-	-	-	-	-	-	-	-	-
~PeCDD, 2,3,7,8-	mg/L	-	-	-	-	-	-	-	-	-
~PeCDF, 1,2,3,7,8-	mg/L	-	-	-	-	-	-	-	-	-
~PeCDF, 2,3,4,7,8-	mg/L	-	-	-	-	-	-	-	-	-
~TCDD, 2,3,7,8-	mg/L	-	-	-	-	-	-	-	-	-
~TCDF, 2,3,7,8-	mg/L	-	-	-	-	-	-	-	-	-
Ethylbenzene	mg/L	3.51E+00	1.58E-01	1.58E-01	7.61E-01	8.07E-02	8.07E-02	3.51E+00	5.09E-02	5.09E-02
Fluoranthene	mg/L	-	-	-	-	-	-	-	-	-
Fluorene	mg/L	4.38E-01	-	4.38E-01	9.95E-02	-	9.95E-02	4.38E-01	-	4.38E-01
Hexachlorobenzene	mg/L	-	-	-	-	-	-	-	-	-
Naphthalene	mg/L	6.61E-01	-	6.61E-01	1.46E-01	-	1.46E-01	6.61E-01	-	6.61E-01
Nitroaniline, 2-	mg/L	3.24E+00	-	3.24E+00	6.95E-01	-	6.95E-01	3.24E+00	-	3.24E+00
Nitroso-di-N-propylamine, N-	mg/L	-	4.47E-03	4.47E-03	-	2.28E-03	2.28E-03	-	1.44E-03	1.44E-03
Phenanthrene ⁱ	mg/L	-	-	-	-	-	-	-	-	-
Polychlorinated Biphenyls, Total ^c	mg/L	-	-	-	-	-	-	-	-	-
~Aroclor 1016	mg/L	-	-	-	-	-	-	-	-	-
~Aroclor 1221	mg/L	-	1.65E-04	1.65E-04	-	8.44E-05	8.44E-05	-	5.32E-05	5.32E-05
~Aroclor 1232	mg/L	-	1.65E-04	1.65E-04	-	8.44E-05	8.44E-05	-	5.32E-05	5.32E-05
~Aroclor 1242	mg/L	-	-	-	-	-	-	-	-	-
~Aroclor 1248	mg/L	-	-	-	-	-	-	-	-	-
~Aroclor 1254	mg/L	-	-	-	-	-	-	-	-	-
~Aroclor 1260	mg/L	-	-	-	-	-	-	-	-	-
Polycyclic aromatic hydrocarbons, Total Carcinogenic ^d	mg/L	-	-	-	-	-	-	-	-	-
~Benz[a]anthracene	mg/L	-	-	-	-	-	-	-	-	-
~Benzo[a]pyrene	mg/L	-	-	-	-	-	-	-	-	-
~Benzo[b]fluoranthene	mg/L	-	-	-	-	-	-	-	-	-
~Benzo[k]fluoranthene	mg/L	-	-	-	-	-	-	-	-	-
~Chrysene	mg/L	-	-	-	-	-	-	-	-	-
~Dibenz[a,h]anthracene	mg/L	-	-	-	-	-	-	-	-	-
~Indeno[1,2,3-cd]pyrene	mg/L	-	-	-	-	-	-	-	-	-

Hazard-based value calculated using target HI of 0.1.

Cancer-based value calculated using target ELCR of 1E-06.

No action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.6. Surface Water No Action Levels for Significant COPCs at PGDP (Continued)

(Values calculated on 11/6/2013 and are based on best available information.)

Analyte	Units	Adult Recreational User Wading ^c			Child Recreational User Wading ^c			Teen Recreational User Wading ^c		
		Hazard	Cancer	No Action	Hazard	Cancer	No Action	Hazard	Cancer	No Action
Pyrene	mg/L	1.43E-01	-	1.43E-01	3.32E-02	-	3.32E-02	1.43E-01	-	1.43E-01
Tetrachloroethylene	mg/L	2.17E-01	8.96E-01	2.17E-01	4.95E-02	4.58E-01	4.95E-02	2.17E-01	2.89E-01	2.17E-01
Trichloroethylene	mg/L	6.50E-02	1.38E-01	6.50E-02	1.39E-02	1.32E-02	1.32E-02	6.50E-02	1.67E-02	1.67E-02
Vinyl Chloride	mg/L	7.75E-01	-	7.75E-01	1.41E-01	2.42E-04	2.42E-04	7.75E-01	-	7.75E-01
Xylene, m-	mg/L	6.51E+00	-	6.51E+00	1.42E+00	-	1.42E+00	6.51E+00	-	6.51E+00
Xylene, Mixture	mg/L	6.93E+00	-	6.93E+00	1.50E+00	-	1.50E+00	6.93E+00	-	6.93E+00
Xylene, o-	mg/L	7.36E+00	-	7.36E+00	1.59E+00	-	1.59E+00	7.36E+00	-	7.36E+00
Xylene, p-	mg/L	7.03E+00	-	7.03E+00	1.52E+00	-	1.52E+00	7.03E+00	-	7.03E+00
Am-241	pCi/L	-	-	-	-	-	-	-	-	-
Cs-137+D	pCi/L	-	-	-	-	-	-	-	-	-
Np-237+D	pCi/L	-	-	-	-	-	-	-	-	-
Pu-238	pCi/L	-	-	-	-	-	-	-	-	-
Pu-239	pCi/L	-	-	-	-	-	-	-	-	-
Pu-240	pCi/L	-	-	-	-	-	-	-	-	-
Tc-99	pCi/L	-	-	-	-	-	-	-	-	-
Th-230	pCi/L	-	-	-	-	-	-	-	-	-
U-234	pCi/L	-	-	-	-	-	-	-	-	-
U-235+D	pCi/L	-	-	-	-	-	-	-	-	-
U-238+D	pCi/L	-	-	-	-	-	-	-	-	-

^a Chromium (Total) utilizes Chromium III hazard values and Chromium VI inhalation risk values (to be conservative in terms of protecting human health).

^b Total dioxins/furans uses values for 2,3,7,8-TCDD, see screening note 9g in the Appendix A introduction, 'Screening Levels' on pages A-3 through A-5.

^c Total PCBs NALs are based on the high risk toxicity values.

^d Total carcinogenic PAHs uses values for benzo(a)pyrene, see screening note 9e in the Appendix A introduction, 'Screening Levels' on pages A-3 through A-5.

^e Wading scenario considers dermal contact only.

^f Acenaphthylene and Phenanthrene use values for Acenaphthene.

Hazard-based value calculated using target HI of 0.1.

Cancer-based value calculated using target ELCR of 1E-06.

No action level value is the lesser of the hazard- and cancer- based values when both are calculated.

Table A.7a. Risk-Based SSLs for Protection of RGA Groundwater for Significant COPCs at PGDP
(Values calculated in November 2013 and are based on best available information.)

CAS Number	Chemical	SSLs for EPA MCL ¹			SSLs PGDP NALs for the Resident (See Table A.5) ¹		
		SSL 1 (mg/kg)	SSL 20 (mg/kg)	GW Conc. (µg/L)	SSL 1 (mg/kg)	SSL 20 (mg/kg)	GW Conc. (µg/L)
7429-90-5	Aluminum	-	-	-	1.57E+03	3.13E+04	1.04E+03
7440-36-0	Antimony (metallic)	2.71E-01	5.42E+00	6.00E+00	1.88E-02	3.75E-01	4.15E-01
7440-38-2	Arsenic, Inorganic	2.92E-01	5.84E+00	1.00E+01	1.11E-03	2.21E-02	3.78E-02
7440-39-3	Barium	8.24E+01	1.65E+03	2.00E+03	8.50E+00	1.70E+02	2.06E+02
7440-41-7	Beryllium and compounds	3.16E+00	6.32E+01	4.00E+00	1.47E+00	2.93E+01	1.86E+00
7440-42-8	Boron And Borates Only	-	-	-	6.65E-01	1.33E+01	2.08E+02
7440-43-9	Cadmium	3.76E-01	7.52E+00	5.00E+00	3.86E-02	7.71E-01	5.13E-01
16065-83-1	Chromium (Total) ^a	1.80E+05	3.60E+06	1.00E+02	2.64E+06	5.28E+07	1.47E+03
18540-29-9	Chromium (III), Insoluble Salts	-	-	-	2.64E+06	5.28E+07	1.47E+03
7440-47-3	Chromium (VI)	-	-	-	5.75E-04	1.15E-02	3.00E-02
7440-48-4	Cobalt	-	-	-	1.42E-02	2.83E-01	3.13E-01
7440-50-8	Copper	4.58E+01	9.15E+02	1.30E+03	1.47E+00	2.93E+01	4.17E+01
7439-89-6	Iron	-	-	-	1.84E+01	3.68E+02	7.29E+02
7439-92-1	Lead	1.35E+01	2.70E+02	1.50E+01	1.35E+01	2.70E+02	1.50E+01
7439-96-5	Manganese	-	-	-	1.60E+00	3.19E+01	2.45E+01
7439-97-6	Mercury, Inorganic Salts	-	-	-	1.62E-02	3.23E-01	3.09E-01
7439-98-7	Molybdenum	-	-	-	1.05E-01	2.10E+00	5.21E+00
7440-02-0	Nickel Soluble Salts	-	-	-	1.36E+00	2.71E+01	2.08E+01
7782-49-2	Selenium	2.60E-01	5.20E+00	5.00E+01	2.71E-02	5.42E-01	5.21E+00
7440-22-4	Silver	-	-	-	4.38E-02	8.75E-01	5.15E+00
7440-28-0	Thallium (Soluble Salts)	1.42E-01	2.85E+00	2.00E+00	7.40E-04	1.48E-02	1.04E-02
NA	Uranium (Soluble Salts)	1.35E+01	2.70E+02	3.00E+01	1.41E+00	2.81E+01	3.13E+00
NA	Vanadium and Compounds	-	-	-	5.10E+00	1.02E+02	5.09E+00
7440-66-6	Zinc and Compounds	-	-	-	1.95E+01	3.89E+02	3.13E+02
83-32-9	Acenaphthene	-	-	-	4.57E-01	9.14E+00	4.46E+01
208-96-8	Acenaphthylene ^c	-	-	-	4.57E-01	9.14E+00	4.46E+01
107-13-1	Acrylonitrile	-	-	-	9.25E-06	1.85E-04	4.25E-02
120-12-7	Anthracene	-	-	-	5.80E+00	1.16E+02	1.76E+02
71-43-2	Benzene	2.56E-03	5.12E-02	5.00E+00	1.94E-04	3.88E-03	3.79E-01
86-74-8	Carbazole	-	-	-	3.67E-02	7.34E-01	1.98E+00
56-23-5	Carbon Tetrachloride	1.95E-03	3.89E-02	5.00E+00	1.49E-04	2.97E-03	3.82E-01
67-66-3	Chloroform	2.22E-02	4.43E-01	8.00E+01	5.25E-05	1.05E-03	1.89E-01
75-35-4	Dichloroethylene, 1,1-	2.52E-03	5.03E-02	7.00E+00	8.20E-03	1.64E-01	2.28E+01
540-59-0	Dichloroethylene, 1,2- (Mixed Isomers)	-	-	-	2.67E-03	5.33E-02	9.06E+00
156-59-2	Dichloroethylene, 1,2- <i>cis</i> -	2.06E-02	4.12E-01	7.00E+01	5.90E-04	1.18E-02	2.01E+00
156-60-5	Dichloroethylene, 1,2- <i>trans</i> -	2.94E-02	5.88E-01	1.00E+02	2.27E-03	4.54E-02	7.72E+00
60-57-1	Dieldrin	-	-	-	7.80E-05	1.56E-03	1.93E-03

Table A.7a. Risk-Based SSLs for Protection of RGA Groundwater for Significant COPCs at PGDP (Continued)
 (Values calculated in November 2013 and are based on best available information.)

CAS Number	Chemical	SSLs for EPA MCL ¹			SSLs PGDP NALs for the Resident (See Table A.5) ¹		
		SSL 1 (mg/kg)	SSL 20 (mg/kg)	GW Conc. (µg/L)	SSL 1 (mg/kg)	SSL 20 (mg/kg)	GW Conc. (µg/L)
1746-01-6	Dioxins/Furans, Total (as TCDD) ^b	1.50E-05	2.99E-04	3.00E-05	2.18E-07	4.35E-06	4.37E-07
37871-00-4	~HpCDD, 2,3,7,8-	-	-	-	1.02E-04	2.03E-03	4.37E-05
38998-75-3	~HpCDF, 2,3,7,8-	-	-	-	5.70E-05	1.14E-03	4.37E-05
34465-46-8	~HxCDD, 2,3,7,8-	-	-	-	6.05E-06	1.21E-04	4.37E-06
55684-94-1	~HxCDF, 2,3,7,8-	-	-	-	3.47E-06	6.94E-05	4.37E-06
3268-87-9	~OCDD	-	-	-	5.65E-03	1.13E-01	1.46E-03
39001-02-0	~OCDF	-	-	-	3.16E-03	6.32E-02	1.46E-03
36088-22-9	~PeCDD, 2,3,7,8-	-	-	-	3.79E-07	7.57E-06	4.37E-07
57117-41-6	~PeCDF, 1,2,3,7,8-	-	-	-	6.80E-06	1.36E-04	1.46E-05
57117-31-4	~PeCDF, 2,3,4,7,8-	-	-	-	6.80E-07	1.36E-05	1.46E-06
1746-01-6	~TCDD, 2,3,7,8-	1.50E-05	2.99E-04	3.00E-05	2.18E-07	4.35E-06	4.37E-07
51207-31-9	~TCDF, 2,3,7,8-	-	-	-	1.22E-06	2.44E-05	4.37E-06
100-41-4	Ethylbenzene	7.85E-01	1.57E+01	7.00E+02	1.48E-03	2.95E-02	1.32E+00
206-44-0	Fluoranthene	-	-	-	4.64E+00	9.27E+01	4.17E+01
86-73-7	Fluorene	-	-	-	4.96E-01	9.92E+00	2.68E+01
118-74-1	Hexachlorobenzene	1.26E-02	2.52E-01	1.00E+00	4.47E-04	8.94E-03	3.55E-02
91-20-3	Naphthalene	-	-	-	4.71E-04	9.42E-03	1.43E-01
88-74-4	Nitroaniline, 2-	-	-	-	4.33E-03	8.65E-02	1.02E+01
621-64-7	Nitroso-di-N-propylamine, N-	-	-	-	6.00E-06	1.20E-04	7.99E-03
85-01-8	Phenanthrene ^f	-	-	-	4.64E+00	9.27E+01	4.17E+01
1336-36-3	Polychlorinated Biphenyls, Total ^c	7.80E-02	1.56E+00	5.00E-01	2.22E-02	4.44E-01	1.42E-01
12674-11-2	~Aroclor 1016	-	-	-	7.00E-03	1.40E-01	7.30E-02
11104-28-2	~Aroclor 1221	-	-	-	8.20E-05	1.64E-03	4.82E-03
11141-16-5	~Aroclor 1232	-	-	-	8.20E-05	1.64E-03	4.82E-03
53469-21-9	~Aroclor 1242	-	-	-	4.44E-03	8.88E-02	2.84E-02
12672-29-6	~Aroclor 1248	-	-	-	4.35E-03	8.70E-02	2.84E-02
11097-69-1	~Aroclor 1254	-	-	-	5.45E-03	1.09E-01	2.09E-02
11096-82-5	~Aroclor 1260	-	-	-	1.99E-02	3.97E-01	2.84E-02
50-32-8	Polycyclic aromatic hydrocarbons, Total Carcinogenic ^d	2.35E-01	4.70E+00	2.00E-01	2.64E-03	5.27E-02	2.24E-03
56-55-3	Benz[a]anthracene	-	-	-	7.95E-03	1.59E-01	2.24E-02
50-32-8	Benzo[a]pyrene	2.35E-01	4.70E+00	2.00E-01	2.64E-03	5.27E-02	2.24E-03
205-99-2	Benzo[b]fluoranthene	-	-	-	2.69E-02	5.38E-01	2.24E-02
207-08-9	Benzo[k]fluoranthene	-	-	-	2.64E-01	5.27E+00	2.24E-01
218-01-9	Chrysene	-	-	-	8.10E-01	1.62E+01	2.24E+00
53-70-3	Dibenz[a,h]anthracene	-	-	-	8.60E-03	1.72E-01	2.24E-03
193-39-5	Indeno[1,2,3-cd]pyrene	-	-	-	1.56E-01	3.11E+00	2.24E-02

Table A.7a. Risk-Based SSLs for Protection of RGA Groundwater for Significant COPCs at PGDP (Continued)
 (Values calculated in November 2013 and are based on best available information.)

CAS Number	Chemical	SSLs for EPA MCL ¹			SSLs PGDP NALs for the Resident (See Table A.5) ¹		
		SSL 1 (mg/kg)	SSL 20 (mg/kg)	GW Conc. (µg/L)	SSL 1 (mg/kg)	SSL 20 (mg/kg)	GW Conc. (µg/L)
129-00-0	Pyrene	-	-	-	1.49E+00	2.98E+01	1.37E+01
127-18-4	Tetrachloroethylene	2.28E-03	4.55E-02	5.00E+00	1.48E-03	2.96E-02	3.26E+00
79-01-6	Trichloroethylene	1.79E-03	3.57E-02	5.00E+00	7.00E-05	1.40E-03	1.95E-01
75-01-4	Vinyl Chloride	6.90E-04	1.38E-02	2.00E+00	3.84E-06	7.68E-05	1.11E-02
108-38-3	Xylene, m-	-	-	-	1.82E-02	3.64E-01	1.87E+01
1330-20-7	Xylene, Mixture	9.85E+00	1.97E+02	1.00E+04	1.84E-02	3.68E-01	1.87E+01
95-47-6	Xylene, o-	-	-	-	1.84E-02	3.68E-01	1.87E+01
106-42-3	Xylene, p-	-	-	-	1.82E-02	3.64E-01	1.87E+01

¹ Values in this table were calculated using the Risk Assessment Information System (RAIS) in November 2013 located at the Web site http://rais.onrl.gov/cgi-bin/prg/PRG_search?select=chem. Prior to using the values in this table in a quantitative risk assessment, a risk assessor must be consulted to determine if any values need to be updated and to verify that the values are being used appropriately. SSL 1 indicates the soil screening level calculated for a dilution attenuation factor (DAF) of 1. SSL 20 indicates the soil screening level calculated for a DAF of 20.

^a Chromium (Total) utilizes chromium III hazard values and chromium VI inhalation risk values, See list of screening levels note 9b (on page A-4).

^b Total dioxins/furans uses values for 2,3,7,8-TCDD, see list of screening levels note 9g (on page A-5).

^c Total PCBs NALs are based on the high risk toxicity values.

^d Total carcinogenic polycyclic aromatic hydrocarbons use values for benzo(a)pyrene, see screening note 9e (page A-4).

^e Values for acenaphthylene use values for acenaphthene, as a surrogate.

^f Values for phenanthrene use values for fluoranthene, as a surrogate.

Only significant COPCs are shown (see Table 2.1). SSLs for other chemicals can be derived using similar methods as needed.

Note: Default parameters from RAIS used are as follows:

Dilution attenuation factor (unitless)	1 or 20
Fraction organic carbon in soil (unitless)	0.002
Water-filled soil porosity (L _{water} /L _{soil})	0.3
Dry soil bulk density (kg/L)	1.5
Soil particle density (kg/L)	2.65

Table A.7b. Risk-Based SSLs for Protection of RGA Groundwater for Radionuclide COPCs at PGDP

(Values calculated in November 2013 and are based on best available information.)

Parameter	Radionuclide	Units	Resident	
			10 ⁻⁶	10 ⁻⁴
14596102	Americium-241	pCi/g	1.83E+01	1.83E+03
10045973	Cesium-137+D	pCi/g	9.25E+00	9.25E+02
13994202	Neptunium-237+D	pCi/g	1.04E+00	1.04E+02
13981163	Plutonium-238	pCi/g	4.21E+00	4.21E+02
15117483	Plutonium-239	pCi/g	4.08E+00	4.08E+02
14119336	Plutonium-240	pCi/g	4.08E+00	4.08E+02
14133767	Technetium-99	pCi/g	1.46E-01	1.46E+01
14269637	Thorium-230	pCi/g	3.53E+01	3.53E+03
13966295	Uranium-234	pCi/g	9.50E-01	9.50E+01
15117961	Uranium-235+D	pCi/g	9.35E-01	9.35E+01
7440611	Uranium-238+D	pCi/g	7.71E-01	7.71E+01

Values in this table were calculated using the best available information in November 2013. Prior to using the values in this table (in a quantitative risk assessment), a risk assessor must be consulted to determine if any values need to be updated, and to verify that the values are being used appropriately.

SSLs calculated using the formula $PRG * DAF * (K_d + (\theta / \rho)) / 1,000$.

Where

PRG is the no action limit (for the 10⁻⁶ column) or action level (for the 10⁻⁴ column) for the resident (see Tables A.5 and A.2, respectively)

DAF is the dilution attenuation factor set at 20

K_d is the chemical-specific distribution coefficient (see below).

θ is the porosity set at 0.3

ρ is the density set at 1.5

K_d values and their references are the following:

Radionuclide	K _d	Reference
Americium-241	1.90E+03	DOE 2012
Cesium-137+D	2.80E+02	DOE 2012
Neptunium-237+D	7.00E+01	DOE 2003
Plutonium-238	5.50E+02	DOE 2003
Plutonium-239	5.50E+02	DOE 2003
Plutonium-240	5.50E+02	DOE 2003
Technetium-99	2.00E-01	DOE 2003
Thorium-230	3.20E+03	DOE 2003
Uranium-234	6.68E+01	DOE 2003
Uranium-235+D	6.68E+01	DOE 2003
Uranium-238+D	6.68E+01	DOE 2003

Table A.8. Dose-Based Soil/Sediment Screening Levels for Site-Related Radionuclides at PGDP
(Values calculated in December 2012 and are based on best available information.)

Parameter	Radionuclide	Units	Excavation Worker			
			1 mrem/yr	15 mrem/yr	25 mrem/yr	100 mrem/yr
14596102	Americium-241	pCi/g	1.41E+01	2.12E+02	3.53E+02	1.41E+03
10045973	Cesium-137	pCi/g	2.17E+00	3.25E+01	5.42E+01	2.17E+02
13994202	Neptunium-237+D	pCi/g	5.78E+00	8.67E+01	1.45E+02	5.78E+02
13981163	Plutonium-238	pCi/g	1.32E+01	1.98E+02	3.31E+02	1.32E+03
15117483	Plutonium-239	pCi/g	1.22E+01	1.82E+02	3.04E+02	1.22E+03
14119336	Plutonium-240	pCi/g	1.22E+01	1.82E+02	3.04E+02	1.22E+03
14133767	Technetium-99	pCi/g	4.45E+03	6.67E+04	1.11E+05	4.45E+05
14269637	Thorium-230	pCi/g	1.45E+01	2.17E+02	3.61E+02	1.45E+03
13966295	Uranium-234	pCi/g	6.19E+01	9.29E+02	1.55E+03	6.19E+03
15117961	Uranium-235+D	pCi/g	7.78E+00	1.17E+02	1.94E+02	7.78E+02
7440611	Uranium-238+D	pCi/g	3.16E+01	4.74E+02	7.90E+02	3.16E+03

Parameter	Radionuclide	Units	Industrial Worker			
			1 mrem/yr	15 mrem/yr	25 mrem/yr	100 mrem/yr
14596102	Americium-241	pCi/g	6.21E+01	9.32E+02	1.55E+03	6.21E+03
10045973	Cesium-137	pCi/g	1.62E+00	2.43E+01	4.05E+01	1.62E+02
13994202	Neptunium-237+D	pCi/g	5.27E+00	7.91E+01	1.32E+02	5.27E+02
13981163	Plutonium-238	pCi/g	9.33E+01	1.40E+03	2.33E+03	9.33E+03
15117483	Plutonium-239	pCi/g	8.56E+01	1.28E+03	2.14E+03	8.56E+03
14119336	Plutonium-240	pCi/g	8.58E+01	1.29E+03	2.15E+03	8.58E+03
14133767	Technetium-99	pCi/g	2.03E+04	3.04E+05	5.07E+05	2.03E+06
14269637	Thorium-230	pCi/g	1.00E+02	1.51E+03	2.51E+03	1.00E+04
13966295	Uranium-234	pCi/g	4.28E+02	6.43E+03	1.07E+04	4.28E+04
15117961	Uranium-235+D	pCi/g	6.46E+00	9.68E+01	1.61E+02	6.46E+02
7440611	Uranium-238+D	pCi/g	4.15E+01	6.23E+02	1.04E+03	4.15E+03

**Table A.8. Dose-Based Soil/Sediment Screening Levels for Site-Related Radionuclides at PGDP
(Continued)**

(Values calculated in December 2012 and are based on best available information.)

Parameter	Radionuclide	Units	Adult Recreator			
			1 mrem/yr	15 mrem/yr	25 mrem/yr	100 mrem/yr
14596102	Americium-241	pCi/g	2.96E+02	4.43E+03	7.39E+03	2.96E+04
10045973	Cesium-137	pCi/g	6.23E+00	9.34E+01	1.56E+02	6.23E+02
13994202	Neptunium-237+D	pCi/g	2.05E+01	3.07E+02	5.12E+02	2.05E+03
13981163	Plutonium-238	pCi/g	5.36E+02	8.04E+03	1.34E+04	5.36E+04
15117483	Plutonium-239	pCi/g	4.91E+02	7.37E+03	1.23E+04	4.91E+04
14119336	Plutonium-240	pCi/g	4.93E+02	7.40E+03	1.23E+04	4.93E+04
14133767	Technetium-99	pCi/g	9.75E+04	1.46E+06	2.44E+06	9.75E+06
14269637	Thorium-230	pCi/g	5.72E+02	8.57E+03	1.43E+04	5.72E+04
13966295	Uranium-234	pCi/g	2.44E+03	3.65E+04	6.09E+04	2.44E+05
15117961	Uranium-235+D	pCi/g	2.49E+01	3.74E+02	6.24E+02	2.49E+03
7440611	Uranium-238+D	pCi/g	1.65E+02	2.47E+03	4.11E+03	1.65E+04

Parameter	Radionuclide	Units	Child Recreator			
			1 mrem/yr	15 mrem/yr	25 mrem/yr	100 mrem/yr
14596102	Americium-241	pCi/g	1.22E+02	1.82E+03	3.04E+03	1.22E+04
10045973	Cesium-137	pCi/g	4.62E+00	6.94E+01	1.16E+02	4.62E+02
13994202	Neptunium-237+D	pCi/g	1.48E+01	2.22E+02	3.69E+02	1.48E+03
13981163	Plutonium-238	pCi/g	1.49E+02	2.23E+03	3.72E+03	1.49E+04
15117483	Plutonium-239	pCi/g	1.39E+02	2.09E+03	3.49E+03	1.39E+04
14119336	Plutonium-240	pCi/g	1.40E+02	2.09E+03	3.49E+03	1.40E+04
14133767	Technetium-99	pCi/g	1.77E+04	2.65E+05	4.42E+05	1.77E+06
14269637	Thorium-230	pCi/g	1.47E+02	2.21E+03	3.68E+03	1.47E+04
13966295	Uranium-234	pCi/g	5.19E+02	7.79E+03	1.30E+04	5.19E+04
15117961	Uranium-235+D	pCi/g	1.81E+01	2.71E+02	4.52E+02	1.81E+03
7440611	Uranium-238+D	pCi/g	1.06E+02	1.60E+03	2.66E+03	1.06E+04

Parameter	Radionuclide	Units	Teen Recreator			
			1 mrem/yr	15 mrem/yr	25 mrem/yr	100 mrem/yr
14596102	Americium-241	pCi/g	2.40E+03	3.60E+04	6.00E+04	2.40E+05
10045973	Cesium-137	pCi/g	5.05E+01	7.58E+02	1.26E+03	5.05E+03
13994202	Neptunium-237+D	pCi/g	1.66E+02	2.49E+03	4.15E+03	1.66E+04
13981163	Plutonium-238	pCi/g	4.55E+03	6.83E+04	1.14E+05	4.55E+05
15117483	Plutonium-239	pCi/g	4.16E+03	6.24E+04	1.04E+05	4.16E+05
14119336	Plutonium-240	pCi/g	4.18E+03	6.26E+04	1.04E+05	4.18E+05
14133767	Technetium-99	pCi/g	6.94E+05	1.04E+07	1.73E+07	6.94E+07
14269637	Thorium-230	pCi/g	4.44E+03	6.66E+04	1.11E+05	4.44E+05
13966295	Uranium-234	pCi/g	1.33E+04	1.99E+05	3.32E+05	1.33E+06
15117961	Uranium-235+D	pCi/g	2.02E+02	3.02E+03	5.04E+03	2.02E+04
7440611	Uranium-238+D	pCi/g	1.30E+03	1.95E+04	3.25E+04	1.30E+05

**Table A.8. Dose-Based Soil/Sediment Screening Levels for Site-Related Radionuclides at PGDP
(Continued)**

(Values calculated in December 2012 and are based on best available information.)

Parameter	Radionuclide	Units	Adult Resident			
			1 mrem/yr	15 mrem/yr	25 mrem/yr	100 mrem/yr
14596102	Americium-241	pCi/g	1.83E+01	2.75E+02	4.59E+02	1.83E+03
10045973	Cesium-137	pCi/g	3.85E-01	5.78E+00	9.64E+00	3.85E+01
13994202	Neptunium-237+D	pCi/g	1.27E+00	1.90E+01	3.17E+01	1.27E+02
13981163	Plutonium-238	pCi/g	3.34E+01	5.01E+02	8.34E+02	3.34E+03
15117483	Plutonium-239	pCi/g	3.06E+01	4.59E+02	7.65E+02	3.06E+03
14119336	Plutonium-240	pCi/g	3.07E+01	4.61E+02	7.68E+02	3.07E+03
14133767	Technetium-99	pCi/g	6.03E+03	9.05E+04	1.51E+05	6.03E+05
14269637	Thorium-230	pCi/g	3.56E+01	5.34E+02	8.90E+02	3.56E+03
13966295	Uranium-234	pCi/g	1.51E+02	2.27E+03	3.78E+03	1.51E+04
15117961	Uranium-235+D	pCi/g	1.54E+00	2.32E+01	3.86E+01	1.54E+02
7440611	Uranium-238+D	pCi/g	1.02E+01	1.53E+02	2.55E+02	1.02E+03

Parameter	Radionuclide	Units	Child Resident			
			1 mrem/yr	15 mrem/yr	25 mrem/yr	100 mrem/yr
14596102	Americium-241	pCi/g	1.01E+01	1.52E+02	2.54E+02	1.01E+03
10045973	Cesium-137	pCi/g	3.85E-01	5.78E+00	9.63E+00	3.85E+01
13994202	Neptunium-237+D	pCi/g	1.23E+00	1.85E+01	3.08E+01	1.23E+02
13981163	Plutonium-238	pCi/g	1.24E+01	1.86E+02	3.11E+02	1.24E+03
15117483	Plutonium-239	pCi/g	1.17E+01	1.75E+02	2.91E+02	1.17E+03
14119336	Plutonium-240	pCi/g	1.17E+01	1.75E+02	2.92E+02	1.17E+03
14133767	Technetium-99	pCi/g	1.47E+03	2.21E+04	3.68E+04	1.47E+05
14269637	Thorium-230	pCi/g	1.23E+01	1.85E+02	3.08E+02	1.23E+03
13966295	Uranium-234	pCi/g	4.34E+01	6.50E+02	1.08E+03	4.34E+03
15117961	Uranium-235+D	pCi/g	1.51E+00	2.26E+01	3.77E+01	1.51E+02
7440611	Uranium-238+D	pCi/g	8.87E+00	1.33E+02	2.22E+02	8.87E+02

Values in this table were calculated using the best available information in December 2012. Prior to using the values in this table (in a quantitative risk assessment), a risk assessor must be consulted to determine if any values need to be updated, and to verify that the values are being used appropriately.

Screening Value = $[\Sigma I / (\text{Pathway-Specific Action Levels})]^{-1}$

Pathways include ingestion, inhalation, and external gamma.

Table A.9. Dose-Based Groundwater Screening Levels for Site-Related Radionuclides at PGDP
(Values calculated in January 2013 and are based on best available information.)

Parameter	Radionuclide	Units	Industrial Worker				
			1	4	15	25	100
			mrem/yr	mrem/yr	mrem/yr	mrem/yr	mrem/yr
14596102	Americium-241	pCi/L	5.41E+00	2.16E+01	8.11E+01	1.35E+02	5.41E+02
10045973	Cesium-137+D	pCi/L	8.32E+01	3.33E+02	1.25E+03	2.08E+03	8.32E+03
13994202	Neptunium-237+D	pCi/L	9.75E+00	3.90E+01	1.46E+02	2.44E+02	9.75E+02
13981163	Plutonium-238	pCi/L	4.70E+00	1.88E+01	7.05E+01	1.18E+02	4.70E+02
15117483	Plutonium-239	pCi/L	4.32E+00	1.73E+01	6.49E+01	1.08E+02	4.32E+02
14119336	Plutonium-240	pCi/L	4.32E+00	1.73E+01	6.49E+01	1.08E+02	4.32E+02
14133767	Technetium-99	pCi/L	1.69E+03	6.76E+03	2.53E+04	4.22E+04	1.69E+05
14269637	Thorium-230	pCi/L	5.15E+00	2.06E+01	7.72E+01	1.29E+02	5.15E+02
13966295	Uranium-234	pCi/L	2.21E+01	8.83E+01	3.31E+02	5.52E+02	2.21E+03
15117961	Uranium-235+D	pCi/L	2.28E+01	9.13E+01	3.42E+02	5.70E+02	2.28E+03
7440611	Uranium-238+D	pCi/L	2.23E+01	8.91E+01	3.34E+02	5.57E+02	2.23E+03

Parameter	Radionuclide	Units	Adult Resident				
			1	4	15	25	100
			mrem/yr	mrem/yr	mrem/yr	mrem/yr	mrem/yr
14596102	Americium-241	pCi/L	1.93E+00	7.72E+00	2.90E+01	4.83E+01	1.93E+02
10045973	Cesium-137+D	pCi/L	2.97E+01	1.19E+02	4.46E+02	7.43E+02	2.97E+03
13994202	Neptunium-237+D	pCi/L	3.48E+00	1.39E+01	5.22E+01	8.71E+01	3.48E+02
13981163	Plutonium-238	pCi/L	1.68E+00	6.71E+00	2.52E+01	4.20E+01	1.68E+02
15117483	Plutonium-239	pCi/L	1.54E+00	6.18E+00	2.32E+01	3.86E+01	1.54E+02
14119336	Plutonium-240	pCi/L	1.54E+00	6.18E+00	2.32E+01	3.86E+01	1.54E+02
14133767	Technetium-99	pCi/L	6.03E+02	2.41E+03	9.05E+03	1.51E+04	6.03E+04
14269637	Thorium-230	pCi/L	1.84E+00	7.35E+00	2.76E+01	4.60E+01	1.84E+02
13966295	Uranium-234	pCi/L	7.88E+00	3.15E+01	1.18E+02	1.97E+02	7.88E+02
15117961	Uranium-235+D	pCi/L	8.15E+00	3.26E+01	1.22E+02	2.04E+02	8.15E+02
7440611	Uranium-238+D	pCi/L	7.95E+00	3.18E+01	1.19E+02	1.99E+02	7.95E+02

Parameter	Radionuclide	Units	Child Resident				
			1	4	15	25	100
			mrem/yr	mrem/yr	mrem/yr	mrem/yr	mrem/yr
14596102	Americium-241	pCi/L	1.91E+00	7.63E+00	2.86E+01	4.77E+01	1.91E+02
10045973	Cesium-137+D	pCi/L	5.36E+01	2.15E+02	8.04E+02	1.34E+03	5.36E+03
13994202	Neptunium-237+D	pCi/L	3.60E+00	1.44E+01	5.39E+01	8.99E+01	3.60E+02
13981163	Plutonium-238	pCi/L	1.66E+00	6.64E+00	2.49E+01	4.15E+01	1.66E+02
15117483	Plutonium-239	pCi/L	1.56E+00	6.24E+00	2.34E+01	3.90E+01	1.56E+02
14119336	Plutonium-240	pCi/L	1.56E+00	6.24E+00	2.34E+01	3.90E+01	1.56E+02
14133767	Technetium-99	pCi/L	2.24E+02	8.95E+02	3.36E+03	5.60E+03	2.24E+04
14269637	Thorium-230	pCi/L	1.66E+00	6.64E+00	2.49E+01	4.15E+01	1.66E+02
13966295	Uranium-234	pCi/L	5.85E+00	2.34E+01	8.78E+01	1.46E+02	5.85E+02
15117961	Uranium-235+D	pCi/L	5.97E+00	2.39E+01	8.96E+01	1.49E+02	5.97E+02
7440611	Uranium-238+D	pCi/L	5.54E+00	2.21E+01	8.30E+01	1.38E+02	5.54E+02

Values in this table were calculated using the best available information in January 2013. Prior to using the values in this table (in a quantitative risk assessment), a risk assessor must be consulted to determine if any values need to be updated and to verify that the values are being used appropriately.

Table A.10. Dose-Based Surface Water Screening Levels for Site-Related Radionuclides at PGDP
(Values calculated in January 2013 and are based on best available information.)

Parameter	Radionuclide	Units	Adult Recreator				
			1 mrem/yr	4 mrem/yr	15 mrem/yr	25 mrem/yr	100 mrem/yr
14596102	Americium-241	pCi/L	2.31E+02	9.24E+02	3.47E+03	5.78E+03	2.31E+04
10045973	Cesium-137+D	pCi/L	3.55E+03	1.42E+04	5.33E+04	8.88E+04	3.55E+05
13994202	Neptunium-237+D	pCi/L	4.17E+02	1.67E+03	6.25E+03	1.04E+04	4.17E+04
13981163	Plutonium-238	pCi/L	2.01E+02	8.03E+02	3.01E+03	5.02E+03	2.01E+04
15117483	Plutonium-239	pCi/L	1.85E+02	7.39E+02	2.77E+03	4.62E+03	1.85E+04
14119336	Plutonium-240	pCi/L	1.85E+02	7.39E+02	2.77E+03	4.62E+03	1.85E+04
14133767	Technetium-99	pCi/L	7.22E+04	2.89E+05	1.08E+06	1.80E+06	7.22E+06
14269637	Thorium-230	pCi/L	2.20E+02	8.80E+02	3.30E+03	5.50E+03	2.20E+04
13966295	Uranium-234	pCi/L	9.43E+02	3.77E+03	1.41E+04	2.36E+04	9.43E+04
15117961	Uranium-235+D	pCi/L	9.75E+02	3.90E+03	1.46E+04	2.44E+04	9.75E+04
7440611	Uranium-238+D	pCi/L	9.52E+02	3.81E+03	1.43E+04	2.38E+04	9.52E+04

Parameter	Radionuclide	Units	Child Recreator				
			1 mrem/yr	4 mrem/yr	15 mrem/yr	25 mrem/yr	100 mrem/yr
14596102	Americium-241	pCi/L	1.71E+02	6.84E+02	2.57E+03	4.28E+03	1.71E+04
10045973	Cesium-137+D	pCi/L	4.81E+03	1.93E+04	7.22E+04	1.20E+05	4.81E+05
13994202	Neptunium-237+D	pCi/L	3.23E+02	1.29E+03	4.84E+03	8.07E+03	3.23E+04
13981163	Plutonium-238	pCi/L	1.49E+02	5.96E+02	2.24E+03	3.73E+03	1.49E+04
15117483	Plutonium-239	pCi/L	1.40E+02	5.60E+02	2.10E+03	3.50E+03	1.40E+04
14119336	Plutonium-240	pCi/L	1.40E+02	5.60E+02	2.10E+03	3.50E+03	1.40E+04
14133767	Technetium-99	pCi/L	2.01E+04	8.03E+04	3.01E+05	5.02E+05	2.01E+06
14269637	Thorium-230	pCi/L	1.49E+02	5.96E+02	2.24E+03	3.73E+03	1.49E+04
13966295	Uranium-234	pCi/L	5.25E+02	2.10E+03	7.88E+03	1.31E+04	5.25E+04
15117961	Uranium-235+D	pCi/L	5.36E+02	2.14E+03	8.04E+03	1.34E+04	5.36E+04
7440611	Uranium-238+D	pCi/L	4.97E+02	1.99E+03	7.45E+03	1.24E+04	4.97E+04

Values in this table were calculated using the best available information in January 2013. Prior to using the values in this table (in a quantitative risk assessment), a risk assessor must be consulted to determine if any values need to be updated and to verify that the values are being used appropriately.

Table A.11. Dose-Based SSLs for Protection of RGA Groundwater for Site-Related Radionuclides at PGDP
(Values calculated in January 2013 and are based on best available information.)

Parameter	Radionuclide	Units	Resident				
			1	4	15	25	100
			mrem/yr	mrem/yr	mrem/yr	mrem/yr	mrem/yr
14596102	Americium-241	pCi/g	4.89E+05	1.96E+06	7.33E+06	1.22E+07	4.89E+07
10045973	Cesium-137 ^a	pCi/g	2.18E+13	8.70E+13	3.26E+14	5.44E+14	2.18E+15
13994202	Neptunium-237+D	pCi/g	2.62E+01	1.05E+02	3.92E+02	6.54E+02	2.62E+03
13981163	Plutonium-238	pCi/g	2.64E+05	1.06E+06	3.96E+06	6.60E+06	2.64E+07
15117483	Plutonium-239	pCi/g	1.91E+08	7.65E+08	2.87E+09	4.78E+09	1.91E+10
14119336	Plutonium-240	pCi/g	7.65E+06	3.06E+07	1.15E+08	1.91E+08	7.65E+08
14133767	Technetium-99	pCi/g	3.35E+01	1.34E+02	5.02E+02	8.36E+02	3.35E+03
14269637	Thorium-230 ^a	pCi/g	5.05E+09	2.02E+10	7.57E+10	1.26E+11	5.05E+11
13966295	Uranium-234	pCi/g	5.79E+01	2.32E+02	8.68E+02	1.45E+03	5.79E+03
15117961	Uranium-235+D	pCi/g	5.78E+01	2.31E+02	8.67E+02	1.45E+03	5.78E+03
7440611	Uranium-238+D	pCi/g	5.55E+01	2.22E+02	8.33E+02	1.39E+03	5.55E+03

^a Calculated using the specific activity constant as defined in the RESRAD User's Manual, Table J.3 (Yu et al. 2001), because there is no limit on the radionuclide concentrations for the specified dose criterion.

Values in this table were calculated using the best available information in January 2013. Prior to using the values in this table (in a quantitative risk assessment), a risk assessor must be consulted to determine if any values need to be updated and to verify that the values are being used appropriately. SSLs estimated using the RESRAD code version 6.5.

Table A.12. Background Concentrations for Surface and Subsurface Soil at PGDP
*Background Levels of Selected Radionuclides and Metals in Soil and Geologic Media
at the Paducah Gaseous Diffusion Plant (DOE 1997)*

Analyte	Background Value ^b	
	Surface	Subsurface
Inorganic Chemicals (mg/kg)^a		
Aluminum	13,000	12,000
Antimony	0.21	0.21
Arsenic	12	7.9
Barium	200	170
Beryllium	0.67	0.69
Cadmium	0.21	0.21
Calcium	200,000	6,100
Chromium (III)	16	43
Chromium (VI) ^d	---	---
Cobalt	14	13
Copper	19	25
Cyanide (CN) ^c	---	---
Iron	28,000	28,000
Lead	36	23
Magnesium	7,700	2,100
Manganese	1,500	820
Mercury	0.2	0.13
Nickel	21	22
Potassium	1,300	950
Selenium	0.8	0.7
Silver	2.3	2.7
Sodium	320	340
Sulfide ^d	---	---
Thallium	0.21	0.34
Tin ^d	---	---
Uranium	4.9	4.6
Vanadium	38	37
Zinc	65	60
Radionuclide (pCi/g)		
Cesium-137	0.49	0.28
Neptunium-237 ^e	0.1	---
Plutonium-238 ^e	0.073	---
Plutonium-239 ^e	0.025	---
Potassium-40	16	16
Radium-226	1.5	1.5
Strontium-90 ^e	4.7	---
Technetium-99	2.5	2.8
Thorium-228	1.6	1.6
Thorium-230	1.5	1.4
Thorium-232	1.5	1.5
Uranium-234	1.2 ^f	1.2 ^f
Uranium-235	0.06 ^f	0.06 ^f
Uranium-238	1.2	1.2

Notes: Cells with "----" indicate data are not available or not applicable.

Values contained in this table have not been approved for all uses by the PGDP Risk Assessment Working Group; therefore, the values presented here are provisional values and subject to change.

^a Includes inorganic chemicals found on Target Analyte List as defined by EPA in 1988 CLP Statement of Work and RCRA Appendix IX list of constituents.

^b Value for use in screening to determine if inorganic chemical or radionuclide detected at naturally occurring concentration in surface or subsurface soil. Details on the derivation of the background concentrations for antimony, beryllium, cadmium, thallium, uranium, and all radionuclides are in DOE 1997. Details on the derivation of the background concentration for all other inorganic chemicals are in DOE 1996.

^c Cyanide is not expected to be naturally occurring in soil at PGDP; background values were not derived.

^d Data are not adequate to calculate a background concentration in soil for this analyte.

^e Concentrations for these radionuclides in subsurface soil were not derived.

^f The values listed for uranium-234 and uranium-235 are not from the 1996 background study, but are derived from the natural isotopic abundance ratio and the uranium-238 values. The values for these radionuclides that appeared in the 2001 version of the Risk Methods Document (DOE 2001) were the UTLs of measured values for the individual isotopes as reported in the PGDP background study (DOE 1997).

Table A.13. Background Concentrations for Groundwater Drawn from the RGA and McNairy Formation at PGDP

Background Concentrations of Naturally Occurring Inorganic Chemicals and Selected Radionuclides in the Regional Gravel Aquifer and McNairy Formation at the Paducah Gaseous Diffusion Plant, Paducah, Kentucky in Feasibility Study for the Groundwater Operable Unit at Paducah Gaseous Diffusion Plant Volume 5(DOE 2000)

Analyte	Over All Observations*		Over Wells*		Comparison Value	
	RGA	McNairy	RGA	McNairy		
Inorganic Chemicals (mg/L)						
Aluminum	2.189	0.687	1.64	0.75	1.04	NAL ^b
Aluminum, Dissolved	0.311	0.579	0.201	0.587	1.04	NAL ^b
Antimony	0.060 ^a	0.060 ^a	0.060 ^a	0.060 ^a	0.006	MCL ^c
Antimony, Dissolved	0.060 ^a	0.060 ^a	0.060 ^a	0.060 ^a	0.006	MCL ^c
Arsenic	0.005 ^a	0.005 ^a	0.005 ^a	0.005 ^a	0.010	MCL ^c
Arsenic, Dissolved	0.005 ^a	0.005 ^a	0.005 ^a	0.005 ^a	0.010	MCL ^c
Barium	0.235	0.296	0.202	0.265	2	MCL ^c
Barium, Dissolved	0.2	0.268	0.179	0.266	2	MCL ^c
Beryllium	0.004 ^a	0.017 ^a	0.004 ^a	0.017 ^a	0.004	MCL ^c
Beryllium, Dissolved	0.004 ^a	0.004 ^a	0.004 ^a	0.004 ^a	0.004	MCL ^c
Cadmium	0.010 ^a	0.010 ^a	0.010 ^a	0.010 ^a	0.005	MCL ^c
Cadmium, Dissolved	0.010 ^a	0.010 ^a	0.010 ^a	0.010 ^a	0.005	MCL ^c
Calcium	41.238	38.858	40	39.47		n/a
Calcium, Dissolved	38.166	38.829	35.8	40.27		n/a
Chloride	91.021	19.708	89.2	20.23		n/a
Chromium	0.144	0.060 ^a	0.134	0.060 ^a	0.1	MCL ^c
Chromium, Dissolved	0.050 ^a	0.050 ^a	0.050 ^a	0.050 ^a	0.1	MCL ^c
Cobalt	0.045 ^a	0.096	0.045 ^a	0.072	0.000313	NAL ^b
Cobalt, Dissolved	0.045 ^a	0.045 ^a	0.045 ^a	0.045 ^a	0.000313	NAL ^b
Copper	0.036	0.057	0.034	0.033	1.3	MCL ^c
Copper, Dissolved	0.02	0.013 ^a	0.018	0.013 ^a	1.3	MCL ^c
Fluoride	0.27	0.33	0.245	0.298	4.0	MCL ^c
Iron	5.03	18.36	3.72	15.83	0.729	NAL ^b
Iron, Dissolved	0.267	12.372	0.164	9.446	0.729	NAL ^b
Lead	0.129	0.050 ^a	0.25	0.050 ^a	0.015	MCL ^c
Lead, Dissolved	0.098	0.050 ^a	0.25	0.050 ^a	0.015	MCL ^c
Magnesium	16.262	13.418	15.7	16.457		n/a
Magnesium, Dissolved	16.215	14.171	15.4	16.533		n/a
Manganese	0.119	0.941	0.082	0.729	0.0245	NAL ^b
Manganese, Dissolved	0.068	0.894	0.048	0.682	0.0245	NAL ^b
Mercury	0.0002 ^a	0.0002 ^a	0.0002 ^a	0.0002 ^a	0.002	MCL ^c
Mercury, Dissolved	0.0002 ^a	0.0002 ^a	0.0002 ^a	0.0002 ^a	0.002	MCL ^c
Molybdenum	0.050 ^a	0.050 ^a	0.050 ^a	0.050 ^a	0.00521	NAL ^b
Molybdenum, Dissolved	0.050 ^a	0.050 ^a	0.050 ^a	0.050 ^a	0.00521	NAL ^b
Nickel	0.682	0.109 ^a	0.530 ^g	0.109 ^a	0.0208	NAL ^b
Nickel, Dissolved	0.305	0.050 ^a	0.305	0.050 ^a	0.0208	NAL ^b
Nitrate as Nitrogen	15.561	1.474	13.5	1.43	10	MCL ^c
Potassium	5.195	55.752	4.47	64.08		n/a
Potassium, Dissolved	4.096	51.205	3.7	58.75		n/a
Selenium	0.005 ^a	0.005 ^a	0.005 ^a	0.005 ^a	0.05	MCL ^c
Selenium, Dissolved	0.005 ^a	0.005 ^a	0.005 ^a	0.005 ^a	0.05	MCL ^c
Silica	26.401	36	21.1	29.4		n/a

Table A.13. Background Concentrations for Groundwater Drawn from the RGA and McNairy Formation at PGDP (Continued)

Background Concentrations of Naturally Occurring Inorganic Chemicals and Selected Radionuclides in the Regional Gravel Aquifer and McNairy Formation at the Paducah Gaseous Diffusion Plant, Paducah, Kentucky in Feasibility Study for the Groundwater Operable Unit at Paducah Gaseous Diffusion Plant Volume 5 (DOE 2000)

Analyte	Over All Observations*		Over Wells*		Comparison	
	RGA	McNairy	RGA	McNairy	Value	
Silver	0.011 ^a	0.050 ^a	0.011 ^a	0.050 ^a	0.00515	NAL ^b
Silver, Dissolved	0.060 ^a	0.050 ^a	0.060 ^a	0.050 ^a	0.00515	NAL ^b
Sodium	59.45	29.2	63.5	24.92		n/a
Sodium, Dissolved	60.433	27.98	65.7	25.9		n/a
Sulfate	19.947	28.9	19.1	27.27		n/a
Thallium	0.056 ^a	0.644	0.056 ^a	0.255	0.002	MCL ^c
Thallium, Dissolved	0.056 ^a	0.056 ^a	0.056 ^a	0.056 ^a	0.002	MCL ^c
Uranium	0.002 ^a	0.001 ^a	0.002 ^a	0.001 ^a	0.03	MCL ^c
Uranium, Dissolved	0.002 ^a	0.001	0.002 ^a	0.001	0.03	MCL ^c
Vanadium	0.134	0.126	0.139	0.119	0.00525	NAL ^b
Vanadium, Dissolved	0.134	0.126	0.131	0.107	0.00525	NAL ^b
Zinc	0.054	0.142	0.025	0.104	0.313	NAL ^b
Zinc, Dissolved	0.049	0.116	0.026	0.08	0.313	NAL ^b
Radionuclides (pCi/L)						
Gross Alpha	5.8	11.9	2.36	5.3		n/a
Gross Beta	13.8	144.5	7.3	125.4		n/a
Neptunium-237	0.8	0.5	0.21	0.13	15	MCL ^d
Plutonium-239	0.1	0.2	0.03	0.04	15	MCL ^d
Radium-226	0.6	1.2	0.1	0.29	5	MCL ^e
Radon-222	626	295	555.3	228.3		n/a
Technetium-99	22.3	20.6	10.8	7.8	18.2	NAL ^b
Thorium-230	1.1	1.5	0.54	0.4	15	MCL ^d
Total Radium	1.3	0.7	0.46	0.36	5	MCL ^e
Uranium-234 ^f	0.7	0.3	0.7	0.3	0.709	NAL ^b
Uranium-235 ^f	0.3	0.2	0.3	0.2	0.698	NAL ^b
Uranium-238 ^f	0.7	0.3	0.7	0.3	0.575	NAL ^b

Values contained in this table have not been approved for all uses by the PGDP Risk Assessment Working Group; therefore, the values presented here are provisional values and subject to change. The issues to be resolved are the data set from which these values were derived and the statistical methods used to analyze the data set.

* For inorganic chemicals, background concentrations were derived for both total and filtered samples over all observations within a group (i.e., both groundwater wells and soil boring data) and over only groundwater wells within a group (i.e., only groundwater wells data). For radionuclides, background concentrations were derived using total sample results only because there were too few results from filtered samples.

For all projects where averages within groundwater wells over time are considered, the values derived for these groundwater wells under the column heading "over wells" should be used. For all other projects, the values shown under the column heading "over all observations" should be used.

^a Background value was derived qualitatively over all observations because analyte was never detected or was detected infrequently at a concentration near the analyte's detection limit.

^b NAL is the no action level for the resident (i.e., the lesser of the child resident HI=0.1 and adult/child ELCR=1E-06, see Table A.5).

^c MCL is the primary maximum contaminant level from "<http://water.epa.gov/drink/contaminants/index.cfm>"; last updated June 3, 2013, (see also Table A.14).

^d See Table A.14 for additional information.

^e MCL is for radium-226 and radium-228 combined.

^f Uranium isotopic concentrations were derived from the mass concentration of uranium.

^g Nickel background value varies from previous Risk Methods Documents due to an error in calculation in the earlier version.

n/a = an NAL or MCL comparison value is not available, as defined in footnotes b and c.

Gray shading. For those background values that were derived qualitatively over all observations, because the analyte never was detected or was detected infrequently at a concentration near the analyte's detection limit (see footnote a), the gray shading indicates that the background value is greater than the comparison value.

Table A.14. Regulatory Action Levels for PGDP
(Values are based on best available information in November 2013.)

Parameter	Chemical	Units	Primary MCLs ^a	Primary MCLGs ^a	Secondary MCLs ^a	State Water Supply WQC ^b	State Fish Consump. WQC ^b	Fed. Combined WQC ^c
7429905	Aluminum	mg/L			0.05 -0.2			
7440360	Antimony	mg/L	0.006	0.006		5.60E-03	6.40E-01	5.60E-03
7440382	Arsenic	mg/L	0.010	0		1.00E-02		1.80E-05
7440393	Barium	mg/L	2	2		1.00E+00		1.00E+00
7440417	Beryllium	mg/L	0.004	0.004		4.00E-03		e
7440428	Boron and Borates Only	mg/L						e
7440439	Cadmium	mg/L	0.005	0.005		5.00E-03		e
7440473	Chromium (Total)	mg/L	0.1	0.1		1.00E-01		e
16065831	Chromium (III)	mg/L						e
18540299	Chromium (VI)	mg/L						e
7440484	Cobalt	mg/L						
7440508	Copper	mg/L	1.3	1.3	1	1.30E+00		1.30E+00
7439896	Iron	mg/L			0.3	3.00E-01		3.00E-01 ^m
7439921	Lead	mg/L	0.015	0		1.50E-02		
7439965	Manganese	mg/L			0.05			5.00E-02
7439976	Mercury (Inorganic)	mg/L	0.002	0.002		2.00E-03	5.10E-05	
7439987	Molybdenum	mg/L						
7440020	Nickel	mg/L				6.10E-01	4.60E+00	6.10E-01
7782492	Selenium	mg/L	0.05	0.05		1.70E-01	4.20E+00	1.70E-01 ^e
7440224	Silver	mg/L			0.1			
7440280	Thallium	mg/L	0.002	0.0005		2.40E-04	4.70E-04	2.40E-04
7440611	Uranium	mg/L	0.03	0				
n/a	Vanadium and Compounds	mg/L						
7440666	Zinc	mg/L			5	7.40E+00	2.60E+01	7.40E+00
83329	Acenaphthene	mg/L				6.70E-01	9.90E-01	6.70E-01
208968	Acenaphthylene	mg/L						
107131	Acrylonitrile	mg/L				5.10E-05	2.50E-04	5.10E-05
120127	Anthracene	mg/L				8.30E+00	4.00E+01	8.30E+00
71432	Benzene	mg/L	0.005	0		2.20E-03	5.10E-02	2.20E-03
86748	Carbazole	mg/L						
56235	Carbon tetrachloride	mg/L	0.005	0		2.30E-04	1.60E-03	2.30E-04
67663	Chloroform	mg/L	0.08 ^d	0.07		5.70E-03	4.70E-01	5.70E-03
75354	Dichloroethylene, 1,1-	mg/L	0.007	0.007		3.30E-01	7.10E+00	3.30E-01
540590	Dichloroethylene, 1,2- (mixed isomers)	mg/L						
156592	Dichloroethylene, <i>cis</i> -1,2-	mg/L	0.07	0.07				
156605	Dichloroethylene, <i>trans</i> -1,2-	mg/L	0.1	0.1		1.40E-01	1.00E+01	1.40E-01 ^e
60571	Dieldrin	mg/L				5.20E-08	5.40E-08	5.20E-08

Table A.14. Regulatory Action Levels for PGDP (Continued)

(Values are based on best available information in November 2013.)

Parameter	Chemical	Units	Primary MCLs ^a	Primary MCLGs ^a	Secondary MCLs ^a	State Water Supply WQC ^b	State Fish Consump. WQC ^b	Fed. Combined WQC ^c
1746016	Dioxins/Furans, Total (as TCDD)	mg/L	0.00000003	0		5.00E-12	5.10E-12	5.00E-12
37871004	~HpCDD, 2,3,7,8-	mg/L						
38998753	~HpCDF, 2,3,7,8-	mg/L						
34465468	~HxCDD, 2,3,7,8-	mg/L						
55684941	~HxCDF, 2,3,7,8-	mg/L						
3268879	~OCDD	mg/L						
39001020	~OCDF	mg/L						
36088229	~PeCDD, 2,3,7,8-	mg/L						
57117416	~PeCDF, 1,2,3,7,8-	mg/L						
57117314	~PeCDF, 2,3,4,7,8-	mg/L						
1746016	~TCDD, 2,3,7,8-	mg/L	0.00000003	0		5.00E-12	5.10E-12	5.00E-12
51207319	~TCDF, 2,3,7,8-	mg/L						
100414	Ethylbenzene	mg/L	0.7	0.7		5.30E-01	2.10E+00	5.30E-01
206440	Fluoranthene	mg/L				1.30E-01	1.40E-01	1.30E-01
86737	Fluorene	mg/L				1.10E+00	5.30E+00	1.10E+00
118741	Hexachlorobenzene	mg/L	0.001	0		2.80E-07	2.90E-07	2.80E-07
91203	Naphthalene	mg/L						
88744	Nitroaniline, 2-	mg/L						
621647	N-Nitrosodi-n-propylamine	mg/L				5.00E-06	5.10E-04	5.00E-06
85018	Phenanthrene	mg/L						
1336363	Polychlorinated biphenyls (PCBs)	mg/L	0.0005	0		6.40E-08	6.40E-08	6.40E-08
12674112	~Aroclor 1016	mg/L						
11104282	~Aroclor 1221	mg/L						
11141165	~Aroclor 1232	mg/L						
53469219	~Aroclor 1242	mg/L						
12672296	~Aroclor 1248	mg/L						
11097691	~Aroclor 1254	mg/L						
11096825	~Aroclor 1260	mg/L						
	Polycyclic aromatic hydrocarbons (cPAH),							
50328	Total Carcinogenic (as BaP)	mg/L	0.0002	0		3.80E-06	1.80E-05	3.80E-06
56553	~Benzo(a)anthracene	mg/L				3.80E-06	1.80E-05	3.80E-06
50328	~Benzo(a)pyrene	mg/L	0.0002	0		3.80E-06	1.80E-05	3.80E-06
205992	~Benzo(b)fluoranthene	mg/L				3.80E-06	1.80E-05	3.80E-06
207089	~Benzo(k)fluoranthene	mg/L				3.80E-06	1.80E-05	3.80E-06
218019	~Chrysene	mg/L				3.80E-06	1.80E-05	3.80E-06
53703	~Dibenz(a,h)anthracene	mg/L				3.80E-06	1.80E-05	3.80E-06
193395	~Indeno(1,2,3-cd)pyrene	mg/L				3.80E-06	1.80E-05	3.80E-06

Table A.14. Regulatory Action Levels for PGDP (Continued)
(Values are based on best available information in November 2013.)

Parameter	Chemical	Units	Primary MCLs ^a	Primary MCLGs ^a	Secondary MCLs ^a	State Water Supply WQC ^b	State Fish Consump. WQC ^b	Fed. Combined WQC ^c
129000	Pyrene	mg/L				8.30E-01	4.00E+00	8.30E-01
127184	Tetrachloroethene	mg/L	0.005	0		6.90E-04	3.30E-03	6.90E-04
79016	Trichloroethene	mg/L	0.005	0		2.50E-03	3.00E-02	2.50E-03
75014	Vinyl chloride	mg/L	0.002	0		2.50E-05	2.40E-03	2.50E-05
1330207	Xylenes, total	mg/L	10	10				
108383	Xylene, m-	mg/L						
95476	Xylene, o-	mg/L						
106423	Xylene, P-	mg/L						
14596102	Am-241	pCi/L	15 ^f					
10045973	Cs-137+D	mrem/yr	4 ^g					
13994202	Np-237+D	pCi/L	15 ^h					
13981163	Pu-238	pCi/L	15 ^{h, i}					
15117483	Pu-239	pCi/L	15 ^{h, i}					
14119336	Pu-240	pCi/L	15 ^{h, i}					
14133767	Tc-99	mrem/yr	4 ^j					
14269637	Th-230	pCi/L	15 ^{h, k}					
n/a	Uranium	pCi/L	1					
13966295	U-234	pCi/L	1					
15117961	U-235	pCi/L	1					
744061	U-238	pCi/L	1					

Notes:

Values in this table were based on the best available information in November 2013. Prior to using the values in this table (in a quantitative risk assessment), a risk assessor must be consulted to determine if any values need to be updated and to verify that the values are being used appropriately. Please see source materials for complete discussions of these values. Only values for water are provided. Values are for planning purposes only.

^a Accessed at “<http://water.epa.gov/drink/contaminants/index.cfm>”; last updated June 3, 2013.

^b From 401 KAR § 10:031 recodified from 401 KAR § 5:031

^c Accessed at “<http://water.epa.gov/scitech/swguidance/standards/criteria/current/index.cfm#hhtable>”; last updated August 22, 2013.

^d MCL is for the sum of the concentrations for trihalomethanes.

^e <http://water.epa.gov/scitech/swguidance/standards/criteria/current/index.cfm#hhtable> indicates more stringent MCL has been issued.

^f “The MCL for alpha particle activity applies to Am-241. The limit is 15 pCi/L alpha particle activity in drinking water,” from “<http://www.epa.gov/radiation/radionuclides/americiam.html>” last updated April 24, 2012.

^g The value derived by the EPA from the 4 mrem/yr MCL for Cs-137 is 200 pCi/L (see “http://www.epa.gov/ogwdw/radionuclides/pdfs/guide_radionuclides_smallsystems_compliance.pdf” dated February 2002; accessed December 12, 2013).

^h Accessed at “http://www.epa.gov/superfund/health/contaminants/radiation/pdfs/9283_1_14.pdf” dated November 6, 2001; accessed December 12, 2013.

ⁱ “EPA has established a maximum contaminant level (MCL) of 15 pCi/L for alpha particle activity, excluding radon and uranium, in drinking water. Plutonium would be covered under this MCL,” from “<http://www.epa.gov/superfund/health/contaminants/radiation/pdfs/plutonium.pdf>” dated February 2002; accessed December 12, 2013.

^j The value derived by the EPA from the 4 mrem/yr MCL for Tc-99 is 900 pCi/L, (see http://www.epa.gov/ogwdw/radionuclides/pdfs/guide_radionuclides_smallsystems_compliance.pdf). An alternate value derived by the EPA from the 4 mrem/yr MCL is 3,790 pCi/L and was proposed in the July 18, 1991 *Federal Register*. See Table A.9 for Tc-99 dose-based groundwater screening levels resulting in a 4 mrem/yr dose based upon more recent dosimetry.

^k “...Thorium would be covered under this MCL,” from “<http://www.epa.gov/superfund/health/contaminants/radiation/pdfs/thorium.pdf>” dated July 2002; accessed December 12, 2013.

^l The uranium MCL is calculated by converting the public drinking water standard of 0.03 mg/L for uranium (chemical toxicity) to 20 pCi/L for total uranium. Isotopic uranium values derived from this standard are 10.24 pCi/L for U-234, 0.466 pCi/L for U-235, and 9.99 pCi/L for U-238, assuming natural occurring uranium at 0.725% uranium-235 and the following ratios:

- Uranium-234/uranium-235 = 21-22 obtained from conversion approximately 21.9
- Uranium-235/uranium-238 = 0.04-0.05 obtained from conversion approximately 0.045

^m Accessed at “http://water.epa.gov/scitech/swguidance/standards/criteria/aqlife/upload/2009_01_13_criteria_goldbook.pdf” dated May 1, 1986; accessed December 12, 2013.

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APPENDIX B

DERIVATION OF PRELIMINARY REMEDIATION GOALS

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B.1 DERIVATION OF RISK-BASED PRELIMINARY REMEDIATION GOALS

This appendix presents the methods used to derive the direct contact risk-based action and no action screening levels [i.e., preliminary remediation goals (PRGs)]. The PRGs presented in Appendix A are taken from a U.S. Environmental Protection Agency (EPA)-sponsored site on the World Wide Web (<http://rais.ornl.gov/>) that maintains a calculator that was used for deriving PRGs using Paducah site-specific parameters. Groundwater protection soil screening levels (SSLs) are taken from an EPA-sponsored site on the World Wide Web (<http://rais.ornl.gov/epa/ssl1.shtml>); methods used to derive these SSLs are discussed on that site.

B.1.1 INTRODUCTION

No action and action direct contact risk-based PRGs may be derived using a modification of methods described in Risk Assessment Guidance for Superfund (RAGS), Part B. In RAGS, Part B, risk-based PRGs are developed by rearranging the equations used to calculate risk or hazard in a risk assessment so that the equations solve for a concentration or activity of an analyte that “yields” a target risk or hazard. To derive the direct contact PRGs, the linear, direct relationship between the concentration or activity of an analyte in an environmental medium and the risk or hazard that exposure to this analyte can present were used. Although this method differs from that in RAGS, Part B, the ultimate results of the modified calculations match those that are received by rearranging the risk or hazard equations.

B.1.2 MATERIALS

In order to derive risk-based PRGs, several pieces of information are required. These are the receptors of interest, the routes through which the receptors may be exposed and equations describing these routes, carcinogenic (cancer) and noncarcinogenic (hazard) toxicity values, and target risk and hazard values. Each of these is discussed in the following subsections.

B.1.2.1 Receptors

Table B.1 provides a matrix showing the medium-receptor combinations for which PRGs were derived. As shown there, over all media, the receptors for which no action and action direct contact risk-based PRGs were derived are the industrial worker, the resident, the recreational user, and the outdoor worker. These receptors were chosen because they represent the most likely current and future receptors for most areas and units at the Paducah Gaseous Diffusion Plant (PGDP). Also, it is believed that the PRGs derived for these receptors yield a range of values that is most useful for determining the cleanup priority for the various areas and units at PGDP. Note: Outdoor worker PRGs (used for surface soil) can be used for a construction/excavation worker (used for surface and subsurface soil); however, because the duration and frequency of exposure for a construction/excavation worker would be markedly less than that for an outdoor worker, scenario-specific PRGs for the construction/excavation worker based on site-specific conditions should be derived, as appropriate.

Table B.1 also includes a series of notes that discusses how the PRGs are to be applied to data during site scoping. These notes should be considered before site scoping is attempted.

**Table B.1. Action and No Action Risk-Based Screening Levels
for Chemicals Derived for PGDP by Medium**

Scenario/Receptor	Medium		
	Groundwater	Surface Water	Soil/Sediment
Outdoor worker	No	Yes	Yes
Industrial Worker	No	Yes	Yes
Adult Recreator	No	Yes	Yes
Teen Recreator	No	Yes	Yes
Child Recreator	No	Yes	Yes
Adult Resident	Yes	No	Yes
Child Resident	Yes	No	Yes

Notes:

1. All groundwater screening is to be performed using the resident. Of the two receptors (i.e., child and adult), use of the child is more conservative (in terms of protecting human health). Note that values for soil deemed protective of groundwater also are available and are based on the resident only.
2. The surface water screening value selected is a location-specific decision. For all areas along effluent ditches or along creeks carrying effluent, the industrial worker screening values are appropriate. Additionally, at areas outside the industrialized areas, use of the recreator values are appropriate. Of the three recreator values available, the child recreator values are most conservative (in terms of protecting human health). Note that two different sets of recreator values are available; these are a set for screening shallow water courses under a wading scenario and a set for screening deeper water courses under a swimming scenario. While which of these two values to use is a location-specific decision, general guidance should be to use the wading values for most areas. If exposure by a resident to surface water is of concern, use of the recreator values is appropriate, because rates of contact for the recreator were selected assuming that the individual would be a local resident.
3. Determining which soil and sediment screening value is appropriate is a location-specific decision. For all locations inside the industrialized area at PGDP where surface soil contamination is of concern, use of the industrial worker values is appropriate. For locations inside the industrialized area at PGDP where contact with surface soil and subsurface soil is of concern (e.g., soil from the surface down to 16 ft bgs), use of the outdoor worker values is appropriate for a construction/excavation worker. (Scenario-specific PRGs for the construction/excavation worker based on site-specific conditions should be derived, as appropriate. See discussion in Section 3.3.4.3.) For locations, outside the industrialized area where surface soil contamination is of concern, screening using the recreator and/or resident values is appropriate. As with the surface water values, the child resident values are the most “conservative (in terms of protecting human health).” Generally, the recreator values are more appropriate for areas along ditches and creeks (i.e., for bank soils), and the resident values are more appropriate for grassy fields. Finally, the outdoor worker values also can be considered for contact with soil in locations outside the industrialized area if this scenario is appropriate for the locations considered. (If screening considers both surface and subsurface soil in locations outside the industrialized area, however, development of scenario-specific PRGs for the outdoor worker based on site-specific conditions is a better approach. See discussion in Section 3.3.4.3.)
4. As mentioned above, values for soil, for protection of groundwater also, are available. These should be used in all areas.

B.1.2.2 Exposure Routes and Equations

The exposure routes considered for the various media-scenario combinations are provided below. Included in this list are the tables from Appendix D that display the equations used to derive chronic daily intake or absorbed dose. The sources for these exposure parameters are provided in the tables in Appendix D. These exposure parameters are summarized in a table following Subsection B.2.3. Since PRGs shown in Appendix A were derived using the Risk Assessment Information System (RAIS) online calculator, equations used for obtaining PGDP PRGs may or may not match the equations for calculating the reasonable maximum exposure (RME) intakes shown in Appendix D. Equations in Appendix D should be used to calculate RME intakes in a PGDP baseline human health risk assessment.

- Residential Scenario (Child and Adult)—Groundwater, Chemicals
Ingestion of water (Table D.1), inhalation of vapors emitted from water during household uses (including showering) (Table D.2), dermal contact with water during showering (Table D.4).
- Residential Scenario (Child and Adult)—Soil and Sediment, Chemicals
Incidental ingestion of contaminated soil or sediment (Table D.5), dermal contact with contaminated soil or sediment (Table D.6), inhalation of particulates emitted from soil or sediment (Table D.7), inhalation of vapors emitted from soil or sediment (Table D.7).

- Residential Scenario (Child and Adult)—Soil and Sediment, Radionuclides
Incidental ingestion of contaminated soil or sediment (Table D.5), inhalation of particulates emitted from soil or sediment (Table D.7), inhalation of vapors emitted from soil or sediment (Table D.7), external exposure to ionizing radiation from soil or sediment (Table D.18).
- Industrial Worker Scenario—Surface Water, Chemicals
Dermal contact with contaminated surface water (Table D.33).
- Industrial Worker Scenario—Soil, Chemicals
Incidental ingestion of contaminated soil (Table D.29), inhalation of particulates emitted from soil (Tables D.31), inhalation of vapors emitted from soil (Table D.31), dermal contact with contaminated soil (Table D.33).
- Industrial Worker Scenario—Soil, Radionuclides
Incidental ingestion of contaminated soil (Table D.29), inhalation of particulates emitted from soil (Table D.31), inhalation of vapors emitted from soil (Table D.31), external exposure to ionizing radiation from soil (Table D.34).
- Outdoor worker Scenario—Surface Water, Chemicals
Dermal contact with contaminated surface water (Table D.36).
- Outdoor worker Scenario—Soil and Sediment, Chemicals
Incidental ingestion of contaminated soil or sediment (Table D.37 or Table D.30), inhalation of particulates emitted from soil or sediment (Tables D.38), inhalation of vapors emitted from soil or sediment (Table D.38), dermal contact with contaminated soil or sediment (Table D.39).
- Outdoor worker Scenario—Soil and Sediment, Radionuclides
Incidental ingestion of contaminated soil or sediment (Table D.37 or Table D.30), inhalation of particulates emitted from soil or sediment (Table D.38), inhalation of vapors emitted from soil or sediment (Table D.38), external exposure to ionizing radiation from soil or sediment (Table D.40).
- Recreational User Scenario (Child, Teen, and Adult)—Sediment, Chemicals
Incidental ingestion of contaminated sediment (Table D.15), dermal contact with contaminated sediment (Table D.16), inhalation of particulates emitted from sediment (Tables D.17), inhalation of vapors emitted from sediment (Table D.17).
- Recreational User Scenario (Child, Teen, and Adult)—Sediment, Radionuclides
Incidental ingestion of contaminated sediment (Table D.15), inhalation of particulates emitted from sediment (Tables D.17), inhalation of vapors emitted from sediment (Table D.17), external exposure to ionizing radiation from soil or sediment (Table D.18).
- Recreational User Scenario (Child, Teen, and Adult)—Surface Water (Swimming), Chemicals
Incidental ingestion of contaminated surface water (Table D.19), dermal contact with surface water (Table D.21).
- Recreational User Scenario (Child, Teen, and Adult)—Surface Water (Wading), Chemicals
Dermal contact with surface water (Table D.20).

It is important to note that PRGs are not derived for industrial use of groundwater. These are not derived because they would not be useful to remedial decision making, as indicated in the following material taken from RAGS, Part B, Section 3.2.1 (EPA 1991).

Once ground water is determined to be suitable for drinking, risk-based concentrations should be based on residential exposures....Similarly, for surface water that is to be used for drinking, the risk-based PRGs should be calculated for residential populations, and not simply worker populations.

Note that the number of exposure routes included in these calculations exceeds that presented in RAGS, Part B, for each scenario. Including exposure routes beyond those discussed in RAGS, Part B, is consistent with material in Section 3.1.1 of RAGS, Part B, where it is stated: "Additional exposure pathways (e.g., dermal absorption) are possible and may be significant at some sites for some contaminants, while perhaps only one exposure pathway (e.g., direct ingestion of water only) may be relevant in others. In any case, the risk-based PRG for each chemical should be calculated by considering all of the relevant exposure pathways."

B.1.2.3 Toxicity Values

The toxicity values used in the derivation of the risk-based concentrations are taken from a variety of sources. The sources of these values are discussed in Section 3.3.5 of the main text. The values are presented in a table following Subsection B.2.3.

B.1.2.4 Values for Volatilization Factors

Derivation of PRGs requires that volatilization factors (VFs) be developed for each chemical based on its physical properties. The soil parameters used in the calculation of VFs and the chemical-specific parameters used in the calculation of VFs and the VF values are presented in tables following Subsection B.2.3.

B.1.2.5 Target Risk and Hazard Values

The target risk and hazard values used when deriving the risk-based concentrations for no action are 1×10^{-6} and 0.1, respectively. The target risk and hazard values used when deriving the risk-based concentrations for action are 1×10^{-4} and 3, respectively. Note, if five or more constituents are detected at a site, it may be appropriate during project scoping to reduce the chemical-specific target risk used to derive the risk-based concentrations for no action.

B.1.3 METHOD OF DERIVATION

Each risk-based PRG is calculated using the same method and generally follows the examples provided by EPA; equations for the derivation for PRGs can be found at the following link:

http://www.epa.gov/reg3hwmd/risk/human/rb-concentration_table/equations.htm

B.2. DERIVATION OF DOSE-BASED PRELIMINARY REMEDIATION GOALS FOR RADIONUCLIDES

The following describes the methods used to derive direct-contact dose-based screening. Methods for deriving the groundwater protection SSLs also are provided for comparison to direct-contact PRGs.

B.2.1 INTRODUCTION

Direct contact dose-based PRGs for radionuclides were derived using a modification of methods described by RAGS, Part B. This modified approach is similar to that used to develop risk-based PRGs for PGDP except for two additional modifications. These are (1) the exposure duration (ED) term was dropped because dose limits are based on annual dose and not lifetime exposure, and (2) slope factors and reference doses were replaced with radiation dose conversion factors (DCFs). Additionally, dose-based SSLs for the protection of groundwater were derived using the Residual Radioactivity Materials Model (RESRAD) computer code, version 6.4. Note that risk-based SSLs were not derived, but were extracted from existing tables provided by the EPA.

B.2.2 MATERIALS

In order to derive dose-based screening levels, several pieces of information are required. These are the receptors of interest, the routes through which the receptors may be exposed and equations describing these routes, activity- or concentration-to-dose conversion factors, and target dose values. Each of these is discussed in the following sections.

B.2.2.1 Receptors

The receptors considered in dose-based screening level calculations are described in the derivation of risk-based PRGs. The description is not repeated here, although it is noted that the ED term is not relevant for dose calculations. This is because dose-based values generally call for yearly rather than lifetime values and are the value that would yield target dose in a given year (e.g., in units of mrem/yr). Direct contact screening levels were derived for the industrial worker, the resident (adult and child), the recreational user (adult, child, and teen), and the outdoor worker. These receptors were chosen because they represent the most likely current and future receptors for most areas and units at PGDP. Also, it is believed that the screening levels derived for these receptors yield a range of values that are most useful for determining the cleanup priority for the various areas and units at PGDP.

Table B.2 lists the media evaluated, by receptor, and includes a series of notes that discuss how the screening levels are to be applied to data during site scoping. These notes should be considered before site scoping is attempted. Table B.2 varies slightly from the version used in nonradiological risk-based PRG development because dermal contact is not a relevant pathway for the radionuclides of interest.

**Table B.2. Action and No Action Risk-Based Screening Levels and SSLs
for Radionuclides Derived for PGDP by Medium**

Scenario/Receptor	Medium		
	Groundwater	Surface Water	Soil/Sediment
Outdoor worker	No	No	Yes
Industrial Worker	No	No	Yes
Adult Recreator	No	Yes	Yes
Teen Recreator	No	Yes	Yes
Child Recreator	No	Yes	Yes
Adult Resident	Yes	No	Yes
Child Resident	Yes	No	Yes

Notes:

1. All groundwater screening is to be performed using the resident. Note that values for soil deemed protective of groundwater also are available and are based on the resident only.
2. Dose-based values for surface water are provided only for recreators.
3. Determining which soil and sediment screening value is appropriate is a location-specific decision. For all locations inside the industrialized area at PGDP where surface soil contamination is of concern, use of the industrial worker values is appropriate. For locations inside the industrialized area at PGDP where contact with surface soil and subsurface soil is of concern (e.g., soil from the surface down to 16 ft bgs), use of the outdoor worker values is appropriate for a construction/excavation worker. (Scenario-specific PRGs for the construction/excavation worker based on site-specific conditions should be derived, as appropriate. See discussion in Section 3.3.4.3.) For locations, outside the industrialized area where surface soil contamination is of concern, screening using the recreator and/or resident values is appropriate. As with the surface water values, the child resident values are the most “conservative (in terms of protecting human health).” Generally, the recreator values are more appropriate for areas along ditches and creeks (i.e., for bank soils), and the resident values are more appropriate for grassy fields. Finally, the outdoor worker values also can be considered for contact with soil in locations outside the industrialized area if this scenario is appropriate for the locations considered. (If screening considers both surface and subsurface soil in locations outside the industrialized area, however, development of scenario-specific PRGs for the outdoor worker based on site-specific conditions is a better approach. See discussion in Section 3.3.4.3.)
4. As mentioned above, values for soil, for protection of groundwater, are also available. These should be used in all areas.

B.2.2.2 Exposure Routes and Equations

As discussed above, the exposure routes and equations used to calculate dose-based screening levels are similar to those used to develop risk-based PRGs. The only pathway-specific difference is that dermal contact is not considered (for radionuclides). Instead, the external gamma pathway is evaluated to account for non-uptake exposures. This being the only difference, the complete list of exposure routes considered for the various media-scenario combinations is not repeated here.

The equations used to calculate dose-based screening levels are similar to those used to develop risk-based values, but with two exceptions. First, dose-based limits are typically for a single year of exposure. Therefore, The ED terms appropriately are dropped from all equations to produce per-year PRG and SSL results. Second, slope factors and reference doses were replaced with DCFs given that the human-health-based limits are radiological doses (in units mrem/yr) rather than carcinogenic risk or non-carcinogenic hazard.

B.2.2.3 Toxicity Values

The toxicity values (i.e., DCFs) used in the derivation of the dose-based concentrations are taken from the latest version of RESRAD output (RESRAD 6.5). DCFs are consistent with International Commission on Radiological Protection (ICRP) Publication 60 and Publication 72. The use of ICRP 60 and 72 is consistent with the requirements established by DOE Order 458.1. These DCFs are given in unit mrem/pCi for the inhalation and ingestion pathways or mrem/yr/pCi/g (i.e., pCi/g in soil/sediment) for the external gamma pathway. The values are provided in Table B.3.

Table B.3. Dose Conversion Factors for Radionuclides of Interest

Radionuclide	Pathway (units)		
	Ingestion ^a (mrem/pCi)	Inhalation ^a (mrem/pCi)	External Gamma ^a (mrem/yr per pCi/g)
Adult			
Americium-241	7.40E-04	3.55E-01	3.72E-02
Cesium-137	4.81E-05	1.44E-04	3.38E+00
Neptunium-237+D	4.10E-04	1.85E-01	1.10E+00
Plutonium-238	8.51E-04	4.07E-01	1.16E-04
Plutonium-239	9.25E-04	4.44E-01	2.64E-04
Plutonium-240	9.25E-04	4.44E-01	1.13E-04
Technetium-99	2.37E-06	4.81E-05	1.08E-04
Thorium-230	7.77E-04	3.70E-01	1.07E-03
Uranium-234	1.81E-04	3.48E-02	3.44E-04
Uranium-235+D	1.75E-04	3.15E-02	8.36E-01
Uranium-238+D	1.74E-04	2.96E-02	1.20E-01
Teen			
Americium-241	7.40E-04	3.40E-01	3.72E-02
Cesium-137	4.81E-05	1.55E-04	3.38E+00
Neptunium-237+D	4.11E-04	1.74E-01	1.01E+00
Plutonium-238	8.14E-04	3.70E-01	1.16E-04
Plutonium-239	8.88E-04	4.07E-01	2.64E-04
Plutonium-240	8.88E-04	4.07E-01	1.13E-04
Technetium-99	3.03E-06	5.55E-05	1.08E-04
Thorium-230	8.14E-04	3.66E-01	1.07E-03
Uranium-234	2.74E-04	3.70E-02	3.44E-04
Uranium-235+D	2.61E-04	3.40E-02	8.36E-01
Uranium-238+D	2.48E-04	3.22E-02	1.20E-01
Child			
Americium-241	9.99E-04	4.44E-01	3.72E-02
Cesium-137	3.55E-05	2.59E-04	3.38E+00
Neptunium-237+D	5.30E-04	2.22E-01	1.01E+00
Plutonium-238	1.15E-03	5.18E-01	1.16E-04
Plutonium-239	1.22E-03	5.55E-01	2.64E-04
Plutonium-240	1.22E-03	5.55E-01	1.13E-04
Technetium-99	8.51E-06	8.88E-05	1.08E-04
Thorium-230	1.15E-03	5.18E-01	1.07E-03
Uranium-234	3.26E-04	7.03E-02	3.44E-04
Uranium-235+D	3.19E-04	6.29E-02	8.36E-01
Uranium-238+D	2.96E-04	6.66E-02	1.20E-01

Notes:

^a From RESRAD version 6.5 output, December 2012. These values are consistent with ICRP 60 and 72.

“+D” indicates all impacts for short-lived decay products (i.e., radioactive decay products with a half-life less than six months) are considered; that is, all short-lived decay products of a principal radionuclide down to, but not including, the next principal radionuclide or the final nonradioactive nuclide in the decay chain).

B.2.2.4 Target Dose Values

The target dose values used when deriving the dose-based concentrations in soil and sediment are 1.0, 15 and 25 mrem/yr. An additional target dose of 4.0 mrem/yr was added for the surface water and groundwater media in consideration of the federal drinking water standard (standards available at <http://water.epa.gov/drink/contaminants/index.cfm>) although these standards are applicable to public drinking water supplies.

B.2.3 METHOD OF DERIVATION

Each dose-based PRG is calculated in the same manner. The general equation used to calculate all PRGs reflects the direct, linear relationship between the environmental concentrations and the dose estimate. This calculation is shown in Eq. 1 to demonstrate the difference in calculation method from that used in developing risk-based PRGs. For this evaluation, PRGs were developed by combining the soil ingestion, dust inhalation, and external gamma pathways. Both surface water and groundwater ingestion were considered separately as these media should be considered on a case-by-case basis.

Dose-based SSLs for protection of RGA groundwater for site-related radionuclides were calculated for each radionuclide of interest used in the RESRAD code version 6.5 and the same exposure parameter values used in risk-based calculations. Other RESRAD input parameters, except for the area of the contamination zone that was set equal to 0.5 acres and the 10,000-year evaluation period, were consistent with those used for establishing single radionuclide soil guidelines at PGDP (ORISE 2012).

Table B.4 includes list of exposure parameters used in calculation of human health PRGs. Tables B.3 and B.5 include the toxicity values and information used in PRG derivation.

$$C_i = \frac{TD}{\sum_{i,j} (DCF_i \times A_{ij})} \quad \text{Eq. 1}$$

where:	C _i	The dose-based concentration for radionuclide “i” (i.e., calculated screening level)
	TD	The target doses (see Section B.2.2.4)
	DCF _i	Dose conversion factor for radionuclide “i” (i.e., in mrem/pCi or mrem/yr per pCi/g)
	A _{ij}	Activity of radionuclide “i” ingested or inhaled (in pCi) or specific activity in soil/sediment (in pCi/g) per unit concentration in medium “j”

Table B.4. Exposure Parameters Used in Calculation of Human Health PRGs

Pathway Variable	Units	Default Industrial Worker	Outdoor Worker	Adult Resident	Child Resident	Adult Recreational User	Teen Recreational User	Child Recreational User
General Parameters Used in All Intake Models (unless otherwise noted)								
Exposure frequency (EF)	days/year	250	185	350	350	104	140	140
Exposure duration (ED)	years	25	25	24	6	12	12	6
Body weight (BW)	kg	70	70	70	15	70	43	15
Averaging time - cancer (AT-C)	days	70 × 365	70 × 365	70 × 365	70 × 365	70 × 365	70 × 365	70 × 365
Averaging time - noncancer (AT-N)	days	365 × 25	365 × 25	365 × 24	365 × 6	365 × 12	365 × 12	365 × 6
Ingestion of Water (Tables D.1, D.26)								
Drinking water ingestion rate (IR)	L/day	NA	NA	2	1.5	NA	NA	NA
Inhalation RGA Groundwater (Table D.2, D.27)								
Indoor inhalation rate	m ³ /hour	0.833	NA	0.833	0.833	NA	NA	NA
Exposure time in the shower (ET _{shower})	hours/day	0.2	NA	0.2	0.2	NA	NA	NA
Time of shower (t1)	hour	0.1	NA	0.1	0.1	NA	NA	NA
Time after shower (t2)	hour	0.1	NA	0.1	0.1	NA	NA	NA
Fraction volatilized while showering (f _{shower})	unitless	0.75	NA	0.75	0.75	NA	NA	NA
Water flow rate (Fw)	L/h	890	NA	890	890	NA	NA	NA
Bathroom volume (Va)	m ³	11	NA	11	11	NA	NA	NA
Averaging time - cancer (AT-C)	hours	24 × 70 × 365	NA	24 × 70 × 365	24 × 70 × 365	NA	NA	NA
Averaging time - noncancer (AT-N)	hours	24 × 365 × 25	NA	24 × 365 × 24	24 × 365 × 6	NA	NA	NA
Exposure time household use (ET _{house})	hours/day	NA	NA	24	24	NA	NA	NA
Exchange rate (ER)	changes/day	NA	NA	10	10	NA	NA	NA
Mixing coefficient (MC)	unitless	NA	NA	0.5	0.5	NA	NA	NA
Fraction volatilized household use (f _{house})	unitless	NA	NA	0.5	0.5	NA	NA	NA
Water flow rate (WHF)	L/day	NA	NA	890	890	NA	NA	NA
House volume (HV)	m ³	NA	NA	450	450	NA	NA	NA
Dermal Contact with RGA Groundwater (showering) (Tables D.4, D.28)								
Body surface area exposed (SA)	m ²	1.815	NA	1.815	0.65	NA	NA	NA
Event time (t _{event})	hour/event	0.2	NA	0.2	0.2	NA	NA	NA
Event frequency (EV)	events/day	1	NA	1	1	NA	NA	NA
Incidental Ingestion of Soil/Sediment (Tables D.5, D.15, D.29, D.30, D.37)								
Incidental ingestion rate (IR)	mg/day	50	480	100	200	100	100	200
Fraction ingested		1	1	1	1	1	1	1
Dermal Contact with Soil/Sediment (Tables D.6, D.16, D.33, D.39)								
Body surface area exposed (SA)	m ² /day	0.47	0.47	0.57	0.28	0.57	0.75	0.28
Soil-to-skin adherence factor (AF)	mg/cm ² -day	1	1	1	1	1	1	1
Inhalation of Vapors and Particulates Emitted from Soil/Sediment (Tables D.7, D.17, D.31, D.32, D.38)								
Total inhalation rate (IR)	m ³ /hour	2.5	2.5	0.833	0.833	2.5	2.5	2.5
Exposure time (ET) (soil)	hours/day	8	8	24	24	5	5	5
Exposure time (ET) (sediment)	hours/day	2.6	NA	NA	NA	NA	NA	NA
Particulate emission factor (PEF)	m ³ /kg	6.20E+08	6.20E+08	9.30E+08	9.30E+08	9.30E+08	9.30E+08	9.30E+08

Table B.4. Exposure Parameters Used in Calculation of Human Health PRGs (Continued)

Pathway Variable	Units	Default Industrial Worker	Outdoor Worker	Adult Resident	Child Resident	Adult Recreational User	Teen Recreational User	Child Recreational User
External Exposure to Ionizing Radiation from Soil/Sediment (Tables D.8, D.18, D.34, D. 35, D.40)								
Exposure frequency (EF)	day/day	250/365	185/365	350/365	350/365	104/365	140/365	140/365
Gamma shielding factor (Se)	unitless	0.2	0.2	0.2	0.2	0	0	0
Gamma exposure time factor (Te)	hr/hr	8/24	8/24	18/24	18/24	5/24	5/24	5/24
Consumption of Home-grown Vegetables (Table D.9)								
Diet fraction (FI)	unitless	NA	NA	0.4	0.4	NA	NA	NA
Ingestion rate (IR)	kg/d	NA	NA	0.72	0.29	NA	NA	NA
Consumption of Beef (Table D.10)								
Diet fraction (FI)	unitless	NA	NA	1	1	NA	NA	NA
Ingestion rate (IR)	kg/d	NA	NA	0.19	0.07	NA	NA	NA
Consumption of Milk (Table D.11)								
Diet fraction (FI)	unitless	NA	NA	1	1	NA	NA	NA
Ingestion rate (IR)	kg/d	NA	NA	1.25	0.9	NA	NA	NA
Consumption of Poultry (Table D.12)								
Diet fraction (FI)	unitless	NA	NA	1	1	NA	NA	NA
Ingestion rate (IR)	kg/d	NA	NA	0.17	0.07	NA	NA	NA
Consumption of Pork (Table D.13)								
Diet fraction (FI)	unitless	NA	NA	1	1	NA	NA	NA
Ingestion rate (IR)	kg/d	NA	NA	0.08	0.03	NA	NA	NA
Consumption of Eggs (Table D.14)								
Diet fraction (FI)	unitless	NA	NA	1	1	NA	NA	NA
Ingestion rate (IR)	kg/d	NA	NA	0.11	0.06	NA	NA	NA
Incidental Ingestion of Surface Water (swimming) (Table D.19)								
Ingestion rate (IR)	L/hr	NA	NA	NA	NA	0.05	0.05	0.05
Exposure time (ET)	hr/day	NA	NA	NA	NA	2.6	2.6	2.6
Exposure frequency (EF)	day/year	NA	NA	NA	NA	45	45	45
Dermal Contact with Surface Water (wading) (Table D.20)								
Body surface area exposed (SA)	m ²	NA	NA	NA	NA	1.06	0.75	0.33
Exposure frequency (EF)	day/year	NA	NA	NA	NA	52	140	140
Exposure time (ET)	hr/day	NA	NA	NA	NA	2.6	2.6	2.6

Table B.4. Exposure Parameters Used in Calculation of Human Health PRGs (Continued)

Pathway Variable	Units	Default Industrial Worker	Outdoor Worker	Adult Resident	Child Resident	Adult Recreational User	Teen Recreational User	Child Recreational User
Dermal Contact with Surface Water (swimming) (Table D.21)								
Body surface area exposed (SA)	m ²	NA	NA	NA	NA	1.815	1.31	0.65
Exposure frequency (EF)	days/year	NA	NA	NA	NA	45	45	45
Exposure time (ET)	hr/day	NA	NA	NA	NA	2.6	2.6	2.6
Event (EV)	event/day	NA	NA	NA	NA	1	1	1
Dermal Contact with Surface Water (Table D.36)								
Body surface area exposed (SA)	m ²	0.47	0.47	NA	NA	NA	NA	NA
Exposure frequency (EF)	days/year	250	20	NA	NA	NA	NA	NA
Exposure time (ET)	hr/day	2.6	8	NA	NA	NA	NA	NA
Consumption of Fish (Table D.22)								
Diet fraction (FI)	Unitless	NA	NA	NA	NA	1	1	1
Ingestion rate (IR)	kg/d	NA	NA	NA	NA	0.029	0.029	0.029
Exposure Frequency (EF)	days/year	NA	NA	NA	NA	365	365	365
Consumption of Venison (Table D.23)								
Diet fraction (FI)	Unitless	NA	NA	NA	NA	1	1	1
Ingestion rate (IR)	kg/d	NA	NA	NA	NA	0.032	0.032	0.007
Consumption of Rabbit (Table D.24)								
Diet fraction (FI)	Unitless	NA	NA	NA	NA	1	1	1
Ingestion rate (IR)	kg/d	NA	NA	NA	NA	0.0165	0.0082	0.0033
Consumption of Quail (Table D.25)								
Diet fraction (FI)	Unitless	NA	NA	NA	NA	1	1	1
Ingestion rate (IR)	kg/d	NA	NA	NA	NA	0.0047	0.0024	0.00094

Information compiled October 2012.

NA = not applicable

Table B.5. Toxicity Values and Information Used in PRG Derivation

Chemical Abstract Number	Analyte	Oral Slope Factor (SFO)	Sfo Ref	Absorbed Dose Slope Factor (SFd)	Inhalation Unit Risk (IUR)	IUR Ref	Oral RfD (RfDo)	RfDo Ref	Absorbed Dose (RfDd)	Inhalation (RfCi)	RfCi Ref
7429905	Aluminum						1.00E+00	P	1.00E+00	5.00E-03	P
7440360	Antimony (metallic)						4.00E-04	I	6.00E-05		
7440382	Arsenic, Inorganic	1.50E+00	I	1.50E+00	4.30E-06	I	3.00E-04	I	3.00E-04	1.50E-05	C
7440393	Barium						2.00E-01	I	1.40E-02	5.00E-04	H
7440417	Beryllium and compounds				2.40E-06	I	2.00E-03	I	1.40E-05	2.00E-05	I
7440428	Boron And Borates Only						2.00E-01	I	2.00E-01	2.00E-02	H
7440439	Cadmium (Diet)				1.80E-06	I	1.00E-03	I	2.50E-05	1.00E-05	A
7440439	Cadmium (Water)				1.80E-06	I	5.00E-04	I	2.50E-05	2.00E-05	C
7440473	Chromium (Total) ^a				9.00E-06	P	1.50E+00		1.95E-02		
16065831	Chromium(III), Insoluble Salts						1.50E+00	I	1.95E-02		
18540299	Chromium(VI)	5.00E-01	J	2.00E+01	8.40E-05	S	3.00E-03	I	7.50E-05	1.00E-04	I
7440484	Cobalt				9.00E-06	P	3.00E-04	P	3.00E-04	6.00E-06	P
7440508	Copper						4.00E-02	H	4.00E-02		
7439896	Iron						7.00E-01	P	7.00E-01		
7439965	Manganese (Diet)						1.40E-01	I	1.40E-01	5.00E-05	I
7439965	Manganese (Non-diet)						2.40E-02	S	9.60E-04	5.00E-05	I
7439976	Mercury, Inorganic Salts						3.00E-04	I	2.10E-05	3.00E-04	S
7439987	Molybdenum						5.00E-03	I	5.00E-03		
7440020	Nickel Soluble Salts				2.60E-07	C	2.00E-02	I	8.00E-04	9.00E-05	A
7782492	Selenium						5.00E-03	I	5.00E-03	2.00E-02	C
7440224	Silver						5.00E-03	I	2.00E-04		
7440280	Thallium (Soluble Salts)						1.00E-05	X	1.00E-05		
N/A	Uranium (Soluble Salts)						3.00E-03	I	3.00E-03	4.00E-05	A
N/A	Vanadium and Compounds						5.04E-03	S	5.04E-03	1.00E-04	A
7440666	Zinc and Compounds						3.00E-01	I	3.00E-01		
83329	Acenaphthene						6.00E-02	I	6.00E-02		
208968	Acenaphthylene ^b						6.00E-02	I	6.00E-02		
107131	Acrylonitrile	5.40E-01	I	5.40E-01	6.80E-08	I	4.00E-02	A	4.00E-02	2.00E-03	I
120127	Anthracene						3.00E-01	I	3.00E-01		
71432	Benzene	5.50E-02	I	5.50E-02	7.80E-09	I	4.00E-03	I	4.00E-03	3.00E-02	I
86748	Carbazole	2.00E-02	H	2.00E-02							
56235	Carbon Tetrachloride	7.00E-02	I	7.00E-02	6.00E-09	I	4.00E-03	I	4.00E-03	1.00E-01	I
67663	Chloroform	3.10E-02	C	3.10E-02	2.30E-08	I	1.00E-02	I	1.00E-02	9.77E-02	A
75354	Dichloroethylene, 1,1-						5.00E-02	I	5.00E-02	2.00E-01	I
540590	Dichloroethylene, 1,2- (Mixed Isomers)						9.00E-03	H	9.00E-03		
156592	Dichloroethylene, 1,2- <i>cis</i> -						2.00E-03	I	2.00E-03		
156605	Dichloroethylene, 1,2- <i>trans</i> -						2.00E-02	I	2.00E-02	6.00E-02	P
60571	Dieldrin	1.60E+01	I	1.60E+01	4.60E-06	I	5.00E-05	I	5.00E-05		
1746016	Dioxins/Furans, Total (as TCDD)	1.30E+05	C	1.30E+05	3.80E-02	C	7.00E-10	I	7.00E-10	4.00E-08	C
37871004	~HpCDD, 2,3,7,8-	1.30E+03	W	1.30E+03	3.80E-04	W	7.00E-08	W		4.00E-06	W
38998753	~HpCDF, 2,3,7,8-	1.30E+03	W	1.30E+03	3.80E-04	W	7.00E-08	W		4.00E-06	W
34465468	~HxCDD, 2,3,7,8-	1.30E+04	W	1.30E+04	3.80E-03	W	7.00E-09	W		4.00E-07	W

Table B.5. Toxicity Values and Information Used in PRG Derivation (Continued)

Chemical Abstract Number	Analyte	Oral Slope Factor (SfO)	Sfo Ref	Absorbed Dose Slope Factor (SFd)	Inhalation Unit Risk (IUR)	IUR Ref	Oral RfD (RfDo)	RfDo Ref	Absorbed Dose (RfDd)	Inhalation (RfCi)	RfCi Ref
55684941	~HxCDF, 2,3,7,8-	1.30E+04	W	1.30E+04	3.80E-03	W	7.00E-09	W		4.00E-07	W
3268879	~OCDD	3.90E+01	W	3.90E+01	1.14E-05	W	2.33E-06	W	3.33E-06	1.33E-04	W
39001020	~OCDF	3.90E+01	W	3.90E+01	1.14E-05	W	2.33E-06	W	3.33E-06	1.33E-04	W
36088229	~PeCDD, 2,3,7,8-	1.30E+05	W	1.30E+05	3.80E-02	W	7.00E-10	W	1.00E-09	4.00E-08	W
57117416	~PeCDF, 1,2,3,7,8-	3.90E+03	W	3.90E+03	1.14E-03	W	2.33E-08	W	3.33E-08	1.33E-06	W
57117314	~PeCDF, 2,3,4,7,8-	3.90E+04	W	3.90E+04	1.14E-02	W	2.33E-09	W	3.33E-09	1.33E-07	W
1746016	~TCDD, 2,3,7,8-	1.30E+05	C	1.30E+05	3.80E-02	C	7.00E-10	I	7.00E-10	4.00E-08	C
51207319	~TCDF, 2,3,7,8-	1.30E+04	W	1.30E+04	3.80E-03	W	7.00E-09	W		4.00E-07	W
100414	Ethylbenzene	1.10E-02	C	1.10E-02	2.50E-09	C	1.00E-01	I	1.00E-01	1.00E+00	I
206440	Fluoranthene						4.00E-02	I	4.00E-02		
86737	Fluorene						4.00E-02	I	4.00E-02		
118741	Hexachlorobenzene	1.60E+00	I	1.60E+00	4.60E-07	I	8.00E-04	I	8.00E-04		
91203	Naphthalene				3.40E-08	C	2.00E-02	I	2.00E-02	3.00E-03	I
88744	Nitroaniline, 2-						1.00E-02	X	1.00E-02	5.00E-05	X
621647	Nitroso-di-N-propylamine, N-	7.00E+00	I	7.00E+00	2.00E-06	C					
85018	Phenanthrene ^c						4.00E-02	I	4.00E-02		
1336363	Polychlorinated Biphenyls (high risk)	2.00E+00	I	2.00E+00	5.71E-07	I					
12674112	~Aroclor 1016	7.00E-02	S	7.00E-02	2.00E-08	S	7.00E-05	I	7.00E-05		
11104282	~Aroclor 1221	2.00E+00	S	2.00E+00	5.71E-07	S					
11141165	~Aroclor 1232	2.00E+00	S	2.00E+00	5.71E-07	S					
53469219	~Aroclor 1242	2.00E+00	S	2.00E+00	5.71E-07	S					
12672296	~Aroclor 1248	2.00E+00	S	2.00E+00	5.71E-07	S					
11097691	~Aroclor 1254	2.00E+00	S	2.00E+00	5.71E-07	S	2.00E-05	I	2.00E-05		
11096825	~Aroclor 1260	2.00E+00	S	2.00E+00	5.71E-07	S					
50328	Polycyclic aromatic hydrocarbons (cPAH), Total Carcinogenic (as BaP)	7.30E+00	I	7.30E+00	1.10E-06	C					
56553	~Benz[a]anthracene	7.30E-01	E	7.30E-01	1.10E-07	C					
50328	~Benzo[a]pyrene	7.30E+00	I	7.30E+00	1.10E-06	C					
205992	~Benzo[b]fluoranthene	7.30E-01	E	7.30E-01	1.10E-07	C					
207089	~Benzo[k]fluoranthene	7.30E-02	E	7.30E-02	1.10E-07	C					
218019	~Chrysene	7.30E-03	E	7.30E-03	1.10E-08	C					
53703	~Dibenz[a,h]anthracene	7.30E+00	E	7.30E+00	1.20E-06	C					
193395	~Indeno[1,2,3-cd]pyrene	7.30E-01	E	7.30E-01	1.10E-07	C					
129000	Pyrene						3.00E-02	I	3.00E-02		
127184	Tetrachloroethylene	2.10E-03	I	2.10E-03	2.60E-10	I	6.00E-03	I	6.00E-03	4.00E-02	I
79016	Trichloroethylene	4.60E-02	I	4.60E-02	4.10E-09	I	5.00E-04	I	5.00E-04	2.00E-03	I
75014	Vinyl Chloride	7.20E-01	I	7.20E-01	4.40E-09	I	3.00E-03	I	3.00E-03	1.00E-01	I
108383	Xylene, m-						2.00E-01	S	2.00E-01	1.00E-01	S
1330207	Xylene, Mixture						2.00E-01	I	2.00E-01	1.00E-01	I
95476	Xylene, o-						2.00E-01	S	2.00E-01	1.00E-01	S
106423	Xylene, P-						2.00E-01	S	2.00E-01	1.00E-01	S

Table B.5. Toxicity Values and Information Used in PRG Derivation (Continued)

Analyte	Volatile Organic?	Mutagen?	GI Absorption Factor (Unitless)	EPA ABS (Unitless)	ABS Ref	PEF Res.	PEF Ind./Comm.	VF Res.	VF Ind./Comm.	KY ABS (Unitless)	Permeability Constant	Perm. Const. Ref
Aluminum			1.00E+00		RAGSE	1.36E+09	1.36E+09			5.00E-02	1.00E-03	RAGSE
Antimony (metallic)			1.50E-01		RAGSE	1.36E+09	1.36E+09			5.00E-02	1.00E-03	RAGSE
Arsenic, Inorganic			1.00E+00	3.00E-02	RAGSE	1.36E+09	1.36E+09			3.00E-02	1.00E-03	RAGSE
Barium			7.00E-02		RAGSE	1.36E+09	1.36E+09			5.00E-02	1.00E-03	RAGSE
Beryllium and compounds			7.00E-03		RAGSE	1.36E+09	1.36E+09			7.00E-03	1.00E-03	RAGSE
Boron And Borates Only			1.00E+00		RAGSE	1.36E+09	1.36E+09			5.00E-02	1.00E-03	RAGSE
Cadmium (Diet)			2.50E-02	1.00E-03	RAGSE	1.36E+09	1.36E+09			1.00E-03	1.00E-03	RAGSE
Cadmium (Water)			5.00E-02	1.00E-03	RAGSE					1.00E-03	1.00E-03	RAGSE
Chromium (Total) ^d			1.30E-02		RAGSE	1.36E+09	1.36E+09			1.30E-02	1.00E-03	RAGSE
Chromium(III), Insoluble Salts			1.30E-02		RAGSE	1.36E+09	1.36E+09			1.30E-02	1.00E-03	RAGSE
Chromium(VI)		YES	2.50E-02		RAGSE	1.36E+09	1.36E+09			2.50E-02	2.00E-03	RAGSE
Cobalt			1.00E+00		RAGSE	1.36E+09	1.36E+09			5.00E-02	4.00E-04	RAGSE
Copper			1.00E+00		RAGSE	1.36E+09	1.36E+09			5.00E-02	1.00E-03	RAGSE
Iron			1.00E+00		RAGSE	1.36E+09	1.36E+09			5.00E-02	1.00E-03	RAGSE
Manganese (Diet)			1.00E+00		RAGSE					5.00E-02	1.00E-03	RAGSE
Manganese (Non-diet)			4.00E-02		RAGSE	1.36E+09	1.36E+09			4.00E-02	1.00E-03	RAGSE
Mercury, Inorganic Salts			7.00E-02		RAGSE	1.36E+09	1.36E+09			5.00E-02	1.00E-03	RAGSE
Molybdenum			1.00E+00		RAGSE	1.36E+09	1.36E+09			5.00E-02	1.00E-03	RAGSE
Nickel Soluble Salts			4.00E-02		RAGSE	1.36E+09	1.36E+09			4.00E-02	2.00E-04	RAGSE
Selenium			1.00E+00		RAGSE	1.36E+09	1.36E+09			5.00E-02	1.00E-03	RAGSE
Silver			4.00E-02		RAGSE	1.36E+09	1.36E+09			4.00E-02	6.00E-04	RAGSE
Thallium (Soluble Salts)			1.00E+00		RAGSE	1.36E+09	1.36E+09			5.00E-02	1.00E-03	RAGSE
Uranium (Soluble Salts)			1.00E+00		RAGSE	1.36E+09	1.36E+09			5.00E-02	1.00E-03	RAGSE
Vanadium and Compounds			2.60E-02		RAGSE	1.36E+09	1.36E+09			5.00E-02	1.00E-03	RAGSE
Zinc and Compounds			1.00E+00		RAGSE	1.36E+09	1.36E+09			5.00E-02	6.00E-04	RAGSE
Acenaphthene	YES		1.00E+00	1.30E-01	RAGSE	1.36E+09	1.36E+09	1.51E+05	1.51E+05	1.30E-01	8.60E-02	EPI
Acenaphthylene ^b	YES		1.00E+00	1.30E-01	RAGSE	1.36E+09	1.36E+09	1.51E+05	1.51E+05	2.50E-01	9.11E-02	EPI
Acrylonitrile	YES		1.00E+00		RAGSE	1.36E+09	1.36E+09	8.27E+03	8.27E+03	2.50E-01	1.16E-03	EPI
Anthracene	YES		1.00E+00	1.30E-01	RAGSE	1.36E+09	1.36E+09	5.63E+05	5.63E+05	1.30E-01	1.42E-01	EPI
Benzene	YES		1.00E+00		RAGSE	1.36E+09	1.36E+09	3.81E+03	3.81E+03	2.50E-01	1.49E-02	EPI
Carbazole			1.00E+00	1.00E-01	RAGSE					1.00E-01	5.36E-02	EPI
Carbon Tetrachloride	YES		1.00E+00		RAGSE	1.36E+09	1.36E+09	1.61E+03	1.61E+03	2.50E-01	1.63E-02	EPI
Chloroform	YES		1.00E+00		RAGSE	1.36E+09	1.36E+09	2.83E+03	2.83E+03	2.50E-01	6.83E-03	EPI
Dichloroethylene, 1,1-	YES		1.00E+00		RAGSE	1.36E+09	1.36E+09	1.24E+03	1.24E+03	2.50E-01	1.17E-02	EPI
Dichloroethylene, 1,2- (Mixed Isomers)	YES		1.00E+00		RAGSE	1.36E+09	1.36E+09	2.70E+03	2.70E+03	2.50E-01	1.10E-02	EPI
Dichloroethylene, 1,2- <i>cis</i> -	YES		1.00E+00		RAGSE	1.36E+09	1.36E+09	2.69E+03	2.69E+03	2.50E-01	1.10E-02	EPI
Dichloroethylene, 1,2- <i>trans</i> -	YES		1.00E+00		RAGSE	1.36E+09	1.36E+09	2.70E+03	2.70E+03	2.50E-01	1.10E-02	EPI
Dieldrin			1.00E+00	1.00E-01	RAGSE	1.36E+09	1.36E+09			1.00E-01	3.26E-02	EPI
Dioxins/Furans, Total (as TCDD)			1.00E+00	3.00E-02	RAGSE	1.36E+09	1.36E+09			3.00E-02	8.08E-01	EPI
~HpCDD, 2,3,7,8-			1.00E+00	3.00E-02	RAGSE						1.33E+00	EPI
~HpCDF, 2,3,7,8-			1.00E+00	1.00E-01	RAGSE						1.45E+00	EPI
~HxCDD, 2,3,7,8-			1.00E+00	3.00E-02	RAGSE						2.86E+00	EPI

Table B.5. Toxicity Values and Information Used in PRG Derivation (Continued)

Analyte	Volatile Organic?	Mutagen?	GI Absorption Factor (Unitless)	EPA ABS (Unitless)	ABS Ref	PEF Res.	PEF Ind./ Comm.	VF Res.	VF Ind./ Comm.	KY ABS (Unitless)	Permeability Constant	Perm. Const. Ref
~HxCDF, 2,3,7,8-			1.00E+00	1.00E-01	RAGSE						1.35E+00	EPI
~OCDD			1.00E+00	3.00E-02	RAGSE					3.00E-02	1.16E+00	EPI
~OCDF			1.00E+00	1.00E-01	RAGSE					1.00E-01	2.63E+00	EPI
~PeCDD, 2,3,7,8-			1.00E+00	3.00E-02	RAGSE					3.00E-02	2.41E-01	EPI
~PeCDF, 1,2,3,7,8-			1.00E+00	1.00E-01	RAGSE					1.00E-01	6.27E-01	EPI
~PeCDF, 2,3,4,7,8-			1.00E+00	1.00E-01	RAGSE					1.00E-01	6.27E-01	EPI
~TCDD, 2,3,7,8-			1.00E+00	3.00E-02	RAGSE	1.36E+09	1.36E+09			3.00E-02	8.08E-01	EPI
~TCDF, 2,3,7,8-			1.00E+00	1.00E-01	RAGSE					1.00E-01	6.57E-01	EPI
Ethylbenzene	YES		1.00E+00		RAGSE	1.36E+09	1.36E+09	6.10E+03	6.10E+03	2.50E-01	4.93E-02	EPI
Fluoranthene			1.00E+00	1.30E-01	RAGSE	1.36E+09	1.36E+09			1.30E-01	3.08E-01	EPI
Fluorene	YES		1.00E+00	1.30E-01	RAGSE	1.36E+09	1.36E+09	3.03E+05	3.03E+05	1.30E-01	1.10E-01	EPI
Hexachlorobenzene			1.00E+00	1.00E-01	RAGSE	1.36E+09	1.36E+09			1.00E-01	2.54E-01	EPI
Naphthalene	YES		1.00E+00	1.30E-01	RAGSE	1.36E+09	1.36E+09	4.99E+04	4.99E+04	1.30E-01	4.66E-02	EPI
Nitroaniline, 2-			1.00E+00	1.00E-01	RAGSE	1.36E+09	1.36E+09			1.00E-01	4.46E-03	EPI
Nitroso-di-N-propylamine, N-			1.00E+00	1.00E-01	RAGSE	1.36E+09	1.36E+09			1.00E-01	2.33E-03	EPI
Phenanthrene ^c			1.00E+00	1.30E-01	RAGSE	1.36E+09	1.36E+09			2.50E-01	1.44E-01	EPI
Polychlorinated Biphenyls (high risk)			1.00E+00	1.40E-01	RAGSE	1.36E+09	1.36E+09			1.40E-01	5.45E-01	EPI
~Aroclor 1016			1.00E+00	1.40E-01	RAGSE	1.36E+09	1.36E+09			1.40E-01	3.05E-01	EPI
~Aroclor 1221	YES		1.00E+00	1.40E-01	RAGSE	1.36E+09	1.36E+09	9.16E+04	9.16E+04	1.40E-01	1.68E-01	EPI
~Aroclor 1232	YES		1.00E+00	1.40E-01	RAGSE	1.36E+09	1.36E+09	9.16E+04	9.16E+04	1.40E-01	1.68E-01	EPI
~Aroclor 1242			1.00E+00	1.40E-01	RAGSE	1.36E+09	1.36E+09			1.40E-01	5.45E-01	EPI
~Aroclor 1248			1.00E+00	1.40E-01	RAGSE	1.36E+09	1.36E+09			1.40E-01	4.75E-01	EPI
~Aroclor 1254			1.00E+00	1.40E-01	RAGSE	1.36E+09	1.36E+09			1.40E-01	7.51E-01	EPI
~Aroclor 1260			1.00E+00	1.40E-01	RAGSE	1.36E+09	1.36E+09			1.40E-01	9.86E-01	EPI
Polycyclic aromatic hydrocarbons (cPAH), Total Carcinogenic (as BaP)		YES	1.00E+00	1.30E-01	RAGSE	1.36E+09	1.36E+09			1.30E-01	7.13E-01	EPI
~Benz[a]anthracene		YES	1.00E+00	1.30E-01	RAGSE	1.36E+09	1.36E+09			1.30E-01	5.52E-01	EPI
~Benzo[a]pyrene		YES	1.00E+00	1.30E-01	RAGSE	1.36E+09	1.36E+09			1.30E-01	7.13E-01	EPI
~Benzo[b]fluoranthene		YES	1.00E+00	1.30E-01	RAGSE	1.36E+09	1.36E+09			1.30E-01	4.17E-01	EPI
~Benzo[k]fluoranthene		YES	1.00E+00	1.30E-01	RAGSE	1.36E+09	1.36E+09			1.30E-01	6.91E-01	EPI
~Chrysene		YES	1.00E+00	1.30E-01	RAGSE	1.36E+09	1.36E+09			1.30E-01	5.96E-01	EPI
~Dibenz[a,h]anthracene		YES	1.00E+00	1.30E-01	RAGSE	1.36E+09	1.36E+09			1.30E-01	9.53E-01	EPI
~Indeno[1,2,3-cd]pyrene		YES	1.00E+00	1.30E-01	RAGSE	1.36E+09	1.36E+09			1.30E-01	1.04E+00	RAGSE
Pyrene	YES		1.00E+00	1.30E-01	RAGSE	1.36E+09	1.36E+09	2.56E+06	2.56E+06	1.30E-01	2.01E-01	EPI
Tetrachloroethylene	YES		1.00E+00		RAGSE	1.36E+09	1.36E+09	2.53E+03	2.53E+03	2.50E-01	3.34E-02	EPI
Trichloroethylene	YES	YES	1.00E+00		RAGSE	1.36E+09	1.36E+09	2.38E+03	2.38E+03	2.50E-01	1.16E-02	EPI
Vinyl Chloride	YES	YES	1.00E+00		RAGSE	1.36E+09	1.36E+09	1.03E+03	1.03E+03	2.50E-01	8.38E-03	EPI
Xylene, m-	YES		1.00E+00		RAGSE	1.36E+09	1.36E+09	5.89E+03	5.89E+03	2.50E-01	5.32E-02	EPI
Xylene, Mixture	YES		1.00E+00		RAGSE	1.36E+09	1.36E+09	6.27E+03	6.27E+03	2.50E-01	5.00E-02	EPI
Xylene, o-	YES		1.00E+00		RAGSE	1.36E+09	1.36E+09	6.95E+03	6.95E+03	2.50E-01	4.71E-02	EPI
Xylene, P-	YES		1.00E+00		RAGSE	1.36E+09	1.36E+09	6.01E+03	6.01E+03	2.50E-01	4.93E-02	EPI

Table B.5. Toxicity Values and Information Used in PRG Derivation (Continued)

Chemical Abstract Number	Analyte	Inhalation Slope Factor (SF _i)	SF _i Ref	Oral Slope Factor for Water (SF _{ow})	SF _{ow} Ref	Oral Slope Factor for Soil (SF _{os})	SF _{os} Ref	Oral Slope Factor for Food (SF _{of})	External Exposure Slope Factor (SF _e)	SF _e Ref	GI Absorption Factor (Unitless)	GI Abs. Ref
14596102	Am-241	2.81E-08	H	1.04E-10	H	2.17E-10	H	1.34E-10	2.76E-08	FGR12	5.00E-04	H
10045973	Cs-137+D	1.19E-11	H	3.04E-11	H	4.33E-11	H	3.74E-11	2.54E-06	FGR12	1.00E+00	H
13994202	Np-237+D	1.77E-08	H	6.74E-11	H	1.62E-10	H	9.10E-11	7.96E-07	FGR12	5.00E-04	H
13981163	Pu-238	3.36E-08	H	1.31E-10	H	2.72E-10	H	1.69E-10	7.22E-11	FGR12	5.00E-04	H
15117483	Pu-239	3.33E-08	H	1.35E-10	H	2.76E-10	H	1.74E-10	2.00E-10	FGR12	5.00E-04	H
14119336	Pu-240	3.33E-08	H	1.35E-10	H	2.77E-10	H	1.74E-10	6.98E-11	FGR12	5.00E-04	H
14133767	Tc-99	1.41E-11	H	2.75E-12	H	7.66E-12	H	4.00E-12	8.14E-11	FGR12	5.00E-01	H
14269637	Th-230	2.85E-08	H	9.10E-11	H	2.02E-10	H	1.19E-10	8.19E-10	FGR12	5.00E-04	H
13966295	U-234	1.14E-08	H	7.07E-11	H	1.58E-10	H	9.55E-11	2.52E-10	FGR12	2.00E-02	H
15117961	U-235+D	1.01E-08	H	7.18E-11	H	1.63E-10	H	9.76E-11	5.19E-07	FGR12	2.00E-02	H
7440611	U-238+D	9.35E-09	H	8.71E-11	H	2.10E-10	H	1.21E-10	1.14E-07	FGR12	2.00E-02	H

Information compiled August 2012.

Note that the toxicity values and information is presented in a split table format.

^a Values for Chromium (Total) use toxicity factors for Chromium III and inhalation unit cancer risk for Chromium VI, consistent with Screening Level note 9b (Appendix A).

^b Values for Acenaphthylene, if not available use toxicity factors for Acenaphthene.

^c Values for Phenanthrene, if not available use toxicity factors for Fluoranthene.

Reference Codes:

- A Agency for Toxic Substances and Disease Registry (ATSDR) minimal risk levels
- C The California EPA Office of Environmental Health Hazard Assessment's (OEHHA) Chronic Reference Exposure Levels (RELS) from December 18, 2008, and the Cancer Potency Values from July 21, 2009
- E Environmental Criteria and Assessment Office
- EPI EPA's Estimation Programs Interface Suite
- FGR12 Federal Guidance Report No. 12
- H HEAST
- I EPA's Integrated Risk Information System (IRIS)
- J New Jersey
- P The Provisional Peer Reviewed Toxicity Values (PPRTVs) derived by EPA's Superfund Health Risk Technical Support Center (STSC) for the EPA Superfund program.
- RAGSE Risk Assessment Guidance for Superfund, Part E.
- S Specific, see EPA's Regional Screening Level (RSL) User's Guide
- W World Health Organization
- X PPRTV Appendix

Notes on Table B.5.

Prior to using the values in this table, a risk assessor must be consulted to determine if any values need to be updated and to verify that the values are being used appropriately.

1. Information used to derive PRGs for COPCs at the PGDP is shown.
2. The "Oral RfD" is the chronic oral reference dose used for ingestion routes of exposure. The units for Oral RfD are mg/(kg x day).

Notes on Table B.5. (Continued)

3. The “Absorbed Dose RfD” calculated by multiplying the Oral RfD by the GI Absorption factor. The units for Absorbed Dose RfD are mg/(kg x day). This value is only applicable to chemical exposures.
4. The “Inhalation RfC” is the chronic inhalation concentration used for inhalation routes of exposure. The units for Inhalation RfC are mg/m³.
5. The “Oral Slope Factor” is the chronic oral slope factor used for the ingestion routes of exposure. The units on this value for chemicals is [mg/(kg x day)]⁻¹. The units on this value for radionuclides is (pCi)⁻¹.
6. “Oral Slope Factor for Water,” “Oral Slope Factor for Soil,” and “Oral Slope Factor for Food” are the indicated values for radionuclides. The units for these factors are (pCi)⁻¹.
7. The “Absorbed Dose Slope Factor” calculated by dividing the Oral Slope Factor by the GI Absorption factor. This value is only applicable to chemical exposures. The units for Absorbed Dose Slope Factor are [mg/(kg x day)]⁻¹.
8. The “Inhalation Unit Risk” is the chronic inhalation factor used for inhalation routes of exposure. The values listed for chemicals is in units of mg/m³, although they are typically expressed in µg/m³. For radionuclides, the inhalation slope factor continues to be used. The units on this value for radionuclides is (pCi)⁻¹.
9. The “External Exposure Slope Factor” is the slope factor used for external exposure to ionizing radiation emitted by radioactive chemicals. The units for external exposure slope factor are [(pCi x year)/g]⁻¹.
10. “Volatile Organic?” is a flag used to specify if the chemical should be assessed as a vapor.
11. The column labeled “Mutagen?” is a flag used to specify if the chemical should be assessed as a mutagen. This assessment is made only when PRGs are developed using the RAIS calculator.
12. The “Particle Emission Factor” is a value used to assess inhalation routes of exposure. The particle emission factor is in units of m³/kg. The values for residential and industrial/commercial scenario listed are taken from the 2002 *Kentucky Risk Assessment Guidance*.
13. The “Volatilization Factor” is a value used to assess inhalation routes of exposure. The volatilization factor is in units of m³/kg. As indicated in the 2002 *Kentucky Risk Assessment Guidance*, the chemical-specific values for residential and industrial/commercial scenario listed here are calculated using Equation (8) of the EPA’s *Soil Screening Level Guidance User’s Guide* (1996).
14. The “EPA ABS” is the dermal absorption value recommended by EPA Region 4 in their guidance material, 2004 *RAGs, Part E*. The dermal absorption value is unitless.
15. The “KY ABS” is the dermal absorption value recommended by the Commonwealth of Kentucky in their guidance material, 2002 *Kentucky Risk Assessment Guidance*. Dermal exposure to soil used default absorption values of 0.25 for volatiles, 0.1 for semivolatiles, and 0.05 for metals. The dermal absorption value is unitless.

In RAGS Part E, 2004, Exhibit 4-1, the following GI absorption efficiencies are listed that are below the 5% dermal absorption KDEP has recommended as a default value for inorganics. For these constituents, the dermal absorption value should be modified from 5% to mimic the GI absorption efficiencies, as follows: Beryllium 0.007 = 0.7%; Chromium III 0.013 = 1.3%; Chromium VI 0.025 = 2.5%; Manganese 0.04 = 4%; Nickel 0.04 = 4%; Silver 0.04 = 4%; Vanadium 0.026 = 2.6%

This is in addition to the chemical-specific dermal absorption fractions listed in RAGS, Part E, Exhibit 3-4, including: Arsenic 0.03 = 3% and Cadmium 0.001=0.1%

16. The “Permeability Constant” is a chemical-specific value used to estimate dermal absorption of chemicals in water. The permeability constant is in units of cm/hr.

For sites for which the concentration in soil exceeds the 400 mg/kg screening level, risks from lead should be analyzed using the Integrated Exposure Uptake Biokinetic (IEUBK) model. The model should be run using the EPA-recommended 10 µg/dl blood lead level cutoff and the site-specific values discussed in the next paragraph. The analysis of risks from lead also should show the probability of exceeding the recommended Commonwealth of Kentucky blood lead level of 2.5 µg/dl (note that this probability distribution can be developed in the IEUBK model from the previous model run by changing the cutoff value in the graph menu). The uncertainty section of the risk assessment should include text indicating that there is no safe level of lead exposure to children and comparing the risks predicted by the IEUBK analyses based on the two cutoff values.

Table B.6 includes parameters that can be used in the IEUBK model to develop more site-specific screening levels for lead. The IEUBK model calculates a blood lead level that includes the contribution from off-site sources such as food in lead and water. To make the model more site-specific, the updated nationwide averages for lead in food can be used in place of the default values in the model. In addition, if regional or site-specific concentrations of lead in food and water are available, the concentration of lead in water can be changed in the model to that value. The PGDP mean value for lead in surface soil from DOE 1995 (17 mg/kg) and the value for lead in RGA groundwater from Appendix A, Table A.13 (0.129 mg/L) should be used in place of the model default value.

Table B.6. Soil Parameters for VF Calculations

Parameter	Definition (units)		Default
Q/C	Inverse the mean conc. at the center of a 0.5-acres square source (g/m ² -s per kg/m ³)	res.	64.177
		ind./com.	43.07
T	Exposure interval (s)		9.50E+08
ρ _b	Dry soil bulk density (g/cm ³)		1.5
θ _a	Air filled soil porosity (L _{air} /L _{soil})		0.28
n	Total soil porosity (L _{pore} /L _{soil})		0.43
θ _w	Water-filled soil porosity (L _{water} /L _{soil})		0.15

Information compiled February 2011.

The revised diet values for the model are available at <http://www.epa.gov/superfund/health/contaminants/lead/ieubkfaq.htm>.

For recreational exposures, the time on-site versus the total time spent outdoors can be included in the model. The model allows only one soil concentration to be entered, but the exposure to on and off-site soil can be incorporated by weighting the soil concentration by the proportion of time spent on and off-site. This method and its limitations are described fully in Appendix A of EPA's review of the human health risk assessment for the Couer d'Alene basin (EPA 2000).

For industrial or outdoor worker scenarios, the Adult Lead Model is used to develop a PRG for soil. This model includes a default blood lead level based on the NHANES survey value for the western United States for all races combined, other measured adult blood lead concentrations from state or regional databases may be used in place of the default value if such values are available. The default soil ingestion value of 50 mg/kg can also be altered if there is a reliable basis for substituting a site-specific value.

Volatilization parameters are provided in Table B.7.

Table B.7. Volatilization Parameters

CAS #	Chemical	D _i (cm ² /s) from RAIS	D _i Ref in RAIS	D _w (cm ² /s) from RAIS	D _w Ref in RAIS	Unitless H' from RAIS	H Ref in RAIS
83329	Acenaphthene	5.06E-02	USEPA 2001	8.33E-06	USEPA 2001	7.52E-03	EPI HenryWin v3.2
208968	Acenaphthylene	4.50E-02	USEPA 2001	6.98E-06	USEPA 2001	4.66E-03	EPI HenryWin v3.2
107131	Acrylonitrile	1.14E-01	USEPA 2001	1.23E-05	USEPA 2001	5.64E-03	EPI HenryWin v3.2
120127	Anthracene	3.90E-02	USEPA 2001	7.85E-06	USEPA 2001	2.27E-03	EPI HenryWin v3.2
71432	Benzene	8.95E-02	USEPA 2001	1.03E-05	USEPA 2001	2.27E-01	EPI HenryWin v3.2
86748	Carbazole	4.17E-02	USEPA 2001	7.45E-06	USEPA 2001	4.74E-06	EPI HenryWin v3.2
56235	Carbon Tetrachloride	5.71E-02	USEPA 2001	9.78E-06	USEPA 2001	1.13E+00	EPI HenryWin v3.2
67663	Chloroform	7.69E-02	USEPA 2001	1.09E-05	USEPA 2001	1.50E-01	EPI HenryWin v3.2
1746016	Dioxins/Furans (Total) (as TCDD)	4.70E-02	USEPA 2001	4.73E-06	USEPA 2001	2.04E-03	EPI
75354	Dichloroethylene, 1,1-	8.63E-02	USEPA 2001	1.10E-05	USEPA 2001	1.07E+00	EPI HenryWin v3.2
540590	Dichloroethylene, 1,2- (Mixed Isomers)	8.97E-02	USEPA 2001	1.12E-05	USEPA 2001	1.67E-01	EPI HenryWin v3.2
156592	Dichloroethylene, 1,2- <i>cis</i> -	8.84E-02	USEPA 2001	1.13E-05	USEPA 2001	1.67E-01	EPI HenryWin v3.2
156605	Dichloroethylene, 1,2- <i>trans</i> -	8.76E-02	USEPA 2001	1.12E-05	USEPA 2001	1.67E-01	EPI HenryWin v3.2
60571	Dieldrin	2.33E-02	USEPA 2001	6.01E-06	USEPA 2001	4.09E-04	EPI HenryWin v3.2
100414	Ethylbenzene	6.85E-02	USEPA 2001	8.46E-06	USEPA 2001	3.22E-01	EPI HenryWin v3.2
206440	Fluoranthene	2.76E-02	USEPA 2001	7.18E-06	USEPA 2001	3.62E-04	EPI HenryWin v3.2
86737	Fluorene	4.40E-02	USEPA 2001	7.89E-06	USEPA 2001	3.93E-03	EPI HenryWin v3.2
118741	Hexachlorobenzene	2.90E-02	USEPA 2001	7.85E-06	USEPA 2001	6.95E-02	EPI HenryWin v3.2
91203	Naphthalene	6.05E-02	USEPA 2001	8.38E-06	USEPA 2001	1.80E-02	EPI HenryWin v3.2
88744	Nitroaniline, 2-	5.19E-02	USEPA 2001	7.41E-06	USEPA 2001	2.41E-06	EPI HenryWin v3.2
621647	Nitroso-di-N-propylamine, N-	5.64E-02	USEPA 2001	7.76E-06	USEPA 2001	2.20E-04	EPI HenryWin v3.2
85018	Phenanthrene	3.45E-02	USEPA 2001	6.69E-06	USEPA 2001	1.73E-03	EPI HenryWin v3.2
1336363	Polychlorinated Biphenyls (high risk)	4.32E-02	USEPA 2001	5.04E-06	USEPA 2001	7.77E-03	EPI HenryWin v3.2
12674112	~Aroclor 1016	4.69E-02	USEPA 2001	5.48E-06	USEPA 2001	8.18E-03	EPI
11104282	~Aroclor 1221	5.78E-02	USEPA 2001	6.75E-06	USEPA 2001	3.01E-02	EPI HenryWin v3.2
11141165	~Aroclor 1232	5.78E-02	USEPA 2001	6.75E-06	USEPA 2001	3.01E-02	EPI HenryWin v3.4
53469219	~Aroclor 1242	4.32E-02	USEPA 2001	5.04E-06	USEPA 2001	7.77E-03	EPI
12672296	~Aroclor 1248	4.32E-02	USEPA 2011	5.04E-06	USEPA 2001	1.80E-02	EPI
11097691	~Aroclor 1254	4.01E-02	USEPA 2001	4.68E-06	USEPA 2001	1.16E-02	EPI
11096825	~Aroclor 1260	3.53E-02	USEPA 2001	4.12E-06	USEPA 2001	1.37E-02	EPI
	Polycyclic aromatic hydrocarbons (cPAH)						
56553	~Benz[a]anthracene	5.09E-02	USEPA 2001	5.94E-06	USEPA 2001	4.91E-04	EPI HenryWin v3.2
50328	~Benzo[a]pyrene	4.76E-02	USEPA 2001	5.56E-06	USEPA 2001	1.87E-05	EPI HenryWin v3.2
205992	~Benzo[b]fluoranthene	4.76E-02	USEPA 2001	5.56E-06	USEPA 2001	2.69E-05	EPI HenryWin v3.2
207089	~Benzo[k]fluoranthene	4.76E-02	USEPA 2001	5.56E-06	USEPA 2001	2.39E-05	EPI HenryWin v3.2
218019	~Chrysene	2.61E-02	USEPA 2001	6.75E-06	USEPA 2001	2.14E-04	EPI HenryWin v3.2
53703	~Dibenz[a,h]anthracene	4.46E-02	USEPA 2001	5.21E-06	USEPA 2001	5.76E-06	EPI HenryWin v3.2
193395	~Indeno[1,2,3-cd]pyrene	4.48E-02	USEPA 2001	5.23E-06	USEPA 2001	6.56E-05	SSL
129000	Pyrene	2.78E-02	USEPA 2001	7.25E-06	USEPA 2001	4.87E-04	EPI HenryWin v3.2
127184	Tetrachloroethylene	5.05E-02	USEPA 2001	9.46E-06	USEPA 2001	7.24E-01	EPI HenryWin v3.2
79016	Trichloroethylene	6.87E-02	USEPA 2001	1.02E-05	USEPA 2001	4.03E-01	EPI HenryWin v3.2
75014	Vinyl Chloride	1.07E-01	USEPA 2001	1.20E-05	USEPA 2001	1.14E+00	EPI HenryWin v3.2
1330207	Xylene, Mixture	8.47E-02	USEPA 2001	9.90E-06	USEPA 2001	2.12E-01	EPI HenryWin v3.2
106423	Xylene, P-	6.82E-02	USEPA 2001	8.42E-06	USEPA 2001	2.82E-01	EPI HenryWin v3.2
108383	Xylene, m-	6.84E-02	USEPA 2001	8.44E-06	USEPA 2001	2.94E-01	EPI HenryWin v3.2
95476	Xylene, o-	6.89E-02	USEPA 2001	8.53E-06	USEPA 2001	2.12E-01	EPI HenryWin v3.2

Table B.7. Volatilization Parameters (Continued)

CAS #	Chemical	K _{oc} (cm ³ /g) from RAIS	K _{oc} Ref in RAIS	K _d (cm ³ /g)* K _{oc} x 0.2%
83329	Acenaphthene	5.03E+03	EPI KOCWIN v2.0	1.01E+01
208968	Acenaphthylene	5.03E+03	EPI KOCWIN v2.0	1.01E+01
107131	Acrylonitrile	8.51E+00	EPI KOCWIN v2.0	1.70E-02
120127	Anthracene	1.64E+04	EPI KOCWIN v2.0	3.28E+01
71432	Benzene	1.46E+02	EPI KOCWIN v2.0	2.92E-01
86748	Carbazole	9.16E+03	EPI KOCWIN v2.0	1.83E+01
56235	Carbon Tetrachloride	4.39E+01	EPI KOCWIN v2.0	8.78E-02
67663	Chloroform	3.18E+01	EPI KOCWIN v2.0	6.36E-02
1746016	Dioxins/Furans (Total) (as TCDD)	2.49E+05	EPI	4.98E+02
75354	Dichloroethylene, 1,1-	3.18E+01	EPI KOCWIN v2.0	6.36E-02
540590	Dichloroethylene, 1,2- (Mixed Isomers)	3.96E+01	EPI KOCWIN v2.0	7.92E-02
156592	Dichloroethylene, 1,2- <i>cis</i> -	3.96E+01	EPI KOCWIN v2.0	7.92E-02
156605	Dichloroethylene, 1,2- <i>trans</i> -	3.96E+01	EPI KOCWIN v2.0	7.92E-02
60571	Dieldrin	2.01E+04	EPI KOCWIN v2.0	4.02E+01
100414	Ethylbenzene	4.46E+02	EPI KOCWIN v2.0	8.92E-01
206440	Fluoranthene	5.55E+04	EPI KOCWIN v2.0	1.11E+02
86737	Fluorene	9.16E+03	EPI KOCWIN v2.0	1.83E+01
118741	Hexachlorobenzene	6.20E+03	EPI KOCWIN v2.0	1.24E+01
91203	Naphthalene	1.54E+03	EPI KOCWIN v2.0	3.08E+00
88744	Nitroaniline, 2-	1.11E+02	EPI KOCWIN v2.0	2.22E-01
621647	Nitroso-di-N-propylamine, N-	2.75E+02	EPI KOCWIN v2.0	5.50E-01
85018	Phenanthrene	1.67E+04	EPI KOCWIN v2.0	3.34E+01
1336363	Polychlorinated Biphenyls (high risk)	7.81E+04	EPI KOCWIN v2.0	1.56E+02
12674112	~Aroclor 1016	4.77E+04	EPI KOCWIN v2.0	9.54E+01
11104282	~Aroclor 1221	8.40E+03	EPI KOCWIN v2.2	1.68E+01
11141165	~Aroclor 1232	8.40E+03	EPI KOCWIN v2.4	1.68E+01
53469219	~Aroclor 1242	7.81E+04	EPI KOCWIN v2.6	1.56E+02
12672296	~Aroclor 1248	7.65E+04	EPI	1.53E+02
11097691	~Aroclor 1254	1.31E+05	EPI KOCWIN v2.10	2.62E+02
11096825	~Aroclor 1260	3.50E+05	EPI KOCWIN v2.12	7.00E+02
	Polycyclic aromatic hydrocarbons (cPAH)			
56553	~Benz[a]anthracene	1.77E+05	EPI KOCWIN v2.0	3.54E+02
50328	~Benzo[a]pyrene	5.87E+05	EPI KOCWIN v2.0	1.17E+03
205992	~Benzo[b]fluoranthene	5.99E+05	EPI KOCWIN v2.0	1.20E+03
207089	~Benzo[k]fluoranthene	5.87E+05	EPI KOCWIN v2.0	1.17E+03
218019	~Chrysene	1.81E+05	EPI KOCWIN v2.0	3.62E+02
53703	~Dibenz[a,h]anthracene	1.91E+06	EPI KOCWIN v2.0	3.82E+03
193395	~Indeno[1,2,3-cd]pyrene	1.95E+06	EPI KOCWIN v2.0	3.90E+03
129000	Pyrene	5.43E+04	EPI KOCWIN v2.0	1.09E+02
127184	Tetrachloroethylene	9.49E+01	EPI KOCWIN v2.0	1.90E-01
79016	Trichloroethylene	6.07E+01	EPI KOCWIN v2.0	1.21E-01
75014	Vinyl Chloride	2.17E+01	EPI KOCWIN v2.0	4.34E-02
1330207	Xylene, Mixture	3.83E+02	EPI KOCWIN v2.0	7.66E-01
106423	Xylene, P-	3.75E+02	EPI KOCWIN v2.0	7.50E-01
108383	Xylene, m-	3.75E+02	EPI KOCWIN v2.0	7.50E-01
95476	Xylene, o-	3.83E+02	EPI KOCWIN v2.0	7.66E-01

* RAIS does not provide K_d values for organic chemicals, therefore, K_d values used in the calculation are taken from PRG Region 9 physical chemical data.

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APPENDIX C

OUTLINE FOR BASELINE HUMAN HEALTH RISK ASSESSMENTS

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OUTLINE FOR BASELINE HUMAN HEALTH RISK ASSESSMENTS

*** Although the following outline can be used for baseline human health risk assessments for both source units and integrator units, not all sections may be relevant to all assessments and additional sections may be needed for some assessments. However, all baseline risk assessments completed for PGDP should include each of the first and second level headers listed below.

*** The document should begin with an introduction that presents the scope and objectives of the baseline human health risk assessment. This should include a description of the general problem at the site and an overview of the design of the baseline human health risk assessment.

1. Results of Previous Studies

*** The section should begin with a brief summary of the previous studies that are relevant to the baseline human health risk assessment. All relevant previous risk evaluations should be summarized.

1.1 Study #1

1.2 Study #2

Etc.

2. Identification of Chemicals of Potential Concern

*** The section should begin with an introduction that describes the purpose of the section and the order in which the material is presented.

2.1 Sources of Data

*** The sources of all data should be listed, and the projects in which the data were collected should be described.

2.2 General Data Evaluation Considerations

*** The eight steps of data evaluation as applied to the baseline risk assessment should be discussed.

2.2.1 Evaluation of Sampling Design

2.2.2 Evaluation of Analytical Methods

2.2.3 Evaluation of Sample Quantitation Limits

2.2.4 Evaluation of Data Qualifiers and Codes

2.2.5 Elimination of Chemicals not Detected

2.2.6 Examination of Toxicity of Detected Analytes

2.2.7 Examination of Essential Nutrients

2.2.8 Comparison of Analyte Concentrations and Activities Detected in Site Samples to Background Concentrations

2.3 Risk Assessment Specific Data Evaluation

*** This section should discuss in detail how the eight steps were applied to identify the chemicals of potential concern under both current and future conditions.

2.3.1 Current Conditions

*** This section should discuss the evaluation of the data set.

2.3.2 Future Conditions

*** This section should discuss any modeling performed to address potential future changes in the identity or concentration of contaminants.

2.4 Evaluation of Data from Other Sources

*** The section should introduce any “special data,” especially nonnumeric data (such as activities of visitors at a site or types of vegetables grown by Kentucky residents) used to develop the exposure assessment that are not used quantitatively in the baseline human health risk assessment. Examples of special data that may be used are found in the survey forms and responses in Appendix E.

2.4.1 Other Source #1

2.4.2 Other Source #2

Etc.

2.5 Summary of Chemicals of Potential Concern

*** This section should present a summary of the quantitative data evaluation and its results.

3. Exposure Assessment

*** This section should begin with a description of the process used in exposure assessment, and the goal of the specific exposure assessment being performed.

3.1 Characterization of Exposure Setting

*** This section should describe either by reference or directly the following:

3.1.1 Surface Features

3.1.2 Meteorology

3.1.3 Geology

3.1.4 Demography and Land Use

3.1.5 Ecology

3.1.6 Hydrology

3.1.7 Hydrogeology

3.2 Identification of Exposure Pathways

*** This section should begin by describing what a pathway is and how a pathway can be complete or incomplete.

3.2.1 Land Use Considerations

*** The land use under current and expected and potential future conditions should be described.

3.2.2 Potential Receptor Populations

*** The potential receptors under both current and future uses should be described and justified.

3.2.3 Delineation of Exposure Points/Exposure Routes

*** All possible exposure routes should be presented and justified. The number of possible exposure routes should be reduced, if possible, so that only probable exposure routes with significant risk or hazard are quantified. The exposure equations used in the assessment to quantify exposure should be presented. Justification for not quantifying a possible route should be presented.

3.2.4 Development of Conceptual Site Models

*** Figures illustrating the pathways of exposure should be presented for each site under investigation. The model for each site should be justified.

3.3 Quantification of Exposure

*** The methods used to quantify exposure (i.e., estimate dose) should be described for each receptor. If modeling is used to determine concentration or activities of chemicals of potential concern in biota, the models should be presented.

3.4 Summary of Exposure Assessment

4. Toxicity Assessment

*** This section should begin by describing the goal and methods used for toxicity assessment. The source of all toxicity values should be discussed. Tables presenting the toxicity information should be presented.

4.1 Inorganics

*** The toxicity of each chemical of potential concern should be profiled. Each profile should include a listing of the carcinogenic and noncarcinogenic toxicity values used in the baseline human health risk assessment.

4.1.1 Chemical 1

4.1.2 Chemical 2

Etc.

4.2 Organics

*** The toxicity of each chemical of potential concern should be profiled. Each profile should include a listing of the toxicity values used in the baseline human health risk assessment.

4.2.1 Chemical 1

4.2.2 Chemical 2

Etc.

4.3 Radionuclides

*** The toxicity of each chemical of potential concern should be profiled. Each profile should include a listing of the toxicity values used in the baseline human health risk assessment.

4.3.1 Radionuclide 1

4.3.2 Radionuclide 2

Etc.

4.4 Chemicals for Which No EPA Toxicity Values Are Available

*** The chemicals of potential concern that fall in this class should be listed. If the baseline human health risk assessment is evaluating multiple units or areas, these chemicals should be listed by unit or area. This section should include the procedure for evaluating potential surrogate chemicals that may be available for some of the chemicals without toxicity values.

4.5 Uncertainties Related to Toxicity Assessment

*** A brief presentation of the uncertainties related to all toxicity assessments and toxicity values should be made.

4.6 Summary

*** The amount of toxicity information for the chemicals of potential concern should be discussed. If the baseline human health risk assessment is evaluating multiple units or areas, this information should be presented by unit or area.

5. Risk Characterization

*** The section should begin with a brief discussion of the purpose and goals of risk characterization and what will result from this step of the assessment.

5.1 Determination of Noncancer Effects

*** The methods used to quantify systemic toxicity for each chemical, both within and across pathways should be presented. If exposure over multiple scenarios or areas is possible, this should be noted.

5.2 Determination of Excess Cancer Risk

*** The methods used to quantify excess lifetime cancer risk for each chemical, both within and across pathways should be presented. If exposure over multiple scenarios or areas is possible, this should be noted.

5.3 Risk Characterization for Current Use Scenario(s)

*** Risk results for each unit or area should be presented in two-way tables and in a narrative summary. If subchronic effects are characterized, they should be presented separately from the chronic effects.

5.3.1 Systemic Toxicity

5.3.2 Excess Lifetime Cancer Risk

5.4 Risk Characterization for Future Use Scenario(s)

*** Risk results for each unit or area should be presented in two-way tables and in a narrative summary. If more than one future time is quantitatively evaluated, the results should be presented for each time period. If subchronic effects are characterized, they should be presented separately from the chronic effects.

5.4.1 Systemic Toxicity

5.4.2 Excess Lifetime Cancer Risk

5.5 Risk Characterization for Lead (if needed)

*** The special problems associated with risk characterization for lead should be discussed. Results from lead modeling and from comparisons against EPA and Kentucky screening values should be presented by unit or area.

5.6 Identification of Use Scenarios, Chemicals, Pathways, and Media of Concern

*** The section should begin with a listing of the rules used to identify use scenarios, chemicals, pathways and media of concern.

5.6.1 Use Scenarios of Concern

*** These should be listed within area or unit under investigation.

5.6.2 Chemicals of Concern

*** These should be listed within area or unit under investigation.

5.6.3 Pathways of Concern

*** These should be listed within area or unit under investigation.

5.6.4 Media of Concern

*** These should be listed within area or unit under investigation

5.7 Summary of Risk Characterization

*** This section should describe and present the risk characterization summary tables.

6. Uncertainty in the Risk Assessment

*** This section should begin with a general discussion of uncertainty. If a qualitative uncertainty analysis is being performed, “small,” “moderate,” and “large” uncertainties should be defined and the following subsections should be included. If a quantitative uncertainty analysis is being performed, the methods and results should be described in detail. Normally, a qualitative analysis, including sensitivity analyses, will be sufficient. Regardless, this section should continue with a discussion of each of the uncertainties affecting the major portions of the risk assessment. (Note, the uncertainties listed below are some of those found in past assessments. The uncertainties to be addressed in future assessments must be determined on a case-by-case basis.)

6.1 Uncertainties Associated with Data

*** The uncertainties to be discussed should be summarized in the introduction of this section. Categories of uncertainties to discuss are presented in the following.

6.1.1 Selection of Chemicals of Potential Concern

6.1.2 Determination of Exposure Point Concentrations—Current Conditions

6.1.3 Determination of Exposure Point Concentrations—Future Conditions

6.1.4 Use of Unfiltered versus Filtered Water Samples

6.2 Uncertainties Associated with Exposure Assessment

*** The uncertainties to be discussed should be summarized in the introduction of this section. Categories of uncertainties to discuss are presented in the following.

6.2.1 Uncertainties in Fate and Transport Modeling

6.2.2 Uncertainties in Use of Reasonable Maximum Exposure (RME) Scenarios

6.2.3 Uncertainties Related to Development of Conceptual Site Models

6.2.4 Uncertainties Related to Use of Default Values When Estimating Dermal Absorbed Dose

6.3 Uncertainties Associated with Toxicity Assessment

*** The uncertainties to be discussed should be summarized in the introduction of this section. Categories of uncertainties to discuss are presented in the following.

6.3.1 Uncertainties Due to Lack of Toxicity Values for Some Chemicals

6.3.2 Uncertainties in Deriving Toxicity Values

6.3.3 Uncertainties Due to Calculation of Absorbed Dose Toxicity Values from Administered Toxicity Values

6.3.4 Uncertainties Due to Use of Toxicity Values for Chronic Exposure for Subchronic Exposure Times

6.4 Uncertainties Associated with Risk Characterization

*** The uncertainties to be discussed should be summarized in the introduction of this section. Categories of uncertainties to discuss are presented in the following.

6.4.1 Uncertainties in Combining Chemical-Specific Risk and Hazard Estimates and Pathway-Specific Risk and Hazard Estimates

6.4.2 Uncertainties in Combining Risk Estimated for Chemical Exposure to those for Risk Estimated for Radioisotope Exposure

6.5 Summary of Uncertainties

*** This section should summarize the uncertainties discussed earlier in the section and present a table reviewing all uncertainties.

7. Conclusions and Summary

*** The purpose of this section is to review the results of the risk assessment without the use of tables and explanations and provide significant observations interpreting the results of the assessment for use by risk managers. When properly presented, it should be possible to insert this section as written into the feasibility study.

7.1 Chemicals of Potential Concern

*** A brief description of the screening process should be provided, and the chemicals of potential concern for each area or unit listed either by name (if the list is short) or by class.

7.2 Exposure Assessment

*** The exposure pathways quantitatively evaluated should be listed for each use scenario

7.3 Toxicity Assessment

*** The amount of available toxicity data for the chemicals of potential concern for each area should be listed. Chemicals of potential concern lacking toxicity values should be highlighted.

7.4 Risk Characterization

*** The use scenarios, chemicals, pathways, and media of concern should be listed for each area or unit, and the rules used to delineate the use scenarios, chemicals, pathways, and media of concern should be presented.

7.5 Observations

*** This section should integrate the risk estimates and the uncertainties to develop a list of salient issues to be considered by risk managers when making decisions in risk management documents. This includes a discussion for each of the chemicals of concern identified in the risk assessment. In addition, the results of the baseline human health risk assessment should be compared to results of previous risk evaluations, if any.

8 Remedial Goal Options

*** This section should present the methods used to derive the remedial goal options and list the remedial goal options for each chemical of concern. Because remedial goal options are medium- and scenario-specific, a separate list should be presented for each area (or unit), scenario, and medium combination.

8.1 Derivation of RGOs

*** This presentation should be as brief as possible.

8.2 Presentation of RGOs

*** These should be presented in tables. Very little narrative beyond directing the reader to the appropriate tables is needed.

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APPENDIX D
EXPOSURE EQUATIONS

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EXPOSURE EQUATIONS

This appendix contains the exposure equations used in environmental human health risk assessments for Department of Energy sites located at the Paducah Gaseous Diffusion Plant (PGDP). It should be noted that the equations shown in this appendix may not be the same as those used in preliminary remediation goal (PRG) calculations. PRG calculations were taken from the Risk Assessment Information System (RAIS) PRG calculator available at <http://rais.ornl.gov/>.

The equations in this appendix are consistent with all Region 4 U.S. Environmental Protection Agency (EPA) and Commonwealth of Kentucky guidance materials. However, the exposure parameters shown are those used to produce daily intakes and absorbed doses used to complete environmental risk assessments performed for PGDP only. These exposure parameters are for a default reasonable maximum exposure (RME). While these exposure parameters generally are consistent with the exposure parameters recommended by Region 4 EPA, they do differ in some cases where Kentucky Department for Environmental Protection (KDEP) values were used. The source of each value is provided below the equation. Equations to complete dose assessments and to derive dose conversion factors are not presented; however, these can be derived from the information provided here.

Chemical-specific values, except for those listed in Appendix B have not been updated for this Risk Methods Document. See Appendix D, Part 2 of *Methods for Conducting Risk Assessments and Risk Evaluations at the Paducah Gaseous Diffusion Plant Paducah, Kentucky Volume 1, Human Health*, DOE/LX/07-0107&D2/R1/V1, for the list (DOE 2011).

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Table D.1. Reasonable Maximum Exposure Assumptions for Ingestion of Water by a Rural Resident^a

Equations:

$$\text{Chemical Intake [mg/(kg} \times \text{day)]} = \frac{C_w \times IR \times EF \times ED}{BW \times AT}$$

$$\text{Radionuclide Intake (pCi)} = A_w \times IR \times EF \times ED$$

Parameter	Units	Value used	References^b
Chemical concentration in water = C_w	mg/L	Chemical-specific	-----
Radiological activity = A_w	pCi/L	Chemical-specific	-----
Ingestion Rate = IR	L/d	2 (adult) 1.5 (child)	[14]
Exposure frequency = EF	d/year	350	[14]
Exposure duration = ED	years	24 (adult) 6 (child)	[14]
Body weight = BW	kg	70 (adult) 15 (child)	[14]
Averaging time = AT	yr × day/yr.	70 × 365 (carcinogen) ED × 365 (noncarcinogen)	[14]

^a Equation from [1].

^b References (noted in brackets []) follow Table D.50.

Table D.2. Reasonable Maximum Exposure Assumptions for Inhalation of Volatile Organic Compounds in Water during Household Use (including Showering) by a Rural Resident^a

Equations:

$$\text{Exposure Concentration } (\mu\text{g}/\text{m}^3) = \frac{[(C_{\text{shower}} \times EF \times ET_{\text{shower}}) + (C_{\text{house}} \times EF \times ET_{\text{house}})] \times ED}{AT} \times CF$$

$$C_{\text{shower}} (\text{mg}/\text{m}^3) = \frac{[(C_{a\text{max}} / 2) \times t_1] + [C_{a\text{max}} \times t_2]}{t_1 + t_2} \qquad C_{a\text{max}} (\text{mg}/\text{m}^3) = \frac{C_{\text{gw}} \times f_{\text{shower}} \times F_w \times t_1}{V_a}$$

$$C_{\text{house}} (\text{mg}/\text{m}^3) = \frac{C_{\text{gw}} \times WHF \times f_{\text{house}}}{HV \times ER \times MC}$$

Parameter	Units	Value used	References ^b
Time-adjusted concentration in shower = C_{shower}	mg/m ³	Chemical-specific	Calculated
Indoor inhalation rate = IR_{air}	m ³ /hour	0.833	[14]
Exposure frequency = EF	day/year	350	[14]
Exposure duration = ED	years	24 (adult) 6 (child)	[14]
Conversion factor = CF	μg/mg	10 ⁻³	-----
Exposure Time = ET_{shower}	hours/day	0.2	[14]
Exposure Time = ET_{house}	hours/day	23.8	[14]
Averaging time = AT	hours/day × yr × day/yr	24 × 70 × 365 (carcinogen) 24 × ED × 365 (noncarcinogen)	[14]
Maximum air concentration = $C_{a\text{max}}$	mg/m ³	Chemical-specific	Calculated
Time of shower = t_1	hour	0.1	[14]
Time after shower = t_2	hour	0.1	[14]
Concentration in groundwater = C_{gw}	mg/L	Chemical-specific	-----
Fraction volatilized = f_{shower}	unitless	0.75	----- ^c
Water flow rate = F_w	L/h	890	[14]
Bathroom volume = V_a	m ³	11	[14]
Concentration in household air = C_{house}	mg/m ³	Chemical-specific	Calculated
Water flow rate = WHF	L/day	890	[14]
Fraction volatilized = f_{house}	unitless	0.5	[14]
House volume = HV	m ³ /change	450	[14]
Exchange rate = ER	changes/day	10	[14]
Mixing coefficient = MC	unitless	0.5	[14]

^a Equations from [1], [37], and [42].

^b References (noted in brackets []) follow Table D.50.

^c Value selected by 2009 Paducah Risk Assessment Working Group because KDEP (2002) does not specify this value for showering.

Table D.3. Reasonable Maximum Exposure Assumptions for Inhalation of Volatile Organic Compounds in Water during Household Use by a Rural Resident

Equation removed. Inhalation intake for multiple environment exposures included in Table D.2.

Table D.4. Reasonable Maximum Exposure Assumptions for Dermal Contact with Water while Showering by a Rural Resident^a

Equation:

$$\text{Absorbed Dose Inorganic [mg/(kg} \times \text{day)]} = \frac{C_w \times SA \times K_p \times CF \times ED \times EF \times ET}{BW \times AT}$$

$$\text{Absorbed Dose Organic [mg/(kg} \times \text{day)]} = \frac{DA_{\text{event}} \times SA \times CF \times CF_1 \times ED \times EF \times EV}{BW \times AT}$$

Parameter	Units	Value used	References ^b
Concentration in water = C_w	mg/L	Chemical-specific	----
Skin surface area exposed = SA^c	m ²	1.815 (adult) 0.65 (child)	[14]
Skin permeability constant = K_p	cm/hr	Chemical-specific	----
Absorbed dose per event = DA_{event}	Mg/cm ² -event	Chemical-specific × C_w	----
Conversion Factor = CF	(L-m)/(cm-m ³)	10	----
Conversion Factor = CF₁	cm ³ /L	1000	----
Exposure duration = ED	years	24 (adult) 6 (child)	[14]
Exposure frequency = EF	days/yr	350	[14]
Exposure time = ET	hrs/bath	0.2	[14]
Event = EV	bath/day	1	[14]
Body weight = BW	kg	70 (adult) 15 (child)	[14]
Averaging time = AT	yr × day/yr	70 × 365 (carcinogen) ED × 365 (noncarcinogen)	[14]

^aEquation from [1].

^bReferences (noted in brackets []) follow Table D.50.

^cEntire surface area of body for both adult and child.

Table D.5. Reasonable Maximum Exposure Assumptions for Incidental Ingestion of Soil by a Rural Resident^a

Equations:

$$\text{Chemical Intake [mg/(kg} \times \text{day)]} = \frac{C_s \times CF \times EF \times FI \times ED \times IR}{BW \times AT}$$

$$\text{Radionuclide Intake (pCi)} = \frac{A_s \times CF_{rad} \times EF \times FI \times ED \times IR \times (1 - e^{-\lambda \times ED})}{ED \times \lambda}$$

Parameter	Units	Value used	References ^b
Chemical concentration in soil = C_s	mg/kg	Chemical-specific	-----
Radiological activity = A_s	pCi/g	Chemical-specific	-----
Conversion factor = CF	kg/mg	10 ⁻⁶	-----
Conversion factor = CF_{rad}	g/mg	10 ⁻³	-----
Exposure frequency = EF	days/yr	350	[14]
Fraction ingested = FI	unitless	1	[14]
Exposure duration = ED	years	24 (adult) 6 (child)	[14]
Ingestion rate of soil = IR	mg/d	100 (adult) 200 (child)	[14]
Body weight = BW	kg	70 (adult) 15 (child)	[14]
Averaging time = AT	yr × day/yr	70 × 365 (carcinogen) ED × 365 (noncarcinogen)	[14]
Decay constant = λ	Unitless	0.693/half-life	-----

^a Equation from [1].

^b References (noted in brackets []) follow Table D.50.

Table D.6. Reasonable Maximum Exposure Assumptions for Dermal Contact with Soil by a Rural Resident^a

Equation:

$$\text{Absorbed Dose [(mg)/(kg} \times \text{day)]} = \frac{C_s \times CF_d \times SA \times AF \times ABS \times EF \times ED}{BW \times AT}$$

Parameter	Units	Value used	References^b
Concentration in soil = C_s	mg/kg	Chemical-specific	-----
Conversion factor = CF_d	(kg-cm ²)/(mg-m ²)	0.01	-----
Surface area ^c = SA	m ² /day	0.57 (adult) 0.28 (child)	[14]
Adherence factor = AF	mg/cm ²	1	[14]
Absorption factor ^d = ABS	unitless	Chemical-specific	[14]
Exposure frequency = EF	day/yr	350	[14]
Exposure duration = ED	years	24 (adult) 6 (child)	[14]
Body weight = BW	kg	70 (adult) 15 (child)	[14]
Averaging time = AT	yr × day/yr	70 × 365 (carcinogen) ED × 365 (noncarcinogen)	[14]

^aEquation from [1].

^bReferences (noted in brackets []) follow Table D.50.

^cIncludes face, forearms, hands and lower legs for adult; face, forearms, hands, lower legs and feet for children.

^dChemical-specific absorption factors available are listed in Table B.5 [38].

Note: Dermal absorbed dose is not applicable to radionuclides per guidance found in [1].

Table D.7. Reasonable Maximum Exposure Assumptions for Inhalation of Vapors and Particulates Emitted from Soil by a Rural Resident^a

Equations:

$$\text{Exposure Concentration } (\mu\text{g}/\text{m}^3) = \frac{C_s \times \text{EF} \times \text{ED} \times \text{ET} \times \left(\frac{1}{\text{VF}} + \frac{1}{\text{PEF}} \right)}{\text{AT}} \times \text{CF}_1$$

$$\text{Radionuclide Intake (pCi)} = \frac{A_s \times \text{EF} \times \text{ED} \times \text{ET} \times \text{CF}_2 \times \left(\frac{1}{\text{PEF}} \right) \times (1 - e^{-\lambda \times \text{ED}})}{\text{ED} \times \lambda}$$

Parameter	Units	Value used	References ^b
Concentration in soil = C_s	mg/kg	Chemical-specific	-----
Activity in soil = A_s	pCi/g	Chemical-specific	-----
Exposure frequency = EF	day/year	350	[14]
Exposure duration = ED	years	24 (adult) 6 (child)	[14]
Exposure time = ET	hours/day	24	[14]
Conversion factor = CF₁	μg/mg	10 ⁻³	-----
Conversion factor = CF₂	g/kg	10 ³	-----
Volatilization factor = VF	m ³ /kg	Chemical-specific	[19]
Particulate emission factor ^c = PEF	m ³ /kg	9.3 × 10 ⁸	[14]
Averaging time = AT	hours/day × yr × day/yr	24 × 70 × 365 (carcinogen) 24 × ED × 365 (noncarcinogen)	[14]
Decay constant = λ	Unitless	0.693/half-life	

^a Equation from [42].

^b References (noted in brackets []) follow Table D.50.

^c PEFs from Kentucky Risk Assessment Guidance use EPA default factors, except for the Q/C value, which is based on the lower 90% confidence interval of the mean dispersion factor of climactic zone VII of Table 3 in the Soil Screening Guidance Technical Background document [41].

Table D.8. Reasonable Maximum Exposure Assumptions for External Exposure to Ionizing Radiation from Soil by a Rural Resident^a

Equation:

$$\text{Absorbed Dose [(pCi} \times \text{year)/g]} = \frac{A_s \times ED \times EF \times (1 - S_e) \times T_e \times (1 - e^{-\lambda \times ED})}{ED \times \lambda}$$

Parameter	Units	Value used	References^b
Activity in soil = A_s	pCi/g	Chemical-specific	-----
Exposure duration = ED	year	24 (adult) 6 (child)	[14]
Exposure frequency = EF	day/day	350/365	[14]
Gamma shielding factor = S_e	unitless	0.2	[20]
Gamma exposure time factor = T_e	hr/hr	18/24	[20]
Decay constant = λ	unitless	0.693/half-life	

^aEquation from [20].

^bReferences (noted in brackets []) follow Table D.50.

Table D.9. Reasonable Maximum Exposure Assumptions for Consumption of Home-Grown Vegetables by a Rural Resident^a

Equations:

$$\text{Chemical Intake [mg/(kg} \times \text{day)]} = \frac{C_{\text{vegetables}} \times FI_v \times IR_v \times EF \times ED}{BW \times AT}$$

$$\text{Radionuclide Intake (pCi)} = A_{\text{vegetables}} \times FI_v \times IR_v \times EF \times ED \times CF$$

Parameter	Units	Value used	References^b
Chemical concentration in vegetables = $C_{\text{vegetables}}$	mg/kg	Chemical-specific	See Table D.42
Radiological activity = $A_{\text{vegetables}}$	pCi/g	Chemical-specific	See Table D.42
Diet fraction = FI_v	unitless	0.4	[21]
Ingestion rate ^c = IR_v	kg/d	0.29 (child) 0.72 (adult)	[23]
Exposure frequency = EF	d/year	350	[14]
Exposure duration = ED	years	6 (child) 24 (adult)	[14]
Body weight (adult) = BW	kg	15 (child) 70 (adult)	[14]
Averaging time = AT	yr \times day/yr	70 \times 365 (carcinogen) ED \times 365 (noncarcinogen)	[14]
Conversion factor = CF	g/kg	1000	-----

^aEquation from [1].

^bReferences (noted in brackets []) follow Table D.50.

^cIngestion values represent the 95th percentile of individuals who consume this food group.

Table D.10. Reasonable Maximum Exposure Assumptions for Consumption of Beef by a Rural Resident^a

Equations:

$$\text{Chemical Intake [mg/(kg} \times \text{day)]} = \frac{C_{\text{beef}} \times FI_b \times IR_b \times EF \times ED}{BW \times AT}$$

$$\text{Radionuclide Intake (pCi)} = A_{\text{beef}} \times FI_b \times IR_b \times EF \times ED$$

Parameter	Units	Value used	References ^b
Chemical concentration in beef = C_{beef}	mg/kg	Chemical-specific	See Table D.46
Radiological activity in beef = A_{beef}	pCi/kg	Chemical-specific	See Table D.46
Beef ingestion rate ^c = IR_b	kg/day	0.07 (child 1–7) 0.19 (adult 8–41)	[23]
Diet fraction = FI_b	unitless	1	[21]
Exposure frequency = EF	d/year	350	[14]
Exposure duration = ED	years	6 (child) 24 (adult)	[14]
Body weight (adult) = BW	kg	15 (child) 70 (adult)	[14]
Averaging time = AT	yr \times day/yr	70 \times 365 (carcinogen) ED \times 365 (noncarcinogen)	[14]

^aEquation from [1].

^bReferences (noted in brackets []) follow Table D.50.

^cIngestion values represent the 95th percentile of individuals who consume this food group.

Table D.11. Reasonable Maximum Exposure Assumptions for Consumption of Milk by a Rural Resident^a

Equations:

$$\text{Chemical Intake [mg/(kg} \times \text{day)]} = \frac{C_{\text{milk}} \times FI_m \times IR_m \times EF \times ED}{BW \times AT}$$

$$\text{Radionuclide Intake (pCi)} = A_{\text{milk}} \times FI_m \times IR_m \times EF \times ED$$

Parameter	Units	Value used	References^b
Chemical concentration in milk = C_{milk}	mg/kg	Chemical-specific	See Table D.47
Radiological activity in milk = A_{milk}	pCi/kg	Chemical-specific	See Table D.47
Milk ingestion rate ^c = IR_m	kg/day	0.9 (child 1 – 7) 1.25(adult 8 – 41)	[23]
Diet fraction = FI_m	unitless	1	[21]
Exposure frequency = EF	d/year	350	[14]
Exposure duration = ED	years	6 (child) 24 (adult)	[14]
Body weight (adult) = BW	kg	15 (child) 70 (adult)	[14]
Averaging time = AT	yr × day/yr	70 × 365 (carcinogen) ED × 365 (noncarcinogen)	[14]

^aEquation from [1].

^bReferences (noted in brackets []) follow Table D.50.

^cIngestion values represent the 95th percentile of individuals who consume this food group.

Table D.12. Reasonable Maximum Exposure Assumptions for Consumption of Poultry by a Rural Resident^a

Equations:

$$\text{Chemical Intake [mg/(kg} \times \text{day)]} = \frac{C_{\text{poultry}} \times FI_p \times IR_p \times EF \times ED}{BW \times AT}$$

$$\text{Radionuclide Intake (pCi)} = A_{\text{poultry}} \times FI_p \times IR_p \times EF \times ED$$

Parameter	Units	Value used	References^b
Chemical concentration in poultry = C_{poultry}	mg/kg	Chemical-specific	See Table D.48
Radiological activity in poultry = A_{poultry}	pCi/kg	Chemical-specific	See Table D.48
Ingestion rate ^c = IR_p	kg/day	0.07 (child 1 – 7) 0.17 (adult 8 – 41)	[23]
Diet fraction = FI_p	unitless	1	[5]
Exposure frequency = EF	day/year	350	
Exposure duration = ED	years	6 (child) 24 (adult)	[14]
Body weight = BW	kg	15 (child) 70 (adult)	[14]
Averaging time = AT	yr × day/yr	70 × 365 (carcinogen) ED × 365 (noncarcinogen)	[14]

^aEquation from [1].

^bReferences (noted in brackets []) follow Table D.50.

^cIngestion values represent the 95th percentile of individuals who consume this food group.

Table D.13. Reasonable Maximum Exposure Assumptions for Consumption of Pork by a Rural Resident^a

Equations:

$$\text{Chemical Intake [mg/(kg} \times \text{day)]} = \frac{C_{\text{pork}} \times FI_{\text{pork}} \times IR_{\text{pork}} \times EF \times ED}{BW \times AT}$$

$$\text{Radionuclide Intake (pCi)} = A_{\text{pork}} \times FI_{\text{pork}} \times IR_{\text{pork}} \times EF \times ED$$

Parameter	Units	Value used	References ^b
Chemical concentration in pork = C_{pork}	mg/kg	Chemical-specific	See Table D.49
Radiological activity in pork = A_{pork}	pCi/kg	Chemical-specific	See Table D.49
Pork ingestion rate ^c = IR_{pork}	kg/day	0.03 (child 1 – 7) 0.08 (adult 8 – 41)	[23]
Diet fraction = FI_{pork}	unitless	1	[21]
Exposure frequency = EF	d/year	350	[14]
Exposure duration = ED	years	6 (child) 24 (adult)	[14]
Body weight (adult) = BW	kg	15 (child) 70 (adult)	[14]
Averaging time = AT	yr \times day/yr	70×365 (carcinogen) $ED \times 365$ (noncarcinogen)	[14]

^aEquation from [1].

^bReferences (noted in brackets []) follow Table D.50.

^cIngestion values represent the 95th percentile of individuals who consume this food group.

Table D.14. Reasonable Maximum Exposure Assumptions for Consumption of Eggs by a Rural Resident^a

Equations:

$$\text{Chemical Intake [mg/(kg} \times \text{day)]} = \frac{C_{egg} \times FI_e \times IR_e \times EF \times ED}{BW \times AT}$$

$$\text{Radionuclide Intake (pCi)} = A_{egg} \times FI_e \times IR_e \times EF \times ED$$

Parameter	Units	Value used	References ^b
Chemical concentration in egg = C_{egg}	mg/kg	Chemical-specific	See Table D.50
Radiological activity in egg = A_{egg}	pCi/kg	Chemical-specific	See Table D.50
Egg ingestion rate ^c = IR_e	kg/day	0.06 (child 1 –7) 0.11 (adult 8 - 41)	[23]
Diet fraction = FI_e	unitless	1	[21]
Exposure frequency = EF	d/year	350	[14]
Exposure duration = ED	years	6 (child) 24 (adult)	[14]
Body weight (adult) = BW	kg	15 (child) 70 (adult)	[14]
Averaging time = AT	yr \times day/yr	70 \times 365 (carcinogen) ED \times 365 (noncarcinogen)	[14]

^aEquation from [1].

^bReferences (noted in brackets []) follow Table D.50.

^cIngestion values represent the 95th percentile of individuals who consume this food group.

Table D.15. Reasonable Maximum Exposure Assumptions for Incidental Ingestion of Sediment by a Recreational User^a

Equations:

$$\text{Chemical Intake [mg/(kg} \times \text{day)]} = \frac{C_{sed} \times CF \times EF \times ED \times IR \times FI}{BW \times AT}$$

$$\text{Radionuclide Intake (pCi)} = \frac{A_{sed} \times CF_{rad} \times EF \times ED \times IR \times FI \times (1 - e^{-\lambda \times ED})}{ED \times \lambda}$$

Parameter	Units	Value used	References ^b
Concentration in sediment = C_{sed}	mg/kg	Chemical-specific	----
Conversion factor = CF	kg/mg	10^{-6}	----
Activity in soil = A_{sed}	pCi/g	Chemical-specific	----
Conversion factor = CF_{rad}	g/mg	10^{-3}	----
Exposure frequency = EF	day/yr	104 (adult) 140 (child and teen)	[14]
Exposure duration = ED	Year	12 (adult) 12 (teen) 6 (child)	[14]
Ingestion rate = IR	mg/day	100 (adult) 100 (teen) 200 (child)	[14]
Fraction ingested = FI	unitless	1	[14]
Body weight = BW	kg	70 (adult) 43 (teen) 15 (child)	[14]
Averaging time = AT	yr \times day/yr	70×365 (carcinogen) $ED \times 365$ (noncarcinogen)	[14]
Decay constant = λ	Unitless	0.693/half-life	

^aEquation after [1].

^bReferences (noted in brackets []) follow Table D.50.

Table D.16. Reasonable Maximum Exposure Assumptions for Dermal Contact with Sediment by a Recreational User^a

Equation:

$$\text{Absorbed Dose [(mg)/(kg} \times \text{day)]} = \frac{C_{sed} \times CF_d \times SA \times AF \times ABS \times EF \times ED}{BW \times AT}$$

Parameter	Units	Value used	References ^b
Concentration in sediment = C_{sed}	mg/kg	Chemical-specific	-----
Conversion factor-dermal = CF_d	(kg-cm ²)/(mg-m ²)	0.01	-----
Surface area ^c = SA	m ² /day	0.57 (adult) 0.75 (teen) 0.28 (child)	[14]
Adherence factor = AF	mg/cm ²	1	[14]
Absorption factor ^d = ABS	unitless	Chemical-specific	[14]
Exposure frequency = EF	day/yr	104 (adult) 140 (teen) 140 (child)	[14]
Exposure duration = ED	years	12 (adult) 12 (teen) 6 (child)	[14]
Body weight = BW	kg	70 (adult) 43 (teen) 15 (child)	[14]
Averaging time = AT	yr × day/yr	70 × 365 (carcinogen) ED × 365 (noncarcinogen)	[14]

^aEquation from [1].

^bReferences (noted in brackets []) follow Table D.50.

^cIncludes face, forearms, lower legs and hands for adults; face, arms, hands, legs, and feet for teens; and face, forearms, hands, lower legs, and feet for children.

^dChemical-specific absorption factors available are listed in Table B.5 [38].

Table D.17. Reasonable Maximum Exposure Assumptions for Inhalation of Vapors or Particulates Emitted from Sediment by a Recreational User^a

Equations:

$$\text{Exposure Concentration } (\mu\text{g}/\text{m}^3) = \frac{C_{\text{sed}} \times EF \times ED \times ET \times \left(\frac{1}{VF} + \frac{1}{PEF} \right)}{AT} \times CF_1$$

$$\text{Radionuclide Intake (pCi)} = \frac{A_{\text{sed}} \times EF \times ED \times ET \times CF_2 \times \left(\frac{1}{PEF} \right) \times (1 - e^{-\lambda \times ED})}{ED \times \lambda}$$

Parameter	Units	Value used	References ^b
Concentration in sediment = C_{sed}	mg/kg	Chemical-specific	-----
Activity in sediment = A_{sed}	pCi/g	Chemical-specific	-----
Exposure frequency = EF	day/year	104 (adult) 140 (teen) 140 (child)	[14]
Exposure duration = ED	years	12 (adult) 12 (teen) 6 (child)	[14]
Exposure time = ET	hour/day	5	[14]
Conversion factor = CF₁	μg/mg	10 ⁻³	-----
Conversion factor = CF₂	g/kg	10 ³	-----
Volatilization factor = VF	m ³ /kg	Chemical-specific	-----
Particulate emission factor ^c = PEF	m ³ /kg	9.3 × 10 ⁸	[14]
Averaging time = AT	hours/day × yr × day/yr	24 × 70 × 365 (carcinogen) 24 × ED × 365 (noncarcinogen)	[14]
Decay constant = λ	unitless	0.693/half-life	

^a Equation from [42].

^b References (noted in brackets []) follow Table D.50.

^c PEFs from Kentucky Risk Assessment Guidance use EPA default factors, except for the Q/C value which is based on the lower 90% confidence interval of the mean dispersion factor of climactic zone VII of Table 3 in the Soil Screening Guidance Technical Background document [41].

Table D.18. Reasonable Maximum Exposure Assumptions for External Exposure to Ionizing Radiation from Sediment by a Recreational User^a

Equation:

$$\text{Absorbed Dose [(pCi}\times\text{year)/g]} = \frac{A_{\text{sed}} \times ED \times EF \times (1 - S_e) \times T_e \times (1 - e^{-\lambda \times ED})}{ED \times \lambda}$$

Parameter	Units	Value used	References ^b
Activity in soil = A_{sed}	pCi/g	Chemical-specific	-----
Exposure duration = ED	year	12 (adult) 12 (teen) 6 (child)	[14]
Exposure frequency = EF	day/day	104/365 (adult) 140/365 (teen) 140/365 (child)	[14]
Gamma shielding factor = S_e	unitless	0.0	[40]
Gamma exposure time factor = T_e	hr/hr	5/24	[20]
Decay constant = λ	unitless	0.693/half-life	

^aEquation from [20].

^bReferences (noted in brackets []) follow Table D.50.

Table D.19. Reasonable Maximum Exposure Assumptions for Incidental Ingestion of Surface Water while Swimming by a Recreational User^a

Equations:

$$\text{Chemical Intake [mg/(kg} \times \text{day)]} = \frac{C_{sw} \times IR \times ET \times EF \times ED}{BW \times AT}$$

$$\text{Radionuclide Intake (pCi)} = A_{sw} \times IR \times ET \times EF \times ED$$

Parameter	Units	Value used	References ^b
Chemical concentration in water = C_{sw}	mg/L	Chemical-specific	-----
Radiological activity = A_{sw}	pCi/L	Chemical-specific	-----
Ingestion Rate = IR	L/hr	0.05	[14]
Exposure time = ET	hr/day	2.6	[14]
Exposure frequency = EF	d/year	45	[14]
Exposure duration = ED	years	12 (adult) 12 (teen) 6 (child)	[14]
Body weight = BW	kg	70 (adult) 43 (teen) 15 (child)	[14]
Averaging time = AT	yr \times day/yr	70 \times 365 (carcinogen) ED \times 365 (noncarcinogen)	[14]

^aEquation from [1].

^bReferences (noted in brackets []) follow Table D.50.

Table D.20. Reasonable Maximum Exposure Assumptions for Dermal Contact with Surface Water (Wading) by a Recreational User^a

Equation:

$$\text{Absorbed Dose Inorganic [mg/(kg} \times \text{day)]} = \frac{C_w \times SA \times K_p \times CF \times ED \times EF \times ET}{BW \times AT}$$

$$\text{Absorbed Dose Organic [mg/(kg} \times \text{day)]} = \frac{DA_{\text{event}} \times SA \times CF \times CF_1 \times ED \times EF \times EV}{BW \times AT}$$

Parameter	Units	Value used	References ^b
Concentration in surface water = C_{sw}	mg/L	Chemical-specific	-----
Adult surface area ^c = SA	m ²	1.06 (adult) 0.75 (teen) 0.33 (child)	[14]
Conversion factor = CF	L/(cm - m ²)	10	-----
Conversion factor 1	cm ³ /L	1000	-----
Skin permeability constant = K_p	cm/hr	Chemical-specific	-----
Absorbed dose per event = DA_{event}	Mg/cm ² -event	Chemical-specific × C_w	-----
Exposure duration = ED	Years	12 (adult) 12 (teen) 6 (child)	[14]
Exposure Frequency = EF	d/yr	52 (adult) 140 (teen) 140 (child)	[14]
Exposure time = ET	hr/day	2.6	[14]
Event = EV	Events/day	1	[14]
Body weight = BW	kg	70 (adult) 43 (teen) 15 (child)	[14]
Averaging time = AT	yr × day/yr	70 × 365 (carcinogen) ED × 365 (noncarcinogen)	[14]

^aEquation from [1].

^bReferences (noted in brackets []) follow Table D.50.

^cIncludes arms, hands, legs, and feet for adult, teen, and child.

Note: Dermal absorbed dose is not applicable to radionuclides per guidance found in [1].

Table D.21. Reasonable Maximum Exposure Assumptions for Dermal Contact with Surface Water (Swimming) by a Recreational User^a

Equation:

$$\text{Absorbed Dose Inorganic [mg/(kg} \times \text{day)]} = \frac{C_w \times SA \times K_p \times CF \times ED \times EF \times ET}{BW \times AT}$$

$$\text{Absorbed Dose Organic [mg/(kg} \times \text{day)]} = \frac{DA_{\text{event}} \times SA \times CF \times CF_1 \times ED \times EF \times EV}{BW \times AT}$$

Parameter	Units	Value used	References ^b
Concentration in surface water = C_{sw}	mg/L	Chemical-specific	----
Surface area ^c = SA	m ²	1.815 (adult) 1.31 (teen) 0.65 (child)	[14]
Conversion factor = CF	L/(cm - m ²)	10	----
Conversion factor 1 = CF₁	Cm ³ /L	1000	----
Skin permeability constant = K_p	cm/hr	Chemical-specific	----
Absorbed dose per event = DA_{event}	Mg/cm ² -event	Chemical-specific × C_w	----
Exposure duration = ED	years	12 (adult) 12 (teen) 6 (child)	[14]
Exposure Frequency = EF	d/yr	45	[14]
Exposure time = ET	hr/day	2.6	[14]
Event = EV	Event/day	1	----
Body weight = BW	kg	70 (adult) 43 (teen) 15 (child)	[14]
Averaging time = AT	yr × day/yr	70 × 365 (carcinogen) ED × 365 (noncarcinogen)	[14]

^aEquation from [1].

^bReferences (noted in brackets []) follow Table D.50.

^cIncludes whole body for adult, teen, and child.

Note: Dermal absorbed dose is not applicable to radionuclides per guidance found in [1].

Table D.22. Reasonable Maximum Exposure Assumptions for Consumption of Fish by a Recreational User^a

Equations:

$$\text{Chemical Intake [mg/(kg} \times \text{day)]} = \frac{C_{\text{fish}} \times FI_f \times IR_f \times EF \times ED}{BW \times AT}$$

$$\text{Radionuclide Intake (pCi)} = A_{\text{fish}} \times FI_f \times IR_f \times EF \times ED$$

Parameter	Units	Value used	References^b
Chemical concentration in fish = C_{fish}	mg/kg	Chemical-specific	See Table D.43
Radiological activity = A_{fish}	pCi/kg	Chemical-specific	See Table D.43
Ingestion rate = IR_f	kg/day	0.029(adult) 0.029 (teen) 0.029 (child)	[39]
Diet fraction = FI_f	unitless	1	[5]
Exposure frequency = EF	days/yr	365	----
Exposure duration = ED	years	12 (adult) 12 (teen) 6 (child)	[14]
Body weight = BW	kg	70 (adult) 43 (teen) 15 (child)	[14]
Averaging time = AT	yr × day/yr	70 × 365 (carcinogen) ED × 365 (noncarcinogen)	[14]

^aEquation from [1].

^bReferences (noted in brackets []) follow Table D.50.

Table D.23. Reasonable Maximum Exposure Assumptions for Consumption of Venison by a Recreational User^a

Equations:

$$\text{Chemical Intake [mg/(kg} \times \text{day)]} = \frac{C_{\text{deer}} \times FI_d \times IR_d \times EF \times ED}{BW \times AT}$$

$$\text{Radionuclide Intake (pCi)} = A_{\text{deer}} \times FI_d \times IR_d \times EF \times ED \times CF$$

Parameter	Units	Value used	References ^b
Chemical concentration in venison = C_{deer}	mg/kg	Chemical-specific	See Table D.41
Radiological activity in venison = A_{deer}	pCi/g	Chemical-specific	See Table D.41
Ingestion rate = IR_d	kg/day	0.032 (adult) 0.032 (teen) 0.007 (child)	See footnote c
Conversion factor = CF	g/kg	1000	-----
Diet fraction = FI_d	unitless	1	[5]
Exposure frequency = EF	day/yr	350	See footnote c
Exposure duration = ED	years	12 (adult) 12 (teen) 6 (child)	[14]
Body weight = BW	kg	70 (adult) 43 (teen) 15 (child)	[14]
Averaging time = AT	yr × day/yr	70 × 365 (carcinogen) ED × 365 (noncarcinogen)	[14]

^aEquation from [1].

^bReferences (noted in brackets []) follow Table D.50.

^cBased on taking 2 deer per year (consistent with regulation in the state of Kentucky), a 50% success rate (Kentucky Department of Fish and Wildlife, 1992, Deer Surveys, Project No: W-45-24.), a dressed weight averaging 108.5 pounds per deer for Ballard and McCracken counties, 60% of venison recovered per deer carcass, 2.5 persons per household in Ballard and McCracken counties, and a child consumption rate 20% of that for adults. Intake values above correspond to 0.467 g/kg bw-day for the child, 0.744 g/kg bw-day for the teen, and 0.457 g/kg bw-day for the adult receptor.

Table D.24. Reasonable Maximum Exposure Assumptions for Consumption of Rabbit by a Recreational User^a

Equations:

$$\text{Chemical Intake [mg/(kg} \times \text{day)]} = \frac{C_{\text{rabbit}} \times FI_r \times IR_r \times EF \times ED}{BW \times AT}$$

$$\text{Radionuclide Intake (pCi)} = A_{\text{rabbit}} \times FI_r \times IR_r \times EF \times ED \times CF$$

Parameter	Units	Value used	References ^b
Chemical concentration in rabbit = C_{rabbit}	mg/kg	Chemical-specific	See Table D.45
Radiological activity in rabbit = A_{rabbit}	pCi/g	Chemical-specific	See Table D.45
Ingestion rate = IR_r	kg/meal	0.0165 (adult) 0.0082 (teen) 0.0033 (child)	See footnote c
Conversion factor = CF	g/kg	1000	-----
Diet fraction = FI_r	unitless	1	[5]
Exposure frequency = EF	meals/yr	350	See footnote c
Exposure duration = ED	years	12 (adult) 12 (teen) 6 (child)	[14]
Body weight = BW	kg	70 (adult) 43 (teen) 15 (child)	[14]
Averaging time = AT	yr × day/yr	70 × 365 (carcinogen) ED × 365 (noncarcinogen)	[14]

^aEquation from [1].

^bReferences (noted in brackets []) follow Table D.50.

^cBased on 20 rabbits bagged per year at WKWMA, a personal communication stating that dressed weight equals 60% of average 1.2 kg rabbit, 2.5 persons per household in Ballard and McCracken counties, a child consumption rate 20% of that for adults, and a teen consumption rate 50% of that for adults. Intake values above correspond to 0.220 g/kg bw-day for the child, 0.191 g/kg bw-day for the teen, and 0.236 g/kg bw-day for the adult receptor.

Table D.25. Reasonable Maximum Exposure Assumptions for Consumption of Quail by a Recreational User^a

Equations:

$$\text{Chemical Intake [mg/(kg} \times \text{day)]} = \frac{C_{\text{quail}} \times FI_q \times IR_q \times EF \times ED}{BW \times AT}$$

$$\text{Radionuclide Intake (pCi)} = A_{\text{quail}} \times FI_q \times IR_q \times EF \times ED \times CF$$

Parameter	Units	Value used	References ^b
Chemical concentration in quail = C_{quail}	mg/kg	Chemical-specific	See Table D.44
Radiological activity in quail = A_{quail}	pCi/g	Chemical-specific	See Table D.44
Ingestion rate ^c = IR_q	kg/meal	0.0047 (adult) 0.0024 (teen) 0.00094 (child)	See footnote c
Conversion factor = CF	g/kg	1000	-----
Diet fraction = FI_q	unitless	1	[5]
Exposure frequency = EF	meals/yr	350	See footnote c
Exposure duration = ED	years	12 (adult) 12 (teen) 6 (child)	[14]
Body weight = BW	kg	70 (adult) 43 (teen) 15 (child)	[14]
Averaging time = AT	yr \times day/yr	70 \times 365 (carcinogen) ED \times 365 (noncarcinogen)	[14]

^aEquation from [1].

^bReferences (noted in brackets []) follow Table D.50.

^cBased on 20 quail bagged per year at WKWMA, personal communication stating dressed weight equals 75% of average 0.183 kg quail, 2.5 persons per household in Ballard and McCracken counties, a child consumption rate 20% of that for adults, and a teen consumption rate 50% of that for adults. Intake values above correspond to 0.063 g/kg bw-day for the child, 0.558 g/kg bw-day for the teen, and 0.067 g/kg bw-day for the adult receptor.

Table D.26. Reasonable Maximum Exposure Assumptions for Ingestion of Water by an Industrial Worker^{a, c}

Equations:

$$\text{Chemical Intake [mg/(kg} \times \text{day)]} = \frac{C_w \times IR_w \times EF \times ED}{BW \times AT}$$

$$\text{Radionuclide Intake (pCi)} = A_w \times IR_w \times EF \times ED$$

Parameter	Units	Value used	References^b
Concentration in groundwater = C_w	mg/L	Chemical-specific	-----
Activity in groundwater = A_w	pCi/L	Chemical-specific	-----
Ingestion rate = IR_w	L/day	1	[14]
Exposure frequency = EF	day/yr	250	[14]
Exposure duration = ED	year	25	[14]
Body weight = BW	kg	70	[14]
Averaging time = AT	yr × day/yr	70 × 365 (carcinogen) ED × 365 (noncarcinogen)	[14]

^aEquation from [1].

^bReferences (noted in brackets []) follow Table D.50.

^cBecause future use of groundwater at the PGDP is uncertain, the industrial worker exposure to groundwater scenario is provided here for informational purposes only. This hypothetical future exposure pathway should represent in most, if not all, locations an incomplete exposure pathway.

Table D.27. Reasonable Maximum Exposure Assumptions for Inhalation of Volatile Organic Compounds in Water while Showering by an Industrial Worker^{a, c}

Equations:

$$\text{Exposure Concentration } (\mu\text{g}/\text{m}^3) = \frac{C_{\text{shower}} \times EF \times ET_{\text{shower}} \times ED}{AT} \times CF$$

$$C_{\text{shower}} (\text{mg}/\text{m}^3) = \frac{[(C_{\text{amax}} / 2) \times t_1] + [C_{\text{amax}} \times t_2]}{t_1 + t_2}$$

$$C_{\text{amax}} (\text{mg}/\text{m}^3) = \frac{C_{\text{gw}} \times f \times F_w \times t_1}{V_a}$$

Parameter	Units	Value used	References ^b
Concentration in shower = C_{shower}	mg/m ³	Chemical-specific	Calculated
Exposure frequency = EF	day/year	250	[14]
Exposure duration = ED	years	25	[14]
Exposure time = ET_{shower}	hours/day	0.2	[14]
Conversion factor = CF	μg/mg	10 ⁻³	----
Averaging time = AT	hours/day × yr × day/yr	24 × 70 × 365 (carcinogen) 24 × ED × 365 (noncarcinogen)	[14]
Maximum concentration = C_{amax}	mg/m ³	Chemical-specific	
Time of shower = t_1	hours	0.1	[14]
Time after shower = t_2	hours	0.1	[14]
Concentration in groundwater = C_{gw}	mg/L	Chemical-specific	
Fraction volatilized = f	unitless	0.75	[14]
Water flow rate = F_w	L/h	890	[14]
Bathroom volume = V_a	m ³	11	[14]

^aEquation after [14] and [37].

^bReferences (noted in brackets []) follow Table D.50.

^cBecause future use of groundwater at the PGDP is uncertain, the industrial worker exposure to groundwater scenario is provided here for informational purposes only. This hypothetical future exposure pathway should represent in most, if not all, locations an incomplete exposure pathway.

Table D.28. Reasonable Maximum Exposure Assumptions for Dermal Contact with Water while Showering by an Industrial Worker^{a, d}

Equation:

$$\text{Absorbed Dose Inorganic [mg/(kg} \times \text{day)]} = \frac{C_w \times SA \times K_p \times CF \times ED \times EF \times ET}{BW \times AT}$$

$$\text{Absorbed Dose Organic [mg/(kg} \times \text{day)]} = \frac{DA_{event} \times SA \times CF \times CF_1 \times ED \times EF \times EV}{BW \times AT}$$

Parameter	Units	Value used	References ^b
Concentration in water = C_w	mg/L	Chemical-specific	----
Skin permeability constant = K_p	cm/hr	Chemical-specific	----
Absorbed dose per event = DA_{event}	Mg/cm ² -event	Chemical-specific × C _w	-----
Skin Surface Area ^c = SA	m ²	1.815	[14]
Exposure frequency = EF	baths/yr	250	[14]
Exposure duration = ED	years	25	[14]
Exposure time = ET	hrs/bath	0.2	[14]
Event = EV	Bath/day	1	[14]
Conversion factor = CF	(L-m)/(cm-m ³)	10	----
Conversion factor = CF₁	Cm ³ /L	1000	----
Body weight = BW	kg	70	[14]
Averaging time = AT	yr × day/yr	70 × 365 (carcinogen) ED × 365 (noncarcinogen)	[14]

^aEquation from [1].

^bReferences (noted in brackets []) follow Table D.50.

^cEntire surface area of body.

^dBecause future use of groundwater at the PGDP is uncertain, the industrial worker exposure to groundwater scenario is provided here for informational purposes only. This hypothetical future exposure pathway should represent in most, if not all, locations an incomplete exposure pathway.

Table D.29. Reasonable Maximum Exposure Assumptions for Incidental Ingestion of Soil by an Industrial Worker^a

Equations:

$$\text{Chemical Intake [mg/(kg} \times \text{day)]} = \frac{C_s \times CF \times EF \times FI \times ED \times IR}{BW \times AT}$$

$$\text{Radionuclide Intake (pCi)} = \frac{A_s \times CF_{rad} \times EF \times FI \times ED \times IR \times (1 - e^{-\lambda \times ED})}{ED \times \lambda}$$

Parameter	Units	Value used	References ^b
Concentration in soil = C_s	mg/kg	Chemical-specific	-----
Activity in soil = A_s	pCi/g	Chemical-specific	-----
Ingestion rate = IR	mg/day	50	[14]
Fraction ingested = FI	unitless	1	[14]
Exposure frequency = EF	day/yr	250	[14]
Exposure duration = ED	year	25	[14]
Conversion factor = CF	kg/mg	10 ⁻⁶	-----
Conversion factor = CF_{rad}	g/mg	10 ⁻³	-----
Body weight = BW	kg	70	[14]
Averaging time = AT	yr × day/yr	70 × 365 (carcinogen) ED × 365 (noncarcinogen)	[14]
Decay constant = λ	unitless	0.693/half-life	

^aEquation from [1].

^bReferences (noted in brackets []) follow Table D.50.

Table D.30. Reasonable Maximum Exposure Assumptions for Incidental Ingestion of Sediment by an Industrial Worker or an Outdoor Worker^a

Equations:

$$\text{Chemical Intake [mg/(kg} \times \text{day)]} = \frac{C_{sed} \times CF \times IR \times EF \times ED}{BW \times AT}$$

$$\text{Radionuclide Intake (pCi)} = \frac{A_{sed} \times CF_{rad} \times IR \times EF \times ED \times (1 - e^{-\lambda \times ED})}{ED \times \lambda}$$

Parameter	Units	Value used	References ^b
Concentration in sediment = C_{sed}	mg/kg	Chemical-specific	-----
Conversion factor = CF	kg/mg	10 ⁻⁶	-----
Activity in sediment = A_{sed}	pCi/g	Chemical-specific	-----
Conversion factor = CF_{rad}	g/mg	10 ⁻³	-----
Ingestion rate = IR	mg/day	50 (indoor) 480 (outdoor)	[14] [14]
Exposure frequency = EF	day/yr	250 (indoor) 185 (outdoor)	[14] [14]
Exposure duration = ED	year	25	[14]
Body weight = BW	kg	70	[14]
Averaging time = AT	yr × day/yr	70 × 365 (carcinogen) ED × 365 (noncarcinogen)	[14]
Decay constant = λ	unitless	0.693/half-life	

^aEquation after [1].

^bReferences (noted in brackets []) follow Table D.50.

Note: ED and EF can be used for a construction/excavation worker. However, when used for the construction/excavation worker scenario, the ED and EF should be reduced and documented on a site-specific basis, based on guidance from the Exposure Factors Handbook or similar RAWG-approved guidance.

Table D.31. Reasonable Maximum Exposure Assumptions for Inhalation of Vapors and Particulates Emitted from Soil by an Industrial Worker^a

Equations:

$$\text{Exposure Concentration } (\mu\text{g}/\text{m}^3) = \frac{C_s \times EF \times ED \times ET \times \left(\frac{1}{VF} + \frac{1}{PEF} \right)}{AT} \times CF_1$$

$$\text{Radionuclide Intake (pCi)} = \frac{A_s \times EF \times ED \times ET \times CF_2 \times \left(\frac{1}{PEF} \right) \times (1 - e^{-\lambda \times ED})}{ED \times \lambda}$$

Parameter	Units	Value used	References ^b
Concentration in soil = C_s	mg/kg	Chemical-specific	-----
Activity in soil or = A_s	pCi/g	Chemical-specific	-----
Conversion factor = CF₁	μg/mg	10 ⁻³	-----
Conversion factor = CF₂	g/kg	10 ³	-----
Exposure frequency = EF	day/year	250	[14]
Exposure duration = ED	years	25	[14]
Exposure time = ET	hour/day	8	[14]
Volatilization factor = VF	m ³ /kg	Chemical-specific	[19]
Particulate emission factor ^c = PEF	m ³ /kg	6.2 × 10 ⁸	[14]
Averaging time = AT	hours/day × yr × day/yr	24 × 70 × 365 (carcinogen) 24 × ED × 365 (noncarcinogen)	[14]
Decay constant = λ	unitless	0.693/half-life	

^aEquation from [42].

^bReferences (noted in brackets []) follow Table D.50.

^cPEFs from Kentucky Risk Assessment Guidance use EPA default factors, except for the Q/C value which is based on the lower 90% confidence interval of the mean dispersion factor of climactic zone VII of Table 3 in the Soil Screening Guidance Technical Background document [41].

Table D.32. Reasonable Maximum Exposure Assumptions for Inhalation of Vapors and Particulates Emitted from Sediment by an Industrial Worker^a

Equations:

$$\text{Exposure Concentration } (\mu\text{g}/\text{m}^3) = \frac{C_s \times EF \times ED \times ET \times \left(\frac{1}{VF} + \frac{1}{PEF} \right)}{AT} \times CF_1$$

$$\text{Radionuclide Intake (pCi)} = \frac{A_{sed} \times EF \times ED \times ET \times CF_2 \times \left(\frac{1}{PEF} \right) \times (1 - e^{-\lambda \times ED})}{ED \times \lambda}$$

Parameter	Units	Value used	References ^b
Concentration in sediment = C_{sed}	mg/kg	Chemical-specific	-----
Activity in sediment = A_{sed}	pCi/g	Chemical-specific	-----
Conversion factor = CF_1	$\mu\text{g}/\text{mg}$	10^{-3}	-----
Conversion factor = CF_2	g/kg	10^3	-----
Exposure frequency = EF	day/year	250	[14]
Exposure duration = ED	years	25	[14]
Exposure time for sediment = ET	hours/day	2.6	[14]
Volatilization factor = VF	m^3/kg	Chemical-specific	[19]
Particulate emission factor ^c = PEF	m^3/kg	6.2×10^8	[14]
Averaging time = AT	hours/day \times yr \times day/yr	$24 \times 70 \times 365$ (carcinogen) $24 \times ED \times 365$ (noncarcinogen)	[14]
Decay constant = λ	unitless	0.693/half-life	

^aEquations after [42].

^bReferences (noted in brackets []) follow Table D.50.

^cPEFs from Kentucky Risk Assessment Guidance use EPA default factors, except for the Q/C value, which is based on the lower 90% confidence interval of the mean dispersion factor of climactic zone VII of Table 3 in the Soil Screening Guidance Technical Background document [41].

Table D.33. Reasonable Maximum Exposure Assumptions for Dermal Contact with Soil or Sediment by an Industrial Worker^a

Equation:

$$\text{Absorbed Dose [(mg)/(kg} \times \text{day)]} = \frac{C_s \times CF_d \times SA \times AF \times ABS \times EF \times ED}{BW \times AT}$$

Parameter	Units	Value used	References ^b
Concentration in soil or sediment = C_s	mg/kg	Chemical-specific	-----
Conversion factor-dermal = CF_d	(kg-cm ²)/(mg-m ²)	0.01	-----
Surface area ^c = SA	m ² /day	0.47	[14]
Adherence factor = AF	mg/cm ²	1	[14]
Absorption factor ^d = ABS	unitless	Chemical-specific	[14]
Exposure frequency = EF	day/yr	250	[14]
Exposure duration = ED	years	25	[14]
Body weight = BW	kg	70	[14]
Averaging time = AT	yr \times day/yr	70 \times 365 (carcinogen) ED \times 365 (noncarcinogen)	[14]

^aEquation after [1].

^bReferences (noted in brackets []) follow Table D.50.

^cArea of hands, arms, and head.

^dChemical-specific absorption factors available are listed in Table B.5 [38].

Note: Dermal absorbed dose is not applicable to radionuclides per guidance found in [1].

Table D.34. Reasonable Maximum Exposure Assumptions for External Exposure to Ionizing Radiation from Soil by an Industrial Worker^a

Equation:

$$\text{Absorbed Dose [(pCi} \times \text{ year)/g]} = \frac{A_s \times ED \times EF \times (1 - S_e) \times T_e \times (1 - e^{-\lambda \times ED})}{ED \times \lambda}$$

Parameter	Units	Value used	References^b
Activity in soil = A_s	pCi/g	Chemical-specific	-----
Exposure frequency = EF	day/day	250/365	[14]
Exposure duration = ED	year	25	[14]
Gamma shielding factor = S_e	unitless	0.2	[20]
Gamma exposure time factor = T_e	hr/hr	8/24	[20]
Decay constant = λ	unitless	0.693/half-life	

^aEquation after [20].

^bReferences (noted in brackets []) follow Table D.50.

Table D.35. Reasonable Maximum Exposure Assumptions for External Exposure to Ionizing Radiation from Sediment by an Industrial Worker^a

Equation:

$$\text{Absorbed Dose [(pCi} \times \text{ year)/g]} = \frac{A_{\text{sed}} \times ED \times EF \times (1 - S_e) \times T_e \times (1 - e^{-\lambda \times ED})}{ED \times \lambda}$$

Parameter	Units	Value used	References^b
Activity in sediment = A_{sed}	pCi/g	Chemical-specific	-----
Exposure frequency = EF	day/day	250/365	[14]
Exposure duration = ED	year	25	[14]
Gamma shielding factor = S_e	unitless	0.2	[20]
Gamma exposure time factor = T_e	hr/hr	2.6/24	[20]
Decay constant = λ	unitless	0.693/half-life	

^aEquation from [20].

^bReferences (noted in brackets []) follow Table D.50.

Table D.36. Reasonable Maximum Exposure Assumptions for Dermal Contact with Surface Water by an Industrial or Outdoor Worker^a

Equation:

$$\text{Absorbed Dose Inorganic [mg/(kg} \times \text{day)]} = \frac{C_w \times SA \times K_p \times CF \times ED \times EF \times ET}{BW \times AT}$$

$$\text{Absorbed Dose Organic [mg/(kg} \times \text{day)]} = \frac{DA_{\text{event}} \times SA \times CF \times CF_1 \times ED \times EF \times EV}{BW \times AT}$$

Parameter	Units	Value used	References ^b
Concentration in surface water = C_{sw}	mg/L	Chemical-specific	----
Adult surface area ^c = SA	m ²	0.47	[14]
Skin permeability constant = K_p	cm/hr	Chemical-specific	----
Absorbed dose per event = DA_{event}	mg/cm ² -event	Chemical-specific × C_w	----
Exposure frequency = EF	day/yr	250 (industrial) 20 (outdoor)	[14] ----
Exposure duration = ED	years	25	[14]
Event = EV	event/day	1	[14]
Exposure Time = ET	hr/day	2.6 (industrial) 8 (outdoor)	---- ----
Conversion factor = CF	L/(cm - m ²)	10	----
Conversion factor = CF	cm ³ /L	1000	----
Body weight = BW	kg	70	[14]
Averaging time = AT	yr × day/yr	70 × 365 (carcinogen) ED × 365 (noncarcinogen)	[14]

^aEquation from [1].

^bReferences (noted in brackets []) follow Table D-50.

^cIncludes area of arms, hands, and head.

Notes:

Dermal absorbed dose is not applicable to radionuclides per guidance found in [1].

Note: ED and EF can be used for a construction/excavation worker. However, when used for the construction/excavation worker scenario, the ED and EF should be reduced and documented on a site-specific basis, based on guidance from the Exposure Factors Handbook or similar RAWG-approved guidance.

Table D.37. Reasonable Maximum Exposure Assumptions for Incidental Ingestion of Soil by an Outdoor Worker^a

Equations:

$$\text{Chemical Intake [mg/(kg} \times \text{day)]} = \frac{C_s \times CF \times EF \times FI \times ED \times IR}{BW \times AT}$$

$$\text{Radionuclide Intake (pCi)} = \frac{A_s \times CF_{rad} \times EF \times FI \times ED \times IR \times (1 - e^{-\lambda \times ED})}{ED \times \lambda}$$

Parameter	Units	Value used	References ^b
Concentration in soil or sediment = C_s	mg/kg	Chemical-specific	-----
Conversion factor = CF	kg/mg	10 ⁻⁶	-----
Activity in soil or sediment = A_s	pCi/g	Chemical-specific	-----
Conversion factor = CF_{rad}	g/mg	10 ⁻³	-----
Ingestion rate = IR	mg/day	480	[14]
Exposure frequency = EF	day/yr	185	[14]
Exposure duration = ED	year	25	[20]
Fraction ingested = FI	unitless	1	[14]
Body weight = BW	kg	70	[14]
Averaging time = AT	yr × day/yr	70 × 365 (carcinogen) ED × 365 (noncarcinogen)	[14]
Decay constant = λ	unitless	0.693/half-life	

^aEquation after [1].

^bReferences (noted in brackets []) follow Table D.50.

Note: ED and EF can be used for a construction/excavation worker. However, when used for the construction/excavation worker scenario, the ED and EF should be reduced and documented on a site-specific basis, based on guidance from the Exposure Factors Handbook or similar RAWG-approved guidance.

Table D.38. Reasonable Maximum Exposure Assumptions for Inhalation of Vapors and Particulates Emitted from Soil by an Outdoor Worker^a

Equations:

$$\text{Exposure Concentration } (\mu\text{g}/\text{m}^3) = \frac{C_s \times EF \times ED \times ET \times \left(\frac{1}{VF} + \frac{1}{PEF} \right)}{AT} \times CF_1$$

$$\text{Radionuclide Intake (pCi)} = \frac{A_s \times EF \times ED \times ET \times CF_2 \times \left(\frac{1}{PEF} \right) \times (1 - e^{-\lambda \times ED})}{ED \times \lambda}$$

Parameter	Units	Value used	References ^b
Concentration in soil or sediment = C_s	mg/kg	Chemical-specific	-----
Activity in soil or sediment = A_s	pCi/g	Chemical-specific	-----
Conversion factor = CF_1	$\mu\text{g}/\text{mg}$	10^{-3}	-----
Conversion factor = CF_2	g/kg	10^3	-----
Exposure frequency = EF	day/yr	185	[14]
Exposure duration = ED	years	25	[20]
Exposure time = ET	hours/day	8	[14]
Volatilization factor = VF	m^3/kg	Chemical-specific	[19]
Particulate emission factor ^c = PEF	m^3/kg	6.2×10^8	[14]
Averaging time = AT	hours/day \times yr \times day/yr	$24 \times 70 \times 365$ (carcinogen) $24 \times ED \times 365$ (noncarcinogen)	[14]
Decay constant = λ	unitless	0.693/half-life	

^aEquation from [42].

^bReferences (noted in brackets []) follow Table D.50.

^cPEFs from Kentucky Risk Assessment Guidance use EPA default factors, except for the Q/C value, which is based on the lower 90% confidence interval of the mean dispersion factor of climactic zone VII of Table 3 in the Soil Screening Guidance Technical Background document [41].

Note: ED and EF can be used for a construction/excavation worker. However, when used for the construction/excavation worker scenario, the ED and EF should be reduced and documented on a site-specific basis, based on guidance from the Exposure Factors Handbook or similar RAWG-approved guidance.

Table D.39. Reasonable Maximum Exposure Assumptions for Dermal Contact with Soil by an Outdoor Worker^a

Equation:

$$\text{Absorbed Dose [(mg)/(kg} \times \text{day)]} = \frac{C_s \times CF_d \times SA \times AF \times ABS \times EF \times ED}{BW \times AT}$$

Parameter	Units	Value used	References ^b
Concentration in soil or sediment = C_s	mg/kg	Chemical-specific	-----
Conversion factor-dermal = CF_d	(kg-cm ²)/(mg-m ²)	0.01	-----
Surface area ^c = SA	m ² /day	0.47	[14]
Adherence factor = AF	mg/cm ²	1	[14]
Absorption factor ^d = ABS	unitless	Chemical-specific	[14]
Exposure frequency = EF	day/yr	185	[14]
Exposure duration = ED	years	25	[20]
Body weight = BW	kg	70	[14]
Averaging time = AT	yr × day/yr	70 × 365 (carcinogen) ED × 365 (noncarcinogen)	[14]

^aEquation from [1].

^bReferences (noted in brackets []) follow Table D.50.

^cIncludes skin area of arms, hands, and head.

^dChemical-specific absorption factors available are listed in Table B.5 [38].

Notes:

Dermal absorbed dose is not applicable to radionuclides per guidance found in [1].

Note: ED and EF can be used for a construction/excavation worker. However, when used for the construction/excavation worker scenario, the ED and EF should be reduced and documented on a site-specific basis, based on guidance from the Exposure Factors Handbook or similar RAWG-approved guidance.

Table D.40. Reasonable Maximum Exposure Assumptions for External Exposure to Ionizing Radiation from Soil by an Outdoor Worker^a

Equation:

$$\text{Absorbed Dose [(pCi} \times \text{year)/g]} = \frac{A_s \times ED \times EF \times (1 - S_e) \times T_e \times (1 - e^{-\lambda \times ED})}{ED \times \lambda}$$

Parameter	Units	Value used	References ^b
Activity in soil or sediment = A_s	pCi/g	Chemical-specific	-----
Exposure frequency = EF	day/day	185/365	[14], [20]
Exposure duration = ED	year	25	[20]
Gamma shielding factor = S_e	unitless	0.2	[20]
Gamma exposure time factor = T_e	hr/hr	8/24	[20]
Decay constant = λ	unitless	0.693/half-life	

^aEquation from [20].

^bReferences (noted in brackets []) follow Table D.50.

Note: ED and EF can be used for a construction/excavation worker. However, when used for the construction/excavation worker scenario, the ED and EF should be reduced and documented on a site-specific basis, based on guidance from the Exposure Factors Handbook or similar RAWG-approved guidance.

Table D.41. Reasonable Maximum Exposure Assumptions for Concentration or Activity of COPCs in Deer^{a, d}

Equations:

$$C_{deer} = F_{deer} \times [(C_{forage} \times AC \times f_s \times Q_f) + (C_s \times AC \times Q_s) + (C_{sw} \times CF_{rad} \times Q_{sw})]$$

$$C_{forage} = (C_s \times R_{upp}) + (C_s \times R_{es})$$

Parameter	Units	Value used	References ^b
Chemical concentration in deer = C_{deer}	mg/kg or pCi/g	Chemical-specific	Calculated
Forage-deer transfer factor = F_{deer}	day/kg	Chemical-specific	-----
Chemical concentration in forage = C_{forage}	mg/kg or pCi/g	Chemical-specific	Calculated
Area of contact ^c = AC	unitless	AS/AD	-----
Area of SWMU = AS	acres	SWMU-specific	-----
Area of deer range = AD	acres	494	[34]
Fraction of deer's food from site when on-site = f_s	unitless	1.0	[5]
Quantity of forage ingested daily by deer = Q_f	kg/day	1.74	[7]
Chemical concentration in soil or sediment = C_s	mg/kg or pCi/g	Chemical-specific	-----
Quantity of soil ingested daily by deer = Q_s	kg/day	0.034	[6]; 2% of forage
Contaminant concentration in surface water = C_{sw}	mg/L or pCi/L	Chemical-specific	-----
Conversion factor for radionuclides = CF_{rad}	kg/g	10^{-3}	-----
Quantity of surface water ingested daily by deer = Q_{sw}	L/day	3.61	[8]
Soil to plant uptake (dry) = R_{upp}	unitless	Chemical-specific or $38 \times K_{ow}^{-0.58}$	[8]
Soil resuspension multiplier = R_{es}	unitless	0.25	[3]

^a Equations after [1], [2], [3], [4].

^b All references (noted in brackets []) follow Table D.50.

^c AC cannot be greater than 1.

^d All ingested water is assumed to be from SWMU or SWMU area.

Table D.42. Reasonable Maximum Exposure Assumptions for Concentration or Activity of COPCs in Home-Grown Vegetables^a

Equations:

$$C_{vegetables} = (C_w \times Irr_{rup} \times CF_{rad}) + (C_s \times AC \times R_{upv}) + (C_w \times Irr_{res} \times CF_{rad}) + (C_s \times AC \times R_{es}) + (C_w \times Irr_{dep} \times CF_{rad})$$

$$Irr_{rup} = \frac{Ir \times F \times Bv_{wet} \times [1 - \exp(-\lambda_B \times t_b)]}{P \times \lambda_B}$$

$$Irr_{dep} = \frac{Ir \times F \times I_f \times T \times [1 - \exp(-\lambda_E \times t_v)]}{Y_v \times \lambda_E}$$

$$Irr_{res} = \frac{Ir \times F \times MLF \times [1 - \exp(-\lambda_B \times t_b)]}{P \times \lambda_B}$$

Parameter	Units	Value used	References ^b
Concentration in vegetable = $C_{vegetables}$	mg/kg or pCi/g	Chemical-specific	Calculated
Concentration in groundwater = C_w	mg/L or pCi/L	Chemical-specific	-----
Root uptake from irrigation = Irr_{rup}	L/kg	Chemical-specific	Calculated
Conversion factor for radionuclides = CF_{rad}	kg/g	10^{-3}	-----
Concentration in soil = C_s	mg/kg or pCi/g	Chemical-specific	-----
Area of contact ^c = AC	unitless	AS/AG	-----
Area of SWMU = AS	acres	SWMU-specific	-----
Area of garden = AG	acres	0.25	[33]
Wet root uptake for leafy vegetables = R_{upv}	kg/kg	Chemical-specific	-----
Resuspension from irrigation = Irr_{res}	L/kg	Chemical-specific	Calculated
Resuspension multiplier = R_{es}	unitless	0.26	[9]
Aerial deposition from irrigation = Irr_{dep}	L/kg	Chemical-specific	Calculated
Irrigation rate = Ir	L/m ² -day	3.62	[10]
Irrigation period = F	unitless	0.25	[10]; 3 months a year
Soil to plant uptake, wet weight = Bv_{wet}	kg/kg	Chemical-specific or $7.7 \times K_{ow}^{-0.58}$	[11]
Effective rate for removal = λ_B	1/day	$\lambda_i + \lambda_{HL}$	[11]
Decay = λ_i	1/day	$0.693/T_r$	[11]
Half-life = T_r	day	Chemical-specific	-----
Soil leaching rate = λ_{HL}	1/day	2.7×10^{-5}	[11]
Long term deposition and build-up = t_b	day	10,950	[2]
Area density for root zone = P	kg/m ²	240	[8], [12], [13]
Plant mass leading factor = MLF	unitless	0.26	[9]
Interception fraction = I_f	unitless	0.42	[7]
Translocation factor = T	unitless	1	[2]
Decay for removal on produce = λ_E	1/day	$\lambda_i + (0.693/t_w)$	[11]
Weathering half-life = t_w	day	14	[2]
Above ground exposure time = t_v	day	60	[2]
Plant yield (wet) = Y_v	kg/m ²	2	[2]

^aEquations after [1], [2], [3], [4].

^bReferences (noted in brackets []) follow Table D.50.

^cAC cannot be greater than 1.

Table D.43. Reasonable Maximum Exposure Assumptions for Concentration or Activity of COPCs in Fish

Equation:

$$C_{fish} = C_{sw} \times BAF_{fish}$$

Parameter	Units	Value used	References
Contaminant concentration in fish = C_{fish}	mg/kg or pCi/kg	Chemical-specific	Calculated
Contaminant concentration in water = C_{sw}	mg/L or pCi/L	Chemical-specific	-----
Bioaccumulation factor = BAF_{fish}	L/kg	Chemical-specific	-----

Table D.44. Reasonable Maximum Exposure Assumptions for Concentration or Activity of COPCs in Quail^{a, d}

Equations:

$$C_{quail} = F_{quail} \times [(C_{forage} \times AC \times f_s \times Q_f) + (C_s \times AC \times Q_s) + (C_{sw} \times CF_{rad} \times Q_{sw}) + (C_i + AC + Q_i)]$$

$$C_{forage} = (C_s \times R_{upp}) + (C_s \times R_{es}) \qquad C_i = (C_s \times BAF_i)$$

Parameter	Units	Value used	References ^b
Chemical concentration in quail = C_{quail}	mg/kg or pCi/g	Chemical-specific	Calculated
Forage-quail transfer factor = F_{quail}	day/kg	Chemical-specific	use $F_{poultry}$ values
Chemical concentration in forage = C_{forage}	mg/kg or pCi/g	Chemical-specific	Calculated
Area of contact ^c = AC	unitless	AS/AQ	-----
Area of SWMU = AS	acres	SWMU-specific	-----
Area of quail range = AQ	acres	15.4	[30]
Fraction of quail's food from site when on-site = f_s	unitless	1.0	-----
Quantity of forage ingested daily by quail = Q_f	kg/day	0.01499	[30] 88.2% of total food
Chemical concentration in invertebrates = C_i	mg/kg or pCi/g	Chemical-specific	-----
Quantity of invertebrates ingested daily by quail = Q_i	kg/day	0.002006	[30] 11.8% of total food
Chemical concentration in soil or sediment = C_s	mg/kg or pCi/g	Chemical-specific	-----
Quantity of soil ingested daily by quail = Q_s	kg/day	0.00158	[32] 9.3% of total food (same as turkey)
Contaminant concentration in surface water = C_{sw}	mg/L or pCi/L	Chemical-specific	-----
Conversion factor for radionuclides = CF_{rad}	kg/g	10^{-3}	-----
Quantity of surface water ingested daily by quail = Q_{sw}	L/day	0.024	[30]
Soil to plant uptake (dry) = R_{upp}	unitless	Chemical-specific or $38 \times K_{ow}^{-0.58}$	[8]
Soil resuspension multiplier = R_{es}	unitless	0.25	[3]

^a Equations after [1], [2], [3], [4].

^b All references (noted in brackets []) follow Table D.50.

^c AC cannot be greater than 1.

^d All ingested water is considered to be from SWMU or SWMU area.

Table D.45. Reasonable Maximum Exposure Assumptions for Concentration or Activity of COPCs in Rabbits^{a, d}

Equations:

$$C_{\text{rabbit}} = F_{\text{rabbit}} \times [(C_{\text{forage}} \times AC \times f_s \times Q_f) + (C_s \times AC \times Q_s) + (C_{\text{sw}} \times CF_{\text{rad}} \times Q_{\text{sw}})]$$

$$C_{\text{forage}} = (C_s \times R_{\text{upp}}) + (C_s \times R_{\text{es}})$$

Parameter	Units	Value used	References ^b
Chemical concentration in rabbit = C_{rabbit}	mg/kg or pCi/g	Chemical-specific	Calculated
Forage-rabbit transfer factor = F_{rabbit}	day/kg	Chemical-specific	use F_{beef} values
Chemical concentration in forage = C_{forage}	mg/kg or pCi/g	Chemical-specific	Calculated
Area of contact ^c = AC	unitless	AS/AR	-----
Area of SWMU = AS	acres	SWMU-specific	-----
Area of rabbit range = AR	acres	3.6	[30]
Fraction of rabbit's food from site when on-site = f_s	unitless	1.0	-----
Quantity of forage ingested daily by rabbit = Q_f	kg/day	0.237	[31]
Chemical concentration in soil or sediment = C_s	mg/kg or pCi/g	Chemical-specific	-----
Quantity of soil ingested daily by rabbit = Q_s	kg/day	0.0149	[31] 6.3% of forage
Contaminant concentration in surface water = C_{sw}	mg/L or pCi/L	Chemical-specific	-----
Conversion factor for radionuclides = CF_{rad}	kg/g	10^{-3}	-----
Quantity of surface water ingested daily by rabbit = Q_{sw}	L/day	0.116	[31]
Soil to plant uptake (dry) = R_{upp}	unitless	Chemical-specific or $38 \times K_{\text{ow}}^{-0.58}$	[8]
Soil resuspension multiplier = R_{es}	unitless	0.25	[3]

^a Equations after [1], [2], [3], [4].

^b All references (noted in brackets []) follow Table D.50.

^c AC cannot be greater than 1.

^d All ingested water is considered to be from SWMU or SWMU area.

Table D.46. Reasonable Maximum Exposure Assumptions for Concentration or Activity of COPCs in Beef^{a, d}

Equations:

$$C_{beef} = F_{beef} \times [(C_{forage} \times AC \times f_s \times Q_f) + (C_s \times AC \times Q_s) + (C_w \times CF_{rad} \times Q_w)]$$

$$C_{forage} = (C_s \times R_{upp}) + (C_s \times R_{es})$$

Parameter	Units	Value used	References ^b
Chemical concentration in beef = C_{beef}	mg/kg or pCi/g	Chemical-specific	Calculated
Forage-beef transfer factor = F_{beef}	day/kg	Chemical-specific	-----
Chemical concentration in pasture = C_{forage}	mg/kg or pCi/g	Chemical-specific	Calculated
Area of contact ^c = AC	unitless	AS/AD	-----
Area of SWMU = AS	acres	SWMU-specific	-----
Area of beef range = AD	acres	2	[29]
Fraction of beef's food from site when on-site = f_s	unitless	1.0	[5]
Quantity of pasture ingested daily by beef = Q_f	kg/day	25	[25]
Chemical concentration in soil or sediment = C_s	mg/kg or pCi/g	Chemical-specific	-----
Quantity of soil ingested daily by beef = Q_s	kg/day	1	[26]
Contaminant concentration in water = C_w	mg/L or pCi/L	Chemical-specific	-----
Conversion factor for radionuclides = CF_{rad}	kg/g	10^{-3}	-----
Quantity of water ingested daily by beef = Q_w	L/day	50	[25]
Soil to plant uptake (dry) = R_{upp}	unitless	Chemical-specific or $38 \times K_{ow}^{-0.58}$	[8]
Soil resuspension multiplier = R_{es}	unitless	0.25	[3]

^aEquations after [1], [2], [3], [4].

^bAll references (noted in brackets []) follow Table D.50.

^cAC cannot be greater than 1.

^dAll ingested water is considered to be from SWMU or SWMU area.

Table D.47. Reasonable Maximum Exposure Assumptions for Concentration or Activity of COPCs in Milk^{a, d}

Equations:

$$C_{milk} = F_{milk} \times [(C_{forage} \times AC \times f_s \times Q_f) + (C_s \times AC \times Q_s) + (C_w \times CF_{rad} \times Q_w)]$$

$$C_{forage} = (C_s \times R_{upp}) + (C_s \times R_{es})$$

Parameter	Units	Value used	References ^b
Chemical concentration in milk = C_{milk}	mg/kg or pCi/g	Chemical-specific	Calculated
Forage-milk transfer factor = F_{milk}	day/kg	Chemical-specific	-----
Chemical concentration in pasture = C_{forage}	mg/kg or pCi/g	Chemical-specific	Calculated
Area of contact ^c = AC	unitless	AS/AD	-----
Area of SWMU = AS	acres	SWMU-specific	-----
Area of dairy range = AD	acres	2	[29]
Fraction of dairy's food from site when on-site = f_s	unitless	1.0	[5]
Quantity of pasture ingested daily by dairy = Q_f	kg/day	25	[25]
Chemical concentration in soil or sediment = C_s	mg/kg or pCi/g	Chemical-specific	-----
Quantity of soil ingested daily by dairy = Q_s	kg/day	1	[26]
Contaminant concentration in water = C_w	mg/L or pCi/L	Chemical-specific	-----
Conversion factor for radionuclides = CF_{rad}	kg/g	10^{-3}	-----
Quantity of water ingested daily by dairy = Q_w	L/day	60	[25]
Soil to plant uptake (dry) = R_{upp}	unitless	Chemical-specific or $38 \times K_{ow}^{-0.58}$	[8]
Soil resuspension multiplier = R_{es}	unitless	0.25	[3]

^a Equations after [1], [2], [3], [4].

^b All references (noted in brackets []) follow Table D.50.

^c AC cannot be greater than 1.

^d All ingested water is considered to be from SWMU or SWMU area.

Table D.48. Reasonable Maximum Exposure Assumptions for Concentration or Activity of COPCs in Poultry^{a, e}

Equations:

$$C_{poultry} = F_{poultry} \times [(C_{forage} \times AC \times f_s \times Q_f) + (C_s \times AC \times Q_s) + (C_w \times CF_{rad} \times Q_w)]$$

$$C_{forage} = (C_s \times R_{upp}) + (C_s \times R_{es})$$

Parameter	Units	Value used	References ^b
Chemical concentration in poultry = $C_{poultry}$	mg/kg or pCi/g	Chemical-specific	Calculated
Forage-poultry transfer factor = $F_{poultry}$	day/kg	Chemical-specific	-----
Chemical concentration in pasture = C_{forage}	mg/kg or pCi/g	Chemical-specific	Calculated
Area of contact ^c = AC	unitless	AS/AD	-----
Area of SWMU = AS	acres	SWMU-specific	-----
Area of poultry range = AD^d	acres	1	[29]
Fraction of poultry's food from site = f_s	unitless	.5	[29] assumes broilers get 50% bought grain
Quantity of pasture ingested daily by poultry = Q_f	kg/day	0.12 (chicken) 0.35 (turkey)	[24] 20 wk old male turkey
Chemical concentration in soil or sediment = C_s	mg/kg or pCi/g	Chemical-specific	-----
Quantity of soil ingested daily by poultry = Q_s	kg/day	0.0024 (chicken) 0.007 (turkey)	[8] same ratio for chicken
Contaminant concentration in water = C_w	mg/L or pCi/L	Chemical-specific	-----
Conversion factor for radionuclides = CF_{rad}	kg/g	10^{-3}	-----
Quantity of water ingested daily by poultry = Q_w	L/day	0.24 (chicken) 1.0 (turkey)	[24] 1:2 ratio of 20 wk old male turkey
Soil to plant uptake (dry) = R_{upp}	unitless	Chemical-specific or $38 \times K_{ow}^{-0.58}$	[8]
Soil resuspension multiplier = R_{es}	unitless	0.25	[3]

^aEquations after [1], [2], [3], [4].

^bAll references (noted in brackets []) follow Table D.50.

^cAC cannot be greater than 1.

^dAssumes 1 acre of pasture for 200 adult birds with a three year rotation.

^eAll ingested water is considered to be from SWMU or SWMU area.

Note: Under this model, poultry raised for use as broilers by subsistence farmers are allowed to forage on pasture where they ingest pasture and soil.

Table D.49. Reasonable Maximum Exposure Assumptions for Concentration or Activity of COPCs in Pork^{a,d}

Equations:

$$C_{pork} = F_{pork} \times [(C_{forage} \times AC \times f_s \times Q_f) + (C_s \times AC \times Q_s) + (C_w \times CF_{rad} \times Q_w)]$$

$$C_{forage} = (C_s \times R_{upp}) + (C_s \times R_{es})$$

Parameter	Units	Value used	References ^b
Chemical concentration in pork = C_{pork}	mg/kg or pCi/g	Chemical-specific	Calculated
Forage-pork transfer factor = F_{pork}	day/kg	Chemical-specific	-----
Chemical concentration in pasture = C_{forage}	mg/kg or pCi/g	Chemical-specific	Calculated
Area of contact ^c = AC	unitless	AS/AD	-----
Area of SWMU = AS	acres	SWMU-specific	-----
Area of swine range = AD	acres	1	[29]
Fraction of swine's food from site = f_s	unitless	0.4	[29]
Quantity of pasture ingested daily by swine = Q_f	kg/day	2.4	[36]
Chemical concentration in soil or sediment = C_s	mg/kg or pCi/g	Chemical-specific	-----
Quantity of soil ingested daily by swine = Q_s	kg/day	0.034	[28]
Contaminant concentration in water = C_w	mg/L or pCi/L	Chemical-specific	-----
Conversion factor for radionuclides = CF_{rad}	kg/g	10^{-3}	-----
Quantity of water ingested daily by swine = Q_w	L/day	6.14	[27] 2.56 to 1, water to feed ratio
Soil to plant uptake (dry) = R_{upp}	unitless	Chemical-specific or $38 \times K_{ow}^{-0.58}$	[8]
Soil resuspension multiplier = R_{es}	unitless	0.25	[3]

^aEquations after [1], [2], [3], [4].

^bAll references (noted in brackets []) follow Table D.50.

^cAC cannot be greater than 1.

^dAll ingested water is considered to be from SWMU or SWMU area.

Note: According to Morrison (1956), subsistence farmers allow 20 to 40 percent of the swine's diet to come from pasture, while the remaining comes from store bought grain.

Table D.50. Reasonable Maximum Exposure Assumptions for Concentration or Activity of COPCs in Egg^{a, c, d}

Equations:

$$C_{egg} = F_{egg} \times (C_w \times CF_{rad} \times Q_w)$$

Parameter	Units	Value used	References ^b
Chemical concentration in egg = C_{egg}	mg/kg or pCi/g	Chemical-specific	Calculated
Forage-egg transfer factor = F_{egg}	day/kg	Chemical-specific	-----
Contaminant concentration in water = C_w	mg/L or pCi/L	Chemical-specific	-----
Conversion factor for radionuclides = CF_{rad}	kg/g	10^{-3}	-----
Quantity of water ingested daily by poultry = Q_w	L/day	0.24 (chicken) 1.0 (turkey)	[24] 1:2 ratio of 20 wk old male turkey

^a Equations after [1], [2], [3], [4].

^b All references (noted in brackets []) follow Table 50.

^c AC cannot be greater than 1.

^d All ingested water is considered to be from SWMU or SWMU area.

Note: Model assumes that laying hens are in a hutch and are not allowed to forage on pasture. Therefore, they eat only store bought grain and are not exposed to pasture or soil. Drinking water is assumed to come from the SWMU or SWMU area.

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APPENDIX E
ADDITIONAL INFORMATION

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APPENDIX E
ADDITIONAL INFORMATION (CD)

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