Risk Assessment Guidance for Superfund: Volume I Human Health Evaluation Manual (Part D, Standardized Planning, Reporting, and Review of Superfund Risk Assessments)

Final

Office of Emergency and Remedial Response U.S. Environmental Protection Agency Washington, DC 20460

NOTICE

This document provides guidance to EPA Regions concerning how the Agency intends to exercise its discretion in implementing one aspect of the CERCLA remedy selection process. The guidance is designed to implement national policy on these issues.

Some of the statutory provisions described in this document contain legally binding requirements. However, this document does not substitute for those provisions or regulations, nor is it a regulation itself. Thus, it cannot impose legally-binding requirements on EPA, States, or the regulated community, and may not apply to a particular situation based upon the circumstances. Any decisions regarding a particular remedy selection decision will be made based on the statute and regulations, and EPA decisionmakers retain the discretion to adopt approaches on a case-by-case basis that differ from this guidance where appropriate.

Interested parties are free to raise questions and objections about the substance of this guidance and the appropriateness of the application of this guidance to a particular situation, and the Agency welcomes public input on the document at any time. EPA may change this guidance in the future.

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DEFINITIONS

These definitions are provided for purposes of this guidance and are intended to be consistent with existing Agency guidance and regualtions.

| Term | Definition |
|---|--|
| Applicable or Relevant and Appropriate Requirements (ARARs) | As defined in the NCP, "Applicable" requirements are those clean-up standards of control, and other substantive environmental protection requirements, criteria, or limitations promulgated under federal or state law that specifically address a hazardous substance, pollutant, contaminant, remedial action, location, or other circumstance at a Comprehensive Environmental Response, Compensation, and Liability Act (CERCLA) site. "Relevant and appropriate" requirements are those clean-up standards which, while not "applicable" at a CERCLA site, address problems or situations sufficiently similar to those encountered at the CERCLA site that their use is well-suited to the particular site. ARARs can be action-specific, location-specific, or chemical-specific. |
| Conceptual Site Model | A "model" of a site developed at scoping using readily available information. Used to identify all potential or suspected sources of contamination, types and concentrations of contaminants detected at the site, potentially contaminated media, and potential exposure pathways, including receptors. This model is also known as "conceptual evaluation model." |
| Deterministic Analysis | Calculation and expression of health risks as single numerical values or "single point" estimates of risk. In risk assessments, the uncertainty and variability are discussed in a qualitative manner. |
| EPA Risk Assessor | The risk assessor responsible for reviewing the risk assessment on behalf of EPA. The individual may be an EPA employee or contractor, a State employee, or some other party, as appropriate for an individual site. |
| Exposure Medium | The contaminated environmental medium to which an individual may be exposed. Includes the transfer of contaminants from one medium to another. |

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| Term | Definition |
|------------------------------|--|
| Exposure Pathway | The course a chemical or radionuclide takes from the source to the exposed individual. An exposure pathway analysis links the sources, locations, and types of environmental releases with population locations and activity patterns to determine the significant pathways of human exposure. Within the Planning Tables, an Exposure Pathway is defined as each unique combination of Scenario Timeframe, Medium, Exposure Medium, Exposure Point, Receptor Population, Receptor Age, and Exposure Route. |
| Exposure Point | An exact location of potential contact between a person and a chemical or radionuclide within an Exposure Medium. |
| Exposure Point Concentration | The value, based on either a statistical derivation of measured data or modeled data, that represents an estimate of the chemical or radionuclide concentration available from a particular Medium or route of exposure. |
| Exposure Route | The way a chemical or radionuclide comes in contact with a person (e.g., by ingestion, inhalation, dermal contact). |
| Interim Deliverables | A series of Planning Tables, Worksheets, and Supporting Information, identified in the Workplan for each site, that should be developed by the risk assessment author, and evaluated by the EPA risk assessor, prior to development of the Draft Baseline Risk Assessment Report. After review and revision, as necessary, these documents should be included in the Baseline Risk Assessment Report. The Planning Tables should be prepared for each site to achieve standardization in risk assessment reporting. The Worksheets and Supporting Information should also be prepared to further improve transparency, clarity, consistency, and reasonableness of risk assessments. |
| Medium | The environmental substance (e.g, air, water, soil) that is a potential source of contaminants in the Exposure Medium. (The Medium will sometimes equal the Exposure Medium.) Usually the Medium is targeted for possible remediation. |

| Term | Definition |
|--------------------------------------|---|
| Preliminary Remediation Goals (PRGs) | Generally, initial cleanup goals that (1) are protective of human health and the environment and (2) comply with ARARs. Pursuant to the NCP, they are developed early in the remedy selection process based on readily available information and should be modified to reflect results of the baseline risk assessment. They also should be used during analysis of remedial alternatives in the remedial investigation/feasibility study (RI/FS). Remedial goals, selected as part of the risk management decision, normally replace PRGs in the Record of Decision. |
| Probabilistic Analysis | Calculation and expression of health risks using multiple risk descriptors to provide the likelihood of various risk levels. Probabilistic risk results approximate a full range of possible outcomes and the likelihood of each, which often are presented as a frequency distribution graph, thus allowing uncertainty or variability to be expressed quantitatively. |
| Risk Assessment Author | The risk assessor responsible for preparing the risk assessment. This individual may be an EPA employee or contractor, a State employee, a PRP employee or contractor, or some other party, as appropriate for an individual site. |
| Receptor Age | The description of the exposed individual as defined by the EPA Region or dictated by the site. |
| Receptor Population | The exposed individual relative to the Exposure Pathway considered. |
| Scenario Timeframe | The time period (current and/or future) being considered for the Exposure Pathway. |

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| Term | Definition |
|------------------------|--|
| Planning Tables | One of the Planning Tools under the RAGS Part D approach. The Planning Tables have been developed to clearly and consistently document important parameters, data, calculations, and conclusions from all stages of human health risk assessment development. Electronic templates for the Planning Tables have been developed in Lotus® and Excel® for ease of use by risk assessors. For each site-specific risk assessment, the Planning Tables, related Worksheets, and Supporting Information should first be prepared as Interim Deliverables for EPA risk assessor review, and should later be included in the Draft and Final Baseline Risk Assessment Reports. The Planning Tables may be found in Appendix A. Use of the Planning Tables will standardize the reporting of human health risk assessments. The Planning Table formats should not be altered (i.e., columns should not be added, deleted, or changed); however, rows and footnotes may be added as appropriate. Standardization of the Tables is needed to achieve Superfund program-wide reporting consistency. |
| Planning Tools | A basic element of the RAGS Part D approach. The Planning Tools have been developed to standardize the planning, reporting, and review of Superfund risk assessments. The three Planning Tools contained in the Part D approach include the Technical Approach for Risk Assessment (TARA), the Planning Tables, and Instructions for the Planning Tables. |
| Supporting Information | Information submissions that substantiate or summarize detailed data analysis, calculations, or modeling and associated parameters and assumptions. Examples of recommended Supporting Information include: derivations of background |

Assessment Report.

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values, exposure point concentrations, modeled intakes, and chemical-specific parameters. Supporting Information should be provided as Interim Deliverables for EPA risk assessor review prior to the development of the Draft Baseline Risk

| DEFINITIONS (Continued) | | |
|---|--|--|
| Term | Definition | |
| Technical Approach for Risk Assessment (TARA) | One of the Planning Tools under the RAGS Part D approach. The TARA is a road map for incorporating continuous involvement of the EPA risk assessor throughout the CERCLA remedial process. Risk-related activities, beginning with scoping and problem formulation, extending through collection and analysis of risk-related data, and supporting risk management decision making and remedial design/remedial action issues are addressed. The TARA should be customized for each site and the requirements identified should be included in project workplans so that risk assessment requirements and approaches are clearly defined. The TARA Schedule Worksheet may be found in Appendix C with the other worksheets. Chapters 2 through 5 of Part D present the TARA. | |
| Worksheets | Formats for documenting assumptions, input parameters, and conclusions regarding complex risk assessment issues. Data Useability, TARA Schedule, Lead, Dermal, Radiation Dose Assessment, and ROD Risk Worksheets are found in Appendix C and should be developed as Interim Deliverables for all risk assessments, as applicable. | |

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ACRONYMS/ABBREVIATIONS

Acronym/

Abbreviation Definition

ARARs Applicable or Relevant and Appropriate Requirements

BRAC Base Realignment and Closure

CERCLA Comprehensive Environmental Response Compensation and

Liability Act

COPCs Chemicals of Potential Concern

CSF Cancer Slope Factor
CT Central Tendency
CWA Clean Water Act
DQOs Data Quality Objectives

EPA U.S. Environmental Protection Agency

EPC Exposure Point Concentration

ESD Explanation of Significant Differences

FS Feasibility Study
FY Fiscal Year

GAO General Accounting Office

HEAST Health Effects Assessment Summary Tables

HI Hazard Index HQ Hazard Quotient

IEUBK Integrated Exposure Uptake Biokinetic Model

IRIS Integrated Risk Information System MCLs Maximum Contaminant Levels

NCEA National Center for Environmental Assessment

NCP National Contingency Plan
NPL National Priorities List
non-TCL non-Target Compound List

OSWER Office of Solid Waste and Emergency Response

PAHs Polynuclear Aromatic Hydrocarbons

PCBs Polychlorinated Biphenyls
PQLs Procedure Quantitation Limits
PRGs Preliminary Remediation Goals
PRP Potentially Responsible Party
QA/QC Quality Assurance/Quality Control
QAPP Quality Assurance Project Plan

RAGS Risk Assessment Guidance for Superfund

RAGS/HHEM Risk Assessment Guidance for Superfund: Volume I --

Human Health Evaluation Manual

RAOs Remedial Action Objectives RfC Reference Concentration

RfD Reference Dose

RI/FS Remedial Investigation/Feasibility Study

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ACRONYMS/ABBREVIATIONS (Continued)

Acronym/ Abbreviation Definition

RI Remedial Investigation

Reasonable Maximum Exposure **RME**

Record of Decision ROD

Remedial Project Manager **RPM** Sampling and Analysis Plan SAP **SDWA** Safe Drinking Water Act

Technical Approach for Risk Assessment **TARA**

Upper Confidence Level **UCL**

URF Unit Risk Factor

UTL **Upper Tolerance Limit**

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ACKNOWLEDGMENTS

This manual was developed by EPA's Office of Emergency and Remedial Response. A large number of EPA regional technical staff participated in the Workgroup that developed the final RAGS Part D approach presented in this manual.

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PREFACE

Risk Assessment Guidance for Superfund: Volume I -- Human Health Evaluation Manual (RAGS/HHEM) Part D is the fourth part in the five-part series of guidance manuals on Superfund human health risk assessment. Part A addresses the baseline risk assessment; Part B addresses the development of risk-based preliminary remediation goals; Part C addresses the human health risk evaluations of remedial alternatives; and Part E addresses dermal exposure. Part D provides guidance on risk assessment planning, reporting, and review throughout the CERCLA remedial process, from scoping through remedy selection and completion and periodic review of the remedial action. Thus, Part D strives for effective and efficient implementation of Superfund risk assessment practice described in Parts A, B, C, and E, and in supplemental Office of Solid Waste and Emergency Response (OSWER) directives and other Agency risk assessment guidance. The potential users of Part D are persons involved in the risk evaluation, remedy selection, and implementation process, including risk assessors, risk assessment reviewers, remedial project managers, and other decisionmakers.

Released in January 1998 as interim guidance, RAGS Part D Revision 0 underwent field testing and evaluation for a 3-year period. This Final guidance considers the comments received from users of the Revision 0 guidance and provides Planning Table format changes as appropriate.

Generally, changes were made to improve useability, transparency, clarity, and/or consistency with other risk guidance (e.g., RAGS Part E dermal guidance [U.S. EPA, 2001], adult lead exposures technical fact sheet [U.S. EPA, 1996d], and Record of Decision guidance [U.S. EPA, 1999a]). These changes may also increase the efficiency of the risk assessor by decreasing the number of versions of each Planning Tables associated with certain sites.

In addition to Planning Table format changes, the Final guidance provides planning formats to document radionuclide and lead risk evaluations, neither of which was addressed in the Revision 0 guidance. The Final guidance also provides more robust and diverse examples than were included in Revision 0. These examples address comments and questions received from users of the Revision 0 guidance and are provided as suggested approaches to address complex situations. In all cases, the EPA regional risk assessor should be consulted to discuss the appropriate approach for a site.

This guidance does <u>not</u> discuss standardization of ecological risk assessments. EPA will provide planning tables for ecological evaluation under separate cover. This guidance does <u>not</u> discuss the risk management decisions that are necessary at a CERCLA site (e.g., selection of final remediation goals).

Upon issuance, RAGS Part D Final will be effective for all new CERCLA risk assessments. Consult the EPA risk assessor for applicability of the final guidance to ongoing risk assessments and non-CERCLA risk assessments. Any updates to this guidance will be posted at the RAGS Part D website at http://www.epa.gov/superfund/programs/risk/ragsd/index.htm.

Comments addressing usefulness, changes, and additional areas where guidance is needed should be addressed to the RAGS Part D website or to:

Senior Process Manager for Risk (RAGS Part D)
U.S. Environmental Protection Agency
Office of Emergency and Remedial Response (5202G)
Ariel Rios Building
1200 Pennsylvania Ave. NW
Washington, DC 20460

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CHAPTER 1

INTRODUCTION

This guidance has been developed by the U.S. Environmental Protection Agency (EPA) to assist remedial project managers (RPMs), risk assessors, site engineers, and others in conducting risk assessment planning, reporting, and review at Comprehensive Environmental Response Compensation and Liability Act (CERCLA) sites. This guidance could also be a useful tool for quantitative risk assessment for non-National Priorities List (Non-NPL), Base Realignment and Closure (BRAC), and Brownfields sites.

This guidance is the fourth part (Part D) in the five-part series Risk Assessment Guidance for Superfund: Volume I -- Human Health Evaluation Manual (RAGS/HHEM) (U.S. EPA, 1989c). Part A of this guidance addresses how to conduct a site-specific baseline risk assessment: the information in Part A is important background for Part D. Part B provides guidance for calculating risk-based concentrations that may be used, along with applicable or relevant and appropriate requirements (ARARs) and other information, to develop preliminary remediation goals (PRGs) during project scoping. PRGs (and final remediation levels set in the Record of Decision [ROD]) can be used throughout the analyses in Part C to assist in evaluating the human health risks of remedial alternatives. Part E provides guidance for evaluation of dermal exposure. Part D complements the guidance provided in Parts A, B, C, and E and presents recommended approaches to standardize risk assessment planning, reporting, and review. Part D guidance spans the CERCLA remedial process from project scoping to periodic review of the implemented remedial action. Exhibit 1-1 illustrates the major correspondence of RAGS/HHEM activities with the steps in the CERCLA remedial process.

The remainder of this chapter:

- presents an overview of Part D, including the background and elements of the Part D approach
- describes the applicability of Part D
- presents the organization of the remainder of

- this document
- describes where to find additional information regarding Part D.

1.1 OVERVIEW OF PART D

1.1.1 BACKGROUND

The March 21, 1995, memorandum on Risk Characterization Policy and Guidance from former EPA Administrator Browner directed improvement in the transparency, clarity, consistency, and reasonableness of risk assessments at EPA. EPA, over the years, has identified opportunities for improvement in presentation of Superfund risk assessments. Furthermore, the General Accounting Office (GAO), members of Congress, and others have called for betterment of Superfund risk The October 1995 Superfund assessments. Administrative Reform #6A directed EPA to: Establish National Criteria to Plan, Report, and Review Superfund Risk Assessments. EPA has developed an approach to respond to these challenges, which is presented in RAGS Part D.

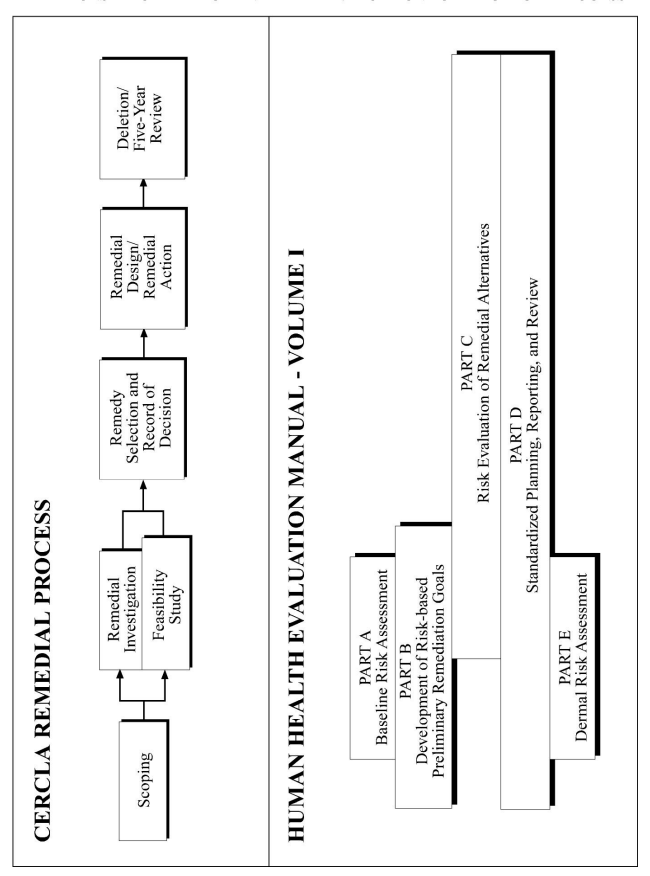
1.1.2 GUIDANCE CHANGES

Released in January 1998 as interim guidance, RAGS Part D Revision 0 underwent field testing and evaluation for a 3-year period. This Final guidance incorporates changes based on the comments received from users of the Revision 0 guidance and provides recommended Planning Table format changes as appropriate.

Generally, changes were made to improve useability, transparency, clarity, or consistency with other risk guidance (e.g., RAGS Part E dermal guidance [U.S. EPA, 2001] and ROD guidance [U.S. EPA, 1999a]). These changes may

also increase the efficiency of the risk assessor by decreasing the number of versions of each Planning Table associated with certain sites.

EXHIBIT 1-1 RELATIONSHIP OF THE HUMAN HEALTH EVALUATION TO THE CERCLA PROCESS



In addition to Planning Table format changes, the Final guidance provides standard formats to document radionuclide and lead risk evaluations, neither of which was addressed in the Revision 0 guidance. This final guidance also provides more robust and diverse examples than were included in Revision 0. These examples address comments and questions received from users of the Revision 0 guidance and are provided as suggested approaches to address complex situations. In all cases, the EPA risk assessor and the RPM (when appropriate) should be consulted to discuss the appropriate approach for a site. Revisions associated with each Planning Table may be found in Exhibit 3-3.

1.1.3 ELEMENTS OF PART DAPPROACH

The Risk Assessment Guidance for Superfund (RAGS) Part D approach consists of three basic elements: Use of Planning Tools, Continuous Involvement of EPA Risk Assessors, and Information Transfer to a National Superfund Risk Data Repository. Brief descriptions of the three components follow:

- Use of Planning Tools The Planning Tools developed by the EPA RAGS Part D Workgroup and refined through regional review include a Technical Approach for Risk Assessment or TARA, Planning Tables, and Instructions for the Planning Tables.
 - The Technical Approach for Risk Assessment (TARA) is a road map for incorporating continuous involvement of the EPA risk assessor throughout the CERCLA remedial process for a particular site. Risk-related activities, beginning with scoping and problem formulation, extending through collection and analysis of risk-related data, and supporting risk management decision making and remedial design/remedial action issues are addressed.

Chapters 2 through 5 of this guidance document present the TARA in the four CERCLA remedial process phases: During Scoping, During the Remedial Investigation, During the Feasibility Study, and After the Feasibility Study. It

- is recommended that the elements identified in the TARA in Chapters 2 through 5 be customized for each site-specific human health risk assessment, as appropriate. These elements should be included in project workplans to better define that risk assessment and facilitate more standardized planning. A planning worksheet that can be used to summarize the TARA for a particular site (the TARA Schedule Worksheet) is found in Appendix C.
- The Planning Tables have been developed to more clearly and consistently document important parameters, data, calculations, and conclusions from all stages of human health risk assessment development. Electronic templates for the Planning Tables have been developed in Lotus® and Excel® for ease of use by risk For site-specific risk assessors. assessments, the Planning Tables, related Worksheets, and Supporting Information should first be prepared as Interim Deliverables for EPA risk assessor review, and should later be included in the Draft and Final Baseline Risk Assessment Reports. The Planning Tables, both a blank set and a fully completed example set, may be found in Appendix A. Additional example scenarios and selected Planning Tables are provided in Appendix D. Use of the Planning Tables will help standardize the reporting of human health risk assessments and improve communication with stakeholders.
- -- Instructions for the Planning Tables have been prepared corresponding to each row and column on each Planning Table. Definitions of each field are supplied in the Glossary and example data or selections for individual data fields are provided. The Instructions should be used to complete and/or review Planning Tables for each site-specific human health risk assessment, where appropriate. The Instructions may be found in Appendix B.
- Continuous Involvement of EPA Risk Assessors - The EPA risk assessor is a critical

participant in the CERCLA remedial process for any site, from scoping through completion and periodic review of the remedial action. EPA risk assessors support reasonable and consistent risk analysis and risk-based decision making. Early and continuous involvement by the EPA risk assessors should include scoping, workplan review, and customization of the TARA for each site to identify all risk-related requirements. The EPA risk assessors should review Interim Deliverables and identify corrections needed prior to preparation of the Draft and Final Baseline Risk Assessment Reports. Participation of the EPA risk assessors in all other phases of the CERCLA remedial process will help ensure human health risk issues are appropriately incorporated in the remedy selection and implementation processes.

 Information Transfer to a Superfund Risk Data Collection - Summary-level site-specific risk information should be contained in a Superfund Risk Data Repository to provide information access and evaluation capabilities to EPA staff.

1.2 APPLICABILITY OF PART D APPROACH

The approach contained in RAGS Part D is strongly recommended for all CERCLA human health risk assessments.

Exhibit 1-2 provides guidelines regarding RAGS Part D applicability as a function of site lead and site type, so that site-specific applicability may be defined by each region.

A brief discussion of the process improvements associated with each RAGS Part D element follows:

- Use of Planning Tools Planning Tools facilitate planning with TARA, reporting with Planning Table formats, and reviewing with Interim Deliverables. The Planning Tools are designed to provide more consistent content and clarity of data, parameters, and assumptions. Transparency for the public and others to understand the risk assessment should be improved by the Planning Tables, and review is facilitated because the basis for conclusions should be more clear. Because Interim Deliverables are integral parts of the baseline risk assessment, their early review and resolution by EPA risk assessors should minimize rework and may reduce project schedules and budgets, while improving consistency.
- Continuous Involvement of EPA Risk Assessor - Involvement of the EPA risk assessor throughout the CERCLA remedial process should result in holistic consideration of risk issues during scoping and helps ensure that appropriate and adequate data are collected. Planning for special evaluations can also be conducted efficiently at project inception rather than at a later point with associated schedule delays and additional Ongoing review of Interim costs. Deliverables by the EPA risk assessor should provide direction regarding reasonable assumptions and should eliminate rework requirements, particularly for those deliverables that build on previous analyses (e.g., the Baseline Risk Assessment Report).

1.3 PROCESS IMPROVEMENTS RESULTING FROM PART D APPROACH

The RAGS Part D approach provides advantages over previous practices in the Superfund program at both the site level and the overall Superfund program level.

Exhibit 1-2 goes here

At later stages of the project (e.g., after the feasibility study), continuous involvement of risk EPA assessor promotes reasonableness and consistency in risk management decision-making by clearly providing risk managers with the information they need. Preparation of draft ROD risk information as an interim deliverable in the format specified in Guide to Preparing Superfund Proposed Plans, Records of Decision, and Other Remedy Selection Decision Documents (U.S. EPA, 1999a) will further support risk managers' efficiency. The ROD Risk Worksheets found in Appendix C match the ROD guidance formats.

• Information Transfer to Superfund Risk Data Collection - Submission of the electronic Planning Tables and Worksheets to the Superfund Risk Data Collection fulfills the review objectives of Superfund Administrative Reform #6A. Use of the information by EPA risk assessors will help improve consistency in future risk assessments.

1.4 ORGANIZATION OF DOCUMENT

The remainder of this guidance is organized into four additional chapters, references, and four appendices as follows:

- Chapter 2: Risk Considerations During Project Scoping;
- Chapter 3: Risk Assessment Data Needs and Tasks During the Remedial Investigation;
- Chapter 4 Risk Evaluations During the Feasibility Study;
- Chapter 5: Risk Evaluations After the Feasibility Study;
- References

- **Planning Tables**
- Appendix C: Worksheets
- Appendix D: Example Scenarios.

In addition, other useful information has been presented in highlight boxes placed throughout the document.

Exhibit 1-3 depicts the continuous involvement of the EPA risk assessor during scoping, during the remedial investigation, and during and after the feasibility study. The various activities the risk assessor conducts are listed, as well as the Part D chapter that addresses that phase.

1.5 ADDITIONAL INFORMATION

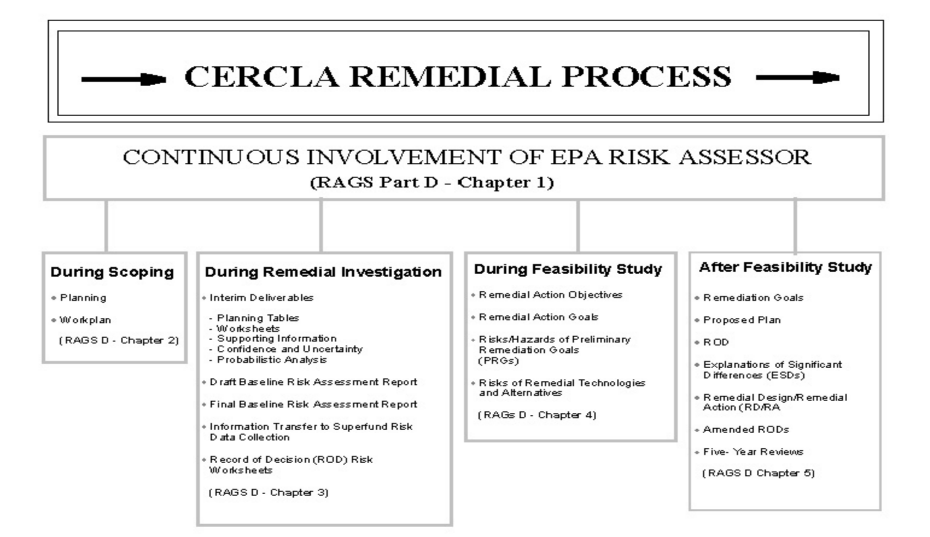
This guidance will be updated periodically in response to user comments and suggestions and to address new human health risk assessment guidance as appropriate.

The Part D guidance and corresponding information may be accessed electronically on the RAGS Part D website, at http://www.epa.gov/superfund/programs/risk/ragsd/index.htm. Updates to Part D will also appear on the website along with an index of the current version of each Chapter or Appendix.

Questions or comments regarding Part D usage for a particular risk assessment should be directed to your EPA risk assessor. General Part D questions or comments should be directed to the RAGS Part D website. Questions or comments received through the website will be considered and a response will be developed and forwarded via telephone or email as appropriate. Frequently asked questions will be assembled and displayed on the website with corresponding responses to provide Part D user support.

- Appendix A: Planning Tables
- Appendix B: Instructions for Completion of

EXHIBIT 1-3 ROLE OF RISK ASSESSOR IN THE CERCLA REMEDIAL PROCESS



CHAPTER 2

RISK CONSIDERATIONS DURING PROJECT SCOPING

The project scoping stage of the remedial investigation (RI) and baseline risk assessment is critical to the success of a Superfund project. The EPA risk assessor should be involved in the project scoping discussions and meetings to help ensure that the planning and workplan development tasks incorporate risk assessment data needs and achieve appropriate standardization in risk assessment planning.

2.1 PLANNING

The following planning activities should be performed at the beginning of the project. These activities should involve the EPA RPM and EPA risk assessor, as decisionmakers, and the risk assessment author and other resources tasked with preparing the Remedial Investigation Report, to support planning. The following pertinent information should be incorporated, as appropriate, into the Remedial Investigation Report or Site Characterization Report and the Baseline Risk Assessment Report:

- Provide site background information, site maps, sample location map; discuss historical site activity and chronology of land use.
- Discuss historical data and data useability, previous studies and actions, and an overview of the nature and extent of contamination.
- Discuss the purpose of the investigation.
- Prepare the preliminary site conceptual model which clearly identifies all known or potential sources of contamination (soil, groundwater, surface water, leachate, air, etc.), release mechanisms, and receptor routes and identifies all potential exposure pathways (including secondary pathways) and the media and receptors associated with each.
- Discuss PRGs and ARARs for the site.

Discuss involvement by the risk assessor in

WHEN PREPARING THE SITE CONCEPTUAL MODEL, CONSIDER THE FOLLOWING:

- Sensitive populations, including but not limited to the elderly, pregnant or nursing women, infants and children, and people suffering from chronic illnesses
- People exposed to particularly high levels of contaminants
- Circumstances where a disadvantaged population is exposed to hazardous materials (i.e., Environmental Justice situations)
- Significant contamination sources
- Potential contaminant release mechanisms (e.g., volatilization, fugitive dust emission, surface runoff/overland flow, leaching to groundwater, tracking by humans/animals, soil gas generation, biodegradation and radioactive decay)
- Contaminant transport pathways such as direct air transport downwind, diffusion in surface water, surface water flow, groundwater flow, soil gas migration, and biomagnification in the food chain
- Cross media transfer effects, such as volatilization to air, wet deposition, dry deposition, groundwater discharge to surface water, groundwater recharge from surface water, and bioaccumulation by aquatic species.

discussions with stakeholders concerning land

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use, groundwater use, and exposure pathways and variables. If possible, the risk assessor should also visit the site.

• Identify interim deliverables for the risk assessment.

INTERIM DELIVERABLES SHOULD INCLUDE THE FOLLOWING:

- Planning Tables 0 through 10
- Worksheets on Data Useability, TARA Schedule, Dermal, Radiation Dose Assessment, and Lead (as applicable)
- Supporting Information (Section 3.1.1)
- Assessment of Confidence and Uncertainty (Section 3.1.2) and Probabilistic Analysis information, as applicable (Section 3.1.3).
- Identify Draft and Final deliverables for the risk assessment. Draft and Final deliverables include the Draft and Final Baseline Risk Assessment Reports, which also incorporate the Interim Deliverables.
- Prepare a preliminary version of Planning Table
 1.
- During project scoping, the EPA RPM and EPA risk assessor may also meet to discuss the potential usefulness of including a Probabilistic Analysis (Monte Carlo) in the RI and the need for a separate Workplan. This preliminary discussion should address whether funds need to be allocated to carry out a Probabilistic Analysis. This decision should be revisited throughout Workplan development and the risk assessment process.

2.2 WORKPLAN DEVELOPMENT

Tasks to be conducted during the remedial investigation/feasibility study (RI/FS) should be identified and documented in several workplans. These usually include the RI/FS Workplan, a Sampling and Analysis Plan (SAP), and a Quality Assurance Project Plan (QAPP). Tasks related to

development of the baseline risk assessment are sometimes presented in a separate Risk Assessment Workplan or incorporated into the RI/FS Workplan.

WHEN EVALUATING WHETHER TO CONDUCT PROBABILISTIC ANALYSIS, CONSIDER THE FOLLOWING:

- Extent of site remediation
- Potential costs of remediation
- Degree of uncertainty associated with the exposure information available for each portion of the site conceptual model

Risk assessment needs should be considered not only in tasks related to development of the baseline risk assessment but also in tasks related to sampling and analysis (i.e., those in the SAP and the QAPP) in the RI and tasks needing risk assessment input in the feasibility study(e.g., development of remedial goals and estimates of potential risk from remediation options).

2.2.1 RI/FS WORKPLAN/BASELINE RISK ASSESSMENT WORKPLAN

The RI/FS Workplan should summarize site background, the current and potential problems posed by site contaminants, and the objectives and scope of the RI/FS. It also should include a description of the tasks to be performed and the information and work products that should be produced from each task. Deliverables for specific tasks should be included. Tasks and deliverables for the baseline risk assessment may be included as a part of the RI/FS Workplan or in a separate Risk Assessment Workplan.

Within these Workplans, it should be clear that risk assessment needs are being considered in the RI/FS objectives. The site-specific objectives and scope of the risk assessment should be included in the Workplan.

This includes information to complete the baseline risk assessment in the RI as well as information for

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the FS, such as that used to develop risk-based preliminary remedial goals (e.g., PRGs), and to assess risks from remediation (e.g., incineration).

These Workplans should also reference the methods (e.g., National guidance such as RAGS/HHEM [U.S. EPA, 1989c]; RAGS Probabilistic Guidance [U.S. EPA, 1997e and g and 2001d.]), used to prepare the Interim, Draft, and Final risk assessment deliverables and define the schedule for submission. These deliverables are described in more detail in Chapter 3. Deliverables related to development of risk-based remedial goals and assessment of risk from remediation should also be included in the Workplan (see Chapter 4).

The EPA risk assessor and EPA RPM may revisit the question of the potential value added by using Probabilistic Analyses in the risk assessment. If these analyses are to be used, the issues concerning the time, expense, and possible benefit associated with the collection of additional exposure information or sampling data should be considered to identify those exposure parameters with the greatest uncertainty, where collection of additional data and/or information may be warranted. A separate Probabilistic Analysis Workplan identifying associated deliverables should be prepared and approved by the EPA RPM and risk assessor.

2.2.2 SAP AND QAPP

Sampling and analysis activities undertaken during the RI should provide adequate data to evaluate all appropriate exposure pathways. Therefore, risk assessors should be involved in the development of the data quality objectives (DQOs) for sampling and analysis and in selecting the types of sampling and analyses that will be done. The DQOs should address the qualitative and quantitative nature of the sampling data in terms of relative quality and intent for use, to ensure that the data collected will be appropriate for the intended objectives. Note that the data quality evaluation should be recorded in the Data Useability Worksheet in Appendix C.

Sampling. The SAP should discuss how the types, numbers, and locations of samples to be collected will be adequate to evaluate each exposure

pathway (both current and future) and medium. The SAP should be accompanied by detailed sampling maps showing the location and type of samples (e.g., grab, composite, or duplicate). It is important to consider how sample results will be used to estimate exposure point concentrations. Background samples should be collected from appropriate areas (e.g., areas proximate to the site, free of potential contamination by site chemicals and similar to the site in topography, geology, meteorology, and other characteristics).

If models will be used to evaluate exposure pathways and estimate exposure point concentrations, these models should be identified in the Workplan. Site-specific data collection needed for these models should also be discussed.

WHEN DEVELOPING THE SAP, CONSIDER THE FOLLOWING:

- How will data from multiple groundwater wells collected over time be used to calculate exposure?
- At what depths will soil samples be taken and how will they be combined to describe exposures for different scenarios (e.g., industrial versus residential) or to characterize hotspots?
- What type of sampling design (e.g., random versus purposive) will be used?
- Are SAPs adequate to distinguish site contamination from background contamination for each medium and for organic and inorganic parameters?

Analysis. Development of the DQOs for analysis should not be limited to concern for the precision, accuracy, representativeness, completeness, and comparability of the data. DQOs that are important for risk assessment should consider: types of laboratory analyses used, sensitivity of detection limits of the analytical techniques (especially for non-Target Compound List [non-TCL] chemicals and non-standard matrices), resulting data quality, and the employment of adequate quality assurance/quality

control (QA/QC) measures.

In some cases, risk assessment data needs may be best supported by additional chemicals, different analytical methods, and/or lower detection limits than are being used for the RI. Based upon the values of the risk-based PRGs calculated during scoping, detection limits may need to be lower than those obtained by the standard Superfund methods. The adequacy of detection limits for conducting the baseline risk assessment and for comparing to PRGs should be evaluated in the Workplan (QAPP). For example, a table listing expected contaminants and comparing the method detection limit or quantitation limit for each compound with the

Analytical data should be evaluated and reviewed in accordance with the criteria to evaluate data (e.g., the National Functional Guidelines). Also refer to your regional Agency office for guidance on data validation and/or other chemical-specific guidance, as applicable.

The Workplan should also discuss how split samples, duplicates, blanks (trip, field, and laboratory), and qualified and rejected data can be used in assessing site risks. The Workplan should describe the analysis for each medium and how the types of analyses were selected based on site history.

appropriate risk-based goal for that chemical could be presented. This information along with issues of cost and other data uses should affect the methods and detection limits finally selected.

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CHAPTER 3

RISK ASSESSMENT DATA AND TASKS DURING THE REMEDIAL INVESTIGATION

Project Management Guidelines. Remedial project managers should establish the schedule of submission for the deliverables for the RI Reports and Baseline Risk Assessment Reports. The schedule may vary from site to site, as appropriate. Interested parties (States, Commonwealths, tribes and other stakeholders) may be involved in the scheduling and review process, as appropriate. Refer to your regional office for guidance regarding the order of the deliverables. These deliverables should also be defined in the Workplan.

General RI Guidelines. Generally, RI guidance should be followed in performing the remedial investigation. The following items are of particular importance to risk assessments. If the risk assessment is being prepared as a stand-alone document, the following items should be included. If, instead, the risk assessment is a section of the RI Report, the items which follow should be addressed in the RI Report and clearly referenced in the Baseline Risk Assessment Report.

- Present a general map of the site depicting boundaries and surface topography, which illustrates site features, such as fences, ponds, structures, as well as geographical relationships between potential receptors and the site
- Discuss historical site activity.
- Discuss chronology of land use (specify agriculture, industry, recreation, waste deposition, and residential development at the site).
- Present an overview of the nature and extent of contamination, including when samples were collected and the kinds of contaminants and media potentially contaminated.
- Describe the analytical and data validation methods used.

 If modeling was used to estimate exposure point concentrations, document the parameters related to soil/sediment, hydrogeology, hydrology, and meteorology either in the risk assessment or the RI Report.

Risk Assessment Guidelines. The risk assessment should be conducted in accordance with all appropriate guidance and policies. Consult with your EPA risk assessor regarding the most appropriate guidance.

Interim Deliverables should be prepared as described in Section 3.1.1 and should ultimately be incorporated into the Baseline Risk Assessment Report. The Interim Deliverables prepared by the risk assessment author should be reviewed by the EPA risk assessor prior to submission of the Baseline Risk Assessment Report. Hazard identification and exposure parameters, among others, may require discussion, refinement, and revision. Review and modification of Interim Deliverables should greatly reduce the Baseline Risk Assessment Report preparation and review time. Discussions of the three categories of risk assessment deliverables (Interim Deliverables, Draft Baseline Risk Assessment Report, and Final Baseline Risk Assessment Report) follow.

3.1 INTERIM DELIVERABLES

This section presents an outline of the Planning Tables, Worksheets, and Supporting Recommended Information that should be prepared as Interim Deliverables for each site. The Workplan discussed in Section 2.2.1 should also describe the Planning Tables, Worksheets, and Supporting Recommended Information for a particular site. Exhibit 3-1 presents a list of recommended Interim Deliverables. Use of these deliverables for each site should improve standardization in risk assessment reporting and

should improve the transparency, clarity, and consistency of risk assessments.

3.1.1 PLANNING TABLES, WORKSHEETS, AND SUPPORTING INFORMATION

More standardized reporting of Superfund human health risk assessments can be achieved through the preparation of Planning Tables, Worksheets, and Supporting Information. These documents should be prepared as Interim Deliverables and reviewed by the EPA risk assessor prior to preparation of the Baseline Risk Assessment Report. After review and revision, as necessary, these documents should be included in the Baseline Risk Assessment Report.

This section describes the Planning Table formats that should be used in EPA CERCLA risk assessments. The Planning Table formats normally should not be altered (i.e., columns should not be added, deleted, or changed); however, rows and footnotes should be added as appropriate. Standardization of the Tables should help to achieve Superfund program-wide reporting consistency. Note that multiple versions of some Planning Tables may be used to address different Media, different Exposure Pathways, or different Exposures (i.e., reasonable maximum exposure [RME] versus central tendency [CT]). Exhibit 3-2 summarizes the relationship between five traditional risk assessment activities and the corresponding Planning Tables that should help standardize risk assessment reporting. The five risk assessment activities follow:

- Data collection
- Data evaluation
- Exposure assessment
- Toxicity assessment
- Risk characterization.

Copies of the blank Planning Tables are provided in both Lotus® and Excel® spreadsheet formats associated with the Part D guidance. Blank Planning Table templates and completed examples of typical Planning Tables are provided in Appendix A. Detailed Instructions for the completion of the Planning Tables are provided in

Appendix B. Additional example scenarios and selected Planning Tables are provided in Appendix D.

In addition to the Planning Tables, six Planning Worksheets are provided in Appendix C. These include Worksheets for Data Useability, TARA Schedule, Dermal, Radiation Dose Assessment, Lead, and ROD Risk. Use of the Worksheets is strongly encouraged to improve transparency, clarity, and consistency.

The Planning Tables and Worksheets document the majority of the data and assumptions used to evaluate risk, as well as the risks and hazards calculated. In most cases, other data and rationale can be used to support the information presented in the Planning Tables. This additional Supporting Information should also be provided to the EPA risk assessor as an Interim Deliverable and later incorporated in the Baseline Risk Assessment Report.

Refer to Exhibit 3-3 for a brief summary of the Revision 1 improvements to the Planning Tables and Worksheets as compared to Revision 0. Descriptions of the RAGS Part D Revision 1 Planning Tables, Worksheets, and Supporting Information follow:

Planning TABLE 0: Site Risk Assessment Identification Information. The purposes of Planning Table 0 are:

- To uniquely identify the risk assessment
- To identify the relevant contacts for the risk assessment.

The information documented in **Planning Table 0** should include:

- Site Information
- Contact information
- Risk assessment document information.

The data elements that should be presented in **Planning Table 0** are listed in the Planning Table 0 highlight box.

KEY DATA ELEMENTS IN PLANNING TABLE 0

Regions should provide the following information: Site Name/OU, Region, EPA ID Number, State, Status, Federal Facility (Y/N), EPA Project Manager, EPA Risk Assessor, Prepared by, Prepared for, Document Title, Document Date, Probabilistic Risk Assessment (Y/N), and Comments.

Regions should perform the following steps associated with the preparation of **Planning Table 0:**

- 1. Provide the identification information for the risk assessment.
- 2. Include Planning Table 0 with the other Planning Tables, Worksheets, and Supporting Information to facilitate tracking of the relevant contacts.

TARA SCHEDULE WORKSHEET. The TARA Schedule of Risk-Related Activities Worksheet (TARA Schedule Worksheet) is the first Worksheet that should be developed for each risk assessment to document the applicability, responsibility, and schedule for each risk-related activity. As the first interim deliverable, the Worksheet documents the plan for a particular site, identifying which Planning Tables, Worksheets, and Supporting Information should be provided as interim deliverables for EPA risk assessor review, and when they are expected to be available. The TARA Schedule Worksheet should be prepared in consultation with the EPA risk assessor assigned to the site.

Regions should perform the following steps associated with the preparation of the TARA Schedule Worksheet:

1. Complete the TARA Schedule Worksheet prior to initiation of any other Planning Tables, Worksheets, or Supporting Information.

2. **Obtain EPA risk assessor consensus** regarding which interim deliverables should be submitted and the schedules for each.

The recommended blank TARA Schedule Worksheet may be found in Appendix C. An example TARA Schedule Worksheet accompanies the Dean Company example in Appendix A.

PLANNING TABLE 1: Selection of Exposure Pathways. The purposes of **Planning Table 1** are:

- To assist in project planning
- To accompany the site conceptual model
- To present possible Receptors, Exposure Routes, and Exposure Pathways
- To present the rationale for selection or exclusion of each Exposure Pathway
- To communicate risk information to interested parties outside EPA
- To establish a framework for the generation of subsequent Planning Tables. All subsequent tables should be built from the information contained in Planning Table 1.

The information that should be documented in **Planning Table 1** includes:

- Exposure Pathways that were examined and excluded from analysis
- Exposure Pathways that are expected to be qualitatively or quantitatively evaluated in the risk assessment.

The data elements that should be presented in **Planning Table 1** are listed in the Planning Table 1 highlight box.

KEY DATA ELEMENTS IN PLANNING TABLE 1

Regions should provide the following information: Scenario Timeframe, Medium, Exposure Medium, Exposure Point, Receptor Population, Receptor Age, Exposure Route, Type of Analysis, Rationale for Selection or Exclusion of Exposure Pathway.

Region should perform the following steps associated with the preparation of **Planning Table 1**:

- Refine site conceptual model which identifies all potential sources of contamination, all potential Exposure Pathways, the Medium associated with each, and the potentially exposed populations (Receptors).
- 2. Select realistic Exposure Pathways for detailed analyses.
- 3. Include rationale for exclusion of potential Exposure Pathways.
- 4. Modify Planning Table 1, where appropriate.
- Planning Table 1 should later be incorporated in the Baseline Risk Assessment Report.

DATA USEABILITY WORKSHEET.

Data quality is an important component of the risk assessment and the evaluation of data quality should be documented. A recommended Data Useability Worksheet is included to address this need.

The Regional EPA risk assessor and the EPA document Guidance for Data Useability in Risk Assessment (Part A, U.S. EPA 1990a), should be consulted before completing the Data Useability Worksheet to define the appropriate level of detail to be reflected in the comment fields in the Worksheet. This Worksheet should be prepared as soon as all data validation reports have been completed for each medium. A medium-specific Data Useability Worksheet should be completed only after the project team (i.e., lead chemist, lead hydrogeologist, risk assessor, etc.) has collectively discussed the data useability criteria. Worksheet should be used to record and identify the impact of data quality issues as they relate to data useability. For example, deviations from approved site Workplans which occurred during sample collection, laboratory analysis, or data review should be assessed. Also, the Worksheet preparer should refer to the Superfund regional office for guidance on data validation when preparing the Worksheet.

Regions should perform the following steps associated with the preparation of the **Data Useability Worksheet**:

- 1. **Complete the** *Data Useability Worksheet* for each Medium prior to screening of chemicals of potential concern (COPCs).
- 2. Incorporate the **Data Useability Worksheet** in the Baseline Risk Assessment Report.

A recommended blank Data Useability Worksheet may be found in Appendix C. An example Data Useability Worksheet accompanies the Dean Company example in Appendix A.

PLANNING TABLE 2: Occurrence, Distribution, and Selection of COPCs. The purposes of Planning Table 2 are:

- To provide information useful for data evaluation of chemicals and radionuclides detected
- To provide adequate information so the user/reviewer gets a sense of the chemicals and radionuclides detected at the site and the potential magnitude of the potential problems at the site
- To provide chemical screening data and rationale for selection of COPCs.

The information documented in **Planning Table 2** should include:

- Statistical information about chemicals and radionuclides detected in each Medium
- The detection limits of chemicals and radionuclides analyzed
- The toxicity screening values for COPC selection
- The chemicals and radionuclides selected and deleted as COPCs.

The data elements presented in **Planning Table 2** are listed in the Planning Table 2 highlight box.

Regions should perform the following steps

associated with the preparation of **Planning Table**2. Refer to the regional office for guidance when performing these steps.

KEY DATA ELEMENTS IN PLANNING TABLE 2

For each unique combination of Scenario Timeframe, Medium, and Exposure Medium, Regions should provide the following information: Exposure Point, CAS Number, Chemical, Minimum Concentration (Qualifier), Maximum Concentration (Qualifier), Units, Location of Maximum Concentration, Detection Frequency, Range of Detection Limits, Concentration Used for Screening, Background Value, Screening Toxicity Value (N/C), Potential ARAR/TBC Value, Potential ARAR/TBC Source, COPC Flag (Y/N), and Rationale for Selection or Deletion.

- 1. Discuss selection criteria for COPCs; including toxicity screening values, frequency of detection, and background comparison, as appropriate.
- Perform screening; select COPCs that will be carried into the risk assessment (include comparison to regulatory standards and criteria where appropriate).
- 3. Submit Supporting Information to substantiate the available Background Value shown for each chemical in Planning Table 2 and to enable verification of those values by EPA. The format of the summary should be determined by each region. The Supporting Information should provide relevant information for each chemical used to determine the background concentration, including (but not limited to) average, maximum, hypothesis testing of equality of the mean, and other information that may be required to fully describe the background selection process.
- 4. Incorporate the Background Supporting Information in the Baseline Risk Assessment Report.

- 5. Complete Planning Table 2 for each combination of Scenario Timeframe, Medium, and Exposure Medium.
- 6. **Incorporate Planning Table 2** in the Baseline Risk Assessment Report.

PLANNING TABLE 3: Exposure Point Concentration Summary. The purposes of Planning Table 3 are:

- To provide the EPCs for measured and modeled values
- To provide statistical information on the derivation of the EPCs.

The information documented in **Planning Table 3** should include:

- Statistical information which was used to calculate the EPCs for chemicals and radionuclides detected in each Medium
- EPCs (RME and/or CT)
- The statistics which were used to make the determinations as well as the rationale for the selection of the statistics for each chemical or radionuclide (i.e., discuss statistical derivation of measured data or approach for modeled data).

The data elements presented in **Planning Table 3** are listed in the Planning Table 3 highlight box.

KEY DATA ELEMENTS IN PLANNING TABLE 3

For each unique combination of Scenario Timeframe, Medium, and Exposure Medium, Regions should provide the following information: Exposure Point, Chemical of Potential Concern, Units, Arithmetic Mean, 95% upper confidence level (UCL), Maximum Concentration (Qualifier), EPC Value, EPC Units, EPC Statistic, and EPC Rationale.

Region should perform the following steps associated with the preparation of Planning Table 3

- 1. Discuss how samples will be grouped (e.g., how hot spots in soil will be considered; how groundwater data will be combined; how temporal and chemical phases will be addressed; how upgradient, downgradient, and cross gradient samples will be addressed).
- 2. Discuss approach to determine how data are distributed (e.g., normal, log-normal).
- 3. Discuss evaluation of lead, total chromium and any other special chemicals.
- 4. Submit Supporting Information document the EPC summary presented in Planning Table 3 and to enable verification of those values by EPA. The format of the summary should be determined by each region. The Supporting Information should discuss EPCs statistically derived from measured data, including identification of the samples used in each calculation, results of distribution testing (Wilk-Shapiro, D'Agostino), mean (transformed if appropriate), maximum (transformed if appropriate), Planning deviation (transformed if appropriate), t- or H-statistic, 95% UCL (including non-parametric methods, where applicable), and other protocols as required. The Supporting Information should also present information for EPCs, including derivation of modeled values, assumptions and values used, statistical derivation of measured values and associated calculations, and other protocols as required.
- 5. Incorporate the **EPC Supporting Information** in the Baseline Risk Assessment Report.
- 6. **Complete Planning Table 3** for each combination of Scenario Timeframe, Medium, Exposure Medium, and Exposure Point. Create separate sets of Planning Table 3 for RME and CT, when appropriate.
- 7. Incorporate **Planning Table 3** in the Baseline

Risk Assessment Report.

Planning TABLE 4: Values Used for Daily Intake Calculations. The purposes of Planning Table 4 are:

- To provide the exposure parameters used for intake calculations for each Exposure Pathway (Scenario Timeframe, Medium, Exposure Medium, Exposure Point, Receptor Population, Receptor Age, and Exposure Route)
- To provide the intake equations or models used for each Exposure Route/Pathway.

The information documented in **Planning Table 4** should include:

- Values used for each intake equation for each Exposure Pathway and the reference/rationale for each
- Intake equation or model used to calculate the intake for each Exposure Pathway.

The data elements presented in **Planning Table 4** are listed in the Planning Table 4 highlight box.

KEY DATA ELEMENTS IN PLANNING TABLE 4

For each unique combination of Scenario Timeframe, Medium, and Exposure Medium, Regions should provide the following information: Exposure Route, Receptor Population, Receptor Age, Exposure Point, Parameter Code, Parameter (Definition, Value, and Units), Rationale/Reference, and Intake Equation/Model Name.

Regions should perform the following steps associated with the preparation of **Planning Table**

1. Provide references for all exposure

parameters.

- 2. Submit Supporting Information to summarize the Modeled Intake Methodology and Parameters used to calculate modeled intake values and to enable verification of those values by EPA. The Supporting Information should be limited to summary level information. The format of the summary should be structured to accommodate the variability and complexity associated with different models.
- 3. Incorporate the **Modeled Intake Supporting Information** in the Baseline Risk Assessment Report.
- 4. Submit **Supporting Information** Chemical-Specific Parameters, which apply to all Planning Tables to be completed for the risk assessment and to enable verification of those values by EPA. The summary should identify and display chemical parameters and constants that are used to calculate risks and hazards, but are not included on Planning Tables. The format of the summary should be determined by each region. The values and constants that are used to calculate risk and hazards, including molecular weight, vapor pressure, K_{oc}, K_{ow}, dermal permeability constant, Henry's Law constant, and other information that the reader would find useful for understanding the risk assessment discussion should be included.
- Incorporate the Chemical-Specific Parameter Supporting Information summary into the Baseline Risk Assessment Report.
- 6. **Complete Planning Table 4** for each combination of Scenario Timeframe, Medium, and Exposure Medium. Create separate sets of Planning Table 4 for RME and CT, where appropriate.
- 7. Incorporate **Planning Table 4** into the Baseline Risk Assessment Report.

DERMAL WORKSHEET. The recommended Dermal Worksheet presents intermediate variables for calculating absorbed dose per event DA (event). A version of this Worksheet should be developed for each medium for which the dermal exposure route will be quantitatively assessed. Available data should be provided for each COPC under evaluation.

Regions should perform the following steps associated with preparation of the **Dermal Worksheet**:

- Complete the Dermal Worksheet prior to calculation of risks and hazards.
- 2. Provide interim deliverables to the EPA risk assessor, as appropriate.
- 3. Incorporate the **Dermal Worksheet** in the Baseline Risk Assessment Report.

A recommended blank Dermal Worksheet may be found in Appendix C. An example Dermal Worksheet accompanies the Dean Company example in Appendix A.

PLANNING TABLES 5 AND 6: Non-Cancer and Cancer Toxicity Data. The purposes of Planning Tables 5.1, 5.2, and 5.3 are:

- To provide information on reference doses (RfDs), reference concentrations (RfCs), Target organs, and adjustment factors for chemicals
- To provide oral to dermal adjustment factors
- To provide RfC to RfD adjustment factors
- To verify references for non-cancer toxicity data
- To provide non-cancer toxicity information for "special-case" chemicals.

KEY DATA ELEMENTS IN PLANNING TABLE 5.1

Region should provide the following information: Chemical of Potential Concern, Chronic/Subchronic, Oral RfD Value and Units, Oral Absorption Efficiency for Dermal, Absorbed RfD for Dermal Value and Units, Primary Target Organ(s), Combined Uncertainty/Modifying Factors, Source(s) RfD: Target Organ(s), and Dates of RfD: Target Organ(s).

The information documented in **Planning Tables 5.1, 5.2, and 5.3** should include:

- The RfDs for each of the COPCs, as well as modifying factors and reference concentration (RfC) to RfD adjustments
- The organ effects of each of the COPCs
- References for RfCs and organ effects.

The data elements presented in **Planning Tables 5.1, 5.2, and 5.3** are listed in the Planning Tables 5.1, 5.2, and 5.3 highlight boxes.

KEY DATA ELEMENTS IN PLANNING TABLE 5.2

Regions should provide the following information: Chemical of Potential Concern, Chronic/Subchronic, Inhalation RfC Value and Units, Extrapolated RfD Value and Units, Primary Target Organ(s), Combined Uncertainty/Modifying Factors, Source(s) of RfC: Target Organ(s), and Date(s) of RfC: Target Organ(s).

KEY DATA ELEMENTS IN PLANNING TABLE 5.3

Regions should provide the following information: Chemical of Potential Concern, Chronic/Subchronic, Parameter Name, Value, and Units), Primary Target Organ(s), Combined Uncertainty/Modifying Factors, Source(s) of Parameter: Target Organ(s), and Date(s) of

The purposes of **Planning Tables 6.1, 6.2, 6.3, and 6.4** are:

- To provide the oral, dermal, and inhalation cancer toxicity information (values and sources of information) for chemicals and radionuclides of potential concern
- To provide the methodology and adjustment factors used to convert oral cancer toxicity values to dermal toxicity values and to convert inhalation unit risks to inhalation cancer slope factors
- To provide weight of evidence/cancer guideline descriptions for each chemical and radionuclide of potential concern
- To provide cancer toxicity information for "special case" chemicals.

The information documented in **Planning Tables 6.1, 6.2, 6.3, and 6.4** should include:

- Oral, dermal, and inhalation toxicity values for chemicals and radionuclides of potential concern
- Weight of evidence/cancer guidelines descriptions for chemicals of potential concern

• The source/reference for each toxicity value.

The data elements presented in **Planning Tables 6.1, 6.2, 6.3, and 6.4** are listed in the Planning Tables 6.1, 6.2, 6.3, and 6.4 highlight box.

KEY DATA ELEMENTS IN PLANNING TABLE 6.1

Regions should provide the following information: Chemical of Potential Concern, Oral Cancer Slope Factor Value and Units, Oral Absorption Efficiency for Dermal, Absorbed Cancer Slope Factor for Dermal Value and Units, Weight of Evidence/Cancer Guideline Description, Source(s) and Date(s) of Oral CSF.

KEY DATA ELEMENTS IN PLANNING TABLE 6.2

Regions should provide the following information: Chemical of Potential Concern, Unit Risk Value and Units, Inhalation Cancer Slope Factor Value and Units, Weight of Evidence/Cancer Guideline Description, Source(s) and Date(s) of Unit Risk: Inhalation CSF.

KEY DATA ELEMENTS IN PLANNING TABLE 6.3

Regions should provide the following information: Chemical of Potential Concern, Parameter (Name, Value, and Units), Source(s), and Dates(s).

KEY DATA ELEMENTS IN PLANNING TABLE 6.4

Regions should provide the following information: Chemical of Potential Concern, Cancer Slope Factor Value and Units, Source(s), and Dates(s).

Regions should perform the following steps associated with the preparation of **Planning Tables 5 and 6.**

1. Refer to the end of Section 3.1.1 for Lead Worksheets.

- 2. Ensure that chronic and subchronic toxicity values are applied correctly based on the duration of exposure. Provide rationale for selection of surrogate toxicity values not in IRIS or HEAST, or provided by NCEA. (EPA may require additional review.)
- 3. Submit Supporting Information regarding **Toxicity Data for Special Case Chemicals** (i.e., those chemicals with cancer risks and non-cancer hazards calculated using methods or toxicity parameters different from those presented on Planning Tables 5.1, 5.2, 6.1, or 6.2). The Supporting Information should be be used to enable verification of those values by EPA. Examples may include selection of potency factors for polychlorinated biphenyls (PCBs), use of relative potencies for polynuclear aromatic hydrocarbons (PAHs) and chlorinated dioxins and furans, and valence species assumptions for metals. Consult the EPA risk assessor regarding the use of these tables.
- Incorporate the Special Case Chemicals Supporting Information in the Baseline Risk Assessment Report.
- 5. Complete Planning Tables 5 and 6 for the exposure routes and chemicals under evaluation.

Planning Table 5.1: Non-Cancer Toxicity Data - Oral/Dermal

Planning Table 5.2: Non-Cancer Toxicity Data - Inhalation

Planning Table 5.3: Non-Cancer Toxicity Data - Special Case Chemicals

Planning Table 6.1: Cancer Toxicity Data - Oral/Dermal

Planning Table 6.2: Cancer Toxicity Data - Inhalation

Planning Table 6.3: Cancer Toxicity Data - Special Case Chemicals

Planning Table 6.4: Cancer Toxicity Data -External (Radiation).

6. Incorporate **Planning Tables 5 and 6** in the Baseline Risk Assessment Report.

PLANNING TABLE 7: Calculation of

Chemical Cancer Risks and Non-Cancer Hazards. The purposes of Planning Table 7 are:

- To provide a summary of the variables used to calculate chemical cancer risks and noncancer hazards
- To show the EPC and intake used in the noncancer hazard and cancer risk calculations
- To present the result of the calculation for each Exposure Route/Pathway for each COPC
- To provide the total hazard index and cancer risks for all Exposure Routes/Pathways for the Scenario Timeframe and Receptor presented in this table.

The information documented in **Planning Table 7** should include:

- The non-cancer hazard quotient (HQ) and cancer risk value for each COPC for each Exposure Route/Pathway
- The values used for EPC, non-cancer intake, cancer intake, reference doses and concentrations, and cancer slope factors for each COPC for each Exposure Route.

The data elements presented in **Planning Table 7** are listed in the Planning Table 7 highlight box.

KEY DATA ELEMENTS IN PLANNING TABLE 7

For each unique combination of Scenario Timeframe, Receptor Population, and Receptor Age, Regions should provide the following information: Medium, Exposure Medium, Exposure Point, Exposure Route, Chemical of Potential Concern, EPC Value and Units, Cancer Risk Calculations (Intake/Exposure Concentration Value and Units, CSF/Unit Risk Value and Units, and Cancer Risk), and Non-Cancer Hazard Calculations (Intake/Exposure Concentration Value and Units, RfD/RfC Value and Units, and Hazard Quotient).

Regions should perform the following steps associated with the preparation of **Planning Table 7.**

- 1. Address non-cancer hazards and cancer risks including the calculations and supporting information by Exposure Route.
- Include RME and CT results in separate tables. Ensure that risks and hazards from multiple chemicals are combined appropriately across Pathways that affect the same individual or population subgroup, for all site-related chemicals.
- Discuss definitions of Planning Tables
 Planning Table 7.n.RME: Calculation of Chemical Cancer Risks and Non-Cancer Hazards (RME)

Planning Table 7.n.CT: Calculation of Chemical Cancer Risks and Non-Cancer Hazards (CT)

- 4. If it is preferred to segregate cancer and noncancer evaluations, see the blank Planning Tables 7.a.1 and 7.b.1 shown in Appendix A as well as Example Scenario 7 in Appendix D.
- 5. Submit Supporting Information that summarizes the approach used to perform Special Chemical Risk and Hazard Calculations and to enable verification of those values by EPA. This summary should address the calculation of non-cancer hazards and cancer risks for chemicals that do not use RfD or cancer slope factor (CSF) values, respectively. The format of the summary should be determined by each region.
- Incorporate the Special Chemical Risk and Hazard Calculations Supporting Information in the Baseline Risk Assessment Report.
- 7. **Complete Planning Table 7** for each combination of Scenario Timeframe, Receptor Population, and Receptor Age.
- 8. Incorporate **Planning Table 7** in the Baseline Risk Assessment Report.

PLANNING TABLE 8: Calculation of Radiation Cancer Risks.

The purposes of **Planning Table 8** are:

- To provide a summary of the variables used to calculate radiation cancer risks
- To show the EPC used in the radiation cancer risk calculations
- To show, based on the documented risk calculation approach, the intake and cancer slope factors
- To present the result of the calculation for each Exposure Route/Pathway for each COPC
- To provide the radiation cancer risks for all Exposure Routes/Pathways for the Scenario Timeframe and Receptor presented in this table.

The information documented in **Planning Table 8** should include:

- The approach for calculating the radiation cancer risk for each COPC for each Exposure Route/Pathway
- The values used for EPC, intake, and cancer slope factor for each COPC for each Exposure Route
- The Cancer risk value for each COPC for each Exposure Route/Pathway
- Total cancer risk values by Exposure Route, Exposure Point, and across all media for the Scenario Timeframe and Receptor presented in this table.

KEY DATA ELEMENTS IN PLANNING TABLE 8

For each unique combination of Scenario Timeframe, Receptor Population, and Receptor Age, Regions should provide the following information: Medium, Exposure Medium, Exposure Point, Exposure Route, Radionuclide of Potential Concern, EPC Value and Units, Risk Calculation Approach, and Cancer Risk Calculations (Intake/Activity Value and Units, CSF Value and Units, and Cancer Risk).

The data elements presented in **Planning Table 8** are listed in the Planning Table 8 highlight box.

Regions should perform the following steps associated with the preparation of **Planning Table 8**.

- 1. Address radiation cancer risks including the calculations and supporting information by Exposure Route.
- 2. Include RME and CT results in separate tables. Ensure that risks from multiple radionuclides are combined appropriately across pathways that affect the same individual or population subgroup, for all site-related radionuclides.
- 3. Discuss definitions of Planning Tables
 Planning Table 8.n.RME: Calculation of
 Cancer Radiation Risks (RME)
 Planning Table 8.n.CT: Calculation of
 Cancer Radiation Risks (CT)
- 4. **Complete Planning Table 8** for each combination of Scenario Timeframe, Receptor Population, and Receptor Age.
- 5. Incorporate **Planning Table 8** in the Baseline Risk Assessment Report.

RADIATION DOSE ASSESSMENT WORKSHEET. The recommended Radiation Dose Assessment Worksheet has been provided to document alternate radionuclide cancer risk

calculations, performed using a dose approach rather than the standard CERCLA risk calculation method.

The Regions should perform the following steps associated with preparation of the **Radiation Dose Assessment Worksheet**, if applicable to the risk assessment:

- Complete the Radiation Dose
 Assessment Worksheet for each Receptor.
- 2. Provide interim deliverables to the EPA risk assessor, as appropriate.

 Incorporate the Radiation Dose Assessment Worksheet in the Baseline Risk Assessment Report.

A recommended blank Radiation Dose Assessment Worksheet may be found in Appendix C. An example Radiation Dose Assessment Worksheet is presented in Appendix D, Example Scenario 11.

PLANNING TABLE 9: Summary of Receptor Risk and Hazards for COPCs.

The purpose of **Planning Table 9** is:

 To provide a summary of cancer risks and non-cancer hazards for each Receptor, by Medium, Exposure Medium, Exposure Route, and Exposure Point.

The information documented in **Planning Table 9** should include:

- The cancer risk and non-cancer hazard to each Receptor for each COPC by Exposure Route and Exposure Point
- The total cancer risk and non-cancer hazard for each Exposure Point, Exposure Medium and Medium across all Exposure Routes
- The total cancer risk and non-cancer hazard for a Receptor across all media
- The primary target organs for noncarcinogenic hazard effects.

The data elements presented in **Planning Table 9** are listed in the Planning Table 9 highlight box.

Regions should perform the following steps associated with the preparation of **Planning Table 9.**

- 1. Address non-cancer hazards and cancer risks including the calculations and supporting information by Exposure Route.
- 2. Include RME and CT results. Ensure that risks and hazards from multiple chemicals are combined appropriately across Pathways that

affect the same individual or population subgroup,

KEY DATA ELEMENTS IN PLANNING TABLE 9

For each unique combination of Scenario Timeframe, Receptor Population, and Receptor Age, Regions should provide the following information: Medium, Exposure Medium, Exposure Point, Chemical of Potential Concern, Carcinogenic Risk (Ingestion, Inhalation, Dermal, External (Radiation) and Exposure Routes Total), and Non-Carcinogenic Hazard Quotient (Primary Target Organ(s), Ingestion, Inhalation, Dermal, and Exposure Routes Total).

for all site-related chemicals.

- Discuss definitions of Planning Tables
 Planning Table 9.n.RME: Summary of Receptor Risks and Hazards for COPCs (RME)
 - **Planning Table 9.n.CT**: Summary of Receptor Risks and Hazards for COPCs (CT)
- 4. **Complete Planning Table 9** for each combination of Scenario Timeframe, Receptor Population, and Receptor Age.
- 5. Incorporate **Planning Table 9** in the Baseline Risk Assessment Report.

PLANNING TABLE 10: Risk Summary. The purpose of **Planning Table 10** is:

 To provide a summary of cancer risks and non-cancer hazards for each Receptor, by Medium, Exposure Medium, Exposure Route, and Exposure Point, that may trigger the need for remedial action.

The information documented in **Planning Table 10** should include:

- The cancer risk and non-cancer hazard to each Receptor for each chemical or radionuclide by Exposure Route and Exposure Point for risk drivers
- The total cancer risk and non-cancer hazard for each Exposure Point, Exposure Medium, and Medium across all Exposure Routes for

risk drivers

- The total cancer risk and non-cancer hazard for a Receptor across all media for risk drivers
- The primary target organs for noncarcinogenic hazard effects for risk drivers.

The data elements presented in **Planning Table 10** are listed in the Planning Table 10 highlight box.

KEY DATA ELEMENTS IN PLANNING TABLE 10

For each unique combination of Scenario Timeframe, Receptor Population, and Receptor Age, Regions should provide the following information: Medium, Exposure Medium, Exposure Point, Chemical, Carcinogenic Risk (Ingestion, Inhalation, Dermal, External (Radiation) and Exposure Routes Total), and Non-Carcinogenic Hazard Quotient (Primary Target Organ(s), Ingestion, Inhalation, Dermal, and Exposure Routes Total).

Regions should perform the following steps associated with the preparation of **Planning Table 10**.

- 1. Address non-cancer hazards and cancer risks including the calculations and supporting information by Exposure Route.
- 2. Include RME and CT results. Ensure that risks and hazards from multiple chemicals are combined appropriately across Pathways that affect the same individual or population subgroup, for all site-related chemicals.
- 3. Discuss definitions of Planning Tables

Planning Table 10.n.RME: Risk Summary (RME)
Planning Table 10.n.CT: Risk Summary (CT)

4. **Complete Planning Table 10** for each combination of Scenario Timeframe, Receptor Population, and Receptor Age.

5. Incorporate **Planning Table 10** in the Baseline Risk Assessment Report.

LEAD WORKSHEETS. Two recommended Lead Worksheets have been provided to document lead risk evaluations performed for young children and adult receptors at a site.

Regions should perform the following steps associated with the preparation of **Lead Worksheets**:

- Complete the Lead Worksheets for Child and Adult. Also attach the appropriate graphs and results from the Integrated Exposure Uptake Biokinetic Model (IEUBK) model (if used) to the Child Worksheet. Also attach results from the adult lead spreadsheet to the Adult Worksheet.
- 2. The **Lead Worksheets** should later be incorporated in the Baseline Risk Assessment Report.

Blank recommended Lead Worksheets may be found in Appendix C. Example Lead Worksheets are presented in Appendix D Example Scenario 10.

3.1.2 ASSESSMENT OF CONFIDENCE AND UNCERTAINTY

Uncertainty assessment is important in risk assessment. Although the risk assessment should indicate sources of variability and uncertainty throughout the process, it will generally be appropriate to include a separate section of the Baseline Risk Assessment Report that also focuses on the uncertainties associated with data evaluation, toxicity assessment, exposure assessment, and risk characterization, as well as overall uncertainty of the final risk numbers. The region may choose to defer presentation of this specific section to the Draft Baseline Risk Assessment Report.

Regions should perform the following steps associated with the **Assessment of Confidence** and Uncertainty:

- 1. Summarize the Assessment of Confidence and Uncertainty.
- 2. Incorporate the **Assessment of Confidence** and Uncertainty in the Baseline Risk Assessment Report.

3.1.3 PROBABILISTIC ANALYSIS INFORMATION

Based upon the results from a deterministic risk characterization calculation (Planning Table 7) a decision should be made if a Probabilistic Analysis will be performed to calculate cancer risks and non-cancer hazards in accordance with Agency policy.

Regions should perform the following steps associated with the **Probabilistic Analysis:**

- 1. Summarize the Probabilistic Analysis (if performed) in a non-standard format. (Planning formats have not been developed to document probabilistic analysis.) Refer to probabilistic analysis guidance (U.S. EPA 1997e, 1997g and 2001d) to determine the information to be documented.
- 2. Incorporate the **Probabilistic Analysis** summary in the Baseline Risk Assessment Report.

3.2 DRAFT BASELINE RISK ASSESSMENT REPORT

Regions should Submit the Draft Baseline Risk Assessment Report after the completion and acceptance of the Interim Deliverables described above. EPA guidance should be consulted in preparing the Draft Baseline Risk Assessment Report. EPA anticipates that this report preparation will be greatly expedited, since it should incorporate the following Interim Deliverables:

- Planning Tables 0 through 10
- Worksheets on Data Useability, Dermal, Radiation Dose Assessments, and Lead, as applicable

- Supporting Information
- The Assessment of Confidence and Uncertainty
- Probabilistic Analysis information (if applicable).

However, the report should not consist exclusively of the Interim Deliverables, because additional narrative should be necessary for a clear and comprehensible Baseline Risk Assessment Report. For example, information such as definition of hazard indices and cancer slope factors, toxicological profiles for COPCs, and other information indicated by risk assessment guidance should be incorporated.

Every risk assessment should contain a Risk Characterization appropriate to the assessment. Risk assessments submitted to the Agency or performed by the Agency should incorporate any current Agency guidance applicable on Risk Characterization (e.g., RAGS/HHEM, EPA 1989c; Memorandum from Carol Browner on Risk Characterization, EPA 1995b).

3.3 FINAL BASELINE RISK ASSESSMENT REPORT

Regions should submit the Final Baseline Risk Assessment Report as a revision of the draft, incorporating review comments as necessary and appropriate.

Regions should Prepare Draft ROD Risk Worksheet (ROD Risk Highlights) as directed by the EPA RPM and EPA risk assessor, upon completion of the Final Baseline Risk Assessment Report. Refer to the ROD guidance (U.S. EPA, 1999a) for human health risk data needs. The draft ROD Risk Worksheets present the Exposure Pathways and Chemicals that help justify the need for remedial action. Regions should prepare these recommended Worksheets when the Final

Baseline Risk Assessment Report is completed, in order to facilitate the EPA risk manager's preparation of the ROD at a later date.

Exhibit 3-4 identifies the RAGS Part D

information sources (Planning Table and column) for ROD Risk Worksheets (Highlights) 6-15, 6-16A, 6-16B, 6-18A, and 6-18B. Blank templates for the five ROD Risk Worksheets (Highlights) may be found in Appendix C

3.4 INFORMATION TRANSFER TO SUPERFUND RISK DATA COLLECTION

Upon the completion of the Final Baseline Risk Assessment Report, provide the Lotus® or Excel® version of the Planning Tables and Worksheets to the EPA risk assessor, who should submit them to the EPA Headquarters Risk Information Manager responsible for the Superfund Risk Data Collection.

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RAGS PART D INFORMATION SOURCES FOR ROD RISK GUIDANCE HIGHLIGHTS

| ROD RISK HIGHLIGHT | PURPOSE OF ROD RISK HIGHLIGHT | ROD FIELDS | ASSOCIATED RAGS D TABLE | ASSOCIATED RAGS D FIELDS |
|-----------------------|--|---------------------------------------|--|---|
| Highlight | Summary of | Scenario Timeframe | Planning Tables 2 & 3 | Scenario Timeframe |
| 6-15 | Chemicals of Concern and | Medium | Planning Tables 2 & 3 | Medium |
| | Medium- Specific | Exposure Medium | Planning Tables 2 & 3 | Exposure Medium |
| | Exposure Point Concentrations | Exposure Point | Planning Tables 2 & 3 | Exposure Point |
| | Concentrations | Chemical of Concern | Significant Chemicals from Planning Table 2 (site specific definition) | Chemical |
| | | Concentration Detected - Min | Planning Table 2 | Minimum Concentration |
| | | Concentration Detected - Max | Planning Table 2 | Maximum Concentration |
| | | Units | Planning Table 2 | Units |
| | | Frequency of Detection | Planning Table 2 | Detection Frequency |
| | | Exposure Point Concentration | Planning Table 3 | Exposure Point Concentration Value |
| | | Exposure Point Concentration Units | Planning Table 3 | Exposure Point Concentration Units |
| | | Statistical Measure | Planning Table 3 | Exposure Point Concentration Statistic |

Notes:

⁻A version of ROD Highlight 6-15 is to be prepared for each combination of Scenario Timeframe, Medium, and Exposure Medium with "significant routes of exposure". The definition of "significant" will be site specific. -Only Exposure Points with "Significant Routes of Exposure" are to be included.

RAGS PART D INFORMATION SOURCES FOR ROD RISK GUIDANCE HIGHLIGHTS (continued)

EXHIBIT 3-4

| ROD RISK HIGHLIGHT | PURPOSE OF ROD RISK HIGHLIGHT | ROD FIELDS | ASSOCIATED RAGS D TABLE | ASSOCIATED RAGS D FIELDS |
|-----------------------|--|--|--|--|
| Highlight 6-16A | Cancer Toxicity Data Summary | Pathway: Ingestion, Dermal | Planning Table 6.1 (Cancer Toxicity Data- Oral/Dermal) | |
| | | Chemical of Concern | Chemicals of Concern from Planning Table 6.1 (site specific definition) | Chemical of Potential Concern |
| | | Oral Cancer Slope Factor | Planning Table 6.1 | Oral Cancer Slope Factor |
| | | Dermal Cancer Slope Factor | Planning Table 6.1 | Absorbed Cancer Slope Factor for Dermal Value |
| | | Slope Factor Units | Planning Table 6.1 | Oral Cancer Slope Factor Units and Absorbed Cancer Slope Factor for Dermal Units |
| | | Weight of Evidence/ Cancer Guideline Description | Planning Table 6.1 | Weight of Evidence/Cancer Guideline Description |
| | | Source | Planning Table 6.1 | Oral CSF Source(s) |
| | | Date | Planning Table 6.1 | Oral CSF Date(s) |
| | | Pathway: Inhalation | Planning Table 6.2 (Cancer Toxicity Data - Inhalation) | |
| | | Chemical of Concern | Chemicals of Concern from Planning Table 6.2 (site specific definition) | Chemical of Potential Concern |
| | | Unit Risk | Planning Table 6.2 | Unit Risk Value |
| | | Units | Planning Table 6.2 | Unit Risk Units |

EXHIBIT 3-4

RAGS PART D INFORMATION SOURCES FOR ROD RISK GUIDANCE HIGHLIGHTS (continued)

| ROD RISK HIGHLIGHT | PURPOSE OF ROD RISK HIGHLIGHT | ROD FIELDS | ASSOCIATED RAGS D TABLE | ASSOCIATED RAGS D FIELDS | |
|-----------------------------|--|---|--|---|------------------------------|
| Highlight 6-16A (continued) | Cancer Toxicity Data Summary | Inhalation Cancer Slope Factor | Planning Table 6.2 | Inhalation Cancer Slope Factor Value | |
| | (continued) | Units | Planning Table 6.2 | Inhalation Cancer Slope Factor Units | |
| | | Weight of Evidence/ Cancer Guideline Description | Planning Table 6.2 | Weight of Evidence/Cancer Guideline Description | |
| | | Source | Planning Table 6.2 | Unit Risk : Inhalation CSF Source(s) | |
| | | Date | Planning Table 6.2 | Unit Risk : Inhalation CSF Date(s) | |
| | | Pathway: External (Radiation) | Planning Table 6.4 (Cancer Toxicity Data - Radiation) | | |
| | | COC | Chemicals of Concern from Planning Table 6.4 (site specific definition) | Chemical of Potential Concern | |
| | | Cancer Slope or Conversion Factor | Planning Table 6.4 | Cancer Slope Factor Value | |
| | | Exposure Route | Planning Table 1 | Exposure Route | |
| | | | Units | Planning Table 6.4 | Cancer Slope Factor Units |
| | | Weight of Evidence/ Cancer Guideline Description | Not Available | Not Available | |
| | | Source | Planning Table 6.4 | Source(s) | |
| | | Date | Planning Table 6.4 | Date(s) | |

Note:

⁻A version of ROD Highlight 6-16A is to be prepared for the Chemicals of Concern. This definition will be site specific.

RAGS PART D INFORMATION SOURCES FOR ROD RISK GUIDANCE HIGHLIGHTS (continued)

EXHIBIT 3-4

| ROD RISK HIGHLIGHT | PURPOSE OF ROD RISK HIGHLIGHT | ROD FIELDS | ASSOCIATED RAGS D TABLE | ASSOCIATED RAGS D FIELDS |
|-----------------------|--|---|--|---|
| Highlight 6-16B | Non-Cancer Toxicity Data Summary | Pathway: Ingestion, Dermal | Planning Table 5.1 (Non-Cancer Toxicity Data - Oral/Dermal) | |
| | | Chemical of Concern | Chemicals of Concern from Planning Table 5.1 (site specific definition) | Chemical of Potential Concern |
| | | Chronic/ Subchronic | Planning Table 5.1 | Chronic/Subchronic |
| | | Oral RfD Value | Planning Table 5.1 | Oral RfD Value |
| | | Oral RfD Units | Planning Table 5.1 | Oral RfD Units |
| | | Dermal RfD | Planning Table 5.1 | Absorbed RfD for Dermal Value |
| | | Dermal RfD Units | Planning Table 5.1 | Absorbed RfD for Dermal Units |
| | | Primary Target Organ | Planning Table 5.1 | Primary Target Organ(s) |
| | | Combined Uncertainty/ Modifying Factors | Planning Table 5.1 | Combined Uncertainty/ Modifying Factors |
| | | Sources of RfD:Target Organ | Planning Table 5.1 | RfD:Target Organ(s) Source(s) |
| | | Dates of RfD:Target Organ | Planning Table 5.1 | RfD:Target Organ(s) Date(s) |
| | | Pathway: Inhalation | Planning Table 5.2 (Non-Cancer Toxicity Data - Inhalation) | |
| | | Chemical of Concern | Chemicals of Concern from Planning Table 5.2 (site specific definition) | Chemical of Potential Concern |

RAGS PART D INFORMATION SOURCES FOR ROD RISK GUIDANCE HIGHLIGHTS (continued)

| ROD RISK HIGHLIGHT | PURPOSE OF ROD RISK HIGHLIGHT | ROD FIELDS | ASSOCIATED RAGS D TABLE | ASSOCIATED RAGS D FIELDS |
|-----------------------|--|---|----------------------------------|---|
| Highlight 6-16B | Non-Cancer Toxicity Data | Chronic/ Subchronic | Planning Table 5.2 | Chronic/ Subchronic |
| (continued) | Summary (continued) | Inhalation RfC | Planning Table 5.2 | Inhalation RfC Value |
| | | Inhalation RfC Units | Planning Table 5.2 | Inhalation RfC Units |
| | | Inhalation RfD | Planning Table 5.2 | Extrapolated RfD Value |
| | | Inhalation RfD Units | Planning Table 5.2 | Extrapolated RfD Units |
| | | Primary Target Organ | Planning Table 5.2 | Primary Target Organ(s) |
| | | Combined Uncertainty/ Modifying Factors | Planning Table 5.2 | Combined Uncertainty/ Modifying Factors |
| | Sources of RfC:RfD: Target Organ | Planning Table 5.2 | RfC:Target Organ(s) Source(s) | |
| | | Dates | Planning Table 5.2 | RfC:Target Organ(s) Date(s) |

Notes

⁻A version of ROD Highlight 6-16B is to be prepared for the Chemicals of Concern. This definition will be site specific.

RAGS PART D INFORMATION SOURCES FOR ROD RISK GUIDANCE HIGHLIGHTS (continued)

| ROD RISK HIGHLIGHT | PURPOSE OF ROD RISK HIGHLIGHT | ROD FIELDS | ASSOCIATED RAGS D TABLE | ASSOCIATED RAGS D FIELDS |
|-----------------------|--|---|--|--|
| Highlight | Risk | Scenario Timeframe | Planning Table 9 or 10 | Scenario Timeframe |
| 6-18A | Characterization Summary - | Receptor Population | Planning Table 9 or 10 | Receptor Population |
| | Carcinogens | Receptor Age | Planning Table 9 or 10 | Receptor Age |
| | | Medium | Planning Table 9 or 10 | Medium |
| | | Exposure Medium | Planning Table 9 or 10 | Exposure Medium |
| | | Exposure Point | Planning Table 9 or 10 | Exposure Point |
| | | Chemical of Concern | Chemicals of Concern from Planning Table 9 or 10 (site specific definition) | Chemical |
| | | Carcinogenic Risk– Ingestion | Planning Table 9 or 10 | Carcinogenic Risk–Ingestion |
| | | Carcinogenic Risk– Inhalation | Planning Table 9 or 10 | Carcinogenic Risk–Inhalation |
| | | Carcinogenic Risk– Dermal | Planning Table 9 or 10 | Carcinogenic Risk-Dermal |
| | Carcinogenic Risk–External (Radiation) | Planning Table 9 or 10 | Carcinogenic Risk–External (Radiation) | |
| | | Carcinogenic Risk Exposure Routes Total | Planning Table 9 or 10 | Carcinogenic Risk - Exposure Routes Total |
| | | Medium Risk Total | Planning Table 9 or 10 | Medium Total (Risk) |
| | | Total Risk | Planning Table 9 or 10 | Receptor Risk Total |

Notes

⁻A version of Highlight 6-18A is to be prepared for each Receptor (combination of Scenario Timeframe, Receptor Population, and Receptor Age) with "Significant Exposure". The definition of "Significant Exposure" will be site specific.

RAGS PART D INFORMATION SOURCES FOR ROD RISK GUIDANCE HIGHLIGHTS (continued)

| ROD RISK HIGHLIGHT | PURPOSE OF ROD RISK HIGHLIGHT | ROD FIELDS | ASSOCIATED RAGS D TABLE | ASSOCIATED RAGS D FIELDS |
|-----------------------|--|---|--|---|
| Highlight | Risk | Scenario Timeframe | Planning Table 9 or 10 | Scenario Timeframe |
| 6-18B | Characterization Summary - | Receptor Population | Planning Table 9 or 10 | Receptor Population |
| | Non- Carcinogens | Receptor Age | Planning Table 9 or 10 | Receptor Age |
| | | Medium | Planning Table 9 or 10 | Medium |
| | | Exposure Medium | Planning Table 9 or 10 | Exposure Medium |
| | | Exposure Point | Planning Table 9 or 10 | Exposure Point |
| | | Chemical of Concern | Chemicals of Concern from Planning Table 9 or 10 (site specific definition) | Chemical |
| | | Primary Target Organ | Planning Table 9 or 10 | Non-Carcinogenic Hazard Quotient - Primary Target Organ(s) |
| | | Non-Carcinogenic Hazard Quotient - Ingestion | Planning Table 9 or 10 | Non-Carcinogenic Hazard Quotient - Ingestion |
| | | Non-Carcinogenic Hazard Quotient - Inhalation | Planning Table 9 or 10 | Non-Carcinogenic Hazard Quotient - Inhalation |
| | | Non-Carcinogenic Hazard Quotient - Dermal | Planning Table 9 or 10 | Non-Carcinogenic Hazard Quotient - Dermal |
| | | Non-Carcinogenic Hazard Quotient - Exposure Routes Total | Planning Table 9 or 10 | Non-Carcinogenic Hazard Quotient - Exposuse Routes Total |

RAGS PART D INFORMATION SOURCES FOR ROD RISK GUIDANCE HIGHLIGHTS (continued)

| ROD RISK HIGHLIGHT | PURPOSE OF ROD RISK HIGHLIGHT | ROD FIELDS | ASSOCIATED RAGS D TABLE | ASSOCIATED RAGS D FIELDS |
|-----------------------------|--|------------------------------|------------------------------------|-----------------------------|
| Highlight 6-18B (continued) | Risk Characterization | Medium Hazard Index Total | Planning Table 9 or 10 | Medium Total (Hazard) |
| | Summary - Non- Carcinogens | Receptor Hazard Index | Planning Table 9 or 10 | Receptor HI Total |
| (continued) | Organ Hazard Index | Planning Table 9 or 10 | Total Organ HI Across All Media | |

Notes

⁻A version of Highlight 6-18B is to be prepared for each Receptor (combination of Scenario Timeframe, Receptor Population, and Receptor Age) with "Significant Exposure". The definition of "Significant Exposure" will be site specific.

CHAPTER 4

RISK EVALUATIONS DURING THE FEASIBILITY STUDY

Continuous involvement of the EPA risk assessor during the FS has numerous the benefits including: 1) supporting the development of remedial action objectives (RAOs) and PRGs, 2) identifying risks and hazards associated with PRGS, and 3) supporting comparison of risks associated with various remedial alternatives. For these reasons, EPA risk assessor involvement in FS preparation and review is strongly encouraged.

4.1 INTRODUCTION

The purpose of the FS generally is to evaluate waste management remedial alternatives. The National Oil and Hazardous Substances Pollution Contingency Plan (NCP) (U.S. EPA, 1990c) provides that a detailed analysis should be performed. The NCP indicates that for screening of remedial alternatives, the long-term and shortterm aspects of three criteria - effectiveness, implementability, and cost - should be used to guide the development and screening of remedial Consideration of effectiveness alternatives. involves evaluating the long-term and short-term human health risks. Long-term risks associated with a remedial alternative are those risks that will remain after the remedy is complete; short-term risks associated with a remedial alternative are generally those risks that occur during implementation of the remedial alternative.

Evaluating long-term risks ideally includes an assessment of the risks associated with treatment of residuals and untreated wastes for a treatment-based remedy, or an evaluation of the remedy's ability to provide protectiveness over time for a containment-based remedy. For short-term human health risks associated with a remedial alternative, a risk assessor may need to evaluate the risks that occur during implementation of the remedial alternative (e.g., risks associated with emissions from an onsite air stripper). Because some remedies may take many years to complete, some "short-term" risks may actually occur over a

period of many years. Populations that may be exposed to chemicals during remedy implementation include people who live and work in the vicinity of the site.

The NCP also provides that RAOs and remediation goals should be developed. These serve as objectives and goals that can be used to identify and assess remedial alternatives at Superfund sites. The remainder of this chapter discusses RAOs and remediation goals. As also discussed in the NCP, final remediation goals are generally not determined until a final remedy for the site is selected in the ROD (see Chapter 5).

4.1.1 REMEDIAL ACTION OBJECTIVES

As discussed in the NCP, RAOs should describe, in general terms, what a remedial action should accomplish in order to be protective of human health and the environment. RAOs are typically narrative statements that specify the contaminants and environmental media of concern, the potential exposure pathways to be addressed by remedial actions, the exposed populations and environmental receptors to be protected, and the acceptable contaminant concentrations or concentration ranges (remediation goals) in each environmental medium.

4.1.2 REMEDIATION GOALS

Remediation goals are normally a subset of the RAOs. They generally provide the acceptable contaminant concentrations in each medium for remedial actions to meet.

As explained in the preamble to the final NCP that remediation goals are generally based on ARARs unless ARARs are not available or are not protective. ARARs do not always exist for all

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SELECTION OF REMEDIATION GOALS

The NCP [U.S. EPA, 1990c; Section 300.430(e) (2)(I)] states that the selection of remediation goals should consider the following:

"...remediation goals shall establish acceptable exposure levels that are protective of human health and the environment and shall be developed considering the following...

ARARs under Federal environmental or State environmental or facility siting laws, if available, and the following factors:

- For systemic toxicants, acceptable exposure levels shall represent concentration levels to which the human population, including sensitive subgroups, may be exposed without adverse effect during a lifetime or part of a lifetime, incorporating an adequate margin of safety;
- 2. For known or suspected carcinogens, acceptable exposure levels are generally concentration levels that represent an excess upper bound lifetime cancer risk to an individual of between 10⁻⁴ and 10⁻⁶ using information on the relationship between dose and response. The 10⁻⁶ risk level shall be used as the point of departure for determining remediation goals for alternatives when ARARs are not available or are not sufficiently protective because of the presence of multiple contaminants at a site or multiple pathways of exposure;
- Factors related to technical limitations such as detection/quantification limits for contaminants;
- 4. Factors related to uncertainty; and
- 5. Other pertinent information."

chemicals and all environmental media.

Therefore, according to the NCP, there are two major sources for determining the acceptable exposure levels used for developing remediation goals: a) concentrations found in Federal and State ARARs and, if these are not available or not

protective, (b) risk-based concentrations that are determined to be protective of human health and the environment. These risk-based concentrations should be calculated using, at a minimum, the criteria sited in numbers 1 and 2 in the Remediation Goals highlight box. Other factors mentioned in the highlight box [i.e., limits of detection (number 3), uncertainty (number 4), and background concentration levels (number 5)] also should be considered.

Risk-based concentrations may need to be developed even if ARARs are available to ensure that these ARARs are protective of human health and the environment.

ARAR-Based Remediation Goals. Potential chemical-specific ARARs include concentration limits set by Federal environmental regulations such as Maximum Contaminant Levels (MCLs) established under the Safe Drinking Water Act (SDWA), ambient water quality criteria established under the Clean Water Act (CWA), and State regulations (e.g., State drinking water laws). Action-specific and location-specific ARARs must also be complied with or waived according to the NCP.

Risk-Based Remediation Goals. In general, remediation goals based on risk-based calculations should be determined using cancer or non-cancer toxicity values with specific exposure assumptions. For chemicals with carcinogenic effects, the NCP has described the development of remediation goals, as a practical matter, as a two-step process [U.S. EPA, 1990c, Section 300.430(e)(2)(I)(D)]. A concentration equivalent to a lifetime cancer risk of 10⁻⁶ is first established as a point of departure. Then, other factors are taken into account to determine where within the acceptable range the remediation goals for a given contaminant at a specific site should be established.

The NCP discusses a generally acceptable risk range of 10^{-4} to 10^{-6} . EPA has further clarified the extent of the acceptable risk range by stating that the upper boundary generally is not a discrete line at $1x10^{-4}$. Risks slightly greater than $1x10^{-4}$ may be considered to be acceptable (i.e., protective) if justified based on site-specific conditions, including any uncertainties about the nature and extent of contamination and associated

risks. [See Role of the Baseline Risk Assessment in Superfund Remedy Selection Decisions (U.S. EPA, 1991d)].

For non-cancer effects, the NCP states that an acceptable exposure level should be defined. (See "Selection of Remediation Goals" highlight box in this section.) According to EPA guidance, generally if the Hazard Index (HI) (Intake/RfD) is above 1 (i.e., the site exposure is estimated to be above the RfD) there may be a concern for potential non-cancer effects [see Role of the Baseline Risk Assessment in Superfund Remedy Selection Decisions (U.S. EPA, 1991d)]. Therefore, in calculating remediation goals at a site to protect for non-cancer effects, remediation goals are generally set at a Hazard Index at or below 1.

4.1.3 PRELIMINARY REMEDIATION GOALS

PRGs for a site are usually established as early in the RI/FS process as possible during project scoping (see Chapter 2). These initial PRGs can then be modified as necessary during the FS, based on site-specific information from the baseline risk assessment. The PRGs should then be used to establish the goals to be met by the remedial alternatives in the FS. The PRGs also should guide the development of the Proposed Plan for remedial action and the selection of remediation levels in the Record of Decision. During the FS, both risk-based and ARAR-based PRGs should be considered. (See Section 4.1.2 for more discussion on ARAR-based PRGs).

Risk-based PRGs (non-ARARs) may be modified within the acceptable risk range during the remedy selection process based on a balancing of the major trade-offs among the alternatives as well as the public and Agency comments on the Proposed Plan (RAGS Part B, U.S. EPA, 1991a). Such balancing among alternatives and consideration of community and State acceptance should establish the specific level of protection the remedy will achieve (i.e., the final remediation levels).

The dialogue begun during Scoping between the EPA risk assessor and the EPA RPM should continue during the FS and beyond to ensure that risk assessment information is used appropriately in the risk management decision process.

The primary guidance on development of the FS is available in "Guidance for Conducting Remedial Investigations and Feasibility Studies Under CERCLA (U.S. EPA, 1988). RAGS Part B (U.S. EPA, 1991a) also presents guidance for the role of risk assessment in the FS. Consult the EPA RPM for guidance.

4.2 DEVELOP REMEDIAL ACTION OBJECTIVES

The risk assessor should be involved in the preparation or review of the following:

- A narrative description of the Medium, Exposure Point and Exposure Routes, and chemicals and radionuclides that will be the focus of the remedial action
- A narrative identifying the remedial action objectives for prevention of exposure and restoration, where appropriate of each contaminated Medium (e.g., restoring groundwater to a potable water source)

A format such as Example Table 1 in Exhibit 4-1 may be a useful approach to present these data for each Medium.

4.3 DEVELOP REMEDIATION GOALS

The risk assessor should be involved in the preparation or review of a short narrative or tables which provide the goals of the remediation. First, all values considered as PRGs should be identified. Then the PRGs selected for each chemical to be used in the FS should be presented.

4.3.1 IDENTIFY VALUES CONSIDERED AS PRELIMINARY REMEDIATION GOALS

The risk assessor should be involved in the following activities:

- Identify which chemicals and/or radionuclides will have PRGs developed.
- · Identify ARAR-based PRGs and associated

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risks/hazards.

- If ARAR-based PRGs are not protective, risk-based PRGs using EPA methods should be calculated.
- Identify other values to consider as PRGs [e.g., background, detection limits, Procedure Quantitation Limits (PQLs)].

A format such as Example Table 2 in Exhibit 4-1 may be a useful approach to present these values, for each Medium and Receptor Population combination.

4.3.2 SELECT PRELIMINARY REMEDIATION GOALS

The risk assessor should be involved in the following activities:

- Select PRG(s) for each chemical from among the values considered (e.g., risk-based for cancer and non-cancer, ARAR-based, other), modifying values as appropriate. Note that the PRG should be ARAR-based unless there is no ARAR available or the ARAR is not protective.
- Provide the rationale for the selected PRG.
 Include the source of the value.

A format such as Example Table 3 in Exhibit 4-1 may be a useful approach to present these values for each Medium and Receptor Population combination.

4.4 SUMMARIZE RISKS AND HAZARDS ASSOCIATED WITH PRELIMINARY REMEDIATION GOALS

The risk assessor should be involved in the preparation or review of a short narrative or tables which summarize the risks and hazards associated with the PRGs. The risk assessor should be involved in the following activities:

• Identify the chemical and/or radionuclide of

- concern, maximum concentration, PRG, basis of PRG, and calculated risks and hazards associated with the PRG for each Medium and Receptor Population.
- Summarize the total risk and total hazard among all chemicals for each Medium and Receptor Population combination.

A format such as Example Table 3 in Exhibit 4-1 may be a useful approach to present these values for each Medium and Receptor Population combination.

4.5 EVALUATE REMEDIAL TECHNOLOGIES AND ALTERNATIVES FOR RISK CONSIDERATIONS

The risk assessor may provide input in the process of evaluating remedial technologies and alternatives for risk considerations beginning in the development and screening stage of the FS and extending into the detailed analysis stage. The major goal for the risk evaluation during these steps is to provide the FS team and the EPA RPM with specific long-term and short-term human health risk information to consider when identifying and screening technologies and alternatives and performing detailed analysis of alternatives.

Generally, the long-term human health risks associated with a remedial technology or alternative are those risks that are expected to remain after the remedy is complete (i.e., residual risks). The risk issues to be considered may include an assessment of the risks associated with treatment residuals, untreated wastes, or contained wastes.

Generally, the short-term human health risks associated with a remedial technology or alternative are those risks that are expected to occur during implementation of the technology or alternative, which may occur over a period of years. Populations to be considered include people who live and work in the vicinity of the site and workers involved in site remediation.

4.5.1 IDENTIFICATION AND

SCREENING OF TECHNOLOGIES AND ALTERNATIVES

The risk assessor may contribute to the identification and screening of technologies and alternatives and focus on evaluating associated short-term and long-term human health risks to ensure that they meet RAOs and PRGs. The goal of the risk assessor is to assist in identifying, and eliminating from further consideration, technologies and/or alternatives with clearly unacceptable risks. This evaluation is typically qualitative, based on simplifying assumptions and professional judgment rather than detailed analysis. The risk assessor's evaluation should be associated with the consideration of effectiveness. one of the NCP's three screening criteria. (Implementability and cost are the other two criteria evaluated at this screening stage, but they do not typically involve risk assessor participation.)

4.5.2 DETAILED ANALYSIS OF ALTERNATIVES

The overall objective of the risk assessor's role in the detailed analysis of alternatives is to support the preparation and evaluation of the risk information needed for RPMs to select a remedial alternative for a site. See the highlight box for the NCP's nine remedial alternatives. The risk assessor should contribute to the analysis of at least three of the nine criteria specified by the NCP:

- Overall Protection of Human Health and the Environment
- Long-term Effectiveness and Permanence
- Short-term Effectiveness.

The detailed analysis of short-term and longterm risks may be qualitative or quantitative depending on the "perceived risk" associated with the alternative based on both professional judgment and community concerns. The risk analysis should follow the same general steps as the baseline risk assessment; however, the steps

will typically not be conducted in the same level of detail for the FS.

NCP CRITERIA FOR EVALUATING REMEDIAL ALTERNATIVES

- 1. Overall Protection of Human Health and Environment
- 2. Compliance with ARARs
- 3. Long-term Effectiveness and Permanence
- 4. Reductions in Toxicity, Mobility, and Volume Through Treatment
- 5. Short-term Effectiveness
- 6. Implementability
- 7. Cost
- 8. State Acceptance
- 9. Community Acceptance.

The detailed analysis of short-term risks should include the following components for each alternative:

- Evaluate short-term exposure
- Evaluate short-term toxicity
- Characterize short-term risks to the community (including people who live or work on or near the site)
- Characterize short-term risks to remediation workers (a qualitative assessment may be appropriate if the risks to remediation workers are addressed adequately in the site-specific Health and Safety Plan).

The detailed analysis of long-term risks includes the following components for each alternative.

- Evaluate residual risk
- Evaluate protectiveness over time.

CHAPTER 5

RISK EVALUATIONS AFTER THE FEASIBILITY STUDY

After completion of the FS, EPA risk assessor involvement in risk evaluations should support the EPA RPM in ensuring that the remedy is protective. While these risk evaluations may not always require a significant level of quantitation, continuous involvement of EPA risk assessors is importantl to ensure consistency in risk evaluation and risk communication. Post-FS activities benefitting from EPA risk assessor involvement typically include the Proposed Plan, the Record of Decision (ROD), the Remedial Design/Remedial Action, and Five-Year Reviews.

5.1 RISK EVALUATION FOR THE PROPOSED PLAN

The Proposed Plan should include sufficient risk assessment information to support the basis for the proposed remedial action. EPA risk assessor support is recommended during the preparation of the Proposed Plan to ensure the consistency of risk information with the Baseline Risk Assessment Report and the FS Report. The level of detail in the Proposed Plan should be appropriate to the needs of the public. Additional EPA risk assessor support at this time may be qualitative or quantitative, typically focusing on refinement of previous analyses, based on newly developed information.

5.2 RISK EVALUATION ASSOCIATED WITH THE RECORD OF DECISION

EPA risk assessor involvement in preparation of the risk evaluation in the ROD is strongly recommended. A summary of the relevant information from the Baseline Risk Assessment Report should be presented in a mixture of text format and table format. In addition, the risks

(short-term and residual) associated with each

alternative should be discussed.

5.2.1 BASELINE RISK SUMMARY IN THE RECORD OF DECISION

To support the preparation of the Record of Decision, the EPA risk assessor should prepare or review a summary of the Baseline Risk Assessment Report which supports the basis for the remedial action. The primary focus should be on those exposure pathways and chemicals of concern found to pose actual or potential threats to human health or the environment. Chemicals included in the risk assessment but determined not to contribute significantly to an unacceptable risk need not be included in the Risk Characterization Summary in the ROD (e.g., chemicals with risk levels less than 1x10⁻⁶ or HQ less than 0.1) unless they are needed to justify a No Action ROD.

Refer to Interim Final Guidance on Preparing Superfund Decision Documents (U.S. EPA, 1989b) and Guide to Preparing Superfund Proposed Plans, Records of Decision, and Other Remedy Selection Decision Documents (U.S. EPA, 1999a) for a recommended format for summarizing human health risk assessment information in the ROD.

Other risk information may also be included in the ROD depending upon the level of detail preferred. Information related to values used for intake calculations and non-cancer and cancer toxicity data and exposure point concentrations are summarized on Planning Tables 4, 5, 6, 7, and 8, which could be placed in appendices to the ROD. Section 3.3 provides recommended ROD Risk Worksheets that correspond to ROD guidance highlights 6-15, 6-16A, 6-16B, 6-18A and 6.18B. Preparation of these recommended

Worksheets previously, as interim deliverables (see Section 3.3), is strongly recommended

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because it should greatly facilitates risk evaluation in the ROD. If these recommended Worksheets were not previously prepared, refer to Exhibit 3-4 for RAGS Part D Planning Table sources for this information.

5.2.2 RISKS ASSOCIATED WITH CLEANUP LEVELS IN THE RECORD OF DECISION

The ROD (except for no-action RODs) should describe how remedial alternatives will reduce risks by achieving cleanup levels through treatment or by eliminating exposures through engineering controls for the contaminated media.

In addition, the risk assessor should prepare/review the following information related to the selected alternative:

- Document short-term risks that may occur during remedy implementation
- Document risks that may remain after completion of the remedy (including residual risk from untreated waste remaining at the site)
- Evaluate the need for five-year reviews.

Refer to the ROD guidance (U.S. EPA, 1999a) for suggestions regarding presentation of risks associated with cleanup levels in the ROD.

5.3 RISK EVALUATION DURING REMEDIAL DESIGN AND REMEDIAL ACTION

The EPA risk assessor's role during remedial design and remedial action may be qualitative or quantitative depending on the site and phase of the project. During the remedial design, short-term and long-term risks may be assessed through refinement of previous analyses and identification of the need for engineering controls or other measures to mitigate risk.

During the remedial action, the EPA risk assessor is more likely to provide quantitative risk evaluation support. Short-term risk evaluation may address impacts to remediation workers and neighboring communities.

Long-term risk evaluations typically focus on the

following:

- Whether cleanup levels specified in the ROD have been attained
- Whether residual risk after completion of the remedy ensures protectiveness.

5.4 RISK EVALUATION ASSOCIATED WITH EXPLANATIONS OF SIGNIFICANT DIFFERENCES (ESDs) AND AMENDED RODS

This may occur when conditions relevant to a site change following the signing of a ROD. It is sometimes necessary to prepare an ESD or amended ROD. Examples of conditions causing this situation may include, but are not limited to, the following:

- · Toxicity values change
- Additional technology performance information becomes available
- ARARs change (e.g., Land Disposal Restrictions).

EPA risk assessor involvement with RPM evaluations of ESDs and Amended RODs should focuses on evaluating: whether cleanup levels are still protective when considering new ARARs; new parameters for risk and hazard calculations; new technology information; and, other new information. Any new information and revised risk evaluations should be thoroughly documented.

5.5 RISK EVALUATION DURING FIVE-YEAR REVIEWS

CERCLA provides for reviews of certain remedies at least every five years to assure that human health and the environment are being protected by the remedial alternative implemented. EPA risk assessor involvement with RPM evaluations during Five-Year Reviews are generally quantitative and should focus on the following three goals:

 Confirm that the remedy remains protective (including any engineering or institutional controls)

- Evaluate whether cleanup levels are still protective by considering new ARARs, new parameters for risk and hazard calculations, and other new information
- Evaluate whether cleanup has reduced risks to levels no longer requiring restricted site use and five-year reviews (U.S. EPA, 2001b).

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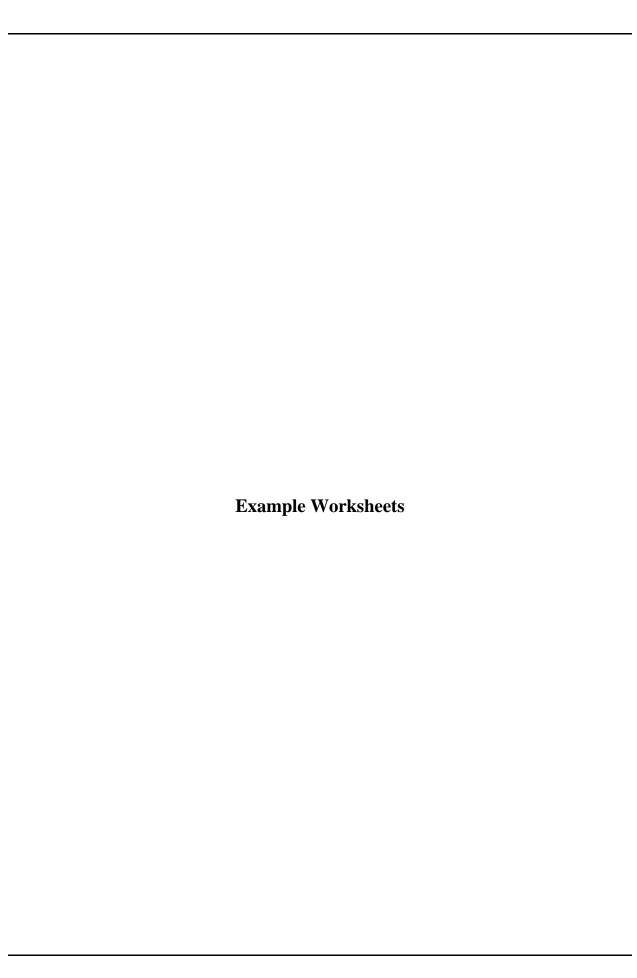
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- * This Reference Section is designed to not only give bibliographic information for documents referred to in the RAGS Part D text, but also to be a source of bibliographic information for documents that are relevant to risk assessment in general.

APPENDIX A

PLANNING TABLES

- -Blank Planning Tables
- -Example Planning Tables

| Blank Planning Tables |
|--|
| |
| |
| The Planning Table formats may not be altered (i.e., columns may be added, deleted, or changed, and rows and footnotes may be added) as appropriate to reflect site-specific conditions. |
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| |





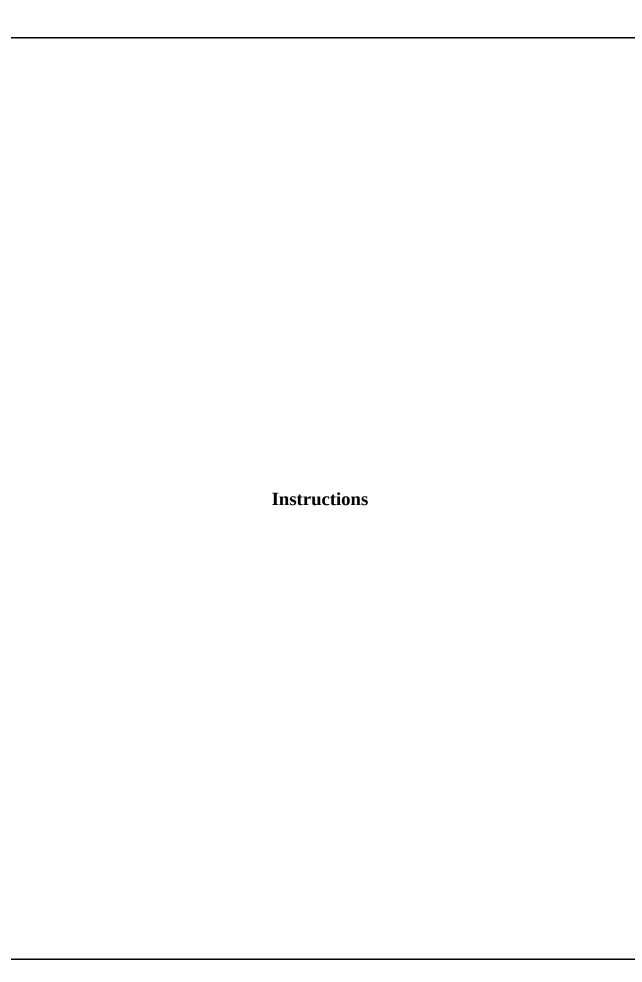
APPENDIX B

INSTRUCTIONS FOR COMPLETION OF THE PLANNING TABLES

- Instructions

-Glossary





APPENDIX C

PLANNING WORKSHEETS

- Data Useability Worksheet
- TARA Schedule Worksheet
- Dermal Worksheet
- Radiation Dose Assessment Worksheet
- Lead Worksheets
- ROD Risk Worksheets

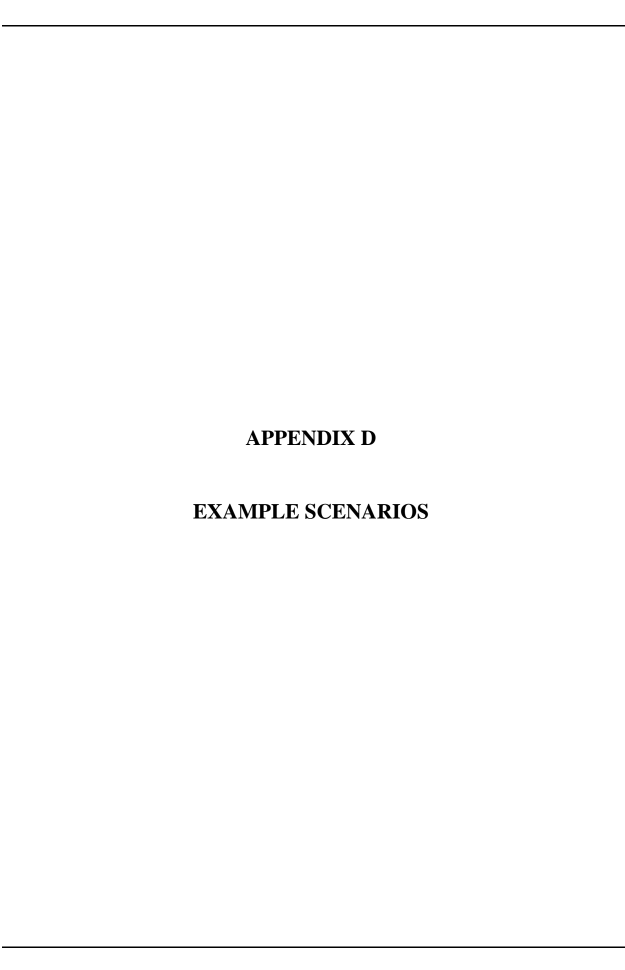
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BLANK PLANNING WORKSHEETS

- -Data Useability Worksheet
- -TARA Schedule Worksheet
- -Dermal Worksheet
- -Radiation Dose Assessment Worksheet
- **-Lead Worksheets**
- -ROD Risk Worksheets

EXAMPLE PLANNING WORKSHEETS

- -Data Useability Worksheet
- -TARA Schedule Worksheet
- **–Dermal Worksheet**
- -Radiation Dose Assessment Worksheet (not included)
- -Lead Worksheets
- -ROD Risk Worksheets
 (not included)



SITE RISK ASSESSMENT IDENTIFICATION INFORMATION

| PURPOSE OF THE TABLE: • To uniquely identify the risk assessment • To identify the relevant contacts for the risk assessment. | | |
|---|--|--|
| INFORMATION DOCUMENTED: • Site information • Contact information . Risk assessment document information. | | |
| TABLE NUMBERING INSTRUCTIONS: Complete one copy of this table for each risk assessment or Set of Planning Tables. Number it Table 0. . | | |
| HOW TO COMPLETE/INTERPRET THE TABLE | | |
| Row 1 - Site Name/OU | | |
| Definition: • The name of the site or operable unit (OU) to which this risk assessment applies. | | |
| Instructions: • Enter the name of the site or operable unit. | | |
| Row 2 - Region | | |
| Definition: • The EPA Region in which the site is located. | | |
| Instructions: • Enter the EPA Region in which the site is located. | | |
| Row 3 - EPA ID Number | | |
| Definition: • The EPA number assigned to identify the site. | | |

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| Ins | structions: | |
|-----|---|--|
| • | Enter the EPA ID Number. The ID can be found either in the site files or in the CERCLIS database. | |
| | | |

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${\bf SITE~RISK~ASSESSMENT~IDENTIFICATION~INFORMATION~(continued)}$

| Row 4 - State | | | |
|--|--------|--|--|
| Definition: • The state in which the site is located. | | | |
| Instructions:Enter the state or commonwealth in which the site is located. | | | |
| Row 5 - Status | | | |
| Definition: • The current status of the site. | | | |
| Instructions: • Enter the site status. | | | |
| Row 6 - Federal Facility (Y/N): | | | |
| Definition: • A flag indicating whether or not the site is a Federal Facility. | | | |
| Instructions:Enter 'Y' if the site is a Federal Facility; enter 'N' otherwise. | Y N | | |
| Row 7 - EPA Project Manager | | | |
| Definition: • The EPA manager responsible for all activity concerning the site. | | | |
| Instructions:Enter the EPA manager responsible for the site. | | | |
| Row 8 - EPA Risk Assessor | | | |
| Definition: • The risk assessor at EPA responsible for this risk assessment. | | | |
| Instructions: • Enter the name of the EPA risk assessor responsible for this risk assessment. | | | |
| Row 9 - Prepared by (Organization): | | | |
| Definition: • The name of the organization that prepared this risk assessment. | | | |
| Instructions:Enter the name of the organization that prepared this risk assessment. | | | |

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${\bf SITE~RISK~ASSESSMENT~IDENTIFICATION~INFORMATION~(continued)}$

| Row 10 - Prepared for (Organization): | | | |
|---|--------|--|--|
| Definition: • The name of the organization for whom this risk assessment was prepared. | | | |
| Instructions: • Enter the name of the organization for whom this risk assessment was prepared | | | |
| Row 11 - Document Title | | | |
| Definition: • The title of this risk assessment document. | | | |
| Instructions:Enter the title of this risk assessment document. | | | |
| Row 12 - Document Date | | | |
| Definition: • The date this risk assessment document was completed or approved. | | | |
| Instructions: Record the date the document was completed or approved in the MM/DD/YYYY format. | | | |
| Row 13 - Probabilistic Risk Assessment (Y/N): | | | |
| Definition: • A flag indicating whether or not a probabilistic risk assessment was done for this risk assessment. | | | |
| Instructions: Enter 'Y' if a probabilistic risk assessment was done; enter 'N' otherwise. | Y N | | |
| Row 14 - Comments | | | |
| Definition: • Any additional information provided about the risk assessment. | | | |
| Instructions: • Enter any additional information about the risk assessment. | | | |

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SELECTION OF EXPOSURE PATHWAYS

| PURPOSE OF THE TABLE: | | |
|--|--|--|
| To assist in project planning To accompany the site conceptual model To present possible Receptors, Exposure Routes, and Exposure Pathways To present the rationale for selection or exclusion of each Exposure Pathway To communicate risk information to interested parties outside EPA To establish a framework for the generation of subsequent Planning Tables. All subsequent tables should be built from the information contained in Table 1. | | |
| INFORMATION DOCUMENTED: Exposure Pathways that were examined and excluded from analysis Exposure Pathways that will be qualitatively and quantitatively evaluated in the risk assessment. | | |
| TABLE NUMBERING INSTRUCTIONS Complete one copy of this table for each risk assessment. Consult the EPA risk assessor to determine if the risk assessment applies to an entire site, a single operable unit, or some other division of the site. Number it Table 1. The table should show each Exposure Pathway considered. | In the Planning Tables, an Exposure Pathway is defined as each unique combination of Scenario Timeframe, Medium, Exposure Medium, Exposure Point, Receptor Population, Receptor Age, and Exposure Route. | |
| HOW TO COMPLETE/INTERPRET THE TABLE | 2 | |
| Column 1 - Scenario Timeframe | | |
| Definition: • The time period (current and/or future) being considered for the Exposure Pathway. | | |
| Instructions: Choose from the picklist to the right. If two Exposure Pathways are identical, Current/Future can be used to describe a future and a current pathway. | Current Future Current/Future Not Documented | |
| Column 2 - Medium | | |
| Definition: The substance (e.g., air, water, soil) that is a potential source of contaminants in the Exposure Medium. (The Medium will sometimes = the Exposure Medium.) Usually, the Medium is that targeted for possible remediation. | | |

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SELECTION OF EXPOSURE PATHWAYS (continued)

| Instructions: • Choose fr | om the picklist to the right. | Groundwater Leachate Sediment Sludge Soil Surface Water Debris Liquid Waste Solid Waste Air Surface Soil Surface Soil Other |
|---------------------------|---|--|
| Column 3 - Exposure | Medium | |
| may be ex | minated environmental medium to which an individual aposed. This includes the transfer of contaminants from aim to another. Contaminants in Groundwater (the Medium) remain in Groundwater (the Exposure Medium) and are available for exposure to receptors. Contaminants in Groundwater (the Medium) may be transferred to Air (the Exposure Medium) and are available for exposure to receptors. Contaminants in Sediment (the Medium) may be transferred to Fish Tissue (the Exposure Medium) and are available for exposure to receptors. | |
| Instructions: | | Groundwater |
| Note: In the o | om the picklist to the right. case of two media transferring contamination to the same Exposure separate Exposure Pathways should be included in Table 1. See nario No. 5. | Leachate Sediment Sludge Soil Surface Water Debris Liquid Waste Solid Waste Air Plant Tissue Animal Tissue Fish Tissue Spring Water Surface Soil Subsurface Soil Particulates Vapors Other |

B1-2 December 2001

SELECTION OF EXPOSURE PATHWAYS (continued)

| Column 4 - Exposure Point | | | | |
|---------------------------|--|--|--|--|
| Def • | An exact location of potential contact between a person and a chemical or radionuclide within an Exposure Medium. | | | |
| | For example: 1) Contaminants are in Groundwater (the Medium and the Exposure Medium) and exposure to Aquifer 1 - Tap Water (the Exposure Point) is evaluated. | | | |
| | 2 Contaminants in Groundwater (the Medium) may be transferred to Air (the Exposure Medium) and exposure to Aquifer 1 - Water Vapors at Showerhead (the Exposure Point) is evaluated. | | | |
| | 3) Contaminants in Sediment (the Medium) may be transferred to Fish Tissue (the Exposure Medium) and Trout from Dean's Creek (the Exposure Point) is evaluated. | | | |
| Ins | Describe the Exposure Point as text in the table. Multiple Exposure Points may be recorded in the same cell/row if all other aspects of their Exposure Pathways (Scenario Timeframe, Medium, Exposure Medium, Receptor Population, Receptor Age, and Exposure Route) are the same. See Example Scenario No. 1. | | | |
| Column 5 - | Receptor Population | | | |
| Def • | inition: The exposed individual relative to the Exposure Pathway considered. | For example, a resident (Receptor Population) who drinks contaminated groundwater. | | |
| Ins | tructions: Choose from the picklist to the right. | Resident Industrial Worker Commercial Worker Construction Worker Other Worker Golfer | | |
| | Note: If there are multiple Trespassers/Visitors of different ages, the use Receptor Age (see Column 6) to distinguish between the different receptors. For example, use Trespasser/Visitor with Adolescent (or Child) to indicate youthful trespassers, and Trespasser/Visitor with Adult for adult visitors. | Jogger Fisher Hunter Fisher/Hunter Swimmer Other Recreational Person Child at School/Daycare/ Playground Trespasser/Visitor Farmer Gardener Gatherer Other | | |

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SELECTION OF EXPOSURE PATHWAYS (continued)

| Column 6 - Receptor Age | | | |
|--|--|--|--|
| Definition: • The description of the exposed individual as defined by the EPA Region or dictated by the site. For example, an adult (Receptor Age) resident (Receptor Population) who drinks contaminated groundwater. | | | |
| Instructions:Choose from the picklist to the right. | Child Adult Adolescents (teens) Pre-Adolescents Not Documented Child/Adult Geriatric Sensitive Other Infant Toddler Pregnant | | |
| Column 7 - Exposure Route | | | |
| Definition: • The way a chemical or radionuclide comes in contact with a person (e.g., by ingestion, inhalation, dermal contact). | | | |
| Instructions:Choose from the picklist to the right. | Inhalation Ingestion Combined (Inhalation and Ingestion) Dermal Not Documented External (Radiation) | | |
| Column 8 - Type of Analysis | | | |
| Definition: • The level of evaluation (quantitative or qualitative) to be performed for the Exposure Pathway based on site-specific analysis. | | | |
| Instructions: • Choose from the picklist to the right. | Quant (Quantitative) Qual (Qualitative) None | | |
| Note: Present pathways that were not further analyzed (Type of Analysis = None) along with the rationale for their exclusion to document that the pathway was considered. | | | |

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SELECTION OF EXPOSURE PATHWAYS (continued)

| Column 9 - Rationale for Selection or Exclusion of Exposure Pathway | | | |
|--|--|--|--|
| Definition: • The reason the Exposure Pathway was selected or not selected for quantitative or qualitative analysis. | | | |
| Instructions: Document the reason for selecting or excluding an Exposure Pathway for analysis. Provide a narrative rationale for each Exposure Pathway. | Consult the EPA risk assessor for the rationale codes. | | |

B1-5 December 2001

OCCURRENCE, DISTRIBUTION AND SELECTION OF CHEMICALS OF POTENTIAL CONCERN

| PURPOSE OF THE TABLE: To provide information useful for data evaluation of chemicals and radionuclides detected To provide adequate information so the user/reviewer gets a sense of the chemicals and radionuclides detected at the site and the potential magnitude of the potential problems at the site To provide chemical screening data and rationale for selection of COPCs. | |
|--|---|
| INFORMATION DOCUMENTED: Statistical information about chemicals and radionuclides detected in each Medium The detection limits of chemicals and radionuclides analyzed The screening toxicity values for COPC selection The chemicals and radionuclides selected or deleted as COPCs. | |
| TABLE NUMBERING AND SUMMARY BOX INSTRUCTIONS: Complete one copy of Table 2 for each unique combination of the following three fields that will be quantitatively evaluated in the risk assessment: Scenario Timeframe, Medium, and Exposure Medium. Enter each combination of these three fields in the Summary Box in the upper left corner of the table. Number each table uniquely, beginning with 2.1 and ending with 2.n, where "n" represents the total number of combinations of the three key fields. | It is possible that some Planning Tables may contain the same data associated with different descriptions in the Summary Box in the upper left corner. Separate tables may be necessary to ensure transparency in data presentation for each Exposure Pathway. Replication of information is readily accomplished using spreadsheet software. Consult the EPA risk assessor for alternatives (e.g., footnotes) to preparing multiple tables with the same data. |
| HOW TO COMPLETE/INTERPRET THE TAI | BLE |
| SUMMARY BOX IN UPPER LEFT CORNER Row 1 - Scenario Timeframe | |
| Definition: The time period (current and/or future) being considered for the exposure pathway. | |
| Instructions:Choose from the picklist to the right. | Current Future Current/Future Not Documented |

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OCCURRENCE, DISTRIBUTION AND SELECTION OF CHEMICALS OF POTENTIAL CONCERN (continued)

| contam someti | bstance (e.g., air, water, soil) that is a potential source of sinants in the Exposure Medium. (The Medium will mes = the Exposure Medium.) Usually, the Medium is that d for possible remediation. | |
|-----------------------|---|--|
| Instructions • Choose | e from the picklist to the right. | Groundwater Leachate Sediment Sludge Soil Surface Water Debris Liquid Waste Solid Waste Air Surface Soil Subsurface Soil |
| - Exposure I | Medium | |
| may be | ntaminated environmental medium to which an individual exposed. Includes the transfer of contaminants from one in to another. | |
| For exan | ple: | |
| 1) | Contaminants in Groundwater (the Medium) remain in Groundwater (the Exposure Medium) and are available for exposure to receptors. | |
| | Contaminants in Groundwater (the Medium) may be transferred to Air (the Exposure Medium) and are available for exposure to | |
| 2) | receptors. | |

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OCCURRENCE, DISTRIBUTION AND SELECTION OF CHEMICALS OF POTENTIAL CONCERN (continued)

| | uctions: Choose from th | ne picklist to the right. | Groundwater Leachate Sediment Sludge Soil Surface Water Debris Liquid Waste Solid Waste Air Plant Tissue Animal Tissue Fish Tissue Spring Water Surface Soil Subsurface Soil Particulates Vapors Other |
|--|---|--|--|
| BODY OF T | HE TABLE | | |
| Column 1 - 1 | Exposure Poin | t | |
| • | For example: 1) Con Mea Poin 2) Con Air Vap 3) Con | on of potential contact between a person and a lionuclide within an exposure medium. taminants are in Groundwater (the Medium and the Exposure lium) and exposure to Aquifer 1 - Tap Water (the Exposure nt) is evaluated. taminants in Groundwater (the Medium) may be transferred to (the Exposure Medium) and exposure to Aquifer 1 - Water ors at Showerhead (the Exposure Point) is evaluated. taminants in Sediment (the Medium) may be transferred to Fish use (the Exposure Medium) and Trout from Dean's Creek (the osure Point) is evaluated. | |
| Instructions: Provide the information as text in the table. | | Exposure Points should be defined the same way as was done in Planning Table 1. | |
| Column 2 - C | CAS Number | | |
| Definition: • The Chemical Abstract Registry Number, a unique standardized number which is assigned to chemicals and radionuclides. | | | |

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OCCURRENCE, DISTRIBUTION AND SELECTION OF CHEMICALS OF POTENTIAL CONCERN (continued)

| Column 3 and consult the EPA risk assessor. | |
|---|---|
| Column 3 - Chemical | L |
| Definition: • The name of the compound detected in samples for the Medium. | |
| Instructions: Provide the names of the chemicals which were detected in the sample for the Medium. | Chemicals can be grouped in the order that the risk assessor prefers. Class descriptions (e.g., PAHs, VOCs, inorganics) can be included as a row before a group of chemicals. |
| Column 4 - Minimum Concentration (Qualifier) | - |
| Minimum Concentration - The lowest detected concentration of the chemical or radionuclide in the medium. Qualifier - The alpha-numeric code assigned to the concentration value by the analytical chemist during data validation for the Minimum Concentration value. | |
| Instructions: Enter the minimum detected concentration for the medium. If there is a detected minimum, enter that as the Minimum Concentration. If the concentration is not detected, enter 'ND' as the Minimum and Maximum Concentrations and record the detection limits in the Range of Detection Limits column. Enter the qualifier associated with the minimum concentration for each chemical or radionuclide in parentheses () after the Minimum Concentration value. Multiple qualifiers should be separated by commas. Provide the definition of each qualifier in the table footnotes. | |

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OCCURRENCE, DISTRIBUTION AND SELECTION OF CHEMICALS OF POTENTIAL CONCERN (continued)

| • | Maximum Concentration - The highest detected concentration of the chemical or radionuclide in the Medium at the current Exposure Point which is above the sample quantitation limit. Qualifier - The alpha-numeric code assigned to the concentration value by the analytical chemist during data validation for the Maximum Concentration value. | |
|-----|--|--|
| Ins | Enter the maximum detected concentration for the medium. Enter the qualifier associated with the Maximum Concentration for each chemical or radionuclide. Provide the definition of each qualifier in the table footnotes. | |

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OCCURRENCE, DISTRIBUTION AND SELECTION OF CHEMICALS OF POTENTIAL CONCERN (continued)

| Definition: • The concentration units for each chemical or radionuclide detected. | |
|---|--|
| Instructions: • Enter the concentration units for each chemical or radionuclide. Units may vary among matrices/media. | Consult with the EPA risk assessor to determine if there is a preference regarding the units used for different matrices (e.g., mg/kg for soil, 2 g/L for groundwater). Choices include: mg/l 2 g/l ng/l pg/l % ppm ppb ppt g/kg mg/kg 2 g/kg ng/kg 2 g/g mg/m³ 2 g/m³ fibers/l fibers/m³ fibers/kg lbs/day 2 g/100cm² mg/cm² 2 Rem/hr Rem/yr pCi/g pCi/kg pCi/m³ pCi/l |
| olumn 7 - Location of Maximum Concentration | Not Documented |
| Definition: • The sample number that identifies the location where the highest concentration sample was taken. | |
| Instructions:Enter the sample identifier which corresponds to the location where the sample was taken. | |
| olumn 8 - Detection Frequency | • |
| Definition: The number of times the chemical or radionuclide was detected versus the number of times it was analyzed, expressed as the "fraction" X/Y. | For example, 5/9 indicates that a chemical was detected in 5 out of 9 samples. |
| Instructions: Indicate the number of times the chemical or radionuclide was detected versus the number of times it was analyzed as the "fraction" X/Y. | Consult the EPA risk assessor for an explanation of how Detection Frequency should be interpreted and applied. |

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OCCURRENCE, DISTRIBUTION AND SELECTION OF CHEMICALS OF POTENTIAL CONCERN (continued)

| De • | finition: The lowest and highest detection limits. | Consult the EPA risk assessor for definitions of detection limits. |
|---------------|---|---|
| Ins • | Enter the lowest and highest detection limit for the chemical or radionuclide in the medium separated by a dash (-). Consult with the EPA risk assessor if detection limits are not reported | |
| Column 10 - C | Concentration Used for Screening | |
| De • | finition: The detected concentration which was used to compare to the screening value. | |
| Ins • | Enter a concentration for each chemical being evaluated for the Medium. Use a footnote to specify the source(s) of the Concentration Used for Screening. | Consult the EPA risk assessor when determining this value. For example, maximum or average. |
| Column 11 - B | Background Value | |
| De • | finition: The background value for the chemical or radionuclide in that Medium as defined by guidance. If a "t-test" or other test which requires backup information is required, this supporting information is should be provided separately. | |
| Ins • • | Enter the numerical value in the column. Specify the source(s)/derivation of the Background Value in table footnotes. For example, literature value, data from a nearby site, statistical tool. | Consult the EPA risk assessor for how background values are determined and whether and how background values are considered for COPC screening. |
| Column 12 - S | Screening Toxicity Value (N/C) | |
| De • | finition: The screening level used to compare detected concentrations of chemicals and radionuclides. Screening Toxicity Values are usually risk-based media concentrations (e.g., RBCs, SSLs, PRGs). | |

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OCCURRENCE, DISTRIBUTION AND SELECTION OF CHEMICALS OF POTENTIAL CONCERN (continued)

| Instructions: Enter the Screening Toxicity Value. Also indicate, with (N) or (C) whether the value is based on non-cancer or cancer effects, respectively. To enter both the cancer and non-cancer screening toxicity values, either (1) record both in the same cell separated by a "/" (e.g., 15C/3.8N), or record one value in Column 12 and one in Column 13. Use a footnote to provide a reference/explanation for the source of the screening values used. | Consult the EPA risk assessor for the source of the screening value and for guidance on comparing the screening value to detected concentrations. |
|--|---|
| Column 13 - Potential ARAR/TBC Value | |
| Definition: • Potential applicable or relevant and appropriate requirements (ARAR) and to be considered (TBC) values. | For example, MCL values, soil cleanup level values, or other values to be considered. |
| Instructions: If multiple values exist, then enter the most conservative ARAR or TBC value. | Consult the EPA risk assessor regarding the requirements for this column. |
| Column 14 - Potential ARAR/TBC Source | • |
| Definition: • The type or source of the ARAR/TBC value entered into the previous column. | For example, MCL or SMCL. |
| Instructions: • Enter the type or source of ARAR/TBC value which corresponds to the value in the previous column. | |
| Column 15 - COPC Flag (Y/N) | |
| Definition: • A code which identifies whether the chemical or radionuclide has been selected as a chemical of potential concern. | |
| Instructions:Enter "Y" or "N" to indicate whether the chemical has been retained as a COPC. | Y N |

B2-8 December 2001

OCCURRENCE, DISTRIBUTION AND SELECTION OF CHEMICALS OF POTENTIAL CONCERN (continued)

| Column 16 - Rationale for Selection or Deletion | | |
|---|--|--|
| Definition: The reason that the chemical or radionuclide was selected or not selected for quantitative or qualitative analysis. | Consult the EPA risk assessor for the rationale codes. | |
| Instructions: Enter the rationale codes for selection/deletion of chemicals of potential concern. Separate multiple codes with commas. Define the codes for the "Rationale for Selection or Deletion" column in a footnote on this table. | The example data table provides rationale codes for example purposes only. | |

B2-9 December 2001

EXPOSURE POINT CONCENTRATION SUMMARY

| PURPOSE OF THE TABLE: To provide the Exposure Point Concentrations (EPCs) for measured and modeled values To provide statistical information on the derivation of the EPCs. | |
|---|--|
| INFORMATION DOCUMENTED: Statistical information which was used to calculate the EPCs for chemicals and radionuclides detected in each Medium Exposure Point Concentrations (RME and/or CT) The statistics which were used to make the determinations as well as the rationale for the selection of the statistics for each chemical or radionuclide (i.e., discuss statistical derivation of measured data or approach for modeled data). | |
| TABLE NUMBERING AND SUMMARY BOX INSTRUCTIONS: Follow the instructions below to create separate sets of Table 3 for RME and CT when appropriate. Complete one copy of Table 3 for each unique combination of the following three fields that will be quantitatively evaluated: Scenario Timeframe, Medium, and Exposure Medium. Enter each combination of these three fields in the Summary Box in the upper left corner of the table. Number each table uniquely, beginning with 3.1 and ending with 3.n, where "n" represents the total number of combinations of the three key fields. Add the extension .RME or .CT to the table number to indicate reasonable maximum exposure or central tendency. Add the line "Reasonable Maximum Exposure" or "Central Tendency" to the table title. | It is possible that some tables may contain the same data associated with different descriptions in the Summary Box in the upper left corner. Separate tables may be necessary to ensure transparency in data presentation for each Exposure Pathway. Replication of information is readily accomplished using spreadsheet software. Consult the EPA risk assessor for alternatives (e.g., footnotes) to preparing multiple tables with the same data. |

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EXPOSURE POINT CONCENTRATION SUMMARY (continued)

GENERAL NOTES/INSTRUCTIONS FOR THIS TABLE:

- Attach supporting documentation regarding how the EPC was calculated.
- Attach an example calculation so the methodology used to develop EPCs is clear to a reviewer.
- Attach supporting information regarding how the concentration term was selected.
- Consult the EPA risk assessor concerning use of decimals or scientific notation for data.
- For certain media, all columns will not be completed.

This information should be of sufficient detail that a reviewer can check and verify the calculations which were performed and obtain the same results as listed in this table.

It is possible that the 95% UCL may not need to be calculated, for example, if only one data point is being considered.

As another example, in some regions, the arithmetic average of concentrations measured from the center of the plume is used as the RME. In this case, the 95% UCL column does not need to be completed.

B3-2 December 2001

EXPOSURE POINT CONCENTRATION SUMMARY (continued)

| HOW TO COMPLETE/INTERPRET THE TABLE SUMMARY BOX IN UPPER LEFT CORNER | |
|--|--|
| | |
| Definition: • The time period (current and/or future) being considered for the exposure pathway. | |
| Instructions:Choose from the picklist to the right. | Current Future Current/Future Not Documented |
| Row 2 - Medium | |
| Definition: The substance (e.g., air, water, soil) that is a potential source of contaminants in the Exposure Medium. (The Medium will sometimes = the Exposure Medium.) Usually, the Medium is that targeted for possible remediation. | |
| Instructions: • Choose from the picklist to the right. | Groundwater Leachate Sediment Sludge Soil Surface Water Debris Other Liquid Waste Solid Waste Air Surface Soil Subsurface Soil |

B3-3 December 2001

EXPOSURE POINT CONCENTRATION SUMMARY (continued)

| v 3 - Exposure Medium | |
|---|--|
| Definition: • The contaminated environmental medium to which an individual may be exposed. Includes the transfer of contaminants from one medium to another. | |
| Contaminants in Groundwater (the Medium) remain in Groundwater (the Exposure Medium) and are available for exposure to receptors. Contaminants in Groundwater (the Medium) may be transferred to Air (the Exposure Medium) and are available for exposure to receptors. Contaminants in Sediment (the Medium) may be transferred to Fish Tissue | |
| Instructions: • Choose from the picklist to the right. | Groundwater Leachate Sediment Sludge Soil Surface Water Debris Other Liquid Waste Solid Waste Air Plant Tissue Animal Tissue Fish Tissue Spring Water Surface Soil Subsurface Soil |

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EXPOSURE POINT CONCENTRATION SUMMARY (continued)

| BODY OF THE TABLE | | |
|---|---|--|
| Column 1 - Exposure Point | | |
| Definition: An exact location of potential contact between a person and a chemical or radionuclide within an Exposure Medium. For example: | | |
| Contaminants are in Groundwater (the Medium and the Exposure Medium) and exposure to Aquifer 1 - Tap Water (the Exposure Point) is evaluated. Contaminants in Groundwater (the Medium) may be transferred to Air (the Exposure Medium) and exposure to Aquifer 1 - Water Vapors at Showerhead (the Exposure Point) is evaluated. | | |
| Contaminants in Sediment (the Medium) may be transferred to Fish Tissue (the Exposure Medium) and Trout from Dean's Creek (the Exposure Point) is evaluated. | | |
| Instructions:Provide the information as text in the table. | Exposure Point should be defined the same way as was done in Planning Table 1. | |
| Column 2 - Chemical of Potential Concern | | |
| Definition: A chemical or radionuclide that is potentially site-related, with data of sufficient quality, that has been retained for quantitative analysis as a result of the screening documented in Table 2. | | |
| Instructions: Enter the names of the chemicals which were selected as COPCs from Table 2. | Chemicals can be grouped in the order that the risk assessor prefers. Class descriptions (e.g., PAHs, VOCs, inorganics) can be included as a row before a group of chemicals. | |
| Column 3 - Units | | |
| Definition: • The concentration units for each chemical and radionuclide detected. | | |

B3-5 December 2001

EXPOSURE POINT CONCENTRATION SUMMARY (continued)

| | ons: er units for each chemical and radionuclide. Units may vary ong matrices/media. | Consult with the EPA risk assessor to determine if there is a preference regarding the units used for different matrices (e.g., mg/kg for soil, zg/L for groundwater). Choices include: mg/l zg/l ng/l pg/l % ppm ppb ppt g/kg mg/kg zg/kg ng/kg zg/g mg/m³ zg/m³ fibers/l fibers/m³ fibers/kg lbs/day zg/100cm² mg/cm² zRem/hr Rem/yr pCi/g pCi/kg pCi/m³ pCi/l pCi/m²/sec Other Not Documented |
|-----------------|--|---|
| Column 4 - Arit | hmetic Mean | _ |
| Defini • | tion: The arithmetic average of detected concentrations. This is the sum of the data divided by the number of data points. | |
| Instru • | ctions: Enter the arithmetic average of detected concentrations. | For duplicate samples, multiple rounds of sampling, and other data evaluation questions, consult the EPA risk assessor. |
| Column 5 - 95% | UCL (Distribution) | |
| Defini • | tion: The statistic for the 95% Upper Confidence Limit on the arithmetic mean, and the type of distribution. | Consult National guidance (Supplemental Guidance to RAGS: Calculating the Concentration Term, OSWER Directive: 9285.7- 08l, May 1992 or most recent updates) and the EPA risk assessor for calculating this term. |
| Instru • • • | Enter the 95% UCL for each COPC. Indicate the distribution of the 95% UCL with (N) or (T) after the value as follows: N is Normal, T is Transformed (lognormal), NP is Nonparametric, O is Other. Define the codes describing the type of distribution in a footnote. Specify any assumptions made in calculating the term in footnotes on this table. Supporting information should be provided in the risk assessment. | For example, for non-detects, ½ the sample quantitation limit is sometimes used as a proxy concentration. For duplicate sample results, the average value is sometimes used in the calculation. |

B3-6 December 2001

EXPOSURE POINT CONCENTRATION SUMMARY (continued)

| Column 6 - Maximum Concentration (Qualifier) | |
|---|--|
| Maximum Concentration - The highest detected concentration of the chemical or radionuclide in the Medium at the current Exposure Point which is above the sample quantitation limit. Maximum Qualifier - The alpha-numeric code assigned to the concentration value by the analytical chemist during data validation for the maximum concentration value. | |
| Instructions: Enter the maximum concentration value. Enter the qualifier associated with the maximum concentration. | Provide the definitions of each qualifier in the table footnotes or in supporting information. |
| Column 7 - Exposure Point Concentration Value | |
| The EPC, based on either a statistical derivation of measured data or modeled data, that represents an estimate of the chemical or radionuclide concentration available from a particular Medium or route of exposure. This EPC value will be used to quantify potential cancer risks and non-cancer hazards. For example, the EPC value may be statistically derived by calculating the 95% UCL of measured groundwater contaminant concentrations from multiple residential wells. Alternatively, the EPC value may be selected as a single measured value, if one data point is used to calculate the risk for each residential well individually. In some cases, the EPC value may be a modeled value (e.g., if upgradient groundwater contaminant concentrations are used to model fish tissue concentrations) | The EPC Value may be calculated, measured, or modeled. |
| Instructions: Enter the value in the column. When using modeled data, enter the Exposure Point, COPC, EPC Value, and EPC Rationale, and include a reference to the location of backup information that show how the data were modeled in the risk assessment document. | Consult the EPA risk assessor concerning how to determine this value. |

B3-7 December 2001

EXPOSURE POINT CONCENTRATION SUMMARY (continued)

| Definition: • The units of the data being used to calculate the EPC. | |
|---|---|
| Instructions: Enter the units for the data being used to calculate the EPC. | Consult the EPA risk assessor for preferences for different media (e.g., ug/L for groundwater; mg/kg for soil). |
| Column 9 - Exposure Point Concentration Statistic | |
| Definition: • The statistic selected to represent the EPC Value based on the distribution of the data, number of data points, etc., and consultation with the EPA risk assessor. | Often, this is 95% UCL of the log- transformed data. |
| Instructions: Enter the statistic used by choosing from the picklist to the right. Define the codes used for the EPC Statistic column in table footnotes. If the statistic used is not on the picklist, enter an abbreviation in Column 9 and provide a description of the statistic in the footnotes of the table. | Max (Maximum) 95% UCL - N (95% UCL of Normal Data) 95% UCL- T (95% UCL of Log-transformed Data) 95% UCL - NP (Mean of Nonparametric Data) Mean - N (Mean of Normal Data) Mean - T (Mean of Log- transformed Data) Mean - NP (Mean of Nonparametric Data) |
| Column 10 - Exposure Point Concentration Rationale | |
| Definition: • The reason the cited statistic was used to represent the EPC. | |
| Instructions: • Enter the rationale for the selection. Footnotes can be used. | |

B3-8 December 2001

VALUES USED FOR DAILY INTAKE CALCULATIONS

| PURPOSE OF THE TABLE: To provide the exposure parameters used for intake calculations for each Exposure Pathway (Scenario Timeframe, Medium, Exposure Medium, Exposure Point, Receptor Population, Receptor Age, and Exposure Route) To provide the intake equations or models used for each Exposure Route/Pathway. | | |
|---|--|--|
| INFORMATION DOCUMENTED: Values used for each intake equation for each Exposure Pathway and the reference/rationale for each Intake equation or model used to calculate the intake for each Exposure Pathway. | | |
| TABLE NUMBERING AND SUMMARY BOX INSTRUCTIONS: Follow the instructions below to create separate sets of Table 4 for RME and CT where appropriate. Complete one copy of Table 4 for each unique combination of the following three fields that will be quantitatively evaluated: Scenario Timeframe, Medium, and Exposure Medium. Enter each combination of these three fields in the Summary Box in the upper left corner of the table. Number each table uniquely, beginning with 4.1 and ending with 4.n, where "n" represents the total number of combinations of the three key fields. Add the line "Reasonable Maximum Exposure" or "Central Tendency" to the table title. Add the extension .RME or .CT to the table number to the line indicate reasonable maximum exposure or central tendency. | Information regarding intake calculations is specific to an Exposure Pathway. Thus, the Summary Box contains the first three identifiers used to specify an exposure pathway: Scenario Timeframe, Medium, and Exposure Medium. It is possible that some tables may contain the same data associated with different descriptions in the Summary Box in the upper left corner. Separate tables may be necessary to ensure transparency in data presentation for each Exposure Pathway. Replication of information is readily accomplished using spreadsheet software. Consult the EPA risk assessor for alternatives (e.g., footnotes) to preparing multiple tables with the same data. | |
| HOW TO COMPLETE/INTERPRET THE TABLE | | |
| SUMMARY BOX IN UPPER LEFT CORNER | | |
| Row 1 - Scenario Timeframe | | |
| Definition:The time period (current and/or future) being considered for the Exposure Pathway. | | |

B4-1 December 2001

VALUES USED FOR DAILY INTAKE CALCULATIONS (continued)

| Instructions: Current Future Current/Future Not Documented |
|--|
|--|

VALUES USED FOR DAILY INTAKE CALCULATIONS (continued)

| Row 2 - Medium | |
|---|--|
| Definition: The substance (e.g., air, water, soil) that is a potential source of contaminants in the Exposure Medium. (The Medium will sometimes = the Exposure Medium.) Usually, the Medium is the targeted for possible remediation. | |
| Instructions: • Choose from the picklist to the right. | Groundwater Leachate Sediment Sludge Soil Surface Water Debris Other Liquid Waste Solid Waste Air Surface Soil Subsurface Soil |
| Row 3 - Exposure Medium | |
| Definition: The contaminated environmental medium to which an individual may be exposed. Includes the transfer of contaminants from or Medium to another. | |
| For example: 1) Contaminants in Groundwater (the Medium) remain in Groundwater (the Exposure Medium) and are available for exposure to receptors. 2) Contaminants in Groundwater (the Medium) may be transferred to Air (the Exposure Medium) and are available for exposure to receptors. 3) Contaminants in Sediment (the Medium) may be transferred to Fish Tissue (the Exposure Medium) and are available for exposure to receptors. | |

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VALUES USED FOR DAILY INTAKE CALCULATIONS (continued)

| Instructions: • Choose from the picklist to the right. | Groundwater Leachate Sediment Sludge Soil Surface Water Debris Other Liquid Waste Solid Waste Air Plant Tissue Animal Tissue Fish Tissue Spring Water Surface Soil |
|---|--|
| | |

B4-4 December 2001

VALUES USED FOR DAILY INTAKE CALCULATIONS (continued)

| BODY OF THE TABLE | | |
|--|---|--|
| Column 1 - Exposure Route | | |
| Definition: • The way a chemical or radionuclide comes in contact with a person (e.g., by ingestion, inhalation, dermal contact). | | |
| Instructions:Choose from the picklist to the right. | Inhalation Ingestion Combined (i.e., Inhalation and Ingestion) Dermal Not Documented External (Radiation) | |
| Column 2 - Receptor Population | | |
| Definition: • The exposed individual relative to the Exposure Pathway considered. | For example, a resident (Receptor Population) who drinks contaminated groundwater. | |
| Instructions: • Choose from the picklist to the right. | Resident Industrial Worker Commercial Worker Construction Worker Other Worker Golfer Jogger Fisher Hunter Fisher/Hunter Swimmer Other Recreational Person Child at School/Daycare/ Playground Trespasser/Visitor Farmer Gardener Gatherer Other | |
| Column 3 - Receptor Age | | |
| Definition: • The description of the exposed individual as defined by the EPA Region or dictated by the site. | For example, a resident (Receptor Population) who drinks contaminated groundwater. | |

B4-5 December 2001

VALUES USED FOR DAILY INTAKE CALCULATIONS (continued)

| Insti • | ructions: Choose from the picklist to the right. | Child Adult Adolescents (teens) Pre-Adolescents Not Documented Child/Adult Geriatric Sensitive Other Infant Toddler Pregnant |
|---------------------------|--|--|
| Column 4 - | Exposure Point | |
| Defi • | nition: An exact location of potential contact between a person and a chemical or radionuclide within an Exposure Medium. For example: 1) Contaminants are in Groundwater (the Medium and the Exposure Medium) and exposure to Aquifer 1 - Tap Water (the Exposure Point) is evaluated. 2) Contaminants in Groundwater (the Medium) may be transferred to Air (the Exposure Medium) and exposure to Aquifer 1 - Water Vapors at Showerhead (the Exposure Point) is evaluated. 3) Contaminants in Sediment (the Medium) may be transferred to Fish Tissue (the Exposure Medium) and Trout in Dean's Creek (the Exposure Point) is evaluated. | |
| Instructions: • | Provide the information as text in the table. Multiple Exposure Points may be recorded in the same cell/row in this table if all other aspects of their Exposure Pathways (Scenario Timeframe, Medium, Exposure Medium, Exposure Route, Receptor Population and Receptor Age) are the same. | Exposure Points should be defined the same way ad was done in Planning Table 1. |
| Column 5 - Parameter Code | | |
| Defi • | nition: The code used for parameters (exposure factors) in the intake equation. | |

B4-6 December 2001

VALUES USED FOR DAILY INTAKE CALCULATIONS (continued)

| Instructions: Enter the appropriate code for the intake parameter from the picklist below. Develop additional intake parameter codes as necessary; be sure that additional codes are unique and defined in this table. | | Do not provide detailed information regarding parameter modeled intakes in this table. This information should be provided separately. Column 10 of this table should list the name of the model or the equation used with a footnote | |
|--|--|---|---|
| Parameter Code | Parameter Definition | Units | referencing supporting information regarding modeled intake development. |
| CS CW IR-W EF ED CF1 BW AT-C AT-N KP ET CF2 SA IN | Chemical Concentration in Soil Chemical Concentration in Water Ingestion Rate of Water Exposure Frequency Exposure Duration Conversion Factor 1 Body Weight Averaging Time (Cancer) Averaging Time (Non-Cancer) Permeability Constant (Dermal for Liquids) Exposure Time Conversion Factor 2 Skin Surface Area Available for Contact Inhalation Rate | mg/kg ug/l liters/day days/year years mg/ug kg days days cm/hr hr/day l/cm3 cm2 m³/hr | |
| IR-SM IR-S DABS SSAF IR-F EF-F | Ingestion Rate (Swimming) Ingestion Rate of Soil Dermal Absorption Factor (Solid) Soil to Skin Adherence Factor Ingestion Rate of Food Exposure Frequency (Food) | l/hr mg/day mg/cm²/event kg/meal meals/year | |
| Definition • The | n: name of the exposure factor (e.g., ing ght) used in the intake equation corresponder entered in Column 5 | | |
| und | ons: er the parameter definition, consistent ver the Parameter Code column. elop additional intake parameter definition | • | Do not provide detailed parameter information regarding modeled intakes in this table. This information should be provided separately. (See instructions for Column 5). |
| Column 7 - Valu | ie | | |
| | n: numeric value of the parameter record the intake calculation. | led in Column 6 used | |

B4-7 December 2001

VALUES USED FOR DAILY INTAKE CALCULATIONS (continued)

Instructions:

- Enter the values used for intake calculations.
- For the CS and CW (chemical concentrations in soil and water, respectively) parameters, refer to Table 3.n or supporting documentation, as appropriate.

Consult the EPA risk assessor for intake parameter values appropriate for each Exposure Pathway.

B4-8 December 2001

VALUES USED FOR DAILY INTAKE CALCULATIONS (CONTINUED)

| Column 8 - Units | |
|---|--|
| Definition: • The units for the parameter code used in the intake equation. | |
| Instructions: Enter the units for each parameter code consistent with the picklist defined under Column 5. Develop additional intake parameter units as necessary. | Consult with the EPA risk assessor to determine if there is a preference regarding the units used for different matrices (e.g., mg/kg for soil, 2 g/L for groundwater). Choices include: |
| | mg/l |
| Column 9 - Rationale/Reference | |
| Definition: • The reason and reference for the parameter value used. | This rationale may be based upon guidance or consultation with the EPA risk assessor. |
| Instructions: Enter the rationale and reference for the value. If the value used is inconsistent with guidance values, provide a detailed explanation of the rationale and a complete reference for the value used. | Provide sufficient detail that the reviewer can easily substantiate the value. |
| Column 10 - Intake Equation/Model Name | |
| Definition: • The calculation, equation, or model used for intake estimates for each Exposure Route. | |
| Instructions: • Enter the intake calculation, equation, and/or model nan • Include a footnote providing a reference to the section the risk assessment where information regarding model intake development is presented. | of the equation used. |

B4-9 December 2001

NON-CANCER TOXICITY DATA - ORAL/DERMAL

| PURPOSE OF THE TABLE: | | |
|--|---|--|
| To provide information on RfDs, target organs, and adjustment | | |
| factors for chemicals | | |
| To provide oral to dermal adjustment factors To verify references for non-cancer toxicity data. | | |
| To verify fereferences for non-cancer toxicity data. | | |
| INFORMATION DOCUMENTED: The RfDs for each of the COPCs, as well as modifying factors and oral to dermal adjustments The organ effects of each of the COPCs References for RfDs and organ effects. | Surrogate toxicity values can also be entered in this table and indicated in the Source(s) column or with a footnote. | |
| TABLE NUMBERING INSTRUCTIONS: Complete one copy of this table only. Number it Table 5.1. The table should contain a row for each COPC considered. | If chronic and subchronic effects are listed for the same COPC, two rows will be required. | |
| GENERAL NOTES/INSTRUCTIONS FOR THIS TABLE: • Table 5.1 does not replace the toxicological profiles for the individual chemicals that will be presented in the risk assessment. | It may be necessary to refer to RAGS, the risk assessment technical approach, and the EPA risk assessor to complete the table. | |
| HOW TO COMPLETE/INTERPRET THE TABLE | E | |
| Column 1 - Chemical of Potential Concern | | |
| Definition: | | |
| Chemicals that are potentially site-related, with data of sufficient quality, that have been retained for quantitative analysis as a result of the screening documented in Table 2. | | |
| Instructions: • Enter the names of the chemicals that were selected as COPCs from Table 2. | Chemicals can be grouped in the order that the risk assessor prefers. Class descriptions (e.g., PAHs, VOCs, inorganics) can be included as a row before a group of chemicals. | |
| Column 2 - Chronic/Subchronic | | |
| Definition: | | |
| Identifies whether the RfD for a particular chemical is for chronic (long-term) and/or subchronic (short-term) exposure. | | |

B5.1-1 December 2001

NON-CANCER TOXICITY DATA - ORAL/DERMAL (continued)

| Instructions: Enter either "Chronic" or "Subchronic" in the field. Both values may be available for an individual COPC. Subchronic values may not be available or necessary for an individual COPC. If that is the case, enter only "Chronic" in Column 2. | Chronic Subchronic | |
|---|---|--|
| Column 3 - Oral RfD Value | | |
| Definition: • The oral RfD value for each of the COPCs. | | |
| Instructions:Enter the value for the chronic and/or subchronic oral RfD (as appropriate). | | |
| Column 4 - Oral RfD Units | | |
| Definition: • The oral RfD units for each COPC. | | |
| Instructions:Enter units for each oral RfD value as necessary. | Consult the EPA risk assessor to determine if there is a preference regarding the units to be used. | |
| Column 5 - Oral Absorption Efficiency Value for Dermal | | |
| Definition: • The adjustment factor used to convert oral RfD values to dermal RfD values. This value is an oral absorption factor. | | |
| Instructions: Enter the adjustment factor in this column. Use a footnote to indicate the source of the Oral Absorption Efficiency for Dermal. Also, specify the section of the risk assessment text where the derivation of the Oral Absorption Efficiency for Dermal can be found. | | |
| Column 6 - Absorbed RfD for Dermal Value | | |
| Definition: • The adjusted RfD for each COPC detected that is derived from the oral RfD. | | |

B5.1-2 December 2001

NON-CANCER TOXICITY DATA - ORAL/DERMAL (continued)

| Instructions: Enter the value that was derived from the adjustment factor in Column 5. In a footnote on this table, reference the section of the risk assessment text where the derivation of the Absorbed RfDs for Dermal can be found. | Derivations of the Absorbed RfD for Dermal should be performed in as directed by the EPA risk assessor. |
|--|--|
| Column 7 - Absorbed RfD for Dermal Units | |
| Definition: • The units associated with the Absorbed RfD for Dermal value for each COPC. | |
| Instructions:Enter units for each Absorbed RfD for Dermal value as necessary. | Consult the EPA risk assessor to determine if there is a preference regarding the units to be used. |
| Column 8 - Primary Target Organ(s) | |
| Definition: The organ(s) most affected (i.e., experiences critical effects) by chronic or subchronic exposure to the specific COPC, and upon which the RfD is based. | |
| Instructions: Enter the name of the most affected organ or organ system in the column. If the critical effect (the one on which the RfD is based) involves multiple target organs, they should be shown, separated by a '/.' Target organs that are affected at higher doses should not be shown. | |
| Column 9 - Combined Uncertainty/Modifying Factors | |
| Definition: • The factors applied to the critical effect level to account for areas of uncertainty inherent in extrapolation from available data. | Refer to IRIS, HEAST, or other source for these values. Examples of uncertainty to be addressed include: - variations in the general population - interspecies variability between humans and animals - use of subchronic data for chronic evaluation - extrapolation from LOAELs to NOAELs. |
| Instructions:Enter number obtained from IRIS, HEAST, or other source. | Refer to IRIS, HEAST, or other source for these values. |

B5.1-3 December 2001

NON-CANCER TOXICITY DATA - ORAL/DERMAL (continued)

| Column 10 - RfD: Target Organ(s) Source(s) | | |
|---|---|--|
| Definition: • The source of the RfD and target organ information. | | |
| Instructions: Enter the source of the RfD and target organ information. Use a colon to delineate multiple sources if the sources of information are different for RfD and target organ. | IRIS HEAST NCEA OTHER | |
| Column 11 - RfD: Target Organ(s) Dates (MM/DD/YYYY) | | |
| Definition: • The date of the source that was consulted for the RfD and target organ information in MM/DD/YYYY format. | The MM/DD/YYYY format refers to month/day/year. | |
| Instructions: Enter the date, in MM/DD/YYYY format, for both RfD and target organ information. Use a colon to delineate multiple dates if the dates of information are different for RfD and target organ. For IRIS references, provide the date IRIS was searched. For HEAST references, provide the date of the HEAST reference. For NCEA references, provide the date of the information provided by NCEA. | For example, the MM/DD/YYYY version of the date March 30, 1995 is 03/30/1995. | |

B5.1-4 December 2001

NON-CANCER TOXICITY DATA - INHALATION

| PURPOSE OF THE TABLE: To provide information on RfCs, RfDs, target organs, and adjustment factors for chemicals To provide RfC to RfD adjustment factors To verify references for non-cancer toxicity data. | |
|---|--|
| INFORMATION DOCUMENTED: The RfDs for each of the COPCs, as well as modifying factors and RfC to RfD adjustments The primary target organ effects of each of the COPCs References for RfCs and organ effects. | Surrogate toxicity values can also be entered in this table and indicated in the Source(s) column or with a footnote. |
| TABLE NUMBERING INSTRUCTIONS: Complete one copy of this table only. Number it Table 5.2. The table should contain a row for each COPC considered. | If chronic and subchronic effects are listed for the same COPC, two rows will be required. |
| GENERAL NOTES/INSTRUCTIONS FOR THIS TABLE: Table 5.2 does not replace the toxicological profiles for the individual chemicals that will be presented in the risk assessment. | It may be necessary to refer to RAGS, the risk assessment technical approach, and EPA Regional guidance to complete the table. |
| HOW TO COMPLETE/INTERPRET THE TABLE | : |
| Column 1 - Chemical of Potential Concern | |
| Definition: Chemicals that are potentially site-related, with data of sufficient quality, that have been retained for quantitative analysis as a result of the screening documented in Table 2. | |
| Instructions: Enter the names of the chemicals that were selected as COPCs from Table 2. | Chemicals can be grouped in the order that the risk assessor prefers. Class descriptions can be included as a row before a group of chemicals. |
| Column 2 - Chronic/Subchronic | |
| Definition: • Identifies whether the RfC or RfD for a particular chemical is for chronic (long-term) and/or subchronic (short-term) exposure. | |
| Instructions: Enter either "Chronic" or "Subchronic" in the field. Both values may be available for an individual chemical. "Subchronic" values may not be available or necessary for an individual COPC. If that is the case, enter "Chronic" in Column 2. | Chronic Subchronic |

B5.2-1 December 2001

NON-CANCER TOXICITY DATA - INHALATION (continued)

| Column 3 - Inhalation RfC Value | | |
|---|---|--|
| Definition: • The RfC value for each of the COPCs. | | |
| Instructions:Enter the value for the chronic and/or subchronic oral RfC (as appropriate). | | |
| Column 4 - Inhalation RfC Units | | |
| Definition: • The RfC units for each chemical detected. | | |
| Instructions:Enter units for each RfC as necessary. | Consult the EPA risk assessor to determine if there is a preference regarding the units to be used. | |
| Column 5 - Extrapolated RfD Value | | |
| Definition: • The inhalation RfD for each COPC that is derived from the RfC value if an RfD is used to calculate risk instead of the RfC. | The derivation of the RfD from an RfC should be performed as directed by the EPA risk assessor. | |
| Instructions: Enter the derived RfD factor in this column. In a footnote on this table, reference the section of the risk assessment text where the derivation of the adjusted RfDs can be found. | The equation to derive the RfD from the RfC is to be included as a footnote in the table. | |
| Column 6 - Extrapolated RfD Units | | |
| Definition: • The Extrapolated RfD units for each COPC. | | |
| Instructions:Enter units for each Extrapolated RfD value as necessary. | Consult the EPA risk assessor to determine if there is a preference regarding the units to be used. | |
| Column 7 - Primary Target Organ(s) | | |
| Definition: The organ that is most affected (i.e., experiences critical effects) by chronic or subchronic exposure to the specific COPC, and upon which the RfD/RfC is based. | | |

B5.2-2 December 2001

NON-CANCER TOXICITY DATA - INHALATION (continued)

| Instructions:Enter the name of the most affected organ or organ system in the | |
|--|--|
| column. If the critical effect (the one on which the RfD/RfC is based) involves multiple target organs, they should all be shown, separated by '/.' Target organs affected at higher doses should not be shown. | |
| Column 8 - Combined Uncertainty/Modifying Factors | |
| Definition: The factors applied to the critical effect level to account for areas of uncertainty inherent in extrapolation from available data. | Refer to IRIS, HEAST, or other source for these values. Examples of uncertainty to be addressed include: - variations in the general population - interspecies variability between humans and animals - use of subchronic data for chronic evaluation - extrapolation from LOAELs to NOAELs. |
| Instructions:Enter number obtained from IRIS, HEAST, or other source. | Refer to IRIS, HEAST, or other source for these values. |
| Column 9 - RfC: Target Organ(s) Source(s) | |
| Definition: • The sources of the RfC and target organ information. | |
| Instructions: Enter the sources of the RfC and target organ information. Use a colon to delineate between multiple information sources if the sources of information are different for RfC and target organ. | IRIS HEAST NCEA OTHER |
| Column 10 - RfC: Target Organ(s) Date(s) (MM/DD/YYYY) | |
| Definition: The dates of the documents that were consulted for the RfC and target organ information in MM/DD/YYYY format. | The MM/DD/YYYY format refers to month/day/year. |

B5.2-3 December 2001

Instructions:

 Enter the dates, in MM/DD/YYYY format, for RfC and target organ information. Use a colon to delineate between multiple dates if the dates of information are different for RfC and target organ. For example, the MM/DD/YYYY version of the date March 30, 1995 is 03/30/1995.

- For IRIS references, provide the date IRIS was searched.
- For HEAST references, provide the date of the HEAST reference.
- For NCEA references, provide the date of the information provided by NCEA.

B5.2-4 December 2001

NON-CANCER TOXICITY DATA - SPECIAL CASE CHEMICALS

| PURPOSE OF THE TABLE: To provide information on toxicity values, target organs, and adjustment factors for unusual chemicals or circumstances or surrogate chemicals that are not covered by Tables 5.1 or 5.2. Table 5.3 is not required if there are not such chemicals or circumstances. To verify references for non-cancer toxicity data. | For example, a toxicity factor derived specifically for an individual risk assessment should be documented in Table 5.3. |
|--|---|
| INFORMATION DOCUMENTED: The toxicity values for each of the COPCs, as well as modifying factors The organ effects of each of the COPCs References for toxicity values and organ effects. | |
| TABLE NUMBERING INSTRUCTIONS: Complete one copy of this table only. Number it Table 5.3. The table should contain a row for each COPC considered. | If chronic and subchronic effects are listed for the same COPC, two rows will be required. |
| GENERAL NOTES/INSTRUCTIONS FOR THIS TABLE: Table 5.3 does not replace the toxicological profiles for the individual chemicals that will be presented in the risk assessment. | Refer to RAGS, the risk assessment technical approach, and the EPA risk assessor to complete the table. |
| HOW TO COMPLETE/INTERPRET THE TABLE | Ε |
| Column 1 - Chemical of Potential Concern | |
| Definition: Chemicals that are potentially site-related, with data of sufficient quality, that have been retained for quantitative analysis as a result of the screening documented in Table 2. | |
| Instructions: Enter the names of the chemicals that were selected as COPCs from Table 2. | Chemicals can be grouped in the order that the risk assessor prefers. Class descriptions (e.g., PAHs, VOCs, inorganics) can be included as a row before a group of chemicals. |
| Column 2 - Chronic/Subchronic | |
| Definition: • Identifies whether the toxicity value for a particular chemical is for chronic (long-term) and/or subchronic (short-term) exposure. | |

B5.3-1 December 2001

NON-CANCER TOXICITY DATA -SPECIAL CASE CHEMICALS (continued)

| Instr | Enter either "Chronic" or "Subchronic" in the field. Both values may be available for an individual COPC. "Subchronic" values may not be available or necessary for an individual chemical. If that is the case, enter only "Chronic" in the column. | Chronic Subchronic | |
|------------|---|---|--|
| Column 3 - | Parameter Name | | |
| Defi • | nition: The name of parameter/toxicity factor being recorded for each COPC. | Toxicity factors derived specifically for an individual risk assessment should be recorded here. | |
| Instr • | ructions: Enter the name of parameter/toxicity factor. | | |
| Column 4 - | Column 4 - Parameter Value | | |
| Defi | nition: The toxicity parameter value for each COPC. | | |
| Instr • | ructions: Enter the value for the chronic and/or subchronic toxicity values (as appropriate). | | |
| Column 5 - | Parameter Units | | |
| Defi | nition: The units associated with the toxicity value for each COPC. | | |
| Instr • | ructions: Enter units for each reference as necessary. | Consult the EPA risk assessor to determine if there is a preference regarding the units to be used. | |
| Column 6 - | Primary Target Organ(s) | | |
| Defi • | nition: The organ(s) most affected (i.e., experiences critical effects) by chronic or subchronic exposure to the specific COPC, and upon which the RfD is based. | | |
| Instr • | Enter the name of the most affected organ or organ system in the column. If the critical effect (the one that the RfD is based on) involves multiple target organs, they should all be shown, separated by a '/.' Target organs affected at higher doses should not be shown. | | |

B5.3-2 December 2001

NON-CANCER TOXICITY DATA -SPECIAL CASE CHEMICALS (continued)

| Column 7 - Combined Uncertainty/Modifying Factors | |
|---|--|
| Definition: • The factors applied to the critical effect level to account for areas of uncertainty inherent in extrapolation from available data. | Refer to IRIS, HEAST, or other source for these values. Examples of uncertainty to be addressed include: - variations in the general population - interspecies variability between humans and animals - use of subchronic data for chronic evaluation - extrapolation from LOAELs to NOAELs. |
| Instructions:Enter number obtained from IRIS, HEAST, or other source. | Refer to IRIS, HEAST, or other source for these values. |
| Column 8 - Parameter: Target Organ(s) Sources | |
| Definition: • The sources of the toxicity and target organ information. | |
| Instructions: Enter the sources of the toxicity and target organ information. Use a colon to delineate multiple sources if the sources of information for toxicity and target organ are different. | IRIS HEAST NCEA OTHER |
| Column 9 - Parameter: Target Organ(s) Date(s) (MM/DD/YYYY) | |
| Definition: • The dates of the sources that were consulted for the toxicity information and the target organ information in MM/DD/YYYY format. | The MM/DD/YYYY format refers to month/day/year. |
| Instructions: Enter the dates, in MM/DD/YYYY format, for the toxicity and target organ information. Use a colon to delineate between multiple dates if the sources of information are different for toxicity and target organ. For IRIS references, provide the date IRIS was searched. | For example, the MM/DD/YYYY version of the date March 30, 1995 is 03/30/1995. |
| For HEAST references, provide the date of the HEAST reference. For NCEA references, provide the date of the information provided by NCEA. | |

B5.3-3 December 2001

CANCER TOXICITY DATA - ORAL/DERMAL

| PURPOSE OF THE TABLE: • To provide the oral and dermal cancer toxicity information (values and sources of information) for chemicals of potential concern To provide the methodal account of interprets for toward to the methodal account of the provide the p | | |
|---|---|--|
| To provide the methodology and adjustment factors used to convert oral cancer toxicity values to dermal toxicity values To provide weight of evidence/cancer guideline descriptions for each chemical of potential concern. | | |
| INFORMATION DOCUMENTED: Oral and dermal toxicity values for chemicals of potential concern Weight of evidence/cancer guidelines descriptions for chemicals of potential concern The source/reference for each toxicity value. | Surrogate toxicity values can also be entered in this table and indicated in the 'Source(s)' column or with a footnote. | |
| GENERAL NOTES/INSTRUCTIONS FOR THIS TABLE: Table 6.1 does not replace toxicological profiles for the individual chemicals that will be presented in the risk assessment. | It may be necessary to refer to RAGS, the risk assessment technical approach, and the EPA risk assessor to complete the table. | |
| HOW TO COMPLETE/INTERPRET THE TABLE | | |
| Column 1 - Chemical of Potential Concern | | |
| Definition: Chemicals that are potentially site-related, with data of sufficient quality, that have been retained for quantitative analysis as a result of the screening documented in Table 2. | | |
| Instructions: Enter the names of the chemicals that were selected as COPCs from Table 2. | Chemicals may be grouped in the order that the risk assessor chooses. Class descriptions can be included as a row before a group of chemicals. | |
| Column 2 - Oral Cancer Slope Factor Value | | |
| Definition: • Cancer slope factor for ingestion. | | |
| Instructions:Enter the oral cancer slope factor value for each of the COPCs. | Refer to IRIS and HEAST. If toxicity information is not available, | |

B6.1-1 December 2001

CANCER TOXICITY DATA - ORAL/DERMAL (continued)

| Column 3 - Oral Cancer Slope Factor Units | |
|---|--|
| Definition: • Units for the cancer slope factor for ingestion. | |
| Instructions:Enter units for each oral cancer slope factor. | Consult the EPA risk assessor to determine if there is a preference regarding the units to be used. |
| Column 4 - Oral Absorption Efficiency for Dermal | |
| Definition: • The absorbed factor used to convert the oral RfD values to dermal RfD values. | |
| Instructions: Enter the oral to dermal adjustment factor. Use a footnote to indicate the source of the Oral Absorption Efficiency for dermal. | |
| Column 5 - Absorbed Cancer Slope Factor for Dermal Value | |
| Definition: The absorbed dermal cancer slope factor for each chemical of potential concern which typically is derived from the oral cancer slope factor. | Derivation of the dermal cancer slope factor should be performed in consultation with the EPA risk assessor. |
| Instructions: Enter the derived dermal cancer slope factor. Use a footnote to specify the section of the risk assessment text where the derivation of the Absorbed Cancer Slope Factor for Dermal can be found. | |
| Column 6 - Absorbed Cancer Slope Factor for Dermal Units | |
| Definition: • The units associated with each Absorbed Cancer Slope Factor for Dermal. | |
| Instructions:Enter the units for the Absorbed Cancer Slope Factors for Dermal. | Typically (mg/kg-day) ⁻¹ . Consult with the EPA risk assessor to determine if there is a preference regarding the units to be used. |
| Column 7 - Weight of Evidence/Cancer Guideline Description | |

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CANCER TOXICITY DATA - ORAL/DERMAL (continued)

| Definition: • An EPA classification system for characterizing the extent to which the available data indicate that an agent is a human carcinogen. | |
|---|---|
| Instructions: Provide the weight of evidence or cancer guideline description. Choose from the categories to the right. | Weight of Evidence: A - Human carcinogen B1 - Probable human carcinogen - indicates that limited human data are available. B2 - Probable human carcinogen - indicates sufficient evidence in animals and inadequate or no evidence in humans. C - Possible human carcinogen D - Not classifiable as a human carcinogen E - Evidence of noncarcinogenicity Cancer Guideline Description: Known/Likely Cannot be Determined Not Likely |
| Column 8 - Oral CSF Source(s) | |
| Definition: • A reference for the oral cancer slope factor. | |
| Instructions: • Enter the reference for the toxicity information. | For example: IRIS HEAST NCEA |
| Column 9 -Oral CSF Date(s) (MM/DD/YYYY) | |
| Definition: • The date of the document that was consulted for the cancer toxicity data in MM/DD/YYYY format. | The MM/DD/YYYY format refers to month/day/year. |
| Instructions: • Enter the date in MM/DD/YYYY format. • For IRIS references, provide the date IRIS was searched. • For HEAST references, provide the date of the HEAST reference. • For NCEA references, provide the date of the information provided by NCEA. | For example, the MM/DD/YYYY version of the date March 30, 1995 is 03/30/1995. |

6.1-3 December 2001

CANCER TOXICITY DATA - INHALATION

| PURPOSE OF THE TABLE: To provide the inhalation cancer toxicity information (values and sources of information) for chemicals of potential concern To provide the methodology and adjustment factors used to convert inhalation unit risks to inhalation cancer slope factors To provide weight of evidence/cancer guideline descriptions for each chemical of potential concern. | |
|--|---|
| INFORMATION DOCUMENTED: Inhalation toxicity values for chemicals of potential concern Weight of evidence/cancer guidelines descriptions for chemicals of potential concern The source/reference for each toxicity value. | Surrogate toxicity values can also be entered in this table and indicated in the 'Source(s)' column or with a footnote. |
| GENERAL NOTES/INSTRUCTIONS FOR THIS TABLE: Table 6.2 does not replace toxicological profiles for the individual chemicals that will be presented in the risk assessment. | It may be necessary to refer to RAGS, the risk assessment technical approach, and the EPA risk assessor to complete the table. |
| HOW TO COMPLETE/INTERPRET THE TABLE | |
| Column 1 - Chemical of Potential Concern | |
| Definition: Chemicals that are potentially site-related, with data of sufficient quality, that have been retained for quantitative analysis as a result of the screening documented in Table 2. | |
| Instructions: Enter the names of the chemicals that were selected as COPCs from Table 2. | Chemicals may be grouped in the order that the risk assessor chooses. Class descriptions (e.g., PAHs, VOCs, inorganics) can be included as a row before a group of chemicals. |
| Column 2 - Unit Risk Value | |
| Definition: Toxicity values for carcinogenic effects expressed in terms of risk per unit concentration of the substance in the medium where human contact occurs. Cancer slope factors can be calculated from unit risk values. | |
| Instructions: • Enter the inhalation unit risk value | Refer to IRIS and HEAST; if toxicity information is not available, contact EPA's National Center for Environmental Assessment (NCEA) office. |

B6.2-1 December 2001

CANCER TOXICITY DATA - INHALATION (continued)

| Column 3 - Unit Risk Units | |
|---|---|
| Definition: • The units used for the unit risk for each chemical detected. | |
| Instructions:Enter the units for the unit risk values. | Consult the EPA risk assessor to determine if there is a preference regarding the units to be used. |
| Column 4 - Inhalation Cancer Slope Factor Value | |
| Definition: A plausible upper-bound estimate of the probability of a response per unit intake of a chemical over a lifetime. | Usually the cancer slope factor is the upper 95th % confidence limit of the dose-response curve for inhalation. |
| Instructions: Enter the Inhalation Cancer Slope Factor if Cancer Slope Factors were used to calculate risk instead of Inhalation Unit Risks. | |
| Column 5 - Inhalation Cancer Slope Factor Units | |
| Definition: • The units used for the Inhalation Cancer Slope Factor for each chemical detected. | |
| Instructions:Enter the units for the Inhalation Cancer Slope Factors. | Consult EPA risk assessor to determine if there is a preference regarding the units to be used. |
| Column 6 - Weight of Evidence/Cancer Guideline Description | |
| Definition: An EPA classification system for characterizing the extent to which the available data indicate that an agent is a human carcinogen. | |

B6.2-2 December 2001

CANCER TOXICITY DATA - INHALATION (continued)

Instructions:

- Provide the weight of evidence or cancer guideline description.
- Choose from the categories to the right.

Weight of Evidence:

- A Human carcinogen
- B1 Probable human carcinogen indicates that limited human data are available.
- B2 Probable human carcinogen indicates sufficient evidence in animals and inadequate or no evidence in humans.
- C Possible human carcinogen
- D Not classifiable as a human carcinogen
- E Evidence of noncarcinogenicity

Cancer Guideline Description: Known/Likely Cannot be Determined Not Likely

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CANCER TOXICITY DATA - INHALATION (continued)

| Column 7 - Unit Risk: Inhalation Cancer Slope Factor Source(s) | | |
|--|---|--|
| Definition: • A reference for the Unit Risk and Inhalation Cancer Slope Factor values. | | |
| Instructions: Enter the reference(s) for Unit Risk and Inhalation Cancer Slope Factor values. Use a colon to delineate multiple sources. | IRIS HEAST NCEA | |
| Column 8 - Unit Risk: Inhalation Cancer Slope Factor Date(s) (MM/DD/YYYY) | | |
| Definition: • The date of the document that was consulted for the cancer toxicity data in MM/DD/YYYY format. | The MM/DD/YYYY format refers to month/day/year. | |
| Instructions: Enter the date in MM/DD/YYYY format. Use a colon to delineate between multiple dates, if multiple sources of information were used. | For example, the MM/DD/YYYY version of the date March 30, 1995 is 03/30/1995. | |
| For IRIS references, provide the date IRIS was searched. For HEAST references, provide the date of the HEAST reference. For NCEA references, provide the date of the information provided by NCEA. | | |

B6.2-4 December 2001

CANCER TOXICITY DATA - SPECIAL CASE CHEMICALS

| PURPOSE OF THE TABLE: To provide cancer toxicity information for unusual chemicals, surrogate chemicals or circumstances that are not covered by Tables 6.1 or 6.2. Table 6.3 (or non-standard tables) can also be used to accommodate threshold carcinogens, if applicable. Table 6.3 is not required if there are no such chemicals or circumstances. | For example, a toxicity factor derived specifically for an individual risk assessment should be documented in Table 6.3. |
|--|--|
| INFORMATION DOCUMENTED: Cancer toxicity information (values and units) for special case chemicals The date and source of the toxicity information. | |
| TABLE NUMBERING INSTRUCTIONS: Complete one copy of this table only. Number it 6.3. The table should contain a row for each COPC considered. | |
| GENERAL NOTES/INSTRUCTIONS FOR THIS TABLE: Table 6.3 does not replace toxicological profiles for the individual chemicals that will be presented in the risk assessment. | It may be necessary to refer to RAGS, the risk assessment technical approach, and consult the EPA risk assessor to complete the table. |
| HOW TO COMPLETE/INTERPRET THE TABLE | |
| Column 1 - Chemical of Potential Concern | |
| Definition: Chemicals that are potentially site-related, with data of sufficient quality, that have been retained for quantitative analysis as a result of the screening documented in Table 2. | |
| Instructions: Enter the names of the chemicals that were selected as COPCs from Table 2. | Chemicals may be grouped in the order that the risk assessor chooses. Class descriptions can be included as a row before a group of chemicals. |
| Column 2 - Parameter Name | |
| Definition: The name of the toxicity parameter being recorded. | |
| Instructions: | |

B6.3-1 December 2001

CANCER TOXICITY DATA - SPECIAL CASE CHEMICALS (continued)

| Column 3 - Parameter Value | |
|--|---|
| Definition: • The toxicity value for each listed parameter for each chemical of potential concern. | |
| Instructions:Enter the toxicity value for each chemical of potential concern. | Refer to IRIS, HEAST, or other source for these valued. |
| Column 4 - Parameter Units | |
| Definition: • The units associated with the toxicity value. | |
| Instructions: • Enter the toxicity units. | Typically (mg/kg-day) ⁻¹ Consult the EPA risk assessor to determine if there is a preference regarding the units to be used. |
| Column 5 -Source(s) | |
| Definition: • A reference for the cancer toxicity information. | |
| Instructions: • Enter the reference for toxicity information. Use a colon to delineate multiple sources. | IRIS HEAST NCEA OTHER |
| Column 6 - Date(s) (MM/DD/YYYY) | |
| Definition: • The date of the document that was consulted for the cancer toxicity data in the MM/DD/YYYY format. | The MM/DD/YYYY format refers to month/day/year. |
| Instructions: Enter the date in MM/DD/YYYY format. Use a comma to delineate between multiple dates, if multiple sources of information were used. | For example, the MM/DD/YYYY version of the date March 30, 1995 is 03/30/1995. |
| For IRIS references, provide the date IRIS was searched. For HEAST references, provide the date of the HEAST reference. For NCEA references, provide the date of the information provided by NCEA. | |

B6.3-2 December 2001

${\bf CANCER\ TOXICITY\ DATA\ -\ EXTERNAL\ (RADIATION)}$

| PURPOSE OF THE TABLE:To provide cancer toxicity information for radionuclides. | |
|--|--|
| INFORMATION DOCUMENTED: Cancer toxicity information (values and units) for radionuclides. The source and date of the toxicity information. | |
| GENERAL NOTES/INSTRUCTIONS FOR THIS TABLE: Table 6.4 does not replace toxicological profiles for the individual radionuclides that will be presented in the risk assessment. | It may be necessary to refer to RAGS, the risk assessment technical approach, and the EPA risk assessor to complete the table. |
| HOW TO COMPLETE/INTERPRET THE TABLE | 2 |
| Column 1 - Chemical of Potential Concern | |
| Definition: Radionuclides that are potentially site-related, with data of sufficient quality, that have been retained for quantitative analysis as a result of the screening documented in Table 2. | |
| Instructions:Enter the names of the radionuclides that were selected as COPCs from Table 2. | Radionuclides may be grouped in the order that the risk assessor chooses. |
| Column 2 - Cancer Slope Factor Value | |
| Definition: A Cancer Slope Factor is an age-averaged lifetime excess cancer incidence rate per unit intake (or unit exposure for external exposure pathways) and is used to convert the intake to a cancer risk. Ingestion and inhalation slope factors are central estimates in a linear model of the age-averaged, lifetime attributable radiation cancer incidence (fatal and nonfatal cancer) risk per unity of activity inhaled or ingested, expressed as risk/picocurie (pCi). External exposure slope factors are central estimates of the lifetime attributable radiation cancer incidence risk for each year of exposure to external radiation from photon-emitting radionuclides distributed uniformly in a thick layer of soil, and are expressed as risk/yr per pCi/gram of soil. | |
| Instructions:Enter the value of the cancer slope factor for each COPC. | |
| Column 3 - Cancer Slope Factor Units | |
| Definition: • The units associated with the Cancer Slope Factor value. | |

B6.4-1 December 2001

${\bf CANCER\ TOXICITY\ DATA\ -\ EXTERNAL\ (RADIATION)\ (continued)}$

| Instructions:Enter the units for the Cancer Slope Factor value. | Consult the EPA risk assessor to determine if there is a preference regarding the units to be used. | |
|--|---|--|
| Column 4 -Source(s) | | |
| Definition: • A reference for the cancer slope or conversion factor value. | | |
| Instructions: Enter the reference(s) for the cancer slope or conversion factor value. Use a colon to delineate multiple sources. | For example: IRIS HEAST NCEA OTHER | |
| Column 5 - Date(s) (MM/DD/YYYY) | | |
| Definition: • The date of the document that was consulted for the cancer slope or conversion factor value in the MM/DD/YYYY format. | The MM/DD/YYYY format refers to month/day/year. | |
| Instructions: Enter the date in MM/DD/YYYY format. Use a colon to delineate between multiple dates, if multiple sources of information were used. | For example, the MM/DD/YYYY version of the date March 30, 1995 is 03/30/1995. | |
| For IRIS references, provide the date IRIS was searched. For HEAST references, provide the date of the HEAST reference. For NCEA references, provide the date of the information provided by NCEA. | | |

B6.4-2 December 2001

CALCULATION OF CHEMICAL CANCER RISKS AND NON-CANCER HAZARDS

| PURPOSE OF THE TABLE: To provide a summary of the variables used to calculate chemical cancer risks and non-cancer hazards To show the EPC and intake used in the non-cancer hazard and cancer risk calculations To present the result of the calculation for each Exposure Route/Pathway for each COPC To provide the total hazard index and cancer risk for all Exposure Routes/Pathways for the Scenario Timeframe and Receptor presented in this table. | |
|---|--|
| INFORMATION DOCUMENTED: The non-cancer hazard quotient and unit risk for each COPC for each Exposure Route/Pathway The values used for EPC, cancer and non-cancer intakes, reference doses, and reference concentrations. | An alternate presentation is also available with cancer information shown on Table 7a and non-cancer information shown on Table 7b. |
| TABLE NUMBERING AND SUMMARY BOX INSTRUCTIONS: Complete one copy of Table 7 for each unique combination of the following three fields that will be quantitatively evaluated (Scenario Timeframe, Receptor Population, and Receptor Age). Enter each combination of these three fields in the Summary Box | It is possible that some tables may contain some of the same data associated with different descriptions in the Summary Box in the upper left corner. |
| in the upper left corner of the table. Note: Each combination of the three key fields and the first four columns should be found as a row in Table 1. Number each table uniquely, beginning with 7.1 and ending with | Separate tables may be necessary to ensure transparency in data presentation for each Exposure Pathway. Replication of information is readily accomplished using spreadsheet software. |

cancer and cancer hazard calculations.

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CALCULATION OF CHEMICAL CANCER RISKS AND NON-CANCER HAZARDS (continued)

TABLE NUMBERING AND SUMMARY BOX INSTRUCTIONS (continued):

- An optional approach is to report cancer and non-cancer values on separate tables as follows:
 - Number non-cancer tables 7.1A.RME 7.nA.RME or 7.1A.CT 7.nA.CT, where "n" represents the total number of combinations of the three key fields.
 - Number cancer tables 7.1B.RME-7.nB RME or 7.1B.CT-7.nB.CT, where "n" represents the total number of combinations of the three key fields.
 - The first seven columns remain the same for both noncancer or cancer tables. Columns 8-12 contain either the Cancer Risk Calculations data or the Non-Cancer Hazard Calculations data.
 - See the blank Planning Tables for an illustration of how Table 7 data can be separated as described above.

When reporting cancer and noncancer values on separate tables, use the column names to identify instructions for completing each column, as the column number will differ after Column 7.

GENERAL NOTES/INSTRUCTIONS FOR THIS TABLE:

- All table entries, with the exception of Intake, Non-Cancer Hazard and Cancer Risk are presented on tables preceding Table
 7.
- With the exception of modeled intakes, the intake value is the result of calculations performed using parameters and equations presented in Table 4 and concentrations presented in Table 3.
- The Total Non-Cancer Hazard is to be summed for each Exposure Route and Exposure Point in the Exposure Route Total and Exposure Point Total rows. The total Non-Cancer Hazard for all Exposure Pathways for a given Receptor is to be presented as the Total of Receptor Hazards Across All Media at the bottom of the table. This value represents the non-cancer hazard of the various exposure routes/pathways combined.
- The total Cancer Risk is to be summed for each Exposure Route and Exposure Point in the Exposure Route Total and Exposure Point Total rows. The Total Cancer Risk for all Exposure Pathways for a given Receptor is to be presented as the Total of Receptor Risks Across All Media at the end of the table. This value represents the cancer risk of the various Exposure Routes/Pathways combined to a given receptor.

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CALCULATION OF CHEMICAL CANCER RISKS AND NON-CANCER HAZARDS (continued)

| HOW TO COMPLETE/INTERPRET THE TABLE SUMMARY BOX IN UPPER LEFT CORNER | |
|--|---|
| | |
| Definition: • The time period (current and/or future) being considered for the Exposure Pathway. | |
| Instructions:Choose from the picklist to the right. | Current Future Current/Future Not Documented |
| Row 2 - Receptor Population | |
| Definition: • The exposed individual relative to the Exposure Pathway considered. | For example, a resident (Receptor Population) who drinks contaminated groundwater. |
| Instructions: • Choose from the picklist to the right. | Resident Industrial Worker Commercial Worker Construction Worker Other Worker Golfer Jogger Fisher Hunter Fisher/Hunter Swimmer Other Recreational Person Child at School/Daycare/ Playground Trespasser/Visitor Farmer Gardener Gatherer Other |
| Row 3 - Receptor Age | |
| Definition:The description of the exposed individual, as defined by the EPA Region or dictated by the site. | For example, an adult (Receptor Age) resident (Receptor Population) who drinks contaminated groundwater. |

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CALCULATION OF CHEMICAL CANCER RISKS AND NON-CANCER HAZARDS (continued)

| Instructions: • Choose from the picklist to the right. | Child Adult Adolescents (teens) Pre-Adolescents Not Documented Child/Adult Geriatric Sensitive Other Infant Toddler Pregnant | |
|---|--|--|
| BODY OF THE TABLE | | |
| Column 1 - Medium | | |
| Definition: The substance (e.g., air, water, soil) that is a potential source of contaminants in the Exposure Medium. (The Medium will sometimes equal the Exposure Medium.) Usually, the Medium is that targeted for possible remediation. | | |
| Instructions: • Choose from the picklist to the right. | Groundwater Leachate Sediment Sludge Soil Surface Water Debris Liquid Waste Solid Waste Air Surface Soil Subsurface Soil Other | |
| Column 2 - Exposure Medium | | |
| Definition: • The contaminated environmental medium to which an individual may be exposed. Includes the transfer of contaminants from one medium to another. For example: 1) Contaminants in Groundwater (the Medium) remain in Groundwater (the Exposure Medium) and are available for exposure to receptors. 2) Contaminants in Groundwater (the Medium) may be transferred to Air (the | | |
| Exposure Medium) and are available for exposure to receptors. 3) Contaminants in Sediment (the Medium) may be transferred to Fish Tissue (the Exposure Medium) and are available for exposure to receptors. | | |

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CALCULATION OF CHEMICAL CANCER RISKS AND NON-CANCER HAZARDS (continued)

| Instruction • Choos | s: e from the picklist to the right. | Groundwater Leachate Sediment Sludge Soil Surface Water Debris Liquid Waste Solid Waste Air Plant Tissue Animal Tissue Fish Tissue Spring Water Surface Soil Subsurface Soil Particulates Vapors Other |
|--------------------------------|---|--|
| Column 3 - Expos | ure Point | |
| chemic For exam 1) 2) | act location of potential contact between a person and a cal or radionuclide within an Exposure Medium. nple: Contaminants are in Groundwater (the Medium and the Exposure Medium) and exposure to Aquifer 1 - Tap Water (the Exposure Point) is evaluated. Contaminants in Groundwater (the Medium) may be transferred to Air (the Exposure Medium) and exposure to Aquifer 1 - Water Vapors at Showerhead (the Exposure Point) is evaluated. Contaminants in Sediment (the Medium) may be transferred to Fish Tissue (the Exposure Medium) and Trout from Dean's Creek (the Exposure Point) is evaluated. | |
| Instruction • Provid | s: le the information as text in the Table. | Exposure Point should be defined in the same way as was done in Planning Table 1. |
| Column 4 - Expo | sure Route | |
| | ray a chemical or radionuclide comes in contact with a n (e.g., by ingestion, inhalation, dermal contact). | |

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CALCULATION OF CHEMICAL CANCER RISKS AND NON-CANCER HAZARDS (continued)

Instructions:

• Enter the Exposure Route considered from the picklist to the right.

Inhalation

Ingestion

Combined (i.e., Inhalation and

Ingestion)

Dermal

Not Documented

External (Radiation)

CALCULATION OF CHEMICAL CANCER RISKS AND NON-CANCER HAZARDS (continued)

| Definition: | |
|---|--|
| • Chemicals that are potentially site-related, with data of sufficient quality, that have been retained for quantitative analysis as a result of the screening documented in Table 2. | |
| Instructions:Enter the COPCs selected from the COPC screening. | Table 2 documents COPC screening. |
| Column 6 - EPC Value | |
| The EPC, based on either a statistical derivation of measured data or modeled data, that represents an estimate of the chemical or radionuclide concentration. The EPC value may be statistically derived by calculating the 95% UCL of measured groundwater contaminant concentrations from multiple residential wells. Alternatively, the EPC value may be selected as a single measured value, if one data point is used to calculate the risk for each residential well individually. In some cases, the EPC value may be a modeled value (e.g., if upgradient groundwater contaminant concentrations are used to model groundwater concentration at a downgradient exposure point, or if sediment concentrations are used to model fish tissue concentrations). | The EPC Value may be calculated, measured, or modeled. |
| Instructions: Enter the EPC value for each COPC. This value should be in Table 3. If an EPC other than the one found in Table 3 is used, indicate it with a footnote and include a reference to supporting information that will show how the data were modeled in the risk assessment. | Table 3 documents EPC calculations for RME and CT. |
| Column 7 - EPC Units | |
| Definition: • The units associated with the EPC value. | |
| Instructions:Enter the units for EPC values. | Consult the EPA risk assessor for unit preferences. |

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CALCULATION OF CHEMICAL CANCER RISKS AND NON-CANCER HAZARDS (continued)

| Definition: Intake is a measure of exposure expressed as the mass of a substance in contact with the exchange boundary per unit body weight per unit time (e.g. mg chemical/kg body weight/day). | Refers to the intake/exposure concentration results using the parameters and equations, calculations and/or models presented in Table 4. |
|---|--|
| Instructions: Enter the result of the intake calculations/modeling or the exposure concentration performed for each COPC and Exposure Route. | The intake equations, calculations, and/or models are documented in Table 4. |
| Column 9 - Cancer Risk Calculations - Intake/Exposure Concentration Uni 7a) | ts (Also Column 9 on Table |
| Definition: • The units for intake or exposure concentration for each COPC and Exposure Route. | |
| Instructions: Enter the units from the intake calculation or exposure concentration for each COPC which corresponds to each Exposure Route. | |
| Column 10 - Cancer Risk Calculations - CSF/Unit Risk Value (Also Column 10 on Table 7a) | |
| Definition: The slope factor is used to estimate an upper-bound probability of an individual developing cancer as a result of a lifetime of exposure to a particular level of potential carcinogen. Unit Risk is a toxicity value for carcinogenic effects expressed in terms of risk per unit concentration of the substance in the medium where human contact occurs. These measures can be calculated from cancer slope factors. | |
| Instructions:Enter the cancer slope factor or unit risk for each COPC which corresponds to each exposure route. | The slope factors and unit risk values for each COPC are presented in Tables 6.1, 6.2, and 6.3. |
| Column 11 - Cancer Risk Calculations - CSF/Unit Risk Units (Also Column 1 | 1 on Table 7a) |
| Definition: • The units for the cancer slope factor or unit risk. | |
| Instructions: • Enter the cancer slope factor or unit risk units for each COPC for each Exposure Route. | |
| Column 12 - Cancer Risk Calculations - Cancer Risk (Also Column 12 on Tabl | e 7a) |

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CALCULATION OF CHEMICAL CANCER RISKS AND NON-CANCER HAZARDS (continued)

| Definition: • The result of the cancer risk calculation for each COPC for each Exposure Route and Exposure Pathway. | |
|---|---|
| Instructions: Enter the cancer risk calculation for each COPC. Sum the cancer risk results for each Exposure Route in the Exposure Route Total row. Sum the cancer risk calculation results for each Exposure Point in the Exposure Route Total row. Sum the total cancer risk results for all Exposure Pathways in the Total of Receptor Risks Across all Media row. | The sum of all Exposure Routes represents the total cancer risk for all Exposure Routes/ Pathways. |
| Column 13 - Non-Cancer Hazard Calculations - Intake/Exposure Concentration Table 7b) | on Value (Also Column 8 |
| Definition: Intake is a measure of exposure expressed as the mass of a substance in contact with the exchange boundary per unit body weight per unit time. | Refers to the intake/exposure concentration results using the parameters and equations/calculations and/or models presented in Table 4. |
| Instructions: Enter the result of the intake calculations/modeling performed for each COPC and Exposure Route. | The intake equations, calculations, and/or models are documented in Table 4. |
| Column 14 - Non-Cancer Hazard Calculations - Intake/Exposure Concentrat on Table 7b) | ion Units (Also Column 9 |
| Definition: • The units for intake for each COPC and Exposure Route. | |
| Instructions:Enter the units from the intake calculation for each COPC which corresponds to each Exposure Route. | |
| Column 15 - Non-Cancer Hazard Calculations - RfD/RfC Value (Also Column | 10 on Table 7b) |
| Definition: RfD is the toxicity value for evaluating non-cancer effects resulting from exposures. RfC is the toxicity value for inhalation. | |

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CALCULATION OF CHEMICAL CANCER RISKS AND NON-CANCER HAZARDS (continued)

Instructions:

- Enter the RfD or RfC value.
- For RfD, enter the reference dose for each COPC which corresponds to each exposure route.
- Enter Oral RfD values for ingestion.
- Enter Adjusted Dermal RfD values for dermal.
- Enter Adjusted Inhalation RfD/RfC values for inhalation.

The reference doses (RfD/RfC) for each COPC are presented in Table

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CALCULATION OF CHEMICAL CANCER RISKS AND NON-CANCER HAZARDS (continued)

| Column 16 - Non-Cancer Hazard Calculations - RfD/RfC Units (Also Column 11 on Table 7b) | | |
|--|---|--|
| Definition: • The units associated with the reference dose or reference concentration. | RfDs are typically reported in mg/kg-day, a dose term, RfCs in mg/m ³ . | |
| Instructions: Enter the units for reference dose or reference concentration for each COPC for each exposure route. RfC is typically reported as a concentration in air (mg/m³) which can be converted to an inhaled dose (mg/kg-day). | | |
| Column 17 - Non-Cancer Hazard Calculations - Hazard Quotient (Also Column 12 on Table 7b) | | |
| Definition: The ratio of a single substance exposure level, over a specified time period, to a reference dose for that substance, derived from a similar exposure period. | | |
| Instructions: Enter the result of the hazard quotient calculation for each COPC. Sum the hazard quotient for each Exposure Route in the Exposure Route Total row. Sum the hazard quotient for each Exposure Point in the Exposure Route Total row. Sum the hazard quotients for all Exposure Pathways in the Total of Receptor Hazards across all Media row. | The Hazard Index represents the total non-cancer hazard for all exposure routes/pathways presented in this table. | |

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CALCULATION OF RADIATION CANCER RISKS

PURPOSE OF THE TABLE:

- To provide a summary of the variables and approaches used to calculate radiation cancer risks
- To show the EPC used in the radiation cancer risk calculations
- To document the radiation risk calculation approach used to calculate radiation cancer risks
- To show, based on the documented risk calculation approach, the intake and cancer slope factors
- To present the result of the calculation for each Exposure Route/Pathway for each COPC
- To provide the total radiation cancer risks for each Exposure Route/Pathway for the Scenario Timeframe, and Receptor presented in this table
- To provide the total radiation cancer risks for each Exposure Point for the Scenario Timeframe and Receptor in this table
- To provide the total radiation cancer risks across all media for the Scenario Timeframe and Receptor in this table

Radiation can be evaluated two ways: 1) Calculate cancer risks.

The evaluation method used needs to be documented in the Planning Tables 2) Compare radiation doses to standards (i.e., EPA NESHAPS or MCLs or DOE/NRC cleanup standards).

Table 8 is used to show the variables and results when using the first method. The Dose Assessment Worksheet can be used to calculate doses which can be compared to radiological dose standards.

INFORMATION DOCUMENTED:

- The approach for calculating the radiation cancer risk for each COPC for each Exposure Route/Pathway
- The values used for EPC, intake and cancer slope factor for each COPC for each Exposure Route
- The cancer risk value for each COPC for each Exposure Route/Pathway
- Total cancer risk values by Exposure Route, Exposure Point, and across all media for the Scenario Timeframe and Receptor presented in this table

TABLE NUMBERING AND SUMMARY BOX INSTRUCTIONS:

- Complete one copy of Table 8 for each unique combination of the following three fields that will be quantitatively evaluated (Scenario Timeframe, Receptor Population, and Receptor Age).
- Enter each combination of these three fields in the Summary Box in the upper left corner of the table.
- Number each table uniquely, beginning with 8.1 and ending with 8.n where "n" represents the total number of combinations of the three key fields.
- Table 8.1.RME through 8.n.RME should be completed for RME cancer risk calculations.

It is possible that some tables may contain the same data associated with different descriptions in the Summary Box in the upper left corner.

Separate tables may be necessary to ensure transparency in data presentation. Replication of information is readily accomplished using spreadsheet software.

Consult the EPA risk assessor for alternatives (e.g., footnotes) to preparing multiple tables with the same data

B8-1 December 2001

CALCULATION OF RADIATION CANCER RISKS (continued)

| GENERAL NOTES/INSTRUCTIONS FOR THIS TABLE: All table entries, with the exception of risk calculation approach, intake, and cancer risk are presented on tables preceding Table 8. With the exception of modeled intakes, the intake value is the result of calculations performed using parameters and equations presented in Table 4 and concentrations presented in Table 3. The total cancer risk for each Exposure Route is to be summed and indicated in the Exposure Route Total row. This value represents the cancer risk of the various Exposure Routes across each Exposure Pathway designated in the table. The total cancer risk for Each Exposure Point is to be summed and presented in the row labeled Exposure Point Total. The total cancer risk for all media is to be summed and presented in the box labeled "Total of Receptor Risks Across All Media". This value represents the total radiation cancer risk to the receptor for the timeframe designated in the table. | ELE | |
|--|--|--|
| SUMMARY BOX IN UPPER LEFT CORNER | | |
| Row 1 - Scenario Timeframe | | |
| Definition: • The time period (current and/or future) being considered for the exposure pathway. | | |
| Instructions:Choose from the picklist to the right. | Current Future Current/Future Not Documented | |
| Row 2 - Receptor Population | | |
| Definition: • The exposed individual relative to the Exposure Pathway considered. | For example, a resident (receptor population) who drinks contaminated groundwater. | |

B8-2 December 2001

CALCULATION OF RADIATION CANCER RISKS (continued)

| Instructions: • Choose from the picklist to the right. | Resident Industrial Worker Commercial Worker Construction Worker Other Worker Golfer Jogger Fisher Hunter Fisher/Hunter Swimmer Other Recreational Person Child at School/Daycare/ Playground Trespasser/Visitor Farmer Gardener Gatherer Other |
|--|---|
| Row 3 - Receptor Age Definition: The description of the exposed individual as defined by the EDA | For example, an adult (Receptor Age) resident (Receptor Population) |
| The description of the exposed individual, as defined by the EPA Region or dictated by the site. | who drinks contaminated groundwater. |
| Instructions: • Choose from the picklist to the right. | Child Adult Adolescents (teens) Pre-Adolescents Not Documented Child/Adult Geriatric Sensitive Infant Toddler Pregnant Other |
| BODY OF THE TABLE | |
| Column 1 - Medium | |
| Definition: The substance (e.g., air, water, soil) that is a potential source of contaminants in the Exposure Medium. (The Medium will sometimes equal the Exposure Medium.) Usually, the Medium is that targeted for possible remediation. | |

B8-3 December 2001

CALCULATION OF RADIATION CANCER RISKS (continued)

| Instructio | ons: sose from the picklist to the right. | Groundwater Leachate Sediment Sludge Soil Surface Water Debris Liquid Waste Solid Waste Air Surface Soil Subsurface Soil Other |
|-------------------|---|--|
| Column 2 - Expo | osure Medium | |
| may Med | contaminated environmental medium to which an individual be exposed. Includes the transfer of contaminants from one dium to another. **Example:* **Contaminants in Groundwater (the Medium) remain in Groundwater (the Exposure Medium) and are available for exposure to receptors. **Contaminants in Groundwater (the Medium) may be transferred to Air (the Exposure Medium) and are available for exposure to receptors. **Contaminants in Sediment (the Medium) may be transferred to Fish Tissue (the Exposure Medium) and are available for exposure to receptors. | |
| Instruction • Cho | ons: cose from the picklist to the right. | Groundwater Leachate Sediment Sludge Soil Surface Water Debris Liquid Waste Solid Waste Air Plant Tissue Animal Tissue Fish Tissue Spring Water Surface Soil Subsurface Soil Particulates Vapors Other |

B8-4 December 2001

CALCULATION OF RADIATION CANCER RISKS (continued)

| יטכו | finition: | |
|----------|---|---|
| • | An exact location of potential contact between a person and a chemical or radionuclide within an Exposure Medium. | |
| | For example: | |
| | 1) Contaminants are in Groundwater (the Medium and the Exposure Medium) and exposure to Aquifer 1 - Tap Water (the Exposure Point) is evaluated. | |
| | Contaminants in Groundwater (the Medium) may be transferred to Air (the Exposure Medium) and exposure to Aquifer 1 - Water Vapors at Showerhead (the Exposure Point) is evaluated. | |
| | 3) Contaminants in Sediment (the Medium) may be transferred to Fish Tissue (the Exposure Medium) and Trout from Dean's Creek (the Exposure Point) is evaluated. | |
| Ins • | tructions: Provide the information as text in the Table. | Exposure Point should be defined in the same way as was done in Planning Table 1. |
| olumn 4 | - Exposure Route | |
| Det • | The way a chemical or radionuclide comes in contact with a person (e.g., by ingestion, inhalation, dermal contact). | |
| Ins | tructions: Enter the Exposure Route considered from the picklist to the right. | Inhalation Ingestion Combined (i.e., Inhalation and Ingestion) Dermal Not Documented External (Radiation) |
| | | |
| olumn 5 | - Radionuclide of Potential Concern | , |
| | - Radionuclide of Potential Concern finition: Radionuclides that are potentially site-related, with data of sufficient quality, that have been retained for quantitative analysis as a result of the screening documented in Table 2. | |

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CALCULATION OF RADIATION CANCER RISKS (continued)

| Definition: • The EPC, based on either a statistical derivation of measured data or modeled data, that represents an estimate of the chemical or radionuclide concentration available from a particular Medium or route of exposure. | The EPC value may be developed from a statistical derivation of measured data or from modeled data. Typically, the EPC units are expressed as activity per mass such as pCi/gram. |
|--|---|
| Instructions: Enter the EPC value for each COPC. If an EPC other than from Table 3 is used, indicate it with a footnote that includes a reference to supporting information that will show how the data were modeled in the risk assessment. | Table 3 documents EPC calculations. |
| Column 7 - EPC Units | |
| Definition: • The units associated with the EPC value. | |
| Instructions: • Enter the units for the EPC values. | The units may vary depending on the medium. |
| Column 8 - Risk Calculation Approach | |
| Definition: • The approach used for calculating radiation cancer risks. | Consult the EPA risk assessor or National guidance for the appropriate risk calculation approach. US EPA RAGS Part A and RESRAD are examples of risk calculation approaches. |
| Instructions:Enter the radiation risk calculation approach used for each COPC. | |
| Column 9 - Cancer Risk Calculations - Intake/Activity Value | |
| Definition:Intake is a measure of exposure expressed in units of activity such as pCi. | Refers to the intake using the parameters and equations/calculations, and/or models presented in Table 4. |
| Instructions: • Enter the result of the intake calculations/modeling performed. | The intake calculations and/or models are documented in Table 4. |
| Column 10 - Cancer Risk Calculations - Intake/Activity Units | |
| Definition: • The units for intake/activity for each COPC and Exposure Route. | |

B8-6 December 2001

CALCULATION OF RADIATION CANCER RISKS (continued)

| Instructions: | |
|---|--|
| Enter the units for the intake/activity for each COPC which | |
| corresponds to each Exposure Route. | |

B8-7 December 2001

CALCULATION OF RADIATION CANCER RISKS (continued)

| Column 11 - Cancer Risk Calculations - CSF Value | |
|---|---|
| Definitions: A cancer slope factor (CSF) is an age-averaged lifetime excess cancer incidence rate per unit intake (or unit exposure for external exposure pathways). Ingestion and inhalation slope factors are central estimates in a linear model of the age-averaged, lifetime attributable radiation cancer incidence (fatal and nonfatal cancer) risk per unity of activity inhaled or ingested, expressed as risk/picocurie (pCi). External exposure slope factors are central estimates of the lifetime attributable radiation cancer incidence risk for each year of exposure to external radiation from photon-emitting radio nuclides distributed uniformly in a thick layer of soil, and are expressed as risk/yr per pCi/gram of soil. | Slope factors presented in Table 6.4 for each radionuclide are the same as those presented here. |
| Instructions:Enter the CSF for each COPC which corresponds to each Exposure Route. | The cancer slope factors for each COPC are presented in Table 6.4. |
| Column 12 - Cancer Risk Calculations - CSF Units | |
| Definition: • The units associated with the cancer slope factor value. | |
| Instructions:Enter the cancer slope factor units for each COPC for each Exposure Route. | Consult the EPA risk assessor to determine if there is a preference regarding the units to be used. |
| Column 13 - Cancer Risk Calculations - Cancer Risk | |
| Definition: The result of the cancer risk calculation for each COPC for each exposure route and pathway. Cancer risk is the incremental probability of an individual's developing cancer over a lifetime as a result of exposure to a potential carcinogen. | |
| Instructions: Enter the cancer risk calculation for each COPC. Sum the cancer risk results for each Exposure Route in the Exposure Route Total row. Sum the cancer risk results for each Exposure Point in the Exposure Point Total row. Sum the total radiation cancer risk results for all media in the bottom right-hand corner box labeled "Total of Receptor Risks Across All Media". | The sum of all Exposure Routes represents the total cancer risk for an Exposure Pathway. The sum of all Exposure Pathways represent the total cancer risk for a medium. The sum of all media represents the "Total of Receptor Risks Across All Media". |

B8-8 December 2001

SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCs

| PURPOSE OF THE TABLE: • To provide a summary of cancer risks and non-cancer hazards for each Receptor by Medium, Exposure Medium, Exposure Route, and Exposure Point INFORMATION DOCUMENTED: The concertrisk and non-cancer bazard to each Recentor for | Table 9 presents cancer risk and non-cancer hazard information for all COPCs and media/exposure points quantitatively evaluated. |
|--|--|
| The cancer risk and non-cancer hazard to each Receptor for each COPC by Exposure Route and Exposure Point The total cancer risk and non-cancer hazard for each Exposure Point, Exposure Medium, and Medium The total cancer risks and non-cancer hazards for a Receptor across all media The primary target organs for non-carcinogenic hazard effects. | |
| TABLE NUMBERING AND SUMMARY BOX INSTRUCTIONS: Complete one copy of Table 9 for each unique combination of the following three fields that will be quantitatively evaluated (Scenario Timeframe, Receptor Population, and Receptor Age). Enter each combination of these three fields in the Summary Box in the upper left corner of the table. Number each table uniquely beginning with 9.1 and ending with 9.n where "n" represents the total number of combinations of the three key fields. Different tables should be prepared to address RME and CT Risk and Hazard summaries. Tables 9.1. RME through 9.n. RME should be completed for RME Risk and Hazard summaries. Table 9.1.CT through 9.n.CT should be completed for CT Risk and Hazard Summaries. | It is possible that some tables may contain the same data associated with different descriptions in the Summary Box in the upper left corner. Separate tables may be necessary to ensure transparency in data presentation. Replication of information is readily accomplished using spreadsheet software. Consult the EPA risk assessor for alternatives (e.g., footnotes) to preparing multiple tables with the same data. |
| • Cancer risk and non-cancer hazard information for all COPCs and media/Exposure Points quantitatively evaluated is to be presented in Table 9. | |
| All table entries are presented on Tables preceding Table 9. Documentation of the non-cancer hazard and carcinogenic risk values for chemicals was presented on Table 7. Documentation of the carcinogenic risk values for radionuclides was presented on Table 8. Total cancer risks and non-cancer hazards associated with each Receptor are to be presented for each Exposure Point, Exposure Medium, and Medium and across all media and all Exposure Routes. | |

B9-1 December 2001

SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCs (continued)

| HOW TO COMPLETE/INTERPRET THE TABLE SUMMARY BOX IN UPPER LEFT CORNER | |
|--|---|
| | |
| Definition:The time period (current and/or future) being considered for the exposure pathway. | |
| Instructions:Choose from the picklist to the right. | Current Future Current/Future Not Documented |
| Row 2 - Receptor Population | |
| Definition: • The exposed individual relative to the Exposure Pathway considered. | For example, a resident (receptor population) who drinks contaminated groundwater. |
| Instructions: • Choose from the picklist to the right. | Resident Industrial Worker Commercial Worker Construction Worker Other Worker Golfer Jogger Fisher Hunter Fisher/Hunter Swimmer Other Recreational Person Child at School/Daycare/ Playground Trespasser/Visitor Gatherer Farmer Gardener Other |
| Row 3 - Receptor Age | |
| Definition:The description of the exposed individual, as defined by the Region or dictated by the site. | For example, an adult (Receptor Age) resident (Receptor Population) who drinks contaminated groundwater. |

B9-2 December 2001

SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCs (continued)

| Instructions: • Choose from the picklist to the right. | Child Adult Adolescents (teens) Pre-Adolescents Not Documented Child/Adult Geriatric Sensitive Other |
|---|--|
| | Infant Toddler Pregnant |
| BODY OF THE TABLE | |
| Column 1 - Medium | |
| Definition: • The substance (e.g., air, water, soil) that is a potential source of contaminants in the Exposure Medium. (The Medium will sometimes equal the Exposure Medium.) Usually, the Medium is that targeted for possible remediation. | |
| Instructions: Choose from the picklist to the right. For each Medium, The last entry in this column should be "Medium Total." In this row, the total risk/HI from each Medium (for all chemicals, Exposure Routes, Exposure Points, and Exposure Media) for the current Receptor is entered in the Exposure Routes Total Column. | Groundwater Leachate Sediment Sludge Soil Surface Water Debris Other Liquid Waste Solid Waste Air Surface Soil Subsurface Soil |
| Column 2 - Exposure Medium | |
| The contaminated environmental medium to which an individual may be exposed. Includes the transfer of contaminants from one medium to another. | |
| For example: | |
| Contaminants in Groundwater (the Medium) remain in Groundwater (the Exposure Medium) and are available for exposure to receptors. Contaminants in Groundwater (the Medium) may be transferred to Air (the | |
| Exposure Medium) and are available for exposure to receptors. 3) Contaminants in Sediment (the Medium) may be transferred to Fish Tissue (the Exposure Medium) and are available for exposure to receptors. | |

B9-3 December 2001

SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCs (continued)

Instructions:

- Choose from the picklist to the right.
- For each Exposure Medium, the last entry in this column should be "Exposure Medium Total." This refers to the total risk/HI from each Exposure Medium (for all chemicals, Exposure Routes and Exposure Points) for the current Receptor. These totals are recorded in the Carcinogenic and Non-Carcinogenic Exposure Routes Total Columns.

Leachate
Sediment
Sludge
Soil
Surface Water
Debris
Other

Groundwater

Liquid Waste Solid Waste

Air

Vapors

Plant Tissue Animal Tissue Fish Tissue Spring Water Surface Soil Subsurface Soil Particulates

B9-4 December 2001

SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCs (continued)

| Column 3 - Exposure Point | | | | | |
|--|--|--|--|--|--|
| Definition: • An exact location of potential contact between a person and a chemical within an Exposure Medium. For example: 1) Contaminants are in Groundwater (the Medium and the Exposure Medium) and exposure to Aquifer 1 - Tap Water (the Exposure Point) is evaluated. 2) Contaminants in Groundwater (the Medium) may be transferred to Air (the Exposure Medium) and exposure to Aquifer 1 - Water Vapors at Showerhead (the Exposure Point) is evaluated. 3) Contaminants in Sediment (the Medium) may be transferred to Fish Tissue (the Exposure Medium) and Trout from Dean's Creek (the Exposure Point) is evaluated. | | | | | |
| Instructions: Provide the information as text in the Table. For each Exposure Point, the last entry in this column should be "Exposure Point Total." This refers to the total risk/HI (for all chemicals and Exposure Routes) for the current Receptor. These totals are recorded in the Carcinogenic and Non-Carcinogenic Exposure Routes Total columns. | Exposure Point should be defined in the same way as was done in Planning Table 1. | | | | |
| Column 4 - Chemical of Potential Concern | | | | | |
| Definition: • The COPCs quantitatively considered in the risk characterization. | | | | | |
| Instructions: Enter the COPCs from previous tables. Enter the term "Chemical Total" at the end of the list of chemicals for each Exposure Point. Use this row to record total risk/HI values from all chemicals at each Exposure Point. Enter the term "Radionuclide Total" at the end of the list of radionuclides for each Exposure Point. Use this row to record total risk/HI values from all radionuclides for each Exposure Point. | | | | | |
| Columns 5, 6, 7, and 8 - Carcinogenic Risk - Ingestion, Inhalation, Dermal | and External (Radiation) | | | | |
| Definition: • The cancer risk value calculated by Receptor for each COPC for each Exposure Route for each Exposure Point. | The value at the bottom of each column presents the total cancer risk by Exposure Route for each Exposure Point. | | | | |

B9-5 December 2001

SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCs (continued)

| Instructions: Enter the cancer risk value calculated by Receptor for each Exposure Route for each Exposure Point. Enter the cancer risk totals for each Exposure Route in the rows labeled "Chemical Total" and "Radionuclide Total." | | | | | | | | | | | | |
|---|---|--|--|--|--|--|--|--|--|--|--|--|
| Column 9 - Carcinogenic Risk - Exposure Routes Total | | | | | | | | | | | | |
| Definition: • The total cancer risk for each COPC across all Exposure Routes at each Exposure Point. | | | | | | | | | | | | |
| Instructions: Enter the sum of the cancer risks across Exposure Routes for each COPC. Enter the sum of the cancer risks in this column for each Exposure Point in the "Exposure Point Total" row. Enter the total cancer risk for each Exposure Medium and individual Medium in the "Exposure Medium Total" and "Medium Total" rows. For each Receptor, enter the total cancer risks across all Media and all Exposure Routes as "Receptor Risk Total." | Consult the EPA risk assessor to determine the appropriate summing of risks. | | | | | | | | | | | |
| Column 10 - Non-Carcinogenic Hazard Quotient - Primary Target Organ | | | | | | | | | | | | |
| Definition:The primary effect reported as a primary target organ effect in IRIS, HEAST, or other source. | | | | | | | | | | | | |
| Instructions: Enter the primary target organ effect as reported in IRIS, HEAST, or other source. | Consult the EPA risk assessor to determine if multiple effects should be provided. | | | | | | | | | | | |
| Columns 11, 12, and 13 - Non-Carcinogenic Hazard Quotient - Ingestion, Inha | alation, Dermal | | | | | | | | | | | |
| Definition: • The non-cancer hazard calculated by Receptor for each COPC for each Exposure Route for each Exposure Point. | The value at the bottom of each column presents the non-cancer hazard by exposure route for each exposure point, for all effects considered together. | | | | | | | | | | | |
| Instructions: Enter the non-cancer hazard value calculated by Receptor for each COPC for each Exposure Route for each Exposure Point. Enter the non-cancer hazard totals for each Exposure Route in the rows labeled "Chemical Total" and "Radionuclide Total." | Consult the EPA risk assessor for summing hazard quotients. | | | | | | | | | | | |

B9-6 December 2001

SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCs (continued)

| Column 14 - Non-Carcinogenic Hazard Quotient - Exposure Routes To | olumn 14 - Non-Carcinogenic Hazard Quotient - Exposure Routes Total | | | | | | | | | | |
|---|---|--|--|--|--|--|--|--|--|--|--|
| Definition: The total non-cancer hazard calculated for each COPC across all Exposure Routes at each Exposure Point. | The Totals in each column present the total non-cancer hazards by Exposure Routes for each Exposure Point. The values beneath the table under this column present hazard quotients for target organs. | | | | | | | | | | |
| Instructions: Enter the sum of non-cancer hazards across the three Exposure Routes in each Exposure Route column. Enter the sum of the non-cancer hazards across Exposure Routes for each COPC and primary target organ. Enter the sum of the non-cancer hazards in this column for each Exposure Point in the "Exposure Point Total" row. Enter the total hazard index for each Exposure Medium and Medium in the "Exposure Medium Total" and "Medium Total" rows. Enter the total hazard index across all media and all Exposure Routes as "Receptor HI Total." Enter the total hazard index for primary target organs. Sum the hazard quotient target organ effects by target organ and enter into the appropriate boxes. | Consult the EPA risk assessor for specific instructions in summing hazard quotients. | | | | | | | | | | |

B9-7 December 2001

RISK SUMMARY

PURPOSE OF THE TABLE:

- To provide a summary for each Receptor by Medium, Exposure Route, and Exposure Point of cancer risks and non-cancer hazards that trigger the need for remedial action.
- The Risk Assessor may consult the Remedial Project Manager and other members of the project team to determine what levels of risk may be actionable at the site and what should be included in Table 10. The risks shown on Table 10 should be based upon the Remedial Project Manager's recommendation. If all risks are below actionable levels, determine with the Remedial Project Manager which chemicals should be shown to document the suitability of a No Action decision.

Table 10 presents cancer risk and non-cancer hazard information for those COPCs and media/exposure points that the Remedial Project Manager determines trigger the need for remedial action (the risk drivers).

INFORMATION DOCUMENTED:

- The cancer risk and non-cancer hazard to each Receptor for each chemical by Exposure Route and Exposure Point for risk drivers
- The cancer risk and non-cancer hazard for each Exposure Point, Exposure Medium, and Medium across all Exposure Routes for risk drivers
- The total cancer risks and non-cancer hazards for a Receptor across all media for risk drivers
- The primary target organs for non-carcinogenic hazard effects for risk drivers.

For the purpose of these instructions, those COPCs determined to trigger the need for cleanup are simply referred to as "Chemicals."

TABLE NUMBERING AND SUMMARY BOX INSTRUCTIONS:

- Complete one copy of Table 10 for each unique combination of the following three fields that will be quantitatively evaluated (Scenario Timeframe, Receptor Population, and Receptor Age).
- Enter each combination of these three fields in the Summary Box in the upper left corner of the table.
- Number each table uniquely beginning with 10.1 and ending with 10.n where "n" represents the total number of combinations of the three key fields.
- Different tables should be prepared to address RME and CT Risk and Hazard summaries.
- Tables 10.1. RME through 10.n. RME should be completed for RME Risk and Hazard summaries.
- Table 10.1 CT through 10.n.CT should be completed for CT Risk and Hazard Summaries.

It is possible that some tables may contain the same data associated with different descriptions in the Summary Box in the upper left corner.

Separate tables may be necessary to ensure transparency in data presentation. Replication of information is readily accomplished using spreadsheet software.

Consult the EPA risk assessor for alternatives (e.g., footnotes) to preparing multiple tables with the same information.

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RISK SUMMARY (continued)

| GENERAL NOTES/INSTRUCTIONS FOR THIS | S TABLE |
|--|--|
| Cancer risk and non-cancer hazard information for only those COPCs and media/exposure points that trigger the need for remedial action (the risk drivers) is to be presented in Table 10. All table entries are presented on Tables preceding Table 10. Documentation of the non-cancer hazard and cancer risk values for chemicals was presented on Table 7. Documentation of the carcinogenic risk values for radionuclides was presented on Table 8. Total cancer risks and non-cancer hazards associated with each Receptor are to be presented for each Exposure Point, Exposure Medium, Medium across all media and all Exposure Routes. | |
| HOW TO COMPLETE/INTERPRET THE TABI | LE |
| UMMARY BOX IN UPPER LEFT CORNER | |
| ow 1 - Scenario Timeframe | |
| Definition: • The time period (current and/or future) being considered for the Exposure Pathway. | |
| Instructions: • Choose from the picklist to the right. | Current Future Current/Future Not Documented |
| ow 2 - Receptor Population | • |
| Definition: • The exposed individual relative to the Exposure Pathway considered. | For example, a resident (receptor population) who drinks contaminated groundwater. |

B10-2 December 2001

RISK SUMMARY (continued)

| Instructions: • Choose from the picklist to the right. | Resident Industrial Worker Commercial Worker Construction Worker Other Worker Golfer Jogger Fisher Hunter Fisher/Hunter Swimmer Other Recreational Person Child at School/Daycare/Playground Trespasser/Visitor Farmer Gatherer |
|---|---|
| | Gatherer Gardener Other |

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RISK SUMMARY (continued)

| Row 3 - Receptor Age | |
|---|--|
| Definition: The description of the exposed individual, as defined by the Region or dictated by the site. | For example, an adult (Receptor Age) resident (Receptor Population) who drinks contaminated groundwater. |
| Instructions: • Choose from the picklist to the right. | Child Adult Adolescents (teens) Pre-Adolescents Not Documented Child/Adult Geriatric Sensitive Other Infant Toddler Pregnant |
| BODY OF THE TABLE | |
| Column 1 - Medium | |
| Definition: The substance (e.g., air, water, soil) that is a potential source of contaminants in the Exposure Medium. (The Medium will sometimes equal the Exposure Medium.) Usually, the Medium is that targeted for possible remediation. | Enter only the Media that have risks or hazards exceeding target levels. |
| Instructions: Choose from the picklist to the right. For each Medium, the last entry in this column should be "Medium Total." This refers to the total risk/HI for each Medium (for all chemicals, Exposure Routes, Exposure Points, and Exposure Media) for the current Receptor. These totals are recorded in th Carcinogenic and Non-Carcinogenic Exposure Routes Total columns. | Groundwater Leachate Sediment Sludge Soil Surface Water Debris Other Liquid Waste Solid Waste Air Surface Soil Subsurface Soil |

B10-4 December 2001

RISK SUMMARY (continued)

Definition:

• The contaminated environmental medium to which an individual may be exposed. Includes the transfer of contaminants from one medium to another.

For example:

- Contaminants in Groundwater (the Medium) remain in Groundwater (the Exposure Medium) and are available for exposure to receptors.
- Contaminants in Groundwater (the Medium) may be transferred to Air (the Exposure Medium) and are available for exposure to receptors.
- Contaminants in Sediment (the Medium) may be transferred to Fish Tissue (the Exposure Medium) and are available for exposure to receptors.

Enter only the Exposure Media that have risks or hazards exceeding target levels.

Instructions:

- Choose from the picklist to the right.
- For each Exposure Medium, the last entry in this column should be "Exposure Medium Total." This refers to the total risk/HI from each Exposure Medium (for all chemicals, Exposure Routes, and Exposure Points) for the current Receptor. These totals are recorded in the Carcinogenic and Non-Carcinogenic Exposure Routes Total columns.

Ground water

Leachate Sediment

Sludge, Soil

Surface Water

Debris Other

Liquid Waste

Solid Waste

Air

Vapors Plant Tissue

Animal Tissue

Fish Tissue

Surface Soil

Subsurface Soil

Particulates

Spring Water

Column 3 - Exposure Point

Definition:

• An exact location of potential contact between a person and a chemical within an Exposure Medium.

For example:

- Contaminants are in Groundwater (the Medium and the Exposure Medium) and exposure to Aquifer 1 - Tap Water (the Exposure Point) is evaluated
- Contaminants in Groundwater (the Medium) may be transferred to Air (the Exposure Medium) and exposure to Aquifer 1 - Water Vapors at Showerhead (the Exposure Point) is evaluated.
- Contaminants in Sediment (the Medium) may be transferred to Fish Tissue (the Exposure Medium) and Trout in Dean's Creek (the Exposure Point) is evaluated.

Enter only the Exposure Points that have risks or hazards exceeding target levels.

B10-5 December 2001

RISK SUMMARY (continued)

| Instructions: Provide the information as text in the Table. For each Exposure Point, the last entry in this column should be "Exposure Point Total." This refers to the total risk/HI from each Exposure Point (for all chemicals, Exposure Routes, and Exposure Points) for the current Receptor. These totals are recorded in the Carcinogenic and Non-Carcinogenic Exposure Routes Total Columns. | Exposure Point should be defined in the same way as was done in the Planning Table 1. |
|--|--|
| Column 4 - Chemical | • |
| Definition: • The COPCs quantitatively considered in the risk characterization. | Enter only the chemicals that have risks exceeding target levels. |
| Instructions: Enter the COPCs from previous tables that exceed target levels. Enter the term "Chemical Total" at the end of the list of chemicals for each Exposure Point. Enter the term "Radionuclide Total" at the end of the list of radionuclides. | |
| Columns 5, 6, 7 and 8 - Carcinogenic Risk - Ingestion, Inhalation, Dermal, | and External (Radiation) |
| Definition: The cancer risk value calculated by Receptor for each chemical for each Exposure Route for each Exposure Point. | Enter only the risks that exceed target levels. The value at the bottom of each column presents the cancer risk from all chemicals by Exposure Route for each Exposure Point. |
| Instructions: Enter the cancer risk value calculated by Receptor for each chemical for each Exposure Route for each Exposure Point that exceeds target levels. Enter the cancer risk totals for each Exposure Route in the last row. | |
| Column 9 - Carcinogenic Risk - Exposure Routes Total | |
| Definition: • The total cancer risk for each chemical across all Exposure Routes at each Exposure Point. | |

B10-6 December 2001

RISK SUMMARY (continued)

| Instructions: Enter the sum of the cancer risks across Exposure Routes for each chemical. Enter the sum of the cancer risks in this column for each Exposure Point in the "Exposure Point Total" row. Enter the total cancer risk for each Exposure Medium and Medium in the "Exposure Medium Total" and "Medium Total" rows. Enter the total cancer risk across all Media and all Exposure Routes as "Receptor Risk Total". | |
|--|--|
| Column 10 - Non-Carcinogenic Hazard Quotient - Primary Target Orga | n |
| Definition: • The primary effect reported as a primary target organ effect in IRIS, HEAST, or other source. | |
| Instructions: Enter the primary target organ effect as reported in IRIS, HEAST, or other source. This target organ should also appear in Table 5. | Consult the EPA risk assessor to determine if multiple effects should be provided. |

B10-7 December 2001

RISK SUMMARY (continued)

| Columns 11, 12, and 13 - Non-Carcinogenic Hazard Quotient - Ingestion, Inh | alation, Dermal |
|---|--|
| Definition: The non-cancer hazard calculated by Receptor for each Chemical for each Exposure Route for each Exposure Point. | Enter only the hazards that exceed target levels. The value at the bottom of each column presents the non-cancer hazard by Exposure Route for each Exposure Point, for all effects considered together. |
| Instructions: Enter the non-cancer hazard value calculated by Receptor for each chemical for each Exposure Route for each Exposure Point that exceeds target levels. Enter the non-cancer hazard totals for each Exposure Route in the last row, corresponding to the term "Chemical Total" in Column 9. | Consult the EPA risk assessor for summing hazard quotients. |
| Column 14 - Non-Carcinogenic Hazard Quotient - Exposure Routes Total | |
| Definition: • The total non-cancer hazard calculated for each chemical across all Exposure Routes at each Exposure Point. | The totals in each column present the total non-cancer hazards across all Exposure Routes for each Exposure Point. |
| | The values at the bottom of this column present hazard quotients for target organs. |
| Instructions: Enter the sum of non-cancer hazards across the three Exposure Routes in Columns 11, 12, and 13. Enter the sum of the non-cancer hazards across Exposure Routes for each chemical and primary target organ. Enter the sum of the non-cancer hazards in this column for each Exposure Point, Exposure Medium, and Medium in the "Exposure Point Total," "Exposure Medium Total," and "Medium Total" rows, respectively. Enter the total hazard index across all Media and all Exposure Routes as "Receptor HI Total." Enter the total hazard index for primary target organs. Sum the hazard quotient target organ effects across all media by target organ and enter into the appropriate boxes below the table. | Consult the EPA risk assessor for specific instructions in summing hazard quotients. |

B10-8 December 2001

TABLE 0

SITE RISK ASSESSMENT IDENTIFICATION INFORMATION

The Dean Company

Site Name/OU: The Dean Company

Region: III

EPA ID Number: PAD123456789

State: PA

Status: Fund Lead Remedial Investigation

Federal Facility (Y/N): N

EPA Project Manager: John Smith

EPA Risk Assessor: Jane Doe

Prepared by (Organization): Eris Consulting Engineers

Prepared for (Organization): EPA

Document Title: Human Health Risk Assessment for the Dean Company Site

Document Date: August 8, 2001

Probabilistic Risk Assessment (Y/N): N

Comments: This site is contaminated with volatile organic compounds, pesticides, and metals. Lead evaluation was conducted.

TABLE 1 SELECTION OF EXPOSURE PATHWAYS Site Name

| Scenario Timeframe | Medium | Exposure Medium | Exposure Point | Receptor Population | Receptor Age | Exposure Route | Type of Analysis | Rationale for Selection or Exclusion of Exposure Pathway |
|-----------------------|--------|--------------------|-------------------|------------------------|-----------------|-------------------|---------------------|---|
| | | | | | | | | |
| | | | | | | | | |
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| | | | | | | | | |

TABLE 2.1

OCCURRENCE, DISTRIBUTION, AND SELECTION OF CHEMICALS OF POTENTIAL CONCERN

The Dean Company

Scenario Timeframe: Future

Medium: Groundwater

Exposure Medium: Groundwater

| Exposure Point | CAS Number | Chemical | Minimum Concentration (Qualifier) | Maximum Concentration (Qualifier) | Units | Location of Maximum Concentration | Detection Frequency | Range of Detection Limits | Concentration Used for Screening (1) | Background Value (2) | Screening Toxicity Value (3) (N/C) | Potential ARAR/TBC Value | Potential ARAR/TBC Source | COPC Flag (Y/N) | Rationale for Selection or Deletion (4) |
|-----------------------|---------------|----------------------------|-----------------------------------|-----------------------------------|-------|---|------------------------|---------------------------------|--------------------------------------|-------------------------|--|--------------------------------|---------------------------------|-----------------------|---|
| Aquifer 1 - Tap Water | 117817 | Bis(2-ethylhexyl)phthalate | 2 J | 5 J | ug/l | GW3D | 4/12 | 3-4 | 5 | NA | 4.8 C | 6 | MCL | Υ | ASL |
| | 67663 | Chloroform | 0.6 J | 9 | ug/l | GW3D | 3/12 | 1 - 1 | 9 | NA | 0.063 C | 100 | MCL | Υ | ASL |
| | 75150 | Carbon Disulfide | 0.3 J | 4.5 | ug/l | GW3D | 3/12 | 1 - 1 | 4.5 | NA | 100 N | NA | NA | N | BSL |
| | 76448 | Heptachlor | 2 J | 33 J | ug/l | GW4D | 6/12 | 0.01 - 0.01 | 33 | NA | 0.015 C | 0.4 | MCL | Y | ASL |
| | 108883 | Toluene | 0.1 J | 0.2 J | ug/l | GW3D | 3/12 | 1 - 1 | 0.2 | NA | 75 N | 1000 | MCL | N | BSL |
| | 7429905 | Aluminum | 134 J | 1340 | ug/l | GW3D | 2/12 | 29 - 38.2 | 1340 | NA | 3700 N | 50 - 200 | SMCL | N | BSL |
| | 7440393 | Barium | 65 J | 489 | ug/l | GW1D | 6/12 | 0.2 - 1 | 489 | NA | 260 N | 2000 | MCL | Y | ASL |
| | 7440417 | Beryllium | 0.2 K | 1.5 K | ug/l | GW2D | 3/12 | 0.1 - 1 | 1.5 | NA | 7.3 N | 4 | MCL | N | BSL |
| | 7439921 | Lead | 6 J | 35 J | ug/l | GW3D | 4/12 | 0.1 - 1 | 35 | NA | 15 | 15 | MCL | Y | ASL |
| | 7439965 | Manganese | 1900 | 12500 | ug/l | GW1D | 6/12 | 0.3 - 1 | 12500 | NA | 73 N | 50 | SMCL | Y | ASL |
| | 7440020 | Nickel | 0.9 J | 1.5 J | ug/l | GW4D | 3/12 | 0.9 - 7 | 1.5 | NA | 73 N | NA | NA | N | BSL |

(1) Maximum concentration used for screening.

(2) To date, no background study has been completed.

(3) All compounds were screened against the Risk-Based Concentration (RBC) Table, U.S. EPA Region III, May 8, 2001 for tap water (cancer benchmark = 1E-06; HQ = 0.1). Lead was screened against the action level of 15 ug/l.

(4) Rationale Codes:

Selection Reason: Above Screening Level (ASL)
Deletion Reason: Below Screening Level (BSL)

Definitions: NA = Not Applicable

MCL = Maximum Contaminant Level

SMCL = Secondary Maximum Contaminant Level

J = Estimated Value

K = Estimated Value - Biased High

C = Carcinogen

N = Noncarcinogen

TABLE 2.2

OCCURRENCE, DISTRIBUTION, AND SELECTION OF CHEMICALS OF POTENTIAL CONCERN

The Dean Company

Scenario Timeframe: Future

Medium: Groundwater

Exposure Medium: Air

| Exposure Point | CAS Number | Chemical | Minimum Concentration (Qualifier) | Maximum Concentration (Qualifier) | Units | Location of Maximum Concentration | Detection Frequency | Range of Detection Limits | Concentration Used for Screening (1) | Background Value (2) | Screening Toxicity Value (3) (N/C) | Potential ARAR/TBC Value | Potential ARAR/TBC Source | COPC Flag (Y/N) | Rationale for Selection or Deletion (4) |
|---------------------------------|---------------|----------------------------|-----------------------------------|-----------------------------------|-------|---|------------------------|---------------------------|--------------------------------------|-------------------------|------------------------------------|--------------------------------|---------------------------------|-----------------------|---|
| Water Vapors from Showerhead | 117817 | Bis(2-ethylhexyl)phthalate | 2 J | 5 J | ug/l | GW3D | 4/12 | 3 - 4 | 5 | NA | 4.8 C | 6 | MCL | Υ | ASL |
| | 67663 | Chloroform | 0.6 J | 9 | ug/l | GW3D | 3/12 | 1 - 1 | 9 | NA | 0.063 C | 100 | MCL | Υ | ASL |
| | 75150 | Carbon Disulfide | 0.3 J | 4.5 | ug/l | GW3D | 3/12 | 1 - 1 | 4.5 | NA | 100 N | NA | NA | N | BSL |
| | 76448 | Heptachlor | 2 J | 33 J | ug/l | GW4D | 6/12 | 0.01 - 0.01 | 33 | NA | 0.015 C | 0.4 | MCL | Υ | ASL |
| | 108883 | Toluene | 0.1 J | 0.2 J | ug/l | GW3D | 3/12 | 1 - 1 | 0.2 | NA | 75 N | 1000 | MCL | N | BSL |
| | 7429905 | Aluminum | 134 J | 1340 | ug/l | GW3D | 2/12 | 29 - 38.2 | 1340 | NA | 3700 N | 50 - 200 | SMCL | N | BSL |
| | 7440393 | Barium | 65 J | 489 | ug/l | GW1D | 6/12 | 0.2 - 1 | 489 | NA | 260 N | 2000 | MCL | Υ | ASL |
| | 7440417 | Beryllium | 0.2 K | 1.5 K | ug/l | GW2D | 3/12 | 0.1 - 1 | 1.5 | NA | 7.3 N | 4 | MCL | N | BSL |
| | 7439921 | Lead | 6 J | 35 J | ug/l | GW3D | 4/12 | 0.1 - 1 | 35 | NA | 15 | 15 | MCL | Υ | ASL |
| | 7439965 | Manganese | 1900 | 12500 | ug/l | GW1D | 6/12 | 0.3 - 1 | 12500 | NA | 73 N | 50 | SMCL | Υ | ASL |
| | 7440020 | Nickel | 0.9 J | 1.5 J | ug/l | GW4D | 3/12 | 0.9 - 7 | 1.5 | NA | 73 N | NA | NA | N | BSL |

(1) Maximum concentration used for screening.

(2) To date, no background study has been completed.

(3) All compounds were screened against the Risk-Based Concentration (RBC) Table, U.S. EPA Region III, May 8, 2001 for tap water (cancer benchmark = 1E-06; HQ = 0.1). Lead was screened against the action level of 15 ug/l.

(4) Rationale Codes:

Selection Reason: Above Screening Level (ASL)
Deletion Reason: Below Screening Level (BSL)

Definitions: NA = Not Applicable

MCL = Maximum Contaminant Level

SMCL = Secondary Maximum Contaminant Level

J = Estimated Value

K = Estimated Value - Biased High

C = Carcinogen

N = Noncarcinogen

TABLE 2.3

OCCURRENCE, DISTRIBUTION, AND SELECTION OF CHEMICALS OF POTENTIAL CONCERN

The Dean Company

Scenario Timeframe: Future

Medium: Soil

Exposure Medium: Soil

| Exposure Point | CAS Number | Chemical | Minimum Concentration (Qualifier) | Maximum Concentration (Qualifier) | Units | Location of Maximum Concentration | Detection Frequency | Range of Detection Limits | Concentration Used for Screening (1) | Background Value (2) | Screening Toxicity Value (3) (N/C) | Potential ARAR/TBC Value | Potential ARAR/TBC Source | COPC Flag (Y/N) | Rationale for Selection or Deletion (4) |
|-------------------|---------------|--------------------|---|---|-------|---|------------------------|---------------------------------|--------------------------------------|-------------------------|--|--------------------------------|---------------------------------|-----------------------|---|
| Soil at Site 1 | 11096825 | Aroclor-1260 | 15 J | 110 J | ug/kg | SS03 | 6 / 29 | 33 - 300 | 110 | NA | 320 C | NA | NA | N | BSL |
| | 56553 | Benzo(a)anthracene | 120 J | 230 J | ug/kg | SS03 | 16 / 29 | 330 - 700 | 230 | NA | 870 C | NA | NA | N | BSL |
| | 50328 | Benzo(a)pyrene | 48 J | 70 J | ug/kg | SS03 | 17 / 29 | 30 - 70 | 70 | NA | 87 C | NA | NA | N | BSL |
| | 75150 | Carbon Disulfide | 2 J | 33 | ug/kg | SB07 | 4 / 29 | 10 - 16 | 33 | NA | 780000 N | NA | NA | N | BSL |
| | 72548 | 4,4'-DDD | 1 J | 4200 | ug/kg | SS09 | 22 / 29 | 3.3 - 1900 | 4200 | NA | 2700 C | NA | NA | Υ | ASL |
| | 72559 | 4,4'-DDE | 0.44 J | 7200 J | ug/kg | SS09 | 28 / 29 | 2.2 - 700 | 7200 | NA | 1900 C | NA | NA | Υ | ASL |
| | 50293 | 4,4'-DDT | 0.69 J | 290000 J | ug/kg | SB08 | 29 / 29 | 3.3 - 700 | 290000 | NA | 1900 C | NA | NA | Υ | ASL |
| | 108883 | Toluene | 1 J | 2 J | ug/kg | SS08 | 2 / 29 | 10 - 16 | 2 | NA | 1600000 N | NA | NA | N | BSL |
| | 7429905 | Aluminum | 1960 | 21700 | mg/kg | SB07 | 29 / 29 | 6.3 - 11 | 21700 | NA | 7800 N | NA | NA | Υ | ASL |
| | 7440417 | Beryllium | 0.1 J | 13.4 | mg/kg | SS06 | 23 / 29 | 0.02 - 0.21 | 13.4 | NA | 16 N | NA | NA | N | BSL |
| | 7439921 | Lead | 56 J | 750 J | mg/kg | SS03 | 16 / 29 | 10 - 16 | 750 | NA | 400 | NA | NA | Υ | ASL |
| | 7439965 | Manganese | 5.9 | 688 | mg/kg | SS03 | 29 / 29 | 0.05 - 0.5 | 688 | NA | 160 N | NA | NA | Υ | ASL |
| | 7782492 | Selenium | 0.53 J | 1 | mg/kg | SS02 | 9 / 29 | 0.43 - 0.75 | 1 | NA | 39 N | NA | NA | N | BSL |
| Soil at Site 2 | 67641 | Acetone | 9 J | 170 | ug/kg | SB01 | 16 / 40 | 10 - 22 | 170 | NA | 780000 N | NA | NA | N | BSL |
| | 56553 | Benzo(a)anthracene | 48 J | 100 J | ug/kg | SS26 | 31 / 40 | 340 - 700 | 100 | NA | 870 C | NA | NA | N | BSL |
| | 50328 | Benzo(a)pyrene | 47 J | 60 J | ug/kg | SS26 | 29 / 40 | 34 - 70 | 60 | NA | 87 C | NA | NA | N | BSL |
| | 75150 | Carbon Disulfide | 2 J | 17 J | ug/kg | SB07 | 13 / 40 | 10 - 22 | 17 | NA | 780000 N | NA | NA | N | BSL |
| | 72559 | 4,4'-DDE | 0.14 J | 4700 J | ug/kg | SS35 | 28 / 40 | 3.3 - 600 | 4700 | NA | 1900 C | NA | NA | Υ | ASL |
| | 50293 | 4,4'-DDT | 0.11 J | 3100 J | ug/kg | SS32 | 27 / 40 | 3.3 - 600 | 3100 | NA | 1900 C | NA | NA | Υ | ASL |
| | 84662 | Diethylphthalate | 30 J | 170 J | ug/kg | SS12 | 10 / 40 | 340 - 3400 | 170 | NA | 6300000 N | NA | NA | N | BSL |
| | 7440417 | Beryllium | 0.08 J | 1.5 J | mg/kg | SB07 | 34 / 40 | 0.02 - 0.36 | 1.5 | NA | 16 N | NA | NA | N | BSL |
| | 7440484 | Cobalt | 0.31 J | 36 | mg/kg | SB02 | 28 / 40 | 0.08 - 2.9 | 36 | NA | 160 N | NA | NA | N | BSL |
| | 7440508 | Copper | 0.9 J | 6470 | mg/kg | SS01 | 26 / 40 | 0.17 - 2.2 | 6470 | NA | 310 N | NA | NA | Υ | ASL |
| | 7439896 | Iron | 371 | 120000 | mg/kg | SS01 | 24 / 40 | 2.7 - 13.5 | 120000 | NA | 2300 N | NA | NA | Υ | ASL |
| | 7782492 | Selenium | 0.49 J | 1.6 J | mg/kg | SS23 | 12 / 40 | 0.4 - 1.1 | 1.6 | NA | 39 N | NA | NA | N | BSL |

(1) Maximum concentration used for screening.

(2) To date, no background study has been completed.

(3) All compounds were screened against the Risk-Based Concentration (RBC) Table, U.S. EPA Region III, May 8, 2001 for residential soil (cancer benchmark = 1E-06; HQ = 0.1). Lead was screened against the U.S. EPA screening value of 400 mg/kg.

(4) Rationale Codes:

Selection Reason: Above Screening Level (ASL)
Deletion Reason: Below Screening Level (BSL)

Definitions: NA = Not Applicable

J = Estimated Value

C = Carcinogen

N = Noncarcinogen

TABLE 3.1.RME

EXPOSURE POINT CONCENTRATION SUMMARY

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future

Medium: Groundwater

Exposure Medium: Groundwater

| Exposure Point | Chemical of Potential Concern | Units | Arithmetic Mean | 95% UCL (Distribution) | Maximum Concentration (Qualifier) | Exposure Poir Value Units | | Point Concentration Statistic | Rationale |
|-----------------------|-------------------------------|-------|--------------------|---------------------------|-----------------------------------|----------------------------|------|-------------------------------|--------------|
| Aquifer 1 - Tap Water | Bis(2-ethylhexyl)phthalate | ug/l | 4 | 5.5 (T) | 5 J | 5 | ug/l | Max | W-Test (1) |
| | Chloroform | ug/l | 1.9 | 14.9 (T) | 9 | 9 | ug/l | Max | W-Test (1) |
| | Heptachlor | ug/l | 27 | 30 (T) | 33 J | 30 | ug/l | 95% UCL - T | W - Test (2) |
| | Barium | ug/l | 224 | 2835 (T) | 489 | 489 | ug/l | Max | W-Test (1) |
| | Lead | ug/l | 21 | 32 (T) | 35 J | 32 | ug/l | 95% UCL - T | W - Test (2) |
| | Manganese | ug/l | 6052 | 33449 (T) | 12500 | 12500 | ug/l | Max | W-Test (1) |

Statistics: Maximum Detected Value (Max); 95% UCL of Transformed Data (95% UCL - T)

T = Transformed

(1) 95% UCL exceeds maximum detected concentration. Therefore, maximum concentration used for EPC.

J = Estimated Value

(2) Shapiro-Wilk W Test indicates data are log-normally distributed.

TABLE 3.2.RME

EXPOSURE POINT CONCENTRATION SUMMARY

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future

Medium: Groundwater Exposure Medium: Air

| Exposure Point | Chemical of | Units | Arithmetic | 95% UCL | Maximum Concentration | Exposure Point Concentration | | | | | |
|-------------------|----------------------------|-------|------------|----------------|--------------------------|------------------------------|-------|-------------|--------------|--|--|
| | Potential Concern | | Mean | (Distribution) | (Qualifier) | Value | Units | Statistic | Rationale | | |
| Water Vapors from | Bis(2-ethylhexyl)phthalate | ug/l | 4 | 5.5 (T) | 5 J | 5 | ug/l | Max | W-Test (1) | | |
| Showerhead | Chloroform | ug/l | 1.9 | 14.9 (T) | 9 | 9 | ug/l | Max | W-Test (1) | | |
| | Heptachlor | ug/l | 27 | 30 (T) | 33 J | 30 | ug/l | 95% UCL - T | W - Test (2) | | |

Statistics: Maximum Detected Value (Max); 95% UCL of Transformed Data (95% UCL - T)

T = Transformed

(1) 95% UCL exceeds maximum detected concentration. Therefore, maximum concentration used for EPC.

J = Estimated Value

(2) Shapiro-Wilk W Test indicates data are log-normally distributed.

TABLE 3.3.RME

EXPOSURE POINT CONCENTRATION SUMMARY

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future

Medium: Soil

Exposure Medium: Soil

| Exposure Point | Chemical of Potential Concern | Units | Arithmetic Mean | 95% UCL (Distribution) | Maximum Concentration (Qualifier) | Exposure Value Units | | e Point Concentration Statistic | Rationale |
|----------------|----------------------------------|-------|--------------------|---------------------------|-----------------------------------|-------------------------|-------|---------------------------------|--------------|
| Soil at Site 1 | 4,4'-DDD | ug/kg | 239 | 452 (T) | 4200 | 452 | ug/kg | 95 % UCL -T | W - Test (2) |
| | 4,4'-DDE | ug/kg | 596 | 6793 (T) | 7200 J | 6793 | ug/kg | 95% UCL - T | W - Test (2) |
| | 4,4'-DDT | ug/kg | 11007 | 28619 (N) | 290000 J | 28619 | ug/kg | 95% UCL - N | W - Test (1) |
| | Aluminum | mg/kg | 7450 | 9964 (T) | 21700 | 9964 | mg/kg | 95% UCL - T | W - Test (2) |
| | Lead | mg/kg | 210 | 345 (T) | 750 J | 345 | mg/kg | 95% UCL - T | W - Test (2) |
| | Manganese | mg/kg | 116 | 201 (T) | 688 | 201 | mg/kg | 95% UCL - T | W - Test (2) |
| Soil at Site 2 | 4,4'-DDE | ug/kg | 230 | 496 | 4700 J | 496 | ug/kg | 95 % UCL - T | W - Test (2) |
| | 4,4'-DDT | ug/kg | 183 | 322 (T) | 3100 J | 322 | ug/kg | 95% UCL - T | W - Test (2) |
| | Copper | mg/kg | 173 | 245 (T) | 6470 | 245 | mg/kg | 95% UCL - T | W - Test (2) |
| | Iron | mg/kg | 19518 | 32230 (T) | 120000 | 32230 | mg/kg | 95% UCL - T | W - Test (2) |

Statistics: 95% UCL of Normal Data (95% UCL - N); 95% UCL of Transformed Data (95% UCL - T)

(1) Shapiro-Wilk W Test indicates data are normally distributed.

(2) Shapiro-Wilk W Test indicates data are log-normally distributed.

N = Normal

T = Transformed

J = Estimated Value

TABLE 4.1.RME

VALUES USED FOR DAILY INTAKE CALCULATIONS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future

Medium: Groundwater

Exposure Medium: Groundwater

| Exposure Route | Receptor Population | Receptor Age | Exposure Point | Parameter | Parameter Definition | Value | Units | Rationale/ | Intake Equation/ |
|----------------|---------------------|--------------|-----------------------|-----------|---|-------------------------------------|-------------|------------------------|---|
| | | | | Code | | | | Reference | Model Name |
| Ingestion | Resident | Adult | Aquifer 1 - Tap Water | cw | Chemical Concentration in Water | See Table 3.1 | mg/l | See Table 3.1 | Chronic Daily Intake (CDI) (mg/kg/day) = |
| | | | | IR-W | Ingestion Rate of Water | 2 | l/day | EPA, 1991 | CW x IR-W x EF x ED x 1/BW x 1/AT |
| | | | | EF | Exposure frequency | 350 | days/year | EPA, 1991 | |
| | | | | ED | Exposure Duration | 24 | years | EPA, 1991 | |
| | | | | BW | Body Weight | 70 | kg | EPA, 1991 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 1989a | |
| | | | | AT-N | Averaging Time - Non-Cancer | 8,760 | days | EPA, 1989a | |
| | | Child | Aquifer 1 - Tap Water | CW | Chemical Concentration in Water | See Table 3.1 | mg/l | See Table 3.1 | CDI (mg/kg/day) = |
| | | | | IR-W | Ingestion Rate of Water | 1 | I/day | EPA, 1989b | CW x IR-W x EF x ED x 1/BW x 1/AT |
| | | | | EF | Exposure frequency | 350 | days/year | EPA. 1991 | |
| | | | | ED | Exposure Duration | 6 | vears | EPA, 1991 | |
| | | | | BW | Body Weight | 15 | kg | EPA, 1991 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 1989a | |
| | | | | AT-N | Averaging Time - Non-Cancer | | , | • | |
| Dames | Desident | A deale | Assistant Ton Mater | 014/ | | 2,190 | days | EPA, 1989a | Describe About ad Dane (DAD) (see the day) |
| Dermal | Resident | Adult | Aquifer 1 - Tap Water | CW | Chemical Concentration in Water | See Table 3.1 | mg/l | See Table 3.1 | Dermally Absorbed Dose (DAD) (mg/kg-day) = |
| | | | | FA Kp | Fraction Absorbed Water Permeability Constant | Chemical Specific Chemical Specific | cm/hr | EPA, 2001 EPA, 2001 | DA-event x EV x ED x EF x SA x 1/BW x 1/AT where for organic compounds, |
| | | | | SA | Skin Surface Area | 18,000 | cm2 | EPA, 2001 | Absorbed Dose per Event (DA-event) (mg/cm2-event) = |
| | | | | tau-event | Lag time per event | Chemical Specific | hours/event | EPA, 2001 | 2 FA x Kp x CW x CF x SQRT{(6 x tau-event x t-event)/pi} |
| | | | | t-event | Event Duration | 0.58 | hours/event | EPA, 2001 | or |
| | | | | В | Ratio of permeability coefficient of a | Chemical Specific | | EPA, 2001 | DA-event = FA x Kp x CW x {(t-event/(1 + B)) + |
| | | | | | compound through the stratum | | | | 2 x tau-event x ((1 + (3 x B) + (3 x B x B))/(1 + B)2)} |
| | | | | | corneum relative to its permeability | | | | and where for inorganic compounds, |
| | | | | | coefficient across the viable | | | | DA-event = Kp x CW x CF x t-event |
| | | | | | epidermis | | | | |
| | | | | EV | Event Frequency | 1 | events/day | EPA, 2001 | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 2001 | |
| | | | | ED | Exposure Duration | 24 | years | EPA, 1991 | |

TABLE 4.1.RME

VALUES USED FOR DAILY INTAKE CALCULATIONS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future Medium: Groundwater

Exposure Medium: Groundwater

| Exposure Route | Receptor Population | Receptor Age | Exposure Point | Parameter Code | Parameter Definition | Value | Units | Rationale/ Reference | Intake Equation/ Model Name |
|--------------------|---------------------|-------------------|-----------------------|-------------------|--|-------------------|-------------|-------------------------|--|
| Dermal (contimued) | Resident (continued | Adult (continued) | Aquifer 1 - Tap Water | CF | Volumetric Conversion Factor for Water | 0.001 | I/cm3 | | |
| | | | | BW | Body Weight | 70 | kg | EPA, 2001 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 2001 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 8,760 | days | EPA, 2001 | |
| | | Child | Aquifer 1 - Tap Water | cw | Chemical Concentration in Water | See Table 3.1 | mg/l | See Table 3.1 | DAD (mg/kg-day) = |
| | | | | FA | Fraction Absorbed Water | Chemical Specific | | EPA, 2001 | DA-event x EV x ED x EF x SA x 1/BW x 1/AT |
| | | | | Кр | Permeability Constant | Chemical Specific | cm/hr | EPA, 2001 | where for organic compounds, |
| | | | | SA | Skin Surface Area | 6,600 | cm2 | EPA, 2001 | DA-event (mg/cm2-event) = |
| | | | | tau-event | Lag time per event | · · | hours/event | EPA, 2001 | 2 FA x Kp x CW x CF x SQRT{(6 x tau-event x t-event)/pi} |
| | | | | t-event | Event Duration | | hours/event | EPA, 2001 | or |
| | | | | В | Ratio of permeability coefficient of a | Chemical Specific | | EPA, 2001 | DA-event = FA x Kp x CW x {(t-event/(1 + B)) + |
| | | | | | compound through the stratum | | | | 2 x tau-event x ((1 + (3 x B) + (3 x B x B))/(1 + B)2)} |
| | | | | | corneum relative to its permeability | | | | and where for inorganic compounds, |
| | | | | | coefficient across the viable | | | | DA-event = Kp x CW x CF x t-event |
| | | | | | epidermis | | | | |
| | | | | EV | Event Frequency | 1 | events/day | EPA, 2001 | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 2001 | |
| | | | | ED | Exposure Duration | 6 | years | EPA, 2001 | |
| | | | | CF | Volumetric Conversion Factor for Water | 0.001 | I/cm3 | | |
| | | | | BW | Body Weight | 15 | kg | EPA, 2001 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 2001 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 2,190 | days | EPA, 2001 | |

EPA 1989a: Risk Assessment Guidance for Superfund. Volume 1: Human Health Evaluation Manual, Part A. OERR EPA/540/1-89/002.

EPA 1989b: Exposure Factors Handbook, July 1989, EPA/600/8-89/043.

EPA 1991: Risk Assessment Guidance for Superfund. Volume 1: Human Health Evaluation Manual - Supplemental Guidance, Standard Default Exposure Factors. Interim Final. OSWER 9285.6-03.

EPA 1992: Dermal Exposure Assessment: Principles and Applications. EPA/600/8-91/011B.

EPA 1997: Exposure Factors Handbook, Volume 1. EPA/600/P-95/002Fa.

EPA 2001: Risk Assessment Guidance for Superfund. Volume 1: Human Health Evaluation Manual (Part E, Supplemental Guidance for Dermal Risk Assessment) Interim.

VALUES USED FOR DAILY INTAKE CALCULATIONS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future

Medium: Groundwater
Exposure Medium: Air

| Exposure Route | Receptor Population | Receptor Age | Exposure Point | Parameter Code | Parameter Definition | Value | Units | Rationale/ | Intake Equation/ Model Name |
|----------------|---------------------|--------------|---------------------------------|-------------------|----------------------|-------|-------|------------|--------------------------------|
| Inhalation (1) | Resident | Adult | Water Vapors from Showerhead | (1) | (1) | (1) | (1) | (1) | Foster and Chrostowski Model |

⁽¹⁾ Refer to the Risk Assessment text for details on the modeled intake methodology and parameters used to calculate modeled intake values for the Foster and Chrostowski Shower Model.

VALUES USED FOR DAILY INTAKE CALCULATIONS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future

Medium: Soil

Exposure Medium: Soil

| Exposure Route | Receptor Population | Receptor Age | Exposure Point | Parameter | Parameter Definition | Value | Units | Rationale/ | Intake Equation/ |
|----------------|---------------------|--------------|----------------|-----------|--------------------------------|---------------|-----------|-----------------------|--|
| | | | | Code | | | | Reference | Model Name |
| Ingestion | Resident | Adult | Soil at Site 1 | cs | Chemical Concentration in Soil | See Table 3.3 | mg/kg | See Table 3.3 | Chronic Daily Intake (CDI) (mg/kg-day) = |
| | | | | IR-S | Ingestion Rate of Soil | 100 | mg/day | EPA, 1991 | CS x IR x FI x EF x ED x CF1 x 1/BW x 1/AT |
| | | | | FI | Fraction Ingested | 1 | | Professional Judgment | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 1991 | |
| | | | | ED | Exposure Duration | 24 | years | EPA, 1991 | |
| | | | | CF1 | Conversion Factor | 1E-06 | kg/mg | | |
| | | | | BW | Body Weight | 70 | kg | EPA, 1991 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 1989 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 8,760 | days | EPA, 1989 | |
| | | | Soil at Site 2 | cs | Chemical Concentration in Soil | See Table 3.3 | mg/kg | See Table 3.3 | CDI (mg/kg-day) = |
| | | | | IR-S | Ingestion Rate of Soil | 100 | mg/day | EPA, 1991 | CS x IR x FI x EF x ED x CF1 x 1/BW x 1/AT |
| | | | | FI | Fraction Ingested | 1 | | Professional Judgment | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 1991 | |
| | | | | ED | Exposure Duration | 24 | years | EPA, 1991 | |
| | | | | CF1 | Conversion Factor | 1E-06 | kg/mg | | |
| | | | | BW | Body Weight | 70 | kg | EPA, 1991 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 1989 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 8,760 | days | EPA, 1989 | |
| | | Child | Soil at Site 1 | cs | Chemical Concentration in Soil | See Table 3.3 | mg/kg | See Table 3.3 | CDI (mg/kg-day) = |
| | | | | IR-S | Ingestion Rate of Soil | 200 | mg/day | EPA, 1991 | CS x IR x FI x EF x ED x CF1 x 1/BW x 1/AT |
| | | | | FI | Fraction Ingested | 1 | | Professional Judgment | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 1991 | |
| | | | | ED | Exposure Duration | 6 | years | EPA, 1991 | |
| | | | | CF1 | Conversion Factor | 1E-06 | kg/mg | | |
| | | | | BW | Body Weight | 15 | kg | EPA, 1991 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 1989 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 2,190 | days | EPA, 1989 | |

VALUES USED FOR DAILY INTAKE CALCULATIONS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future

Medium: Soil

Exposure Medium: Soil

| Exposure Route | Receptor Population | Receptor Age | Exposure Point | Parameter | Parameter Definition | Value | Units | Rationale/ | Intake Equation/ |
|-----------------------|----------------------|-------------------|----------------|-----------|---|-------------------|--------------|-----------------------|---|
| | | | | Code | | | | Reference | Model Name |
| Ingestion (continued) | Resident (continued) | Child (continued) | Soil at Site 2 | CS | Chemical Concentration in Soil | See Table 3.3 | mg/kg | See Table 3.3 | CDI (mg/kg-day) = |
| | | | | IR-S | Ingestion Rate of Soil | 200 | mg/day | EPA, 1991 | CS x IR x FI x EF x ED x CF1 x 1/BW x 1/AT |
| | | | | FI | Fraction Ingested | 1 | | Professional Judgment | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 1991 | |
| | | | | ED | Exposure Duration | 6 | years | EPA, 1991 | |
| | | | | CF1 | Conversion Factor | 1E-06 | kg/mg | | |
| | | | | BW | Body Weight | 15 | kg | EPA, 1991 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 1989 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 2,190 | days | EPA, 1989 | |
| Dermal | Resident | Adult | Soil at Site 1 | CS | Chemical Concentration in Soil | See Table 3.3 | mg/kg | See Table 3.3 | Dermal Absorbed Dose (DAD) (mg/kg-day) = |
| | | | | CF | Conversion Factor | 1E-06 | kg/mg | | DA-event x EF x ED x EV x SA X 1/BW x 1/AT |
| | | | | SA | Skin Surface Area Available for Contact | 5,700 | cm2 | EPA, 2001 | where |
| | | | | AF | Soil to Skin Adherence Factor | 0.07 | mg/cm2-event | EPA, 2001 | Absorbed Dose per Event (DA-event) (mg/cm2-event) = |
| | | | | ABS-d | Dermal Absorption Factor | chemical-specific | unitless | EPA, 2001 | CS x CF x AF x ABS-d |
| | | | | EV | Event Frequency | 1 | events/day | EPA, 2001 | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 2001 | |
| | | | | ED | Exposure Duration | 24 | years | EPA, 1991 | |
| | | | | BW | Body Weight | 70 | kg | EPA, 2001 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 2001 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 8,760 | days | EPA, 2001 | |

VALUES USED FOR DAILY INTAKE CALCULATIONS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future

Medium: Soil

Exposure Medium: Soil

| Exposure Route | Receptor Population | Receptor Age | Exposure Point | Parameter Code | Parameter Definition | Value | Units | Rationale/ | Intake Equation/ Model Name |
|--------------------|----------------------|-------------------|----------------|-------------------|---|-------------------|--------------|---------------|--|
| Dermal (continued) | Resident (continued) | Adult (continued) | Soil at Site 2 | CS | Chemical Concentration in Soil | See Table 3.3 | mg/kg | See Table 3.3 | DAD (mg/kg-day) = |
| | | | | CF | Conversion Factor | 1E-06 | kg/mg | | DA-event x EF x ED x EV x SA X 1/BW x 1/AT |
| | | | | | | | | | |
| | | | | SA | Skin Surface Area Available for Contact | 5,700 | cm2 | EPA, 2001 | where |
| | | | | AF | Soil to Skin Adherence Factor | 0.07 | mg/cm2-event | EPA, 2001 | DA-event (mg/cm2-event) = |
| | | | | ABS-d | Dermal Absorption Factor | chemical-specific | unitless | EPA, 2001 | CS x CF x AF x ABS-d |
| | | | | EV | Event Frequency | 1 | events/day | EPA, 2001 | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 2001 | |
| | | | | ED | Exposure Duration | 24 | years | EPA, 1991 | |
| | | | | BW | Body Weight | 70 | kg | EPA, 2001 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 2001 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 8,760 | days | EPA, 2001 | |
| | | Child | Soil at Site 1 | CS | Chemical Concentration in Soil | See Table 3.3 | mg/kg | See Table 3.3 | DAD (mg/kg-day) = |
| | | | | CF | Conversion Factor | 1E-06 | kg/mg | | DA-event x EF x ED x EV x SA X 1/BW x 1/AT |
| | | | | SA | Skin Surface Area Available for Contact | 2,800 | cm2 | EPA, 2001 | where |
| | | | | AF | Soil to Skin Adherence Factor | 0.2 | mg/cm2-event | EPA, 2001 | DA-event (mg/cm2-event) = |
| | | | | ABS-d | Dermal Absorption Factor | chemical-specific | unitless | EPA, 2001 | CS x CF x AF x ABS-d |
| | | | | EV | Event Frequency | 1 | events/day | EPA, 2001 | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 2001 | |
| | | | | ED | Exposure Duration | 6 | years | EPA, 2001 | |
| | | | | BW | Body Weight | 15 | kg | EPA, 2001 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 2001 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 2,190 | days | EPA, 2001 | |

VALUES USED FOR DAILY INTAKE CALCULATIONS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future

Medium: Soil

Exposure Medium: Soil

| Exposure Route | Receptor Population | Receptor Age | Exposure Point | Parameter Code | Parameter Definition | Value | Units | Rationale/ Reference | Intake Equation/ Model Name |
|--------------------|----------------------|-------------------|----------------|-------------------|---|-------------------|--------------|-------------------------|--|
| Dermal (continued) | Resident (continued) | Child (continued) | Soil at Site 2 | CS | Chemical Concentration in Soil | See Table 3.3 | mg/kg | See Table 3.3 | DAD (mg/kg-day) = |
| | | | | CF | Conversion Factor | 1E-06 | kg/mg | | DA-event x EF x ED x EV x SA X 1/BW x 1/AT |
| | | | | SA | Skin Surface Area Available for Contact | 2,800 | cm2 | EPA, 2001 | where |
| | | | | AF | Soil to Skin Adherence Factor | 0.2 | mg/cm2-event | EPA, 2001 | DA-event (mg/cm2-event) = |
| | | | | ABS-d | Dermal Absorption Factor | chemical-specific | unitless | EPA, 2001 | CS x CF x AF x ABS-d |
| | | | | EV | Event Frequency | 1 | events/day | EPA, 2001 | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 2001 | |
| | | | | ED | Exposure Duration | 6 | years | EPA, 2001 | |
| | | | | BW | Body Weight | 15 | kg | EPA, 2001 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 2001 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 2,190 | days | EPA, 2001 | |

EPA 1989: Risk Assessment Guidance for Superfund. Volume 1: Human Health Evaluation Manual, Part A. OERR EPA/540/1-89/002.

EPA 1991: Risk Assessment Guidance for Superfund. Volume 1: Human Health Evaluation Manual - Supplemental Guidance, Standard Default Exposure Factors. Interim Final. OSWER 9285.6-03.

EPA 1995: Assessing Dermal Exposure from Soil, Technical Guidance Manual, Region III, EPA/903-K-95-003.

EPA 1997: Exposure Factors Handbook, Volume 1. EPA/600/P-95/002Fa.

EPA 2001: Risk Assessment Guidance for Superfund. Volume 1: Human Health Evaluation Manual (Part E, Supplemental Guidance for Dermal Risk Assessment) Interim.

TABLE 5.1

NON-CANCER TOXICITY DATA -- ORAL/DERMAL

The Dean Company

| Chemical of Potential | Chronic/ Subchronic | Ora | l RfD | Oral Absoprtion Absorbed RfD for Dermal (2) Efficiency for Dermal (1) | | Primary Target | Combined Uncertainty/Modifying | RfD:Tar | get Organ(s) | |
|----------------------------|------------------------|----------|-----------|--|----------|-------------------|--------------------------------|---------|--------------|--------------|
| Concern | | Value | Units | | Value | Units | Organ(s) | Factors | Source(s) | Date(s) |
| | | | | | | | | | | (MM/DD/YYYY) |
| 4,4'-DDD | NA | NA | NA | 1 | NA | NA | NA | NA | NA | NA |
| 4,4'-DDE | NA | NA | NA | 1 | NA | NA | NA | NA | NA | NA |
| 4,4'-DDT | Chronic | 5.0E-004 | mg/kg/day | 1 | 5.0E-004 | mg/kg/day | Liver | 100 | IRIS | 06/21/2001 |
| 4,4'-DDT | Subchronic | 5.0E-004 | mg/kg/day | 1 | 5.0E-004 | mg/kg/day | Liver | 100 | HEAST | 07/01/1997 |
| Bis(2-ethylhexyl)phthalate | Chronic | 2.0E-02 | mg/kg/day | 1 | 2.0E-02 | mg/kg/day | Liver | 1000 | IRIS | 06/21/2001 |
| Bis(2-ethylhexyl)phthalate | Subchronic | 2.0E-02 | mg/kg/day | 1 | 2.0E-02 | mg/kg/day | Liver | 1000 | HEAST | 07/01/1997 |
| Chloroform | Chronic | 1.0E-02 | mg/kg/day | 1 | 1.0E-02 | mg/kg/day | Liver | 1000 | IRIS | 06/21/2001 |
| Chloroform | Subchronic | 1.0E-02 | mg/kg/day | 1 | 1.0E-02 | mg/kg/day | Liver | 1000 | HEAST | 07/01/1997 |
| Heptachlor | Chronic | 5.0E-04 | mg/kg/day | 1 | 5.0E-04 | mg/kg/day | Liver | 300 | IRIS | 06/21/2001 |
| Heptachlor | Subchronic | 5.0E-04 | mg/kg/day | 1 | 5.0E-04 | mg/kg/day | Liver | 300 | HEAST | 07/01/1997 |
| Aluminum | Chronic | 1.0E+00 | mg/kg/day | 1 | 1.0E+00 | mg/kg/day | Central Nervous System | 100 | NCEA | 06/21/2001 |
| Barium | Chronic | 7.0E-02 | mg/kg/day | 0.07 | 4.9E-03 | mg/kg/day | Heart | 3 | IRIS | 02/02/2001 |
| Barium | Subchronic | 7.0E-02 | mg/kg/day | 0.07 | 4.9E-03 | mg/kg/day | Heart | 3 | HEAST | 07/01/1997 |
| Copper | Chronic | 3.7E-02 | mg/kg/day | 1 | 3.7E-02 | mg/kg/day | Gastrointestinal | NA | HEAST | 07/01/1997 |
| Copper | Subchronic | 3.7E-02 | mg/kg/day | 1 | 3.7E-02 | mg/kg/day | Gastrointestinal | NA | HEAST | 07/01/1997 |
| Iron | Chronic | 3.0E-01 | mg/kg/day | 1 | 3.0E-01 | mg/kg/day | Gastrointestinal | 1 | NCEA | 06/21/2001 |
| Lead | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| Manganese (nonfood) | Chronic | 2.0E-02 | mg/kg/day | 0.04 | 8.0E-04 | mg/kg/day | Central Nervous System | 1 | IRIS | 06/21/2001 |

(1) Source: Risk Assessment Guidance for Superfund. Volume 1: Human Health Evaluation Manual (Part E, Supplemental Guidance for Dermal Risk Assessment) Interim. Section 4.2 and Exhibit 4-1.

(2) See Risk Assessment text for the derivation of the "Absorbed RfD for Dermal".

Definitions: NA = Not Available

IRIS = Integrated Risk Information System

HEAST = Health Effects Assessment Summary Table, July 1997

NCEA = National Center for Environmental Assessment

TABLE 5.2

NON-CANCER TOXICITY DATA -- INHALATION

The Dean Company

| Chemical of Potential | Chronic/ Subchronic | Inhalat | ion RfC | Extrapolat | ed RfD (1) | Primary Target | Combined Uncertainty/Modifying | RfC : Target Organ(s) | |
|----------------------------|------------------------|---------|---------|------------|------------|------------------------|--------------------------------|-----------------------|--------------|
| Concern | | Value | Units | Value | Units | Organ(s) | Factors | Source(s) | Date(s) |
| | | | | | | | | | (MM/DD/YYYY) |
| 4,4'-DDD | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| 4,4'-DDE | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| 4,4'-DDT | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| Bis(2-ethylhexyl)phthalate | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| Chloroform | Chronic | 3.0E-04 | mg/m3 | 8.6E-05 | mg/kg/day | Nasal | 1000 | NCEA | 06/21/2001 |
| Chloroform | Subchronic | 3.0E-03 | mg/m3 | 8.6E-4 | mg/kg/day | Nasal | 100 | NCEA | 06/21/2001 |
| Heptachlor | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| Aluminum | Chronic | 5.0E-03 | mg/m3 | 1.4E-03 | mg/kg/day | Central Nervous System | 300 | NCEA | 06/21/2001 |
| Barium | Chronic | 5.0E-04 | mg/m3 | 1.4E-04 | mg/kg/day | Fetus | 1000 | HEAST | 07/01/1997 |
| Barium | Subchronic | 5.0E-03 | mg/m3 | 1.4E-03 | mg/kg/day | Fetus | 100 | HEAST | 07/01/1997 |
| Copper | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| Iron | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| Lead | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| Manganese (nonfood) | Chronic | 5.0E-05 | mg/m3 | 1.4E-05 | mg/kg/day | Central Nervous System | 1000 | IRIS | 06/21/2001 |

(1) See Risk Assessment text for the derivation of the "Extrapolated RfD".

Definitions: NA = Not Available

IRIS = Integrated Risk Information System

HEAST = Health Effects Assessment Summary Table, July 1997

NCEA = National Center for Environmental Assessment

TABLE 5.3 NON-CANCER TOXICITY DATA -- SPECIAL CASE CHEMICALS The Dean Company

| Chemical of Potential | Chronic/ Subchronic | Parameter Primary Target Organ(s) | | | Combined Uncertainty/Modifying | Parameter:1 | Parameter:Target Organ(s) | |
|-----------------------|------------------------|-----------------------------------|-------|-------|--------------------------------|-------------|---------------------------|-------------------------|
| Concern | | Name | Value | Units | | Factors | Source(s) | Date(s) (MM/DD/YYYY) |
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There are no special case chemicals in this risk assessment. As a result, the table is blank.

TABLE 6.1

CANCER TOXICITY DATA -- ORAL/DERMAL

The Dean Company

| Chemical of Potential | Oral Cancer Slope Factor | | Oral Cancer Slope Factor | | Oral Absorption Efficiency for Dermal (1) | | cer Slope Factor | Weight of Evidence/ | C | oral CSF |
|----------------------------|--------------------------|-------------|--------------------------|----------|---|-------------|------------------|----------------------|---|----------|
| Concern | Value | Units | | Value | Units | Description | Source(s) | Date(s) (MM/DD/YYYY) | | |
| 4,4'-DDD | 2.4E-01 | 1/mg/kg/day | 1 | 2.4E-01 | 1/mg/kg/day | B2 | IRIS | 06/21/2001 | | |
| 4,4'-DDE | 3.4E-01 | 1/mg/kg/day | 1 | 3.4E-01 | 1/mg/kg/day | B2 | IRIS | 06/21/2001 | | |
| 4,4'-DDT | 3.4E-001 | 1/mg/kg/day | 1 | 3.4E-001 | 1/mg/kg/day | B2 | IRIS | 06/21/2001 | | |
| Bis(2-ethylhexyl)phthalate | 1.4E-02 | 1/mg/kg/day | 1 | 1.4E-02 | 1/mg/kg/day | B2 | IRIS | 06/21/2001 | | |
| Chloroform | 6.1E-03 | 1/mg/kg/day | 1 | 6.1E-03 | 1/mg/kg/day | B2 | IRIS | 06/21/2001 | | |
| Heptachlor | 4.5E+00 | 1/mg/kg/day | 1 | 4.5E+00 | 1/mg/kg/day | B2 | IRIS | 06/21/2001 | | |
| Aluminum | NA | NA | 1 | NA | NA | NA | NA | NA | | |
| Barium | NA | NA | 0.07 | NA | NA | NA | NA | NA | | |
| Copper | NA | NA | 1 | NA | NA | NA | NA | NA | | |
| Iron | NA | NA | 1 | NA | NA | NA | NA | NA | | |
| Lead | NA | NA | NA | NA | NA | NA | NA | NA | | |
| Manganese (nonfood) | NA | NA | 0.04 | NA | NA | NA | NA | NA | | |

(1) Source: Risk Assessment Guidance for Superfund. Volume 1: Human Health Evaluation Manual (Part E, Supplemental Guidance for Dermal Risk Assessment) Interim. Section 4.2 and Exhibit 4-1.

(2) See Risk Assessment text for the derivation of the "Absorbed Cancer Slope Factor for Dermal".

Definitions: NA = Not Available

IRIS = Integrated Risk Information System

B2 = Probable Human Carcinogen - indicates sufficient evidence

in animals and inadequate or no evidence in humans

TABLE 6.2

CANCER TOXICITY DATA -- INHALATION

The Dean Company

| Chemical of Potential | Unit | Risk | Inhalation Cand | eer Slope Factor | Weight of Evidence/ | | | |
|----------------------------|----------|---------|-----------------|------------------|---------------------|-----------|--------------|--|
| Concern | Value | Units | Value | Units | Description | Source(s) | Date(s) | |
| | | | | | | | (MM/DD/YYYY) | |
| 4,4'-DDD | NA | NA | NA | NA | NA | NA | NA | |
| 4,4-DDE | NA | NA | NA | NA | NA | NA | NA | |
| 4,4'-DDT | 9.7E-005 | 1/ug/m3 | 3.4E-001 | 1/mg/kg/day | B2 | IRIS | 06/21/2001 | |
| Bis(2-ethylhexyl)phthalate | NA | NA | NA | NA | NA | NA | NA | |
| Chloroform | 2.3E-05 | 1/ug/m3 | 8.1E-02 | 1/mg/kg/day | B2 | IRIS | 06/21/2001 | |
| Heptachlor | 1.3E-03 | 1/ug/m3 | 4.5E+00 | 1/mg/kg/day | B2 | IRIS | 06/21/2001 | |
| Aluminum | NA | NA | NA | NA | NA | NA | NA | |
| Barium | NA | NA | NA | NA | NA | NA | NA | |
| Copper | NA | NA | NA | NA | NA | NA | NA | |
| Iron | NA | NA | NA | NA | NA | NA | NA | |
| Lead | NA | NA | NA | NA | NA | NA | NA | |
| Manganese (nonfood) | NA | NA | NA | NA | NA | NA | NA | |
| Thallium | NA | NA | NA | NA | NA | NA | NA | |

Definitions: NA = Not Available

IRIS = Integrated Risk Information System

B2 = Probable Human Carcinogen - indicates sufficient evidence

in animals and inadequate or no evidence in humans

TABLE 6.3 CANCER TOXICITY DATA -- SPECIAL CASE CHEMICALS The Dean Company

| Chemical of Potential | | Parameters | | Source(s) | Date(s) (MM/DD/YYYY) | |
|-----------------------|------|-------------|------|-----------|-------------------------|--|
| Concern | Name | Value Units | | | | |
| | | Not Applica | able | | | |

There are no special case chemicals in this risk assessment. As a result, this table is blank.

$\label{eq:table 6.4}$ Cancer toxicity data -- external (radiation)

The Dean Company

| Chemical of Potential | | ope Factor | Source(s) | Date(s) (MM/DD/YYYY) |
|-----------------------|-------|------------|-----------|-------------------------|
| Concern | Value | Units | | |
| | No | t Applicak | ole | |

There are no radionuclides in this risk assessment. As a result, this table is blank.

TABLE 7.1.RME

CALCULATION OF CHEMICAL CANCER RISKS AND NON-CANCER HAZARDS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

| Medium | Exposure Medium | Exposure Point | Exposure Route | Chemical of | EF | PC | | С | ancer Risk Calculat | ions | | | Non- | Cancer Hazard Cal | culations | |
|-------------------|-----------------------|-----------------------|------------------|------------------------------------|---------------|--------|---------------|------------------|---------------------|-------------------|-------------|---------------|------------------|-------------------|-----------------|-----------------|
| | | | | Potential Concern | Value | Units | Intake/Exposu | re Concentration | CSF/L | Init Risk | Cancer Risk | Intake/Exposu | re Concentration | RfD | D/RfC | Hazard Quotient |
| | | | | | | | Value | Units | Value | Units | | Value | Units | Value | Units | |
| Groundwater | Groundwater | Aquifer 1 - Tap Water | Ingestion | Bis(2-ethylhexyl)phthalate | 0.005 | mg/l | 4.7E-05 | mg/kg/day | 1.4E-02 | 1/mg/kg/day | 7E-07 | 1.4E-04 | mg/kg/day | 2.0E-02 | mg/kg/day | 0.007 |
| | | | | Chloroform | 0.009 | mg/l | 8.5E-05 | mg/kg/day | 6.1E-03 | 1/mg/kg/day | 5E-07 | 2.5E-04 | mg/kg/day | 1.0E-02 | mg/kg/day | 0.03 |
| | | | | Heptachlor | 0.03 | mg/l | 2.8E-04 | mg/kg/day | 4.5E-00 | 1/mg/kg/day | 1E-03 | 8.1E-04 | mg/kg/day | 5.0E-04 | mg/kg/day | 2 |
| | | | | Barium | 0.489 | mg/l | 4.6E-03 | mg/kg/day | NA | NA | NA | 1.3E-02 | mg/kg/day | 7.0E-02 | mg/kg/day | 0.2 |
| | | | | Lead (1) | | | | | | | | | | | | |
| | | | | Manganese | 12.5 | mg/l | 1.2E-01 | mg/kg/day | NA | NA | NA | 3.4E-01 | mg/kg/day | 2.0E-02 | mg/kg/day | 17 |
| | | | Exp. Route Total | | • | • | | • | • | • | 1E-03 | | • | • | • | 19 |
| | | | Dermal | Bis(2-ethylhexyl)phthalate | 0.005 | mg/l | 7.2E-05 | mg/kg/day | 1.4E-02 | 1/mg/kg/day | 1E-06 | 2.1E-04 | mg/kg/day | 2.2E-02 | mg/kg/day | 0.01 |
| | | | | Chloroform | 0.009 | mg/l | 1.7E-04 | mg/kg/day | 6.1E-03 | 1/mg/kg/day | 1E-06 | 4.9E-04 | mg/kg/day | 1.0E-02 | mg/kg/day | 0.05 |
| | | | | Heptachlor | 0.03 0.489 | mg/l | 1.3E-04 NA | mg/kg/day NA | 4.5E-00 NA | 1/mg/kg/day NA | 6E-04 NA | 3.9E-04 NA | mg/kg/day NA | 5.0E-04 NA | mg/kg/day NA | 0.8 NA |
| | | | | Barium | 0.469 | mg/l | INA | INA | INA | INA | INA | INA | | INA | INA | INA |
| | | | | Lead (1) | 12.5 | mg/l | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | Exp. Route Total | Manganese | 12.5 | IIIg/I | INA | INA | IVA | IVA | 6E-04 | INA | INA | INA | INA | 0.9 |
| | | Exposure Point Total | ' ' ' ' | | | | | | | | 2E-03 | | | | | 20 |
| | Exposure Medium Total | | | | | | | | | | 2E-03 | | | | | 20 |
| | Air | Water Vapors from | Inhalation | Dis (O subsulta see District state | 0.005 | mg/l | 2.3E-06 | mg/kg/day | NA | NA | NA NA | 3.6E-06 | mg/kg/day | NA | NA | NA NA |
| | | Showerhead | | Bis(2-ethylhexyl)phthalate | 0.009 | mg/l | 1.3E-04 | mg/kg/day | 8.1E-02 | 1/mg/kg/day | 1E-05 | 3.9E-04 | mg/kg/day | 8.6E-05 | mg/kg/day | 5 |
| | | | | Chloroform | 0.03 | mg/l | 2.6E-04 | mg/kg/day | 4.5E-00 | 1/mg/kg/day | 1E-03 | 7.7E-04 | mg/kg/day | NA. | NA | NA NA |
| | | | Exp. Route Total | Heptachlor | | | | | | 9.19 / | 1E-03 | | 99, | | <u> </u> | 5 |
| | | Exposure Point Total | | | | | | | | | 1E-03 | | | | | 5 |
| | Exposure Medium Total | | | | | | | | | | 1E-03 | | | | | 5 |
| Groundwater Total | 1 | | | | | | | | | | 3E-03 | | | | | 25 |
| Soil | Soil | Soil at Site 1 | Ingestion | 4.4'-DDD | 0.452 | mg/kg | 2.1E-07 | mg/kg/day | 2.4E-01 | 1/mg/kg/day | 5E-08 | 6.2E-07 | mg/kg/day | NA | NA | NA |
| | | | | * | 6.8 | mg/kg | 3.2E-06 | mg/kg/day | 3.4E-01 | 1/mg/kg/day | 1E-06 | 9.3E-06 | mg/kg/day | NA | NA. | NA NA |
| | | | | 4,4'-DDE | 28.6 | mg/kg | 1.3E-05 | mg/kg/day | 3.4E-01 | 1/mg/kg/day | 5E-06 | 3.9E-05 | mg/kg/day | 5.0E-04 | mg/kg/day | 0.08 |
| | | | | 4,4'-DDT Aluminum | 9964 | mg/kg | 4.7E-03 | mg/kg/day | NA | NA | NA | 1.4E-02 | mg/kg/day | 1.0E+00 | mg/kg/day | 0.01 |
| | | | | | | | | | | | | | | | | |
| | | | | Lead (1) | 201 | mg/kg | 9.5E-05 | mg/kg/day | NA | NA. | NA | 2.8E-04 | mg/kg/day | 1.4E-01 | mg/kg/day | 0.002 |
| | | | Exp. Route Total | Manganese | | | | 3 3 1 7 | <u> </u> | <u> </u> | 6E-06 | | 0 0 11 | ļ | 3 3 , | 0.09 |
| | | | Dermal | 4 # 000 | 0.452 | mg/kg | NA | NA NA | NA | NA NA | NA | NA | NA | NA | NA | NA NA |
| | | | Soma | 4,4'-DDD 4.4'-DDE | 6.8 | mg/kg | NA NA | NA NA | NA NA | NA NA | NA NA | NA NA | NA NA | NA NA | NA NA | NA NA |
| | | | | 4,4'-DDT | 28.6 | mg/kg | 1.6E-06 | mg/kg/day | 3.4E-01 | 1/mg/kg/day | 5E-07 | 4.7E-06 | mg/kg/day | 5.0E-04 | mg/kg/day | 0.009 |
| | | | | Aluminum | 9964 | mg/kg | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | | Lead (1) | | | | | | | | | | | | |
| | | | | Manganese | 201 | mg/kg | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | Exp. Route Total | | | | | | | | 5E-07 | | | | | 0.009 |
| | | Exposure Point Total | | | | | | | | | 7E-06 | | | | | 0.1 |
| | | | | | | | | | | | | | | | | |

TABLE 7.1.RME

CALCULATION OF CHEMICAL CANCER RISKS AND NON-CANCER HAZARDS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

| Medium | Exposure Medium | Exposure Point | Exposure Route | Chemical of | Е | PC | | С | ancer Risk Calculati | ions | | | Non- | Cancer Hazard Cal | culations | |
|------------------|-----------------------|----------------------|------------------|-------------------|-------|-------|---------------|-----------------|----------------------|------------------|-------------|----------------|------------------|-------------------|------------------|-----------------|
| | | | | Potential Concern | Value | Units | Intake/Exposu | e Concentration | CSF/U | Jnit Risk | Cancer Risk | Intake/Exposur | re Concentration | RfE | D/RfC | Hazard Quotient |
| | | | | | | | Value | Units | Value | Units | | Value | Units | Value | Units | |
| Soil (continued) | Soil (continued) | Soil at Site 2 | Ingestion | 4,4'-DDE | 0.496 | mg/kg | 2.3E-07 | mg/kg/day | 3.4E-01 | 1/mg/kg/day | 8E-08 | 6.8E-07 | mg/kg/day | NA | NA | NA |
| | | | İ | 4,4'-DDT | 0.322 | mg/kg | 1.5E-07 | mg/kg/day | 3.4E-01 | 1/mg/kg/day | 5E-08 | 4.4E-07 | mg/kg/day | 5.0E-04 | mg/kg/day | 0.0009 |
| | | | | Copper | 245 | mg/kg | 1.2E-04 | mg/kg/day | NA | NA | NA | 3.4E-04 | mg/kg/day | 3.7E-02 | mg/kg/day | 0.009 |
| | | | | Iron | 32230 | mg/kg | 1.5E-02 | mg/kg/day | NA | NA | NA | 4.4E-02 | mg/kg/day | 3.0E-01 | mg/kg/day | 0.1 |
| | | | Exp. Route Total | | 1 | 1 | | | Į. | • | 1E-07 | | • | | * | 0.1 |
| | | | Dermal | 4,4'-DDE | 0.496 | mg/kg | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | | 4,4'-DDT | 0.322 | mg/kg | 1.8E-08 | mg/kg/day | 3.4E-01 | 1/mg/kg/day | 6E-09 | 5.3E-08 | mg/kg/day | 5.0E-04 | mg/kg/day | 0.0001 |
| | | | | Copper | 245 | mg/kg | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | | Iron | 32230 | mg/kg | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | Exp. Route Total | | | • | | • | • | | 6E-09 | | • | • | • | 0.0001 |
| | | Exposure Point Total | | | | | | | | | 1E-07 | | | | | 0.1 |
| | Exposure Medium Total | | | | | | | | • | | 7E-06 | | • | | | 0.2 |
| Soil Total | | | | | | | | | | | 0.2 | | | | | |
| | | | | | | | | Tota | of Receptor Risks | Across All Media | 3E-03 | | Total of | Receptor Hazards | Across All Media | 25 |

⁽¹⁾ Lead is evaluated for the resident using the IEUBK model. See Risk Assessment text for discussion of results and appendix for the lead modeling run results.

TABLE 7.2.RME

CALCULATION OF CHEMICAL CANCER RISKS AND NON-CANCER HAZARDS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future
Receptor Population: Resident

| Medium | Exposure Medium | Exposure Point | Exposure Route | Chemical of | E | PC | | Ca | ncer Risk Calculat | ions | | | Non- | Cancer Hazard Ca | lculations | |
|-------------------|-----------------------|--|------------------|----------------------------|-------|-------|---------------|-------------------|--------------------|-------------|-------------|---------------|------------------|------------------|------------|-----------------|
| | | | | Potential Concern | Value | Units | Intake/Exposu | ire Concentration | CSF/L | Init Risk | Cancer Risk | Intake/Exposu | re Concentration | Rfl | D/RfC | Hazard Quotient |
| | | | | | | | Value | Units | Value | Units | | Value | Units | Value | Units | |
| Groundwater | Groundwater | Aquifer 1 - Tap Water | Ingestion | Bis(2-ethylhexyl)phthalate | 0.005 | mg/l | 2.7E-05 | mg/kg/day | 1.4E-02 | 1/mg/kg/day | 4E-07 | 3.2E-04 | mg/kg/day | 2.0E-02 | mg/kg/day | 0.02 |
| | | | | Chloroform | 0.009 | mg/l | 4.9E-05 | mg/kg/day | 6.1E-03 | 1/mg/kg/day | 3E-07 | 5.8E-04 | mg/kg/day | 1.0E-02 | mg/kg/day | 0.06 |
| | | | | Heptachlor | 0.03 | mg/l | 1.6E-04 | mg/kg/day | 4.5E-00 | 1/mg/kg/day | 7E-04 | 1.9E-03 | mg/kg/day | 5.0E-04 | mg/kg/day | 4 |
| | | | | Barium | 0.489 | mg/l | 2.7E-03 | mg/kg/day | NA | NA | NA | 3.1E-02 | mg/kg/day | 7.0E-02 | mg/kg/day | 0.4 |
| | | | | Lead (1) | | | | | | | | | | | | |
| | | | | Manganese | 12.5 | mg/l | 6.8E-02 | mg/kg/day | NA | NA | NA | 8.0E-01 | mg/kg/day | 2.0E-02 | mg/kg/day | 40 |
| | | | Exp. Route Total | | • | • | | • | | • | 7E-04 | | • | | • | 44 |
| | | | Demal | Bis(2-ethylhexyl)phthalate | 0.005 | mg/l | 3.1E-05 | mg/kg/day | 1.4E-02 | 1/mg/kg/day | 4E-07 | 3.6E-04 | mg/kg/day | 2.2E-02 | mg/kg/day | 0.02 |
| | | | | Chloroform | 0.009 | mg/l | 7.2E-05 | mg/kg/day | 6.1E-03 | 1/mg/kg/day | 4E-07 | 8.4E-04 | mg/kg/day | 1.0E-02 | mg/kg/day | 0.08 |
| | | | | Heptachlor | 0.03 | mg/l | 5.7E-05 | mg/kg/day | 4.5E-00 | 1/mg/kg/day | 3E-04 | 6.7E-04 | mg/kg/day | 5.0E-04 | mg/kg/day | 1 |
| | | | | Barium | 0.489 | mg/l | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | | Lead (1) | | | | | | | | | | | | |
| | | | | Manganese | 12.5 | mg/l | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | Exp. Route Total | | • | • | | • | | • | 3E-04 | | • | | | 1 |
| | | Exposure Point Total | | | | | | | | | 1E-03 | | | | | 45 |
| l [r | Exposure Medium Total | • | | | | | | | | | 1E-03 | | | | | 45 |
| Groundwater Total | | | | | | | | | | | 1E-03 | | | | | 45 |
| Soil | Soil | Soil at Site 1 | Ingestion | 4,4'-DDD | 0.452 | mg/kg | 5.0E-07 | mg/kg/day | 2.4E-01 | 1/mg/kg/day | 1E-07 | 5.8E-06 | mg/kg/day | NA | NA | NA |
| | | | i | 4,4'-DDE | 6.8 | mg/kg | 7.4E-06 | mg/kg/day | 3.4E-01 | 1/mg/kg/day | 3E-06 | 8.7E-05 | mg/kg/day | NA | NA | NA |
| | | | | 4,4'-DDT | 28.6 | mg/kg | 3.1E-05 | mg/kg/day | 3.4E-01 | 1/mg/kg/day | 1E-05 | 3.7E-04 | mg/kg/day | 5.0E-04 | mg/kg/day | 0.7 |
| | | | | Aluminum | 9964 | mg/kg | 1.1E-02 | mg/kg/day | NA | NA | NA | 1.3E-01 | mg/kg/day | 1.0E-00 | mg/kg/day | 0.1 |
| | | | | Lead (1) | | | | | | | | | | | | |
| | | | | Manganese | 201 | mg/kg | 2.2E-04 | mg/kg/day | NA | NA | NA | 2.6E-03 | mg/kg/day | 1.4E-01 | mg/kg/day | 0.02 |
| | | | Exp. Route Total | | • | • | | • | | | 1E-05 | İ | • | | | 0.8 |
| | | | Dermal | 4,4'-DDD | 0.452 | mg/kg | NA NA | NA. | NA | NA | NA | NA. | NA | NA | NA | NA |
| | | | | 4,4'-DDE | 6.8 | mg/kg | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | | 4,4'-DDT | 28.6 | mg/kg | 2.6E-06 | mg/kg/day | 3.4E-01 | 1/mg/kg/day | 9E-07 | 3.1E-05 | mg/kg/day | 5.0E-04 | mg/kg/day | 0.06 |
| | | | | Aluminum | 9964 | mg/kg | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | | Lead (1) | | | | | | | | | | | | |
| | | | | Manganese | 201 | mg/kg | NA | NA | NA | NA | NA | NA | NA | MA | NA | NA |
| | | | Exp. Route Total | | | • | | • | - | • | 9E-07 | | • | - | · | 0.06 |
| 1 | l l | Exp. Route Total Exposure Point Total | | | | | | | | | 1F-05 | | | | | 0.9 |

TABLE 7.2.RME

CALCULATION OF CHEMICAL CANCER RISKS AND NON-CANCER HAZARDS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

| Medium | Exposure Medium | Exposure Point | Exposure Route | Chemical of | E | PC | | Ca | ncer Risk Calculat | ions | | | Non- | Cancer Hazard Ca | Iculations | |
|--------------------|-----------------------|----------------------|------------------|-------------------|-------|-------|----------------|------------------|--------------------|------------------|-------------|----------------|-----------------|------------------|------------------|-----------------|
| | | | | Potential Concern | Value | Units | Intake/Exposur | re Concentration | CSF/L | Init Risk | Cancer Risk | Intake/Exposur | e Concentration | RfE | D/RfC | Hazard Quotient |
| | | | | | | | Value | Units | Value | Units | | Value | Units | Value | Units | |
| Soil (continued) | Soil (continued) | Soil at Site 2 | Ingestion | 4,4'-DDE | 0.496 | mg/kg | 5.4E-07 | mg/kg/day | 3.4E-01 | 1/mg/kg/day | 2E-07 | 6.3E-06 | mg/kg/day | NA | NA | NA |
| | | | İ | 4,4'-DDT | 0.322 | mg/kg | 3.5E-07 | mg/kg/day | 3.4E-01 | 1/mg/kg/day | 1E-07 | 4.1E-06 | mg/kg/day | 5.0E-04 | mg/kg/day | 0.008 |
| | | | | Copper | 245 | mg/kg | 2.7E-04 | mg/kg/day | NA | NA | NA | 3.1E-03 | mg/kg/day | 3.7E-02 | mg/kg/day | 0.08 |
| | | | | Iron | 32230 | mg/kg | 3.5E-02 | mg/kg/day | NA | NA | NA | 4.1E-01 | mg/kg/day | 3.0E-01 | mg/kg/day | 1 |
| | | | Exp. Route Total | | • | • | | | | • | 3E-07 | | | | • | 1 |
| | | | Dermal | 4,4'-DDE | 0.496 | mg/kg | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | | 4,4'-DDT | 0.322 | mg/kg | 3.0E-08 | mg/kg/day | 3.4E-04 | 1/mg/kg/day | 1E-08 | 3.5E-007 | mg/kg/day | 5.0E-004 | mg/kg/day | 0.0007 |
| | | | | Copper | 245 | mg/kg | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | | Iron | 32230 | mg/kg | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | Exp. Route Total | | • | | | | | | 1E-08 | | | | | 0.0007 |
| | | Exposure Point Total | | | | | | | | | 3E-07 | | | | | 1 |
| | Exposure Medium Total | | | | | | | | | | 1E-05 | | | | | 2 |
| Soil Total 1E-05 2 | | | | | | | | | 2 | | | | | | | |
| | | <u> </u> | | <u> </u> | | | | Total | of Receptor Risks | Across All Media | 1E-03 | | Total of | Receptor Hazards | Across All Media | 47 |

⁽¹⁾ Lead is evaluated for the resident using the IEUBK model. See Risk Assessment text for discussion of results and appendix for the lead modeling run results.

TABLE 8.1.RME CALCULATION OF RADIATION CANCER RISKS The Dean Company

| Scenario Timeframe: | |
|----------------------|--|
| Receptor Population: | |
| Receptor Age: | |

| Medium | Exposure Medium | Exposure Point | Exposure Route | Radionuclide of Potential Concern | EF | PC . | Risk Calculation | | | Cancer Risk Calcula | ations | |
|--------|-----------------|----------------------|------------------|-----------------------------------|-------|-------|------------------|--------|-----------|----------------------|------------------|-------------|
| | | | | | Value | Units | Approach | Intake | /Activity | | SF | Cancer Risk |
| | | | | | | | | Value | Units | Value | Units | |
| | | | | | | | | | | | | |
| | | | | | | | | | | | | |
| | | | | | | | | | | | | |
| | | | | | | | | | | | | |
| | | | | Ī | | | | | | | | |
| | | | Exp. Route Total | | | | | | | | | |
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| | | | | | | | | | | | | |
| | | | | | | | | | | | | |
| | | | Exp. Route Total | • | | | | | | | | |
| | ı | Exposure Point Total | , | | | | | | | | ļ | |
| | | | | | | | | | | | | |
| | | | | Not Appli | aabla | | | | | | | |
| | | | | Not Appli | Lable | | | | | | | |
| | | | | | | | | | | | | |
| | | | Exp. Route Total | Ī | | | | | | | | |
| | ı | Exposure Point Total | Exp. Route Total | | | | | | | | | |
| | | Exposure Form Total | ļ . | | 1 | 1 | | 1 | 1 | | | |
| | | | | | | | | | | | | |
| | | | | | | | | | | | | |
| | | | | | | | | | | | | |
| | | | | | | | | | | | | |
| | | | Exp. Route Total | | | | | | | | | |
| | | | | | | | | | | | | |
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| | | | | | | | | | | | | |
| | | | | | | | | | | | | |
| | | | | | | | | | | | | |
| |] | | Exp. Route Total | | | | | | | | | |
| | | Exposure Point Total | | | | | | | | | | |
| | | • | ·· | | | | ·· | | Tot | al of Receptor Risks | Across All Media | |

There are no radionuclides in this risk assessment. As a result, this table is blank.

TABLE 9.1.RME SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCs REASONABLE MAXIMUM EXPOSURE

The Dean Company

| Medium | Exposure Medium | Exposure Point | Chemical of Potential | | | Carcinogen | ic Risk | | | Non-Carcinoge | enic Hazard Quo | otient | |
|----------------|--------------------|--|----------------------------|-----------|------------|------------|-------------|--------------|------------------------|---------------|-----------------|--------|--------------|
| | | | Concern | Ingestion | Inhalation | Dermal | External | Exposure | Primary | Ingestion | Inhalation | Dermal | Exposure |
| | | | | | | | (Radiation) | Routes Total | Target Organ(s) | | | | Routes Total |
| Groundwater | Groundwater | Aquifer 1 - Tap Water | Bis(2-ethylhexyl)phthalate | 7E-07 | | 1E-06 | | 2E-06 | Liver | 0.007 | | 0.01 | 0.02 |
| | | | Chloroform | 5E-07 | | 1E-06 | | 2E-06 | Liver | 0.03 | | 0.05 | 0.08 |
| | | | Heptachlor | 1E-03 | | 6E-04 | | 2E-03 | Liver | 2 | | 0.8 | 3 |
| | | | Barium | | | | | | Heart | 0.2 | | | 0.2 |
| | | | Lead (1) | | | | | | | | | | |
| | | | Manganese | | | | | | Central Nervous System | 17 | | | 17 |
| | | | Chemical Total | 1E-03 | | 6E-04 | | 2E-03 | | 19 | | 0.9 | 20 |
| | | | | | | | | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | |
| | | Exposure Point Total | • | | | | | 2E-03 | | • | • | | 20 |
| | Exposure Medium To | tal | | | | | | 2E-03 | | | | | 20 |
| | Air | Water Vapors from | Bis(2-ethylhexyl)phthalate | | | | | | | | | | |
| | | Showerhead | Chloroform | | 1E-05 | | | 1E-05 | Liver | | 5 | | 5 |
| | | | Heptachlor | | 1E-03 | | | 1E-03 | | | | | |
| | | | Barium | | | | | | | | | | |
| | | | Lead (1) | | | | | | | | | | |
| | | | Manganese | | | | | | | | | | |
| | | | Chemical Total | | 1E-03 | | | 1E-03 | | | 5 | | 5 |
| | | | | | | | | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | |
| | | Exposure Point Total | - | | • | | • | 1E-03 | | | • | | 5 |
| | Exposure Medium To | · · · · · · · · · · · · · · · · · · · · | | | | | | 1E-03 | | | | | 5 |
| Groundwater To | tal | | | | | | | 3E-03 | | | | | 25 |

TABLE 9.1.RME SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCs REASONABLE MAXIMUM EXPOSURE The Dean Company

| Medium | Exposure Medium | Exposure Point | Chemical of Potential | | | Carcinogen | ic Risk | | | Non-Carcinoge | enic Hazard Quo | otient | |
|----------------|--------------------|----------------------|-----------------------|-----------|------------|------------|-------------|--------------|------------------------|---------------|-----------------|--------|--------------|
| | | | Concern | Ingestion | Inhalation | Dermal | External | Exposure | Primary | Ingestion | Inhalation | Dermal | Exposure |
| | | | | | | | (Radiation) | Routes Total | Target Organ(s) | | | | Routes Total |
| Soil | Soil | Soil at Site 1 | 4,4'-DDD | 5E-08 | | | | 5E-08 | | | | | |
| | | | 4,4'-DDE | 1E-06 | | | | 1E-06 | | | | | |
| | | | 4,4'-DDT | 5E-06 | | 5E-07 | | 6E-06 | Liver | 0.08 | | 0.009 | 0.09 |
| | | | Aluminum | | | | | | Central Nervous System | 0.01 | | | 0.01 |
| | | | Lead (1) | | | | | | | | | | |
| | | | Manganese | | | | | | Central Nervous System | 0.002 | | | 0.002 |
| | | | Chemical Total | 6E-06 | | 5E-07 | | 7E-06 | | 0.09 | | 0.009 | 0.1 |
| | | | | | | | | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | |
| | | Exposure Point Total | | | | | | 7E-06 | | | | | 0.1 |
| | | Soil at Site 2 | 4,4'-DDE | 8E-08 | | | Ï | 8E-08 | | | | | |
| | | | 4,4'-DDT | 5E-08 | | 6E-09 | | 6E-08 | Liver | 0.0009 | | 0.0001 | 0.001 |
| | | | Copper | | | | | | Gastrointestinal | 0.009 | | | 0.009 |
| | | | Iron | | | | | | Gastrointestinal | 0.1 | | | 0.1 |
| | | | Chemical Total | 1E-07 | | 6E-09 | | 1E-07 | | 0.1 | | 0.0001 | 0.1 |
| | | | | | | | | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | |
| | | Exposure Point Total | | | | | | 1E-07 | | | | | 0.1 |
| | Exposure Medium To | otal | | | | | | 7E-06 | | | | | 0.2 |
| Soil Total | | | | | | | | 7E-06 | | | | | 0.2 |
| Receptor Total | | | | | | | | 3E-03 | | | | | 26 |

| _ | | _ | |
|-------------------------------|-------|-----------------------------------|----|
| Total Risk Across All Media = | 3E-03 | Total Hazard Across All Media | 26 |
| _ | | | |
| | | Total Liver HI Across All Media = | 8 |

TABLE 9.1.RME

SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCs

REASONABLE MAXIMUM EXPOSURE

The Dean Company

| Medium | Exposure Medium | Exposure Point | Chemical of Potential | | | Carcinogen | ic Risk | | | Non-Carcinoge | enic Hazard Quo | otient | |
|-------------------|---------------------------|-------------------------------|-----------------------------------|------------------|-----------------|----------------|----------------------|--------------------------|-------------------------|----------------------|-----------------|-----------------|--------------------------|
| | | | Concern | Ingestion | Inhalation | Dermal | External (Radiation) | Exposure Routes Total | Primary Target Organ(s) | Ingestion | Inhalation | Dermal | Exposure Routes Total |
| (1) Lead is evalu | Lated for the resident us | I Ina the IEUBK model. See Ri | sk Assessment text for discussion | of results and a | ppendix for the | e lead modleii | | Noutes Total | | L Central Nervous | Svstem HI Acre | oss All Media = | |

TABLE 9.2.RME SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCS REASONABLE MAXIMUM EXPOSURE The Dean Company

| Medium | Exposure Medium | Exposure Point | Chemical of Potential | | | Carcinogen | ic Risk | | | Non-Carcinoge | enic Hazard Quo | otient | |
|----------------|--------------------|-----------------------|----------------------------|-----------|------------|------------|-------------|--------------|------------------------|---------------|-----------------|--------|--------------|
| | | | Concern | Ingestion | Inhalation | Dermal | External | Exposure | Primary | Ingestion | Inhalation | Dermal | Exposure |
| | | | | | | | (Radiation) | Routes Total | Target Organ(s) | | | | Routes Total |
| Groundwater | Groundwater | Aquifer 1 - Tap Water | Bis(2-ethylhexyl)phthalate | 4E-07 | | 4E-07 | | 8E-07 | Liver | 0.02 | | 0.02 | 0.04 |
| | | | Chloroform | 3E-07 | | 4E-07 | | 7E-07 | Liver | 0.06 | | 0.08 | 0.1 |
| | | | Heptachlor | 7E-04 | | 3E-04 | | 1E-03 | Liver | 4 | | 1 | 5 |
| | | | Barium | | | | | | Heart | 0.4 | | | 0.4 |
| | | | Lead (1) | | | | | | | | | | |
| | | | Manganese | | | | | | Central Nervous System | 40 | | | 40 |
| | | | Chemical Total | 7E-04 | | 3E-04 | | 1E-03 | | 44 | | 1 | 45 |
| | | | | | | | | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | |
| | | Exposure Point Total | | | | | | 1E-03 | | | | | 45 |
| | Exposure Medium To | tal | | | | | | 1E-03 | | | | | 45 |
| Groundwater To | tal | | | | | | | 1E-03 | | | | | 45 |
| Soil | Soil | Soil at Site 1 | 4,4'-DDD | 1E-07 | | | | 1E-07 | | | | | |
| | | | 4,4'-DDE | 3E-06 | | | | 3E-06 | | | | | |
| | | | 4,4'-DDT | 1E-05 | | 9E-07 | | 1E-05 | Liver | 0.7 | | 0.06 | 0.8 |
| | | | Aluminum | | | | | | Central Nervous System | 0.1 | | | 0.1 |
| | | | Lead (1) | | | | | | | | | | |
| | | | Manganese | | | | | | Central Nervous System | 0.02 | | | 0.02 |
| | | | Chemical Total | 1E-05 | | 9E-07 | | 1E-05 | | 0.8 | | 0.06 | 0.9 |
| | | | | | | | | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | |
| | | Exposure Point Total | | | | | | 1E-05 | | | | | 0.9 |

TABLE 9.2.RME

SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCs

REASONABLE MAXIMUM EXPOSURE

The Dean Company

| Medium | Exposure Medium | Exposure Point | Chemical of Potential | | | Carcinogen | ic Risk | | | Non-Carcinoge | nic Hazard Quo | otient | |
|------------------|-----------------------|----------------------|-----------------------|-----------|------------|------------|-------------|--------------|------------------|---------------|----------------|--------|--------------|
| | | | Concern | Ingestion | Inhalation | Dermal | External | Exposure | Primary | Ingestion | Inhalation | Dermal | Exposure |
| | | | | | | | (Radiation) | Routes Total | Target Organ(s) | | | | Routes Total |
| Soil (continued) | Soil (continued) | Soil at Site 2 | 4,4'-DDE | 2E-07 | | | | 2E-07 | | | | | |
| | | | 4,4'-DDT | 1E-07 | | 1E-08 | | 1E-07 | Liver | 0.008 | | 0.0007 | 0.008 |
| | | | Copper | | | | | | Gastrointestinal | 0.08 | | | 0.08 |
| | | | Iron | | | | | | Gastrointestinal | 1 | | | 1 |
| | | | Chemical Total | 3E-07 | | 1E-08 | | 3E-07 | | 1 | | 0.0007 | 1 |
| | | | | | | | | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | |
| | | Exposure Point Total | | | | | | 3E-07 | | | • | | 1 |
| | Exposure Medium Total | | | | | | | 1E-05 | | | • | | 2 |
| Soil Total | | | | | | | | 1E-05 | | | • | | 2 |
| Receptor Total | | | | | | | | 1E-03 | | | | | 47 |

| 47 | Total Hazard Across All Media | 1E-03 | Total Risk Across All Media = |
|----|--|-------|-------------------------------|
| | | | |
| 6 | Total Liver HI Across All Media = | | |
| 40 | Total Central Nervous System HI Across All Media = | | |
| 1 | Total Gastrointestinal HI Across All Media = | | |

TABLE 10.1.RME RISK SUMMARY REASONABLE MAXIMUM EXPOSURE The Dean Company

Scenario Timeframe: Future Receptor Population: Resident Receptor Age: Adult

| Medium | Exposure Medium | Exposure Point | Chemical | Carcinogenic Risk Non-Carcinogenic Hazard Quotient | | | | itient | | | | | |
|----------------------------|--------------------|-----------------------|--|--|--------------|----------------|----------------------|--------------------------|---------------------------------|----------------|------------|--------------|--------------------------|
| | | | | Ingestion | Inhalation | Dermal | External (Radiation) | Exposure Routes Total | Primary Target Organ(s) | Ingestion | Inhalation | Dermal | Exposure Routes Total |
| Groundwater | Groundwater | Aquifer 1 - Tap Water | Bis(2-ethylhexyl)phthalate Chloroform | 7E-07 5E-07 | | 1E-06 1E-06 | | 2E-06 2E-06 | Liver Liver | 0.007 0.03 | | 0.01 0.05 | 0.02 0.08 |
| | | | Heptachlor Manganese | 1E-03 | | 6E-04 | | 2E-03 | Liver Central Nervous System | 2 17 | | 0.8 | 3 17 |
| | | | Chemical Total | 1E-03 | | 6E-04 | | 2E-03 | | 19 | | 0.8 | 20 |
| | | Exposure Point Total | | | | | | 2E-03 | | | | | 20 |
| | Exposure Medium To | tal | | | | | | 2E-03 | | | | | 20 |
| | Air | Water Vapors from | Chloroform | | 1E-05 | | | 1E-05 | Liver | | 5 | | 5 |
| | | Showerhead | Heptachlor | | 1E-03 | | | 1E-03 | | | _ | | _ |
| | | Exposure Point Total | Chemical Total | | 1E-03 | | ļ. | 1E-03 1E-03 | | | 5 | | 5 |
| | Exposure Medium To | | | | | | | 1E-03 | | | | | 5 |
| Groundwater Tot | | tai | | | | | | 3E-03 | | | | | 25 |
| Soil | Soil | Soil at Site 1 | 4,4'-DDE | 1E-06 | | | | 1E-06 | | | | | 25 |
| Con | Oon | Son at Site 1 | 4,4'-DDT | 5E-06 | | 5E-07 | | 6E-06 | | | | | |
| | | | Chemical Total | 6E-06 | | 5E-07 | | 7E-06 | | | | | |
| | | Exposure Point Total | | | | | | 7E-06 | | | | | |
| | Exposure Medium To | tal | | | | | | 7E-06 | | | | | |
| Soil Total | | | | | | | | 7E-06 | | | | | |
| Receptor Total | | | | | | | | 3E-03 | | | | | 25 |
| Total Risk Across All Medi | | | | | ss All Media | 3E-03 | | T | Total Hazard Ac | ross All Media | 25 | | |

The information in this example table is for illustration only. The site screening threshold was determined by the RPM.

Total Liver HI Across All Media = 8

Total Central Nervous System HI Across All Media = 17

TABLE 10.2:RME RISK SUMMARY REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future
Receptor Population: Resident
Receptor Age: Child

| Medium | Exposure Medium | Exposure Point | · | | Carcinogenic Risk | | | | | Non-Carcinoge | enic Hazard Quo | otient | |
|-------------------------------|--------------------|-----------------------|----------------|-----------|-------------------|-----------------|----------------|--------------|------------------------|---------------|-----------------|--------|--------------|
| | | | | Ingestion | Inhalation | Dermal | External | Exposure | Primary | Ingestion | Inhalation | Dermal | Exposure |
| | | | | | | | (Radiation) | Routes Total | Target Organ(s) | | | | Routes Total |
| Groundwater | Groundwater | Aquifer 1 - Tap Water | Heptachlor | 7E-04 | | 3E-04 | | 1E-03 | Liver | 4 | | 1 | 5 |
| | | | Manganese | | | | | | Central Nervous System | 40 | | | 40 |
| | | | Chemical Total | 7E-04 | | 3E-04 | | 1E-03 | | 44 | | 1 | 45 |
| | | Exposure Point Total | | | | | | 1E-03 | | | | | 45 |
| | Exposure Medium To | tal | | | | | | 1E-03 | | | | | 45 |
| Groundwater Tot | al | | | | | | | 1E-03 | | | | | 45 |
| Soil | Soil | Soil at Site 1 | 4,4'-DDE | 3E-06 | | | | 3E-06 | | | | | |
| | | | 4,4'-DDT | 1E-05 | | 9E-07 | | 1E-05 | | | | | |
| | | | Chemical Total | 1E-05 | | 9E-07 | | 1E-05 | | | | | |
| | | Exposure Point Total | | | | | | 1E-05 | | | | | |
| | | Soil at Site 2 | Iron | | | | | | Gastrointestinal | 1 | | | 1 |
| | | | Chemical Total | | | | | | | 1 | | | 1 |
| | | Exposure Point Total | • | | • | | | | | • | • | | 1 |
| | Exposure Medium To | tal | | | | | | 1E-05 | | | | | 1 |
| Soil Total | | | | 1E-05 | | | | | 1 | | | | |
| Receptor Total | | | | | | | | 1E-03 | | | | | 46 |
| . Total Risk Across All Media | | | 1E-03 | | 7 | Total Hazard Ac | ross All Media | 46 | | | | | |

The information in this example table is for illustration only. The site screening threshold was determined by the RPM.

Total Liver HI Across All Media = 5

Total Central Nervous System HI Across All Media = 40

Total Gastrointestinal HI Across All Media = 1

DATA USEABILITY WORKSHEET

| Activity | Comment | | | | |
|---|--|--|--|--|--|
| Field Sampling | | | | | |
| Discuss sampling problems and field conditions that affect data useability. | Groundwater samples were collected from 12 monitoring wells located onsite. There were no apparent problems reported from the field collection program that could affect data useability. | | | | |
| Are samples representative of receptor exposure for this medium (e.g. sample depth, grab vs composite, filtered vs unfiltered, low flow, etc.)? | Groundwater samples submitted for organic and inorganic analyses were non-filtered samples collected using low flow purging and sampling techniques. These samples are representative of receptor exposure. | | | | |
| Assess the effect of field QC results on data useability. | A few of the metals in the samples were qualified "B" due to the presence of the metals in blank samples. | | | | |
| Summarize the effect of field sampling issues on the risk assessment, if applicable. | There are no field sampling issues that should affect the risk assessment. | | | | |
| Analytical | Techniques | | | | |
| Were the analytical methods appropriate for quantitative risk assessment? | Yes. Groundwater samples were analyzed for organic compounds according to Contract Laboratory Program (CLP) Statement of Work (SOW) for Organic Analysis, Multi-Media, Multi-Concentration, OLM04.2. Inorganic groundwater samples were analyzed according to CLP SOW for Inorganic Analysis, Multi-Media, Multi-Concentration, ILM04.1. | | | | |
| Were detection limits adequate? | Yes. The method detection and quantitation limit were less than the associated risk-based concentration (RBC) values, except for chloroform and thallium. For these two compounds, no available methods can achieve the RBC as a quantitation limit. For all non-detected chemicals in groundwater, the method detection and quantitation limits were less than the associated RBC values. Recommend no changes to the data set. | | | | |
| Summarize the effect of analytical technique issues on the risk assessment, if applicable. | There are no analytical technique issues that should affect the risk assessment. | | | | |

| Activity | Comment | | | | |
|--|--|--|--|--|--|
| Data Quality Objectives | | | | | |
| Precision - How were duplicates handled? | Relative percent differences (RPDs) were calculated for one pair of duplicate samples. The RPDs were less than the EPA-approved RPD of 20%. The highest concentration of a compound detected in the samples was used in the risk assessment. | | | | |
| Accuracy - How were split samples handled? | Split samples were not collected. | | | | |
| Representativeness - Indicate any problems associated with data representativeness (e.g., trip blank or rinsate blank contamination, chain of custody problems, etc.). | Analytes qualified with a "B" due to blank contamination will be considered as non-detects during the risk assessment. | | | | |
| Completeness - Indicate any problems associated with data completeness (e.g., incorrect sample analysis, incomplete sample records, problems with field procedures, etc.). | No problems were associated with data completeness. | | | | |
| Comparability - Indicate any problems associated with data comparability. | No problems have been associated with data comparability. | | | | |
| Were the DQOs specified in the QAPP satisfied? | Yes, the DQOs identified in the Sampling and Analysis Plan were satisfied. | | | | |
| Summarize the effect of DQO issues on the risk assessment, if applicable. | There are no DQO issues that should affect the risk assessment. | | | | |

| Activity | Comment | | | | |
|---|---|--|--|--|--|
| Data Validation and Interpretation | | | | | |
| What are the data validation requirements? | For organic samples, validators were required to check the following items: holding times, instrument performance checks, initial and continuing calibrations, blanks, system monitoring compounds, matrix spike/matrix spike duplicates, regional QA/QC, internal standards, target compound identification, contract required quantitation limits, tentatively identified compounds, system performance, and overall assessment of data. For inorganic samples, validators were required to check holding times, calibration, blanks, interference checks, laboratory control samples, duplicate samples, matrix spike samples, furnace atomic absorption QC, ICP Serial Dilution, sample result verification, field duplicates, and perform an overall assessment of the data. | | | | |
| What method or guidance was used to validate the data? | Region III modifications to "Laboratory Data Validation Functional Guidelines for Validating Organic (and Inorganic) Analyses", USEPA 9/94 (and 4/93). | | | | |
| Was the data validation method consistent with guidance? Discuss any discrepancies. | Yes. The data validation method was consistent with regional guidance. | | | | |
| Were all data qualifiers defined? Discuss those which were not. | Yes. All data qualifiers were defined. | | | | |
| Which qualifiers represent useable data? | B, J, L, U, UJ, and UL | | | | |
| Which qualifiers represent unuseable data? | R | | | | |
| How are tentatively identified compounds handled? | Only TICs that were determined not to be laboratory or field artifacts were reported. All TICs were reported with an "N" and/or a "J" qualifier. "N" qualified data indicates that the analyte is tentatively identified. "J" qualified data indicates that the analyte is present but reported value is estimated. TICs will be evaluated qualitatively in the risk assessment. | | | | |

| Activity | Comment |
|--|---|
| Summarize the effect of data validation and interpretation issues on the risk assessment, if applicable. | Unusable data qualified with an "R" will not be used in the risk assessment. All other data, both qualified and unqualified, will be used in the risk assessment. |
| Additional notes: | None. |

DATA USEABILITY WORKSHEET

The Dean Company Medium: Soil

| Activity | Comment | | | | | |
|---|---|--|--|--|--|--|
| Field Sampling | | | | | | |
| Discuss sampling problems and field conditions that affect data useability. | There were no apparent problems that could affect data useability. | | | | | |
| Are samples representative of receptor exposure for this medium (e.g. sample depth, grab vs composite, filtered vs unfiltered, low flow, etc.)? | Yes. Soil samples are representative of receptor exposure for this medium. | | | | | |
| Assess the effect of field QC results on data useability. | Overall, the trip, field, and rinsate blanks were generally non-detect for VOCs and SVOCs with the exception of low levels of commonly reported laboratory contaminants. Several of the metals in the samples were qualified "B" due to the presence of the metals in blank samples. | | | | | |
| Summarize the effect of field sampling issues on the risk assessment, if applicable. | There are no field sampling issues that should affect the risk assessment. | | | | | |
| Analytical | Techniques | | | | | |
| Were the analytical methods appropriate for quantitative risk assessment? | Yes. Samples were analyzed for organic compounds according to Contract Laboratory Program (CLP) Statement of Work (SOW) for Organic Analysis, Multi-Media, Multi-Concentration, OLM04.2. Inorganic soil samples were analyzed according to CLP SOW for Inorganic Analysis, Multi-Media, Multi-Concentration, ILM04.1. | | | | | |
| Were detection limits adequate? | Yes. The method detection and quantitation limit were less than the associated risk-based concentration (RBC) values. | | | | | |
| Summarize the effect of analytical technique issues on the risk assessment, if applicable. | There are no analytical technique issues that should affect the risk assessment. | | | | | |

The Dean Company Medium: Soil

| Activity | Comment | | | | |
|--|--|--|--|--|--|
| Data Quality Objectives | | | | | |
| Precision - How were duplicates handled? | Relative percent differences (RPDs) were calculated for one pair of duplicate samples. The RPDs were less than the EPA-approved RPD of 35%. The highest concentration of a compound detected in the samples was used in the risk assessment. | | | | |
| Accuracy - How were split samples handled? | Split samples were not collected. | | | | |
| Representativeness - Indicate any problems associated with data representativeness (e.g., trip blank or rinsate blank contamination, chain of custody problems, etc.). | Analytes qualified with a "B" due to blank contamination will be considered as non-detects during the risk assessment. | | | | |
| Completeness - Indicate any problems associated with data completeness (e.g., incorrect sample analysis, incomplete sample records, problems with field procedures, etc.). | No problems were associated with data completeness. | | | | |
| Comparability - Indicate any problems associated with data comparability. | No problems have been associated with data comparability. | | | | |
| Were the DQOs specified in the QAPP satisfied? | Yes, the DQOs identified in the Sampling and Analysis Plan were satisfied. | | | | |
| Summarize the effect of DQO issues on the risk assessment, if applicable. | There are no DQO issues that should affect the risk assessment. | | | | |

The Dean Company Medium: Soil

| Activity | Comment | | | | |
|---|---|--|--|--|--|
| Data Validation a | and Interpretation | | | | |
| What are the data validation requirements? | For organic samples, validators were required to check the following items: holding times, instrument performance checks, initial and continuing calibrations, blanks, system monitoring compounds, matrix spike/matrix spike duplicates, regional QA/QC, internal standards, target compound identification, contract required quantitation limits, tentatively identified compounds, system performance, and overall assessment of data. For inorganic samples, validators were required to check holding times, calibration, blanks, interference checks, laboratory control samples, duplicate samples, matrix spike samples, furnace atomic absorption QC, ICP serial dilution, sample result verification, field duplicates, and perform an overall assessment of the data. | | | | |
| What method or guidance was used to validate the data? | Region III modifications to "Laboratory Data Validation Functional Guidelines for Validating Organic (and Inorganic) Analyses", USEPA 9/94 (and 4/93). | | | | |
| Was the data validation method consistent with guidance? Discuss any discrepancies. | Yes. The data validation method was consistent with regional guidance. | | | | |
| Were all data qualifiers defined? Discuss those which were not. | Yes. All data qualifiers were defined. | | | | |
| Which qualifiers represent useable data? | B, J, K, L, U, UJ, and UL | | | | |
| Which qualifiers represent unuseable data? | R | | | | |
| How are tentatively identified compounds handled? | Only TICs that were determined not to be laboratory or field artifacts were reported. All TICs were reported with an "N" and/or a "J" qualifier. "N" qualified data indicates that the analyte is tentatively identified. "J" qualified data indicates that the analyte is present but the reported value is estimated. TICs will be evaluated qualitatively in the risk assessment. | | | | |

The Dean Company Medium: Soil

| Activity | Comment |
|--|---|
| Summarize the effect of data validation and interpretation issues on the risk assessment, if applicable. | Unusable data qualified with an "R" will not be used in the risk assessment. All other data, both qualified and unqualified, will be used in the risk assessment. |
| Additional notes: | None. |

EXAMPLE TECHNICAL APPROACH TO RISK ASSESSMENT (TARA) SCHEDULE WORKSHEET

The Dean Company

| Activity - RAGS Part D Reference ⁽¹⁾ | Comments ⁽²⁾ |
|---|-------------------------|
| PROJECT SCOPING | |
| Preliminary site conceptual model - Section 2.1 | November 30, 2000 |
| Site visit - Sec 2.1 | November 4, 2000 |
| Scoping meeting - Sec 2.1 | November 2, 2000 |
| PRGs and ARARs (initial discussion) - Sec 2.1 | November 2, 2000 |
| Identification of deliverables - Sec 2.1 | November 30, 2000 |
| Planning Table 1 (preliminary version) - Sec 2.1 | November 30, 2000 |
| Probabilistic Analysis (preliminary consideration) - Sec 2.1 | November 30, 2000 |
| RI/FS Workplan (consideration of risk assessment objectives) - Sec 2.2 | November 30, 2000 |
| Baseline Risk Assessment Workplan (consideration of risk assessment objectives) - Sec 2.2 | November 30, 2000 |
| Probabilistic Analysis (additional consideration and Workplan as appropriate) - Sec 2.2.1 | November 30, 2000 |
| REMEDIAL INVESTIGATION | |
| Planning Table 0 - Sec. 3.1.1 | August 30, 2001 |
| TARA Schedule Worksheet - Sec. 3.1.1 and Appendix C | August 30, 2001 |
| Planning Table 1 - Sec 3.1.1 | August 30, 2001 |
| Data Useability Worksheet - Sec 3.1.1 and Appendix C | August 30, 2001 |
| Supporting information for background value for Planning Table 2 - Sec 3.1.1 | August 30, 2001 |
| Planning Table 2 - Sec 3.1.1 | August 30, 2001 |
| Supporting information for EPC for Planning Table 3 - Sec 3.1.1 | August 30, 2001 |
| Planning Table 3 -Sec 3.1.1 | August 30, 2001 |

Notes:

¹Add other activities as appropriate for the site.

²Use this column to identify the applicability, schedule, and responsibility for each activity. Activities that are not required for a particular site can be noted as NA (not applicable). It is recommended that the responsibility and schedule for both the preparation and review of each activity be noted.

EXAMPLE TECHNICAL APPROACH TO RISK ASSESSMENT (TARA) SCHEDULE WORKSHEET

The Dean Company

| Activity - RAGS Part D Reference ⁽¹⁾ | Comments ⁽²⁾ | | | | | | |
|---|-------------------------|--|--|--|--|--|--|
| REMEDIAL INVESTIGATION (continued) | | | | | | | |
| Supporting information on modeled intake methodology and parameters for Planning Table 4 - Sec 3.1.1 | August 30, 2001 | | | | | | |
| Supporting information on chemical-specific parameters for Planning Table 4 - Sec 3.1.1 | August 30, 2001 | | | | | | |
| Dermal Worksheet - Sec 3.1.1 and Appendix C | August 30, 2001 | | | | | | |
| Planning Table 4 - Sec 3.1.1 | August 30, 2001 | | | | | | |
| Supporting information on toxicity data for special case chemicals on Planning Tables 5/6 - Sec 3.1.1 | August 30, 2001 | | | | | | |
| Planning Table 5 - Sec 3.1.1 | August 30, 2001 | | | | | | |
| Planning Table 6 - Sec 3.1.1 | August 30, 2001 | | | | | | |
| Supporting information on special chemical risk and hazard calculations for Planning Tables 7/8 - Sec 3.1.1 | October 21, 2001 | | | | | | |
| Planning Table 7 - Sec 3.1.1 | October 21, 2001 | | | | | | |
| Planning Table 8 - Sec. 3.1.1 | October 21, 2001 | | | | | | |
| Radiation Dose Assessment Worksheet - Sec 3.1.1 and Appendix C | October 21, 2001 | | | | | | |
| Planning Table 9 - Sec 3.1.1 | October 21, 2001 | | | | | | |
| Planning Table 10 - Sec 3.1.1 | October 21, 2001 | | | | | | |
| Lead Worksheets - Sec 3.1.1 and Appendix C | October 21, 2001 | | | | | | |
| Assessment of Confidence and Uncertainty - Sec 3.1.2 | October 21, 2001 | | | | | | |
| Summary of Probabilistic Analysis - Sec 3.1.3 | October 21, 2001 | | | | | | |
| Draft Baseline Risk Assessment - Sec 3.2 | October 21, 2001 | | | | | | |
| Final Baseline Risk Assessment - Sec 3.3 | January 15, 2001 | | | | | | |

Notes:

¹Add other activities as appropriate for the site.

²Use this column to identify the applicability, schedule, and responsibility for each activity. Activities that are not required for a particular site can be noted as NA (not applicable). It is recommended that the responsibility and schedule for both the preparation and review of each activity be noted.

EXAMPLE TECHNICAL APPROACH TO RISK ASSESSMENT (TARA) SCHEDULE WORKSHEET

The Dean Company

| Activity - RAGS Part D Reference ⁽¹⁾ | Comments ⁽²⁾ | | | | |
|---|-------------------------|--|--|--|--|
| REMEDIAL INVESTIGATION (continued) | • | | | | |
| Draft ROD Risk Worksheets - Sec 3.3 and Appendix C | January 15, 2001 | | | | |
| FEASIBILITY STUDY | | | | | |
| Remedial Action Objectives - Sec 4.2 | January 15, 2001 | | | | |
| Remediation Goals - Sec 4.2 | January 15, 2001 | | | | |
| Risks and hazards associated with PRGs - Sec 4.4 | January 15, 2001 | | | | |
| Risk considerations of remedial technologies and alternatives - Sec 4.5 | January 15, 2001 | | | | |
| AFTER THE FEASIBILITY STUDY | | | | | |
| Risk evaluation for the Proposed Plan - Sec 5.1 | To be determined | | | | |
| Documentation of risks in the Record of Decision - Sec 5.2 | To be determined | | | | |
| Revise ROD Risk Worksheets - Sec 5.2 and Appendix C | To be determined | | | | |
| Risk evaluation during remedial design and remedial action - Sec 5.3 | To be determined | | | | |
| Risk evaluation associated with explanations of significant differences - Sec 5.4 | To be determined | | | | |
| Risk evaluations during five-year review - Sec 5.5 | To be determined | | | | |
| Public meeting participation | To be determined | | | | |

Notes:

3 of 3 December 2001

¹Add other activities as appropriate for the site.

²Use this column to identify the applicability, schedule, and responsibility for each activity. Activities that are not required for a particular site can be noted as NA (not applicable). It is recommended that the responsibility and schedule for both the preparation and review of each activity be noted.

Dermal Worksheet Intermediate Variables for Calculating DA(event) The Dean Company

| Chemical of | Medium | Dermal Absorption | FA | Кр | | T(event) | | Tau | | T* | | В |
|-------------------|-------------|-------------------|---------|-----------|---------|----------|------------|---------|---------|---------|---------|---------|
| Potential Concern | | Fraction (soil) | Value | Value | Units | Value | Units | Value | Units | Value | Units | Value |
| phthalate | Groundwater | | 0.8 | 2.50E-002 | cm/hour | 0.58 | hour/event | 16.27 | hour | 39.05 | hour | 0.2 |
| Chloroform | Groundwater | | 1 | 1.50E-001 | cm/hour | 0.58 | hour/event | 0.49 | hour | 1.18 | hour | 0 |
| Heptachlor | Groundwater | | 0.8 | 8.70E-003 | cm/hour | 0.58 | hour/event | 12.99 | hour | 31.16 | hour | 0.1 |
| Barium * | Groundwater | | | | | | | | | | | |
| Manganese * | Groundwater | | | | | | | | | | | |
| Thallium * | Groundwater | | | | | | | | | | | |
| 4,4'-DDD * | Soil | | | | | | | | | | | |
| 4,4'-DDE * | Soil | | | | | | | | | | | |
| 4,4-DDT | Soil | 0.03 | No data | No data | No data | No data | No data | No data | No data | No data | No data | No data |
| Aluminum * | Soil | | | | | | | | | | | |
| Copper * | Soil | | | | | | | | | | | |
| Iron * | Soil | | | | | | | | | | | |
| Manganese * | Soil | | | | | | | | | | | |
| Thallium * | Soil | | | | - | | | | | | | |

FA = Fraction Absorbed Water

Kp = Dermal Permeability Coefficient of

Compound in Water

T(event) = Event Duration Tau = Lag Time T* = Time to Reach Steady-State

B = Dimensionless Ratio of the Permeability Coefficient of a Compound Through the Stratum Corneum Relative to its Permeability Coefficient Across the Viable Epidermis

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^{* =} Dermal assessment not recommended based on RAGS Part E, Appendix B-3 screening table.

TABLE X (RAGS D IEUBK LEAD WORKSHEET)

Site Name: <SITE and OU>

Receptor: <Receptor> (Age <X> Months) Exposure to Media as Described

1. Lead Screening Questions

| Medium | Lead Cond Used in M | | Basis for Lead Concentration Used | Lead Sc Concent | _ | Basis for Lead Screening | |
|--------|------------------------|-------|--------------------------------------|--------------------|-------|--|--|
| | Value | Units | For Model Run | Value | Units | Level | |
| Soil | <x></x> | mg/kg | Average Detected Value | 400 | mg/kg | Recommended Soil Screening Level | |
| Water | <x></x> | ug/L | Average Detected Value | 15 | ug/L | Recommended Drinking Water Action Level | |

2. Lead Model Questions

| Question | Response for Residential Lead Model |
|---|---|
| What lead model (version and date) was used? | <model> <version and="" date=""></version></model> |
| Where are the input values located in the risk assessment report? | Located in Appendix <x> <ieubkwin output=""></ieubkwin></x> |
| What range of media concentrations were used for the model? | <refer data="" sampling="" table="" to=""></refer> |
| What statistics were used to represent the exposure concentration terms and where are the data on concentrations in the risk assessment that support use of these statistics? | <statistic used=""> Data are Located in Appendix <x></x></statistic> |
| Was soil sample taken from top 2 cm? If not, why? | <yes no=""></yes> |
| Was soil sample sieved? What size screen was used? If not sieved, provide rationale. | <yes no=""> Mesh size <x> um</x></yes> |
| What was the point of exposure/location? | <describe></describe> |
| Where are the output values located in the risk assessment report? | Located in Appendix X <ieubkwin output=""></ieubkwin> |
| Was the model run using default values only? | <yes no=""></yes> |
| Was the default soil bioavailability used? | <yes no=""> Default is 30%</yes> |
| Was the default soil ingestion rate used? | <yes no=""> Default values for 7 age groups are 85, 135, 135, 100, 090, and 85 mg/day</yes> |
| If non-default values were used, where are the rationale for the values located in the risk assessment report? | Located in Appendix X <ieubkwin output=""></ieubkwin> |

3. Final Result

| Medium | Result | Comment/PRG 1 |
|-------------------|---|--|
| <medium></medium> | Input value of <x> (units) in <medium> results in YYY% of <receptor> above a blood lead level of 10 ug/dL. Geometric mean blood lead = ZZZ ug/dL. This exceeds the blood lead goal as described in the 1994 OSWER Directive of no more than 5% of children exceeding 10 ug/dL blood lead.</receptor></medium></x> | Based on site conditions, a PRG of X (units) is indicated for <medium>.</medium> |

 $^{1. \} Attach \ the \ IEUBK \ text \ output \ file \ and \ graph \ upon \ which \ the \ PRG \ was \ based \ as \ an \ appendix. \ For \ additional \ information, see \ \underline{www.epa.gov/superfund/programs/lead}$

TABLE Y (RAGS D ADULT LEAD WORKSHEET)

Site Name: Example Site, Slag Pile 2

Receptor: Adult Worker, Exposure to Media as Described

1. Lead Screening Questions

| Medium | Lead Concentration used in Model Run | | Basis for Lead Concentration Used | Lead Sc Concent | | Basis for Lead Screening Level | |
|--------|--------------------------------------|-------|--------------------------------------|--------------------|-------|----------------------------------|--|
| | Value | Units | For Model Run | Value | Units | Ü | |
| Soil | 2000 | mg/kg | Average Detected Value | 750 | mg/kg | Recommended Soil Screening Level | |

2. Lead Model Questions

| Question | Response |
|---|---|
| What lead model was used? Provide reference and version | EPA Interim Adult Lead Model (1996) |
| If the EPA Adult Lead Model (ALM) was not used provide rationale for model selected. | n/a |
| Where are the input values located in the risk assessment report? | Located in Appendix 5 |
| What statistics were used to represent the exposure concentration terms and where are the data on concentrations in the risk assessment that support use of these statistics? | Mean soil concentration. Data are Located in Appendix 2 |
| What was the point of exposure and location? | OU 3 Slag pile area |
| Where are the output values located in the risk assessment report? | Located in Appendix 5 |
| What GSD value was used? If this is outside the recommended range of 1.8-2.1, provide rationale in Appendix <y>.</y> | 1.8 |
| What baseline blood lead concentration (PbB $_0$) value was used? If this is outside the default range of 1.7 to 2.2 provide rationale in Appendix $<$ Y $>$. | 2.0 |
| Was the default exposure frequency (EF; 219 days/year) used? | Yes |
| Was the default BKSF used (0.4 ug/dL per ug/day) used? | Yes |
| Was the default absorption fraction (AF; 0.12) used? | Yes |
| Was the default soil ingestion rate (IR; 50 mg/day) used? | Yes |
| If non-default values were used for any of the parameters listed above, where are the rationale for the values located in the risk assessment report? | Located in Appendix 5 |

3. Final Result

| Medium | Result | Comment/RBRG 1 |
|--------|--|----------------|
| Soil | 2000 ppm lead in soil results in >5% of receptors above a blood lead level of 10 ug/d and geometric mean blood lead = 11.6 ug/dL. This exceeds the blood lead goal as described in the 1994 OSWER Directive of no more than 5% of children (fetuses of exposed women) exceeding 10 ug/dL blood lead. | 1500 ppm |

^{1.} Attach the ALM spreadsheet output file upon which the Risk Based Remediation Goal (RBRG) was based and description of rationale for parameters used. For additional information, see www.epa.gov/superfund/programs/lead

APPENDIX D

EXAMPLE SCENARIOS

- 1. Duplicate Exposure Information for Different Exposure Points
- 2. Modeled Inhalation from Showering
- 3. Measured Data and Subsequent Ingestion
- 4. Modeled Data and Subsequent Ingestion
- 5. Modeled Data
- **6.** Multiple Source Exposures
- 7. Possible Summing Options on Planning Tables 9 and 10
- 8. Child/Adult Lifetime Cancer Risk
- 9. Transfer of Contaminants Through Multiple Media
- 10. Lead Data Example
- 11. Radiation Data Example

Example Scenario No. 1 Duplicate Exposure Information for Different Exposure Points (with Planning Tables 1 and 4)

<u>Scenario Description</u>: Data are available for several exposure points that are to be evaluated separately in the risk assessment. In this risk assessment, data will be evaluated separately for ingestion and dermal contact from three different slag piles (Slag Piles 1, 2, and 3) for the same scenario timeframe, medium, and exposure medium.

Planning Table Issues Associated with this Scenario:

The primary issue with this scenario is whether or how to show the exposure points on Planning Tables 1 and 4. Note that the exposure parameter values used for daily intake calculations are identical for each individual pathway, i.e. the values presented on Planning Table 4 are the same for all exposure points for each type of exposure route.

1. How will Planning Table 1 show the three separate exposure points?

Planning Table 1 will need to show the three separate exposure points since each data set will be evaluated separately in the risk assessment. Planning Table 1 needs to show:

Medium: Solid Waste

Exposure Medium: Solid Waste Exposure Point: Slag Pile 1

Medium: Solid Waste

Exposure Medium: Solid Waste Exposure Point: Slag Pile 2

Medium: Solid Waste

Exposure Medium: Solid Waste Exposure Point: Slag Pile 3

2. Do the values used for daily intake calculations need to be shown three separate times on Planning Table 4 for each exposure point even though the values and intake equations are identical?

There are two options that can be followed:

Option 1: Complete Planning Table 4 according to the RAGS Part D instructions. For this example, Planning Table 4 would have three sets of identical values and intake equations, one for each exposure point.

Option 2: Complete Planning Table 4 using only one set of values and intake equations and indicate on the table that these values are identical for all three different exposure points. This can be accomplished by including "Slag Piles 1, 2, and 3" in the Exposure

Example Scenario No. 1 (continued) Duplicate Exposure Information for Different Exposure Points (with Planning Tables 1 and 4)

Point column and footnoting that these values and intake equations are the same for all three exposure points.

Option 1 is provided in the Example Tables in Appendix A. Option 2, consisting of a revised example Planning Table 4, is illustrated in the accompanying table.

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Example Scenario No. 2 Modeled Inhalation from Showering (with Planning Tables 1, 2, 3, 4, and 7)

<u>Scenario Description</u>: Individuals may be exposed to chemicals of potential concern in air by inhalation of chemicals through showering. The inhalation pathway is modeled using an EPA-accepted inhalation model. For this example scenario, a model accepted by EPA regions, such as the Foster and Chrostowski Shower Model, is used to evaluate *future adult resident inhalation exposure to groundwater*. See Example Scenario 4 for illustrations of how to present modeled data.

Planning Table Issues Associated with this Scenario:

1. How will use of an inhalation model affect Planning Table 1?

Planning Table 1 can accommodate this easily. Planning Table 1 can be completed to include an exposure medium (e.g., Water Vapors at Showerhead) and include the inhalation exposure route for all applicable scenarios. For this scenario example, Planning Table 1 would include a row that would describe this inhalation exposure pathway.

2. What data will be included in Planning Table 2 -- modeled air concentrations or measured groundwater concentrations?

In this example, Planning Table 2 will show measured groundwater concentrations. The data will be screened against tap water screening values.

- 3. What data will be included in Planning Table 3?
 - In this example, Planning Table 3 will show measured groundwater statistics.
- 4. How will the inhalation model parameters be shown on Planning Table 4?

For this example, the upper left hand corner Summary Box and the exposure route, receptor population, receptor age, and exposure point fields should be completed. However, exposure parameters and intake equations do not need to be entered into the table if there are space limitations. In the exposure route column, enter "Inhalation" with a footnote. Include the footnote explanation beneath the table that describes the model to be used and the section of the risk assessment text where information regarding modeled intake development can be found. Supporting information that summarizes the modeled intake methodology and parameters used to calculate modeled intake values should be included in the Baseline Risk Assessment Report as an attachment. Non-standard tables may also be used to display modeled information. Refer to the Risk Assessment text for details on the modeled intake methodology, the parameters used to calculate modeled intake values, and the modeled air concentrations predicted by the model.

Example Scenario No. 2 Modeled Inhalation from Showering (with Planning Tables 1, 2, 3, 4, and 7)

5. How are the modeled results displayed on Planning Table 7?

For this example, EPC values are calculated using measured groundwater data. They can be found on Planning Table 3. Intake/Exposure concentration values are values that are generated using the inhalation model. These values need to be included on this table. The risks and hazards will be calculated using the "Intake / Exposure concentration values" based on modeling and appropriate toxicity information.

Example Scenario No. 3 Measured Data and Subsequent Ingestion (Planning Tables 1, 2 and 3)

<u>Scenario Description</u>: Measured fish tissue data are available for evaluation in the risk assessment. The data are available for a specific species: trout. The measured data will be used in the risk assessment to determine the potential for adverse effects from ingestion of fish. This scenario is based upon fish tissue to show how to include measured data in the tables, but it can be applied to other exposure media.

Planning Table Issues Associated with this Scenario:

1. How will Planning Table 1 show fish tissue exposure?

In this situation, it is assumed that the source of exposure for the fish was the sediment, Planning Table 1 will need to show a specific exposure point for the trout as follows:

Medium: Sediment

Exposure Medium: Fish Tissue

Exposure Point: Trout

- 2. What data will be included in Planning Table 2 measured fish tissue data or sediment data? Planning Table 2 will show measured trout analytical data. The data will be screened against fish tissue screening values.
- 3. What data will be included in Planning Table 3?

 Planning Table 3 will show measured fish tissue statistics for the trout.

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Example Scenario No. 4 Modeled Data and Subsequent Ingestion (Planning Tables 1 and 2)

<u>Scenario Description</u>: Modeled fish tissue data are available for evaluation in the risk assessment based on concentrations of contaminants in the sediment. The modeled data will be used in the risk assessment to determine the potential for adverse effects from ingestion of the fish. This scenario is based upon fish tissue to show how to include modeled data in the tables, but it can be applied to other exposure media.

Planning Table Issues Associated with this Scenario:

The primary issue with this scenario is what data to show on Planning Table 2 and subsequent tables (modeled fish tissue or measured sediment data). There are two options for data presentation.

Option 1 (Modeled Fish Tissue Concentrations): The modeled fish tissue concentrations could appear on Planning Table 2 in the Concentration Used for Screening column. These modeled concentrations would be screened against fish tissue screening values. The methodology used to develop the modeled concentrations should be referenced on the tables. This option should be used when screening on fish tissue concentrations.

Option 2 (Measured Sediment Concentrations): Measured sediment concentrations could be presented on Planning Table 2. The measured concentrations are the values used as input in the model to determine predicted fish tissue concentrations. The modeling methodology could be discussed in the text and referenced on Planning Table 4. The model results would be used for intake calculations in Planning Table 7. This option should be used when screening on sediment concentrations.

1. How will Planning Table 1 show fish tissue exposure?

Assuming the source of exposure for the fish is sediment, Planning Table 1 will need to show a specific exposure point for the fish as follows:

Medium: Sediment

Exposure Medium: Fish Tissue

Exposure Point: Trout

2. What data will be included in Planning Table 2 - measured sediment data or modeled fish tissue data?

See discussion of options, above, and footnotes on Planning Table 2.

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Example Scenario No. 5 Modeled Data (Planning Table 1)

<u>Scenario Description:</u> The risk assessment uses data that have been modeled to evaluate potential risks. The modeling results are for spatial changes, temporal changes, and transfer between media.

Planning Table Issues Associated with this Scenario:

The issue associated with this scenario is how to identify and evaluate each different modeled data set. In this temporal change example, groundwater data have been modeled to represent concentrations in future years (1 year, 2 years, and 5 years in the future). This evaluation can be accommodated by assigning a separate exposure point to each future year.

1. How will Planning Table 1 be completed?

Planning Table 1 could show temporal changes using the exposure point column, as shown on the accompanying table.

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Example Scenario No. 6 Multiple Source Exposures (Planning Table 1)

<u>Scenario Description</u>: The risk assessment is evaluating the ingestion of fish tissue affected by both contaminated surface water <u>and</u> sediment.

<u>Planning Table Issues Associated with this Scenario:</u>

1. How will the medium, exposure medium, and exposure point be represented in Planning Table 1 for fish tissue?

The exposure point for fish tissue ingestion can be presented in two different ways, as described in the options below:

Option 1

Medium: Surface Water/Sediment Exposure Medium: Fish Tissue

Exposure Point: Trout - contaminant uptake from surface water and sediment

This option should be used if screening will be performed against measured or modeled fish tissue data.

Option 2

Medium: Surface Water Exposure Medium: Fish Tissue

Exposure Point: Trout - contaminant uptake from surface water

AND

Medium: Sediment

Exposure Medium: Fish Tissue

Exposure Point: Trout - contaminant uptake from sediment

This option should be used if screening will be performed against measured surface water or sediment data.

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Example Scenario No. 7 Possible Summing Options (Planning Tables 9 and 10)

<u>Scenario Description</u>: The risk assessment is evaluating several different exposure points for a particular set of media and exposure media. The EPA risk assessor for the site may allow the risk assessor to use abridged versions of Planning Tables 9 and 10 which do not require the same level of summation as the version of Planning Tables 9 and 10 shown in Appendix A.

<u>Planning Table Issues Associated with this Scenario:</u>

1. How will the risk data be summed on Planning Tables 9 and 10 for medium, exposure medium, exposure point, and receptor (combination of scenario timeframe, receptor population, and receptor age)?

The summing of risk for these exposure pathway elements can be presented in two different ways, as described in the options below. The EPA risk assessor will determine the type of summing that is appropriate for a particular site.

Option 1

Summing will occur in the standard fashion at four levels: medium, exposure medium, exposure point, and receptor.

Option 1 is shown in the accompanying tables and in Appendix A

Option 2

Summing will occur at fewer levels only: e.g., for exposure point and receptor only. Consult the EPA risk assessor to determine the appropriate procedure to follow. *Option 2 is shown in the accompanying tables*.

Example Scenario No. 8 Child/Adult Lifetime Cancer Risk (Planning Tables 1, 4, 7, 9)

<u>Scenario Description</u>: For this risk assessment the lifetime risk will be evaluated. Lifetime risk evaluates the combined risk from childhood through adulthood.

<u>Planning Table Issues Associated with this Scenario:</u>

In some regions, lifetime cancer risks are calculated by adding child and adult risk estimates together. In other regions, age-adjusted exposure factors are used to calculate lifetime cancer risk.

- 1. How should lifetime cancer risk be presented on Planning Table 1?

 For the "receptor age" column, choose from the picklist and enter "Adult", "Child", and "Child/Adult"
- 2. How should the other Planning Tables be completed? *Two options are presented:*

Option 1–Child/Adult calculated through summing cancer risks for separate Child and Adult receptors

Planning Tables 1, 4, and 7 would have separate Child and Adult receptor ages.

Planning Table 1 would also show a Child/Adult receptor to indicate that the Child/Adult analyses will be performed. Planning Table 4s would be developed for Child and Adult receptors with appropriate exposure factor values. A Planning Table 4 would also be shown for the Child/Adult receptor with no exposure factor values provided. Instead, a note would indicate that Child/Adult cancer risks will be calculated based upon the sum of Child cancer risk and Adult cancer risk.

Planning Table 7s and 9s would then be developed for three receptor ages: Child, Adult, and Child/Adult (a version of Planning Tables 7 and 9 combining the Child and the Adult cancer risk data into a single Child/Adult table with a note that the data on the table was derived from summing the Child and Adult data).

Option 2–Child/Adult calculated using age-adjusted exposure factors

As in Option 1, Planning Tables 1, 4, and 7 in Option 2 would show separate Child and Adult receptor ages as well as the Child/Adult receptor age. For the Option 2 Planning Table 4, the Child/Adult receptor age would be shown with age-adjusted exposure factor values. For the Option 2 Planning Tables 7 and 9, the Child/Adult cancer risks would be calculated using age-adjusted exposure factor values.

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Example Scenario No. 9 Transfer of Contaminants Through Multiple Media (Planning Table 1)

<u>Scenario Description:</u> The risk assessment evaluates the potential adverse effects from contaminants in soil that is taken up by plants and then taken up by an animal that is then ingested by human receptors.

Planning Table Issues Associated with this Scenario:

1. How can Planning Table 1 accommodate this three-way transfer?

Planning Table 1 can accommodate this scenario as follows:

Medium: Soil

Exposure Medium: Animal Tissue

Exposure Point: Beef from cattle grazing in field

This example scenario assumes that only the first and last media are of interest and no evaluation is needed for intermediate media. Consult with the EPA Risk Assessor to determine if screening is to be conducted on intermediate media (e.g., in an exposure scenario in which a contaminant moves from soil to plant tissue to animal tissue, whether an evaluation should be conducted for the intermediate plant tissue step).

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Example Scenario No. 10 Lead Data Example (Lead Worksheets)

<u>Scenario Description:</u> Lead is present in site soil and the child and adult lead models were used to evaluate blood lead levels. The standard tables do not accommodate lead model results.

Planning Table Issues Associated with this Scenario:

1. Since there are no standard tables that accommodate lead, how should lead results be presented?

The Lead Worksheets should be completed to demonstrate the evaluation performed and the results of analysis.

Examples of completed Lead Worksheets follow.

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Example Scenario No. 11 Radiation Data Example

<u>Scenario Description:</u> The site has radiological and chemical waste associated with it and radiological and chemical analyses were performed as part of the investigation. Potential adverse health effects will be evaluated in the risk assessment.

Planning Table Issues Associated with this Scenario:

Since radiological risk assessment uses different methodologies and terminologies than chemical risk assessment, how can the radiological risk assessment data be shown in the Planning Tables?

Planning Table 6.4 (Cancer Toxicity Data - External (Radiation)) and Planning Table 8 (Calculation of Radiation Cancer Risks) were developed by the Workgroup. The carcinogenic risk sections of Planning Tables 9 and 10 were expanded to include an External (Radiation) column. The following radiological risk example includes these Planning Tables.

Note: Many of the Example Planning Tables (i.e., those Example Planning Tables that do not specifically address radionuclides) provided for this Example Scenario are identical to those from Appendix A.

TABLE 1 SELECTION OF EXPOSURE PATHWAYS The Dean Company

| Scenario Timeframe | Medium | Exposure Medium | Exposure Point | Receptor Population | Receptor Age | Exposure Route | Type of Analysis | Rationale for Selection or Exclusion of Exposure Pathway |
|-----------------------|-------------|--------------------|-------------------|------------------------|-----------------|-------------------|---------------------|--|
| Future | Solid Waste | Solid Waste | Slag Pile 1 | Receptor Population | Age 1 | Ingestion | Quant | Rationale |
| | | | | | | Dermal | Quant | Rationale |
| | | | Slag Pile 2 | Receptor Population | Age 1 | Ingestion | Quant | Rationale |
| | | | | | | Dermal | Quant | Rationale |
| | | | Slag Pile 3 | Receptor Population | Age 1 | Ingestion | Quant | Rationale |
| | | | | | | Dermal | Quant | Rationale |

EXAMPLE SCENARIO 1 Option 2

TABLE 4.1.RME

VALUES USED FOR DAILY INTAKE CALCULATIONS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future Medium: Solid Waste

Exposure Medium: Solid Waste

| Exposure Route | Receptor Population | Receptor Age | Exposure Point | Parameter Code | Parameter Definition | Value | Units | Rationale/ | Intake Equation/ Model Name |
|----------------|---------------------|--------------|------------------------|-------------------|---|-------------------|--------------|-----------------------|---|
| | | | | i | | | <u> </u> | | |
| Ingestion | Receptor Population | Age 1 | Slag Piles 1, 2, 3 (1) | i | Chemical Concentration in Slag | See Table 3.1 | mg/kg | See Table 3.1 | Chronic Daily Intake (CDI) (mg/kg-day) = |
| | | | | IR | Ingestion Rate of Slag | 100 | mg/day | EPA, 1991 | CS x IR x FI x EF x ED x CF1 x 1/BW x 1/AT |
| | | | | FI | Fraction Ingested | 1 | | Professional Judgment | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 1991 | |
| | | | | ED | Exposure Duration | 24 | years | EPA, 1991 | |
| | | | | CF1 | Conversion Factor | 1E-06 | kg/mg | | |
| | | | | BW | Body Weight | 70 | kg | EPA, 1991 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 1989 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 8,760 | days | EPA, 1989 | |
| Dermal | Receptor Population | Age 1 | Slag Piles 1, 2, 3 (1) | cs | Chemical Concentration in Slag | See Table 3.1 | mg/kg | See Table 3.1 | Dermal Absorbed Dose (DAD) (mg/kg-day) = |
| | | | | CF1 | Conversion Factor | 1E-06 | kg/mg | | DA-event x EF x ED x EV x SA X 1/BW x 1/AT |
| | | | | SA | Skin Surface Area Available for Contact | 5,700 | cm2 | EPA, 2001 | where |
| | | | | AF | Soil to Skin Adherence Factor | 0.19 | mg/cm2-event | EPA, 2001 | Absorbed Dose per Event (DA-event) (mg/cm2-event) = |
| | | | | ABS-d | Absorption Factor | chemical-specific | unitless | EPA, 2001 | CS x CF1 x AF x ABS-d |
| | | | | EV | Event Frequency | 1 | events/day | EPA, 2001 | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 2001 | |
| | | | | ED | Exposure Duration | 24 | years | EPA, 1991 | |
| | | | | BW | Body Weight | 70 | kg | EPA, 2001 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 2001 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 8,760 | days | EPA, 2001 | |

⁽¹⁾ Parameters for Slag Piles 2 and 3 are identical to Slag Pile 1, and are therefore not repeated.

EPA 1989: Risk Assessment Guidance for Superfund. Volume 1: Human Health Evaluation Manual, Part A. OERR EPA/540/1-89/002.

EPA 1991: Risk Assessment Guidance for Superfund. Volume 1: Human Health Evaluation Manual - Supplemental Guidance, Standard Default Exposure Factors. Interim Final. OSWER 9285.6-03.

EPA 1995: Assessing Dermal Exposure from Soil, Technical Guidance Manual, Region III, EPA/903-K-95-003.

EPA 1997: Exposure Factors Handbook, Volume 1. EPA/600/P-95/002Fa.

EPA 2001: Risk Assessment Guidance for Superfund. Volume 1: Human Health Evaluation Manual (Part E, Supplemental Guidance for Dermal Risk Assessment) Interim.

NA = Not Available

TABLE 1

SELECTION OF EXPOSURE PATHWAYS

The Dean Company

| Scenario Timeframe | Medium | Exposure Medium | Exposure Point | Receptor Population | Receptor Age | Exposure Route | Type of Analysis | Rationale for Selection or Exclusion of Exposure Pathway |
|-----------------------|-------------|--------------------|-----------------------|------------------------|-----------------|-------------------|---------------------|--|
| Future | Groundwater | Groundwater | Aquifer 1 - Tap Water | Resident | Adult | Dermal | Quant | Future onsite residents may rely on domestic wells drawing from Aquifer 1. |
| | | | | | | Ingestion | Quant | Future onsite residents may rely on domestic wells drawing from Aquifer 1. |
| | | | | | Child | Dermal | Quant | Future onsite residents may rely on domestic wells drawing from Aquifer 1. |
| | | | | | | Ingestion | Quant | Future onsite residents may rely on domestic wells drawing from Aquifer 1. |
| | | Air | Water Vapors at | Resident | Adult | Inhalation | Quant | Future onsite residents may rely on domestic wells drawing from Aquifer 1. |
| | | | Showerhead | | Child | Inhalation | None | Children are assumed not to shower. |

TABLE 2.2

OCCURRENCE, DISTRIBUTION, AND SELECTION OF CHEMICALS OF POTENTIAL CONCERN

The Dean Company

Scenario Timeframe: Future

Medium: Groundwater

Exposure Medium: Air

| Exposure Point | CAS Number | Chemical | Minimum (1) Concentration (Qualifier) | Maximum (1) Concentration (Qualifier) | Units | Location of Maximum Concentration | Detection Frequency | Range of Detection Limits | Concentration Used for Screening (2) | Background Value (3) | Screening Toxicity Value (4) (N/C) | Potential ARAR/TBC Value | Potential ARAR/TBC Source | COPC Flag (Y/N) | Rationale for Selection or Deletion (5) |
|-------------------|---------------|----------------------------|---------------------------------------|---------------------------------------|-------|---|------------------------|---------------------------------|--------------------------------------|-------------------------|------------------------------------|--------------------------------|---------------------------------|-----------------------|---|
| Water Vapors | 117817 | Bis(2-ethylhexyl)phthalate | 2 J | 5 J | ug/l | GW3D | 4/12 | 7 - 11 | 5 | NA | 4.8 C | 6 | MCL | Υ | ASL |
| at | 67663 | Chloroform | 0.6 J | 9 | ug/l | GW3D | 3/12 | 1 - 1 | 9 | NA | 0.063 C | 100 | MCL | Υ | ASL |
| Showerhead | 75150 | Carbon Disulfide | 0.3 J | 4.5 | ug/l | GW3D | 3/12 | 1 - 1 | 4.5 | NA | 100 N | NA | NA | N | BSL |
| | 76448 | Heptachlor | 2 J | 33 J | ug/l | GW4D | 6/12 | 0.05 - 0.05 | 33 | NA | 0.015 C | 0.4 | MCL | Υ | ASL |
| | 108883 | Toluene | 0.1 J | 0.2 J | ug/l | GW3D | 3/12 | 1 - 1 | 0.2 | NA | 75 N | 1000 | MCL | N | BSL |

(1) Measured groundwater concentrations.

(2) Maximum concentration used for screening.

(3) To date, no background study has been completed.

 $\hbox{ (4) All compounds are screened against the Risk-Based Concentration (RBC) Table, U.S. EPA Region III, } \\$

October 5, 2000 for tap water (cancer benchmark = 1E-06; HQ = 0.1).

(5) Rationale Codes:

Selection Reason: Above Screening Level (ASL)
Deletion Reason: Below Screening Level (BSL)

Definitions: NA = Not Applicable

COPC = Chemical of Potential Concern

ARAR/TBC = Applicable or Relevant and Appropriate Requirement/To Be Considered

MCL = Maximum Contaminant Level

J = Estimated Value
C = Carcinogen

N = Noncarcinogen

TABLE 3.2.RME

EXPOSURE POINT CONCENTRATION SUMMARY

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future

Medium: Groundwater

Exposure Medium: Air

| Exposure Point | Chemical of Potential Concern | Units | Arithmetic Mean | 95% UCL (N/T) | Maximum Concentration (Qualifier) | Exposure P Value Units | | e Point Concentration Statistic | Rationale |
|-----------------|-------------------------------|-------|--------------------|------------------|-----------------------------------|-------------------------|------|---------------------------------|--------------|
| | | | | | | | | | |
| Water Vapors at | Bis(2-ethylhexyl)phthalate | ug/l | 4 | 5.5 T | 5 J | 5 | ug/l | Max | W-Test (1) |
| Showerhead | Chloroform | ug/l | 1.9 | 14.9 T | 9 | 9 | ug/l | Max | W-Test (1) |
| | Heptachlor | ug/l | 27 | 30 T | 33 J | 30 | ug/l | 95% UCL - T | W - Test (2) |

Note: Measured groundwater concentrations used to calculate EPC values.

Statistics: Maximum Detected Value (Max); 95% UCL of Transformed Data (95% UCL - T)

(1) 95% UCL exceeds maximum detected concentration. Therefore, maximum concentration used for EPC.

(2) Shapiro-Wilk W Test indicates data are lognormally transformed.

N = Normal

T = Transformed

J = Estimated Value

TABLE 4.2.RME

VALUES USED FOR DAILY INTAKE CALCULATIONS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future Medium: Groundwater

Exposure Medium: Air

| Exposure Route | Receptor Population | Receptor Age | Exposure Point | Parameter | Parameter Definition | Value | Units | Rationale/ | Intake Equation/ | |
|----------------|---------------------|--------------|-----------------|-----------|----------------------|-------|-------|------------|------------------------------|--|
| | | | | Code | | | | Reference | Model Name | |
| Inhalation (1) | Resident | Adult | Water Vapors at | (1) | (1) | (1) | (1) | (1) | Foster and Chrostowski Model | |
| | | | Showerhead | | | | | | | |

⁽¹⁾ Refer to the Risk Assessment text for details on the modeled intake methodology, the parameters used to calculate modeled intake values, and the modeled air concentrations predicted by the Foster and Chrostowski Shower Model.

TABLE 7.1.RME

CALCULATION OF CHEMICAL CANCER RISKS AND NON-CANCER HAZARDS REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future
Receptor Population: Resident
Receptor Age: Adult

| Medium | Exposure Medium | Exposure Point | Exposure Route | Chemical of | Е | PC | | Cancer R | isk Calculati | ons | | | Non-Cance | r Hazard Ca | alculations | |
|-------------|-----------------|-----------------------|------------------|----------------------------|-------|----------|---------------|------------------|---------------|---------------|-------------|----------------|-------------------|--------------|--------------|-----------------|
| | | | | Potential Concern | Value | Units | Intake/Exposu | re Concentration | CSF/ | Unit Risk | Cancer Risk | Intake/Exposur | e Concentration | RfD |)/RfC | Hazard Quotient |
| | | | | | | | Value | Units | Value | Units | | Value | Units | Value | Units | |
| Groundwater | Groundwater | Aquifer 1 - Tap Water | Ingestion | Bis(2-ethylhexyl)phthalate | 0.005 | mg/l | 4.7E-005 | mg/kg/day | 1.4E-002 | 1/mg/kg/day | 7E-007 | 1.4E-004 | mg/kg/day | 2.0E-002 | mg/kg/day | 0.007 |
| | | | | Chloroform | 0.009 | mg/l | 8.5E-005 | mg/kg/day | 6.1E-003 | 1/mg/kg/day | 5E-007 | 2.5E-004 | mg/kg/day | 1.0E-002 | mg/kg/day | 0.03 |
| | | | | Heptachlor | 0.03 | mg/l | 2.8E-004 | mg/kg/day | 4.5E+000 | 1/mg/kg/day | 1E-003 | 8.1E-004 | mg/kg/day | 5.0E-004 | mg/kg/day | 2 |
| | | | Exp. Route Total | ĺ | | | | | | | 1E-003 | | | | | 2 |
| | | | Dermal | Bis(2-ethylhexyl)phthalate | 0.005 | mg/l | 3.9E-006 | mg/kg/day | 2.5E-002 | 1/mg/kg/day | 1E-007 | 1.1E-005 | mg/kg/day | 1.1E-002 | mg/kg/day | 0.001 |
| | | | | Chloroform | 0.009 | mg/l | 1.9E-006 | mg/kg/day | 6.1E-003 | 1/mg/kg/day | 1E-008 | 5.5E-006 | mg/kg/day | 1.0E-002 | mg/kg/day | 0.0006 |
| | | | | Heptachlor | 0.03 | mg/l | 7.6E-006 | mg/kg/day | 9.0E+000 | 1/mg/kg/day | 7E-005 | 2.2E-005 | mg/kg/day | 2.5E-004 | mg/kg/day | 0.09 |
| | | | Exp. Route Total | | | | | | | | 7E-005 | | | | | 0.09 |
| | | Exposure Point Total | • | • | | • | | • | • | • | 1E-003 | | • | • | | 2 |
| | Air | Water Vapors at | Inhalation | Bis(2-ethylhexyl)phthalate | 0.005 | mg/l (1) | 2.3E-006 | mg/kg/day | NA | NA | NA | 3.6E-006 | mg/kg/day | NA | NA | NA |
| | | Showerhead | | Chloroform | 0.009 | mg/l (1) | 1.3E-004 | mg/kg/day | 8.1E-002 | 1/mg/kg/day | 1E-005 | 3.9E-004 | mg/kg/day | 8.6E-005 | mg/kg/day | 5 |
| | | | | Heptachlor | 0.03 | mg/l (1) | 2.6E-004 | mg/kg/day | 4.5E+000 | 1/mg/kg/day | 1E-003 | 7.7E-004 | mg/kg/day | NA | NA | NA |
| | | | Exp. Route Total | | | | | | | | 1E-003 | | | | | 5 |
| | | Exposure Point Total | | | • | • | | • | • | | 1E-003 | | • | | | 5 |
| | | | | | • | | | Total of Recept | or Risks Acr | oss All Media | 2E-003 | Tot | al of Receptor Ha | azards Acros | ss All Media | 7 |

⁽¹⁾ EPC values are shown as measured groundwater values and are found on Table 3.2.RME.

TABLE 1 SELECTION OF EXPOSURE PATHWAYS The Dean Company

| Scenario Timeframe | Medium | Exposure Medium | Exposure Point | Receptor Population | Receptor Age | Exposure Route | Type of Analysis | Rationale for Selection or Exclusion of Exposure Pathway |
|-----------------------|----------|--------------------|-------------------|------------------------|-----------------|-------------------|---------------------|--|
| Future | Sediment | Sediment | Pond 1 | Receptor Population | Age 1 | Route 1 | Quant | Rationale |
| | | | | | | Route 2 | Quant | Rationale |
| | | | | | Age 2 | Route 1 | Quant | Rationale |
| | | | | | | Route 2 | Quant | Rationale |
| | | Fish Tissue | Trout | Receptor Population | Age 1 | Route 1 | Quant | Rationale |
| | | | | | Age 2 | Route 1 | Quant | Rationale |

TABLE 2.1

OCCURRENCE, DISTRIBUTION, AND SELECTION OF CHEMICALS OF POTENTIAL CONCERN

The Dean Company

Scenario Timeframe: Future

Medium: Sediment

Exposure Medium: Fish Tissue

| Exposure Point | CAS Number | Chemical | Minimum (1) Concentration (Qualifier) | Maximum (1) Concentration (Qualifier) | Units | Location of Maximum Concentration | Detection Frequency | Range of Detection Limits | Concentration Used for Screening (1) | Background Value (2) | Screening Toxicity Value (3) (N/C) | Potential ARAR/TBC Value | Potential ARAR/TBC Source | COPC Flag (Y/N) | Rationale for Selection or Deletion (4) |
|-------------------|---------------|----------------------------------|---------------------------------------|---------------------------------------|-------|---|------------------------|---------------------------------|--------------------------------------|-------------------------|--|--------------------------------|---------------------------------|-----------------------|---|
| Trout | 11096825 | Arochlor 1260 | 0.0002 J | 0.005 J | mg/kg | Trout - 1 | 3 / 10 | 0.0001 - 0.0001 | 0.005 | NA | 0.0016 C | NA | NA | Y | ASL |
| | 7439921 | Lead | 0.004 J | 0.007 J | mg/kg | Trout - 3 | 5 / 10 | 0.001 - 0.001 | 0.007 | NA | NA | NA | NA | Υ | NTX |
| | 1746016 | 2,3,7,8-Tetrachlorodibenzodioxin | 0.00000001 J | 0.00000005 J | mg/kg | Trout - 1 | 4 / 10 | 0.00000001 - 0.00000001 | 0.0000005 | NA | 0.000000021 C | NA | NA | Υ | ASL |

(1) Measured fish tissue concentrations. Maximum measured fish tissue concentrations used for screening.

(2) Background values are not available.

(3) All compounds were screened against the Risk-Based Concentration (RBC) Table, U.S. EPA Region III, May 8, 2001 for fish tissue (cancer benchmark = 1E-06; HQ = 0.1).

(4) Rationale Codes:

Selection Reason: Above Screening Level (ASL)

No Toxicity Infomation (NTX)

Definitions: NA = Not Applicable

COPC = Chemical of Potential Concern

ARAR/TBC = Applicable or Relevant and Appropriate Requirement/To Be Considered

J = Estimated Value

C = Carcinogen

N = Noncarcinogen

TABLE 3.1.RME

EXPOSURE POINT CONCENTRATION SUMMARY

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future

Medium: Sediment

Exposure Medium: Fish Tissue

| Exposure Point | Chemical of | Units | Arithmetic | 95% UCL | Maximum Concentration | | Exposure | Point Concentration | |
|----------------|----------------------------------|----------------|----------------|-------------------------------|--------------------------|------------------|----------------|----------------------------|------------------------------|
| | Potential Concern | | Mean | (N/T) | (Qualifier) | Value | Units | Statistic | Rationale |
| Trout | Arochlor 1260 Lead | mg/kg mg/kg | 0.003 0.005 | 0.0035 (T) 0.0063 (T) | 0.005 J 0.007 J | 0.0035 0.0063 | mg/kg mg/kg | 95% UCL - T 95% UCL - T | W - Test (1) W - Test (1) |
| | 2,3,7,8-Tetrachlorodibenzodioxin | mg/kg | 0.00000002 | 0.00000(T) 0.000000047 (T) | 0.00000005 J | 0.00000047 | mg/kg | 95% UCL -T | W - Test (1) |

Statistics: 95% UCL of Transformed Data (95% UCL - T)

(1) Shapiro-Wilk W Test indicates data are log-normally distributed.

Note: Measured fish tissue concentrations used to calculate EPC values.

N = Normal

T = Transformed

J = Estimated Value

The Dean Company

TABLE 1 SELECTION OF EXPOSURE PATHWAYS

| Scenario Timeframe | Medium | Exposure Medium | Exposure Point | Receptor Population | Receptor Age | Exposure Route | Type of Analysis | Rationale for Selection or Exclusion of Exposure Pathway |
|-----------------------|----------|--------------------|-------------------|------------------------|-----------------|-------------------|---------------------|--|
| Timeframe | Sediment | Fish Tissue | Trout | Population 1 | Age 1 | Route 1 | Quant | Rationale |
| | | | | | Age 2 | Route 1 | Quant | Rationale |
| | | | | Population 2 | Age 1 | Route 1 | Quant | Rationale |
| | | | | | Age 2 | Route 1 | Quant | Rationale |

EXAMPLE SCENARIO 4 Option 1

TABLE 2.1

OCCURRENCE, DISTRIBUTION, AND SELECTION OF CHEMICALS OF POTENTIAL CONCERN

The Dean Company

Scenario Timeframe: Future Medium: Sediment Exposure Medium: Fish Tissue

| Exposure Point | CAS Number | Chemical | Minimum Concentration (1) (Qualifier) | Maximum Concentration (1) (Qualifier) | Units | Location of Maximum Concentration | Detection Frequency | Range of Detection Limits | Concentration Used for Screening (2) | Background Value (3) | Screening Toxicity Value (4) (N/C) | Potential ARAR/TBC Value | Potential ARAR/TBC Source | COPC Flag (Y/N) | Rationale for Selection or Deletion (5) |
|-------------------|---------------|--|---------------------------------------|---------------------------------------|----------------|---|------------------------|---------------------------------|--------------------------------------|-------------------------|------------------------------------|--------------------------------|---------------------------------|-----------------------|---|
| Trout | | Arochlor 1260 | 0.6 J | 5.5 J | mg/kg | SD01 | 3 / 10 | 0.1 - 0.2 | 0.005 | NA | 0.0016 (C) | NA | NA | Y | ASL |
| | 1400021 | Lead 2,3,7,8-Tetrachlorodibenzodioxin | 210 J 0.000001 J | 500 J 0.00005 J | mg/kg mg/kg | SD03 SD01 | 5 / 10 4 / 10 | 10 - 16 0.000001 - 0.000001 | 0.007 0.00000005 | NA NA | NA 0.000000021 (C) | NA NA | NA NA | Y | NTX ASL |

- (1) Measured sediment concentrations.
- (2) Concentrations used for screening are fish tissue values derived from the X model. Refer to the risk assessment text for details on the model methodology.
- (3) To date, no background study has been completed.
- (4) All compounds were screened against the Risk-Based Concentration (RBC) Table, U.S. EPA Region III,

May 8, 2001 for fish tissue (cancer benchmark = 1E-06; HQ = 0.1).

(5) Rationale Codes:

Selection Reason:

Above Screening Level (ASL)

No Toxicity Infomation (NTX)

EXAMPLE SCENARIO 4 Option 2

TABLE 2.1

OCCURRENCE, DISTRIBUTION, AND SELECTION OF CHEMICALS OF POTENTIAL CONCERN

The Dean Company

Scenario Timeframe: Future
Medium: Sediment
Exposure Medium: Fish Tissue

| Exposure Point | CAS Number | Chemical | Minimum Concentration (1) (Qualifier) | Maximum Concentration (1) (Qualifier) | Units | Location of Maximum Concentration | Detection Frequency | Range of Detection Limits | Concentration Used for Screening (1) | Background Value (2) | Screening Toxicity Value (3) (N/C) | Potential ARAR/TBC Value | Potential ARAR/TBC Source | COPC Flag (Y/N) | Rationale for Selection or Deletion (4) |
|-------------------|---------------|---|---------------------------------------|---------------------------------------|-------------------------|---|----------------------------|---|--------------------------------------|-------------------------|------------------------------------|--------------------------------|---------------------------------|-----------------------|---|
| Trout | 7439921 | Arochlor 1260 Lead 2,3,7,8-Tetrachlorodibenzodioxin | 0.6 J 210 J 0.000001 J | 5.5 J 500 J 0.00005 J | mg/kg mg/kg mg/kg | SD01 SD03 SD01 | 3 / 10 5 / 10 4 / 10 | 0.1 - 0.2 10 - 16 0.000001 - 0.000001 | 5.5 500 0.00005 | NA NA NA | 3.2 (C) 400 0.000043 (C) | NA NA NA | NA NA NA | Y Y Y | ASL ASL ASL |

(1) Measured sediment concentrations are shown and maximum concentrations are used for screening. These data will be used as input in the X model to predict fish tissue concentrations. Refer to the risk assessment text for details on the model methodology.

(2) To date, no background study has been completed.

(3) All compounds were screened against the Risk-Based Concentration (RBC) Table, U.S. EPA Region III, May 8, 2001 for 10 times the residential soil value (cancer benchmark = 10 x 1E-06; HQ = 10 x 0.1). Lead was screened against the U.S. EPA screening value of 400 mg/kg.

(4) Rationale Codes:

Selection Reason: Above Screening Level (ASL)

Definitions: NA = Not Applicable

COPC = Chemical of Potential Concern

ARAR/TBC = Applicable or Relevant and Appropriate Requirement/To Be Considered

J = Estimated Value C = Carcinogen

N = Noncarcinogen

TABLE 1 SELECTION OF EXPOSURE PATHWAYS Site Name

| Scenario Timeframe | Medium | Exposure Medium | Exposure Point | Receptor Population | Receptor Age | Exposure Route | Type of Analysis | Rationale for Selection or Exclusion of Exposure Pathway |
|-----------------------|-------------|--------------------|--|------------------------|-----------------|-------------------|---------------------|--|
| Future | Groundwater | Groundwater | Groundwater - Modeled 1 year into the future | Resident | Adult | Ingestion | Quant | Rationale |
| | | | | | | Dermal | Quant | Rationale |
| | | | Groundwater - Modeled 2 Years into the Future | Resident | Adult | Ingestion | Quant | Rationale |
| | | | | | | Dermal | Quant | Rationale |
| | | | Groundwater - Modeled 5 Years into the Future | Resident | Adult | Ingestion | Quant | Rationale |
| | | | | | | Dermal | Quant | Rationale |

EXAMPLE SCENARIO 6 OPTION 1

TABLE 1

SELECTION OF EXPOSURE PATHWAYS

The Dean Company

| Scenario Timeframe | Medium | Exposure Medium | Exposure Point | Receptor Population | Receptor Age | Exposure Route | Type of Analysis | Rationale for Selection or Exclusion of Exposure Pathway |
|-----------------------|------------------------|--------------------|---|------------------------|-----------------|-------------------|---------------------|--|
| Future | Surface Water/Sediment | Fish Tissue | TroutContaminant Uptake from Surface Water and Sediment | Receptor Population | Age 1 | Ingestion | | Rationale |
| | | | | | Age 2 | Ingestion | Quant | Rationale |

EXAMPLE SCENARIO 6 OPTION 2

TABLE 1

SELECTION OF EXPOSURE PATHWAYS

The Dean Company

| Scenario Timeframe | Medium | Exposure Medium | Exposure Point | Receptor Population | Receptor Age | Exposure Route | Type of Analysis | Rationale for Selection or Exclusion of Exposure Pathway |
|-----------------------|---------------|--------------------|--|------------------------|-----------------|-------------------|---------------------|--|
| Future | Surface Water | Fish Tissue | TroutContaminant Uptake from Surface Water | Receptor Population | Age 1 | Ingestion | Quant | Rationale |
| | | | | | Age 2 | Ingestion | Quant | Rationale |
| | Sediment | Fish Tissue | TroutContaminant Uptake from Sediment | Receptor Population | Age 1 | Ingestion | Quant | Rationale |
| | | | | | Age 2 | Ingestion | Quant | Rationale |

Option 1

TABLE 9.1.RME

SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCs REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future
Receptor Population: Resident
Receptor Age: Adult

| Medium | Exposure Medium | Exposure Point | Chemical of Potential | | | Carcinogen | c Risk | | | Non-Carcinoge | enic Hazard Quo | | |
|------------------|---------------------|-----------------------|----------------------------|-----------|------------|------------|-------------|--------------|-----------------|---------------|-----------------|--------|--------------|
| | | | Concern | Ingestion | Inhalation | Dermal | External | Exposure | Primary | Ingestion | Inhalation | Dermal | Exposure |
| | | | | | | | (Radiation) | Routes Total | Target Organ(s) | | | | Routes Total |
| Groundwater | Groundwater | Aquifer 1 - Tap Water | Bis(2-ethylhexyl)phthalate | 7E-07 | | 1E-07 | | 8E-07 | Liver | 0.007 | | 0.001 | 0.008 |
| | | | Chloroform | 5E-07 | | 1E-08 | | 5E-07 | Liver | 0.03 | | 0.0006 | 0.03 |
| | | | Chemical Total | 1E-06 | | 1E-07 | | 1E-06 | | 0.03 | | 0.002 | 0.04 |
| | | | | | | | | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | |
| | | Exposure Point Total | • | | • | | • | 1E-06 | | • | • | | 0.04 |
| | Exposure Medium Tot | tal | | | | | | 1E-06 | | | | | 0.04 |
| | Air | Water Vapors from | Bis(2-ethylhexyl)phthalate | | 3E-08 | | | 3E-08 | | | | | |
| | | Showerhead | Chloroform | | 1E-05 | | | 1E-05 | Liver | | 5 | | 5 |
| | | | Chemical Total | | 1E-05 | | | 1E-05 | | | 5 | | 5 |
| | | | | | | | | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | |
| | | Exposure Point Total | | | • | | | 1E-05 | | | | | 5 |
| | Exposure Medium Tot | tal | | | | | | 1E-05 | | | | | 5 |
| Groundwater Tota | al | · | | · | | | 1E-05 | | | | · | 5 | |

Option 1

TABLE 9.1.RME

SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCs

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future
Receptor Population: Resident
Receptor Age: Adult

| Medium | Exposure Medium | Exposure Point | Chemical of Potential | | | Carcinogen | ic Risk | | | Non-Carcinoge | enic Hazard Quo | tient | |
|----------------|--------------------|----------------------|---------------------------------------|-----------|------------|------------|-------------|--------------|-----------------|---------------|-----------------|--------|--------------|
| | | | Concern | Ingestion | Inhalation | Dermal | External | Exposure | Primary | Ingestion | Inhalation | Dermal | Exposure |
| | | | | | | | (Radiation) | Routes Total | Target Organ(s) | | | | Routes Total |
| Soil | Soil | Soil at Site 1 | 4,4'-DDE | 1E-06 | | 1E-06 | | 2E-06 | | | | | |
| | | | 4,4'-DDT | 5E-06 | | 5E-006 | | 1E-005 | Liver | 0.08 | | 0.08 | 0.2 |
| | | | Chemical Total | 6E-06 | | 6E-06 | | 1E-05 | | 0.08 | | 0.08 | 0.2 |
| | | | | | | | | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | |
| | | Exposure Point Total | • | | • | • | · | 1E-05 | | • | • | | 0.2 |
| | | Soil at Site 2 | 4,4'-DDE | 8E-08 | | 8E-08 | 1 | 2E-07 | | | | | |
| | | | 4,4'-DDT | 5E-08 | | 5E-08 | | 1E-07 | Liver | 0.0009 | | 0.0009 | 0.002 |
| | | | Chemical Total | 1E-07 | | 1E-07 | | 3E-07 | | 0.0009 | | 0.0009 | 0.002 |
| | | | | | | | | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | |
| | | Exposure Point Total | · · · · · · · · · · · · · · · · · · · | | | | | 3E-07 | | | | | 0.002 |
| | Exposure Medium To | tal | | | | | | 1E-05 | | | | | 0.002 |
| Soil Total | Total | | | | | | | 1E-05 | | | | | 0.002 |
| Receptor Total | | | _ | | | • | | 2E-05 | | | | • | 5 |

| Total Risk Across All Media | 2E-05 | Total Hazard Across All Media | 5 |
|-----------------------------|-------|-----------------------------------|---|
| | | | |
| | | Total Liver HI Across All Media = | 5 |

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Option 2

TABLE 9.1.RME

SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCs

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future
Receptor Population: Resident
Receptor Age: Adult

| Medium | Exposure Medium | Exposure Point | Chemical of Potential | | | Carcinogen | ic Risk | | Non-Carcinogenic Hazard Quotient | | | | |
|-------------|----------------------|-----------------------|----------------------------|-----------|------------|------------|----------------------|--------------------------|----------------------------------|-----------|------------|--------|--------------------------|
| | | | Concern | Ingestion | Inhalation | Dermal | External (Radiation) | Exposure Routes Total | Primary Target Organ(s) | Ingestion | Inhalation | Dermal | Exposure Routes Total |
| Groundwater | Groundwater | Aquifer 1 - Tap Water | Bis(2-ethylhexyl)phthalate | 7E-07 | | 1E-07 | | 8E-07 | Liver | 0.007 | | 0.001 | 0.008 |
| | | | Chloroform | 5E-07 | | 1E-08 |] | 5E-07 | Liver | 0.03 | | 0.0006 | 0.03 |
| | | | Chemical Total | 1E-06 | | 1E-07 | | 1E-06 | | 0.03 | | 0.002 | 0.04 |
| | | | | | | | | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | |
| | | Exposure Point Total | | | | | | 1E-06 | | | | | 0.04 |
| | Air | Water Vapors from | Bis(2-ethylhexyl)phthalate | | 3E-08 | | | 3E-08 | | | | | |
| | | Showerhead | Chloroform | | 1E-05 | | | 1E-05 | Liver | | 5 | | 5 |
| | | | Chemical Total | | 1E-05 | | | 1E-05 | | | 5 | | 5 |
| | | | | | | |] | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | |
| | | Exposure Point Total | | | | | | 1E-05 | | | | | 5 |
| Soil | Soil | Soil at Site 1 | 4,4'-DDE | 1E-06 | | 1E-06 | | 2E-06 | | | | | |
| | | | 4,4'-DDT | 5E-06 | | 5E-006 | | 1E-005 | Liver | 0.08 | | 0.08 | 0.2 |
| | | | Chemical Total | 6E-06 | | 6E-06 | | 1E-05 | | 0.08 | | 0.08 | 0.2 |
| | | | | | | | | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | |
| | | Exposure Point Total | | | 1 | | | 1E-05 | | 1 | | 1 | 0.2 |
| | | Soil at Site 2 | 4,4'-DDE | 8E-08 | | 8E-08 | | 2E-07 | | | | | |
| | | | 4,4'-DDT | 5E-08 | | 5E-08 | | 1E-07 | Liver | 0.0009 | | 0.0009 | 0.002 |
| | | | Chemical Total | 1E-07 | | 1E-07 | | 3E-07 | | 0.0009 | | 0.0009 | 0.002 |
| | | | | | | | | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | |
| | Exposure Point Total | | | | | | | 3E-07 | | | | | 0.002 |

| Total Risk Across All Media | 2E-05 | Total Hazard Across All Media = | 5 |
|-----------------------------|-------|-----------------------------------|---|
| | | | |
| | | Total Liver HI Across All Media = | 5 |

Option 1

TABLE 10.1.RME
RISK SUMMARY
REASONABLE MAXIMUM EXPOSURE
The Dean Company

Scenario Timeframe: Future Receptor Population: Resident Receptor Age: Adult

| Medium | Exposure Medium | Exposure Point | Chemical of Potential | | | Carcinogen | c Risk | | Non-Carcinogenic Hazard Quotient | | | | |
|----------------------|-----------------------|---------------------------------|-----------------------|-----------|------------|------------|-------------|--------------|----------------------------------|-----------|------------|--------|--------------|
| | | | Concern | Ingestion | Inhalation | Dermal | External | Exposure | Primary | Ingestion | Inhalation | Dermal | Exposure |
| | | | | | | | (Radiation) | Routes Total | Target Organ(s) | | | | Routes Total |
| Groundwater | Air | Water Vapors from Showerhead | Chloroform | | 1E-05 | | | 1E-05 | Liver | | 5 | | 5 |
| | | | Chemical Total | | 1E-05 | | | 1E-05 | | | 5 | | 5 |
| | | | | | | | | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | |
| | Exposure Point Total | | | | | • | | 1E-05 | | | | | 5 |
| | Exposure Medium To | tal | | | | | | 1E-05 | | | | | 5 |
| Groundwater Tota | al | | | | | | | 1E-05 | | | | | 5 |
| Soil | Soil | Soil at Site 1 | 4,4'-DDE | 1E-06 | | 1E-06 | | 2E-06 | | | | | |
| | | | 4,4'-DDT | 5E-06 | | 5E-06 | | 1E-05 | | | | | |
| | | | Chemical Total | 6E-06 | | 6E-06 | | 1E-05 | | | | | |
| | | | | | | | | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | |
| Exposure Point Total | | | | | | | 1E-05 | | | | | | |
| | Exposure Medium Total | | | | | | | | | | | | |
| Soil Total | Soil Total | | | | | | | 1E-05 | | | | | |
| Receptor Total | | • | | | | • | | 2E-05 | | | | | 5 |

Total Risk Across All Media 2E-05 Total Hazard Across All Media 5

Cancer risks presented are those greater than 1E-06; Non-cancer risks presented are those greater than 1.

Total Liver HI Across All Media = 5

Option 2

TABLE 10.1.RME
RISK SUMMARY
REASONABLE MAXIMUM EXPOSURE
The Dean Company

Scenario Timeframe: Future
Receptor Population: Resident
Receptor Age: Adult

| Medium | Exposure Medium | Exposure Point | Chemical of Potential | | | Carcinogen | ic Risk | | Non-Carcinogenic Hazard Quotient | | | | |
|-------------|----------------------|---------------------------------|-----------------------|-----------|------------|------------|----------------------|--------------------------|----------------------------------|-----------|------------|--------|--------------------------|
| | | | Concern | Ingestion | Inhalation | Dermal | External (Radiation) | Exposure Routes Total | Primary Target Organ(s) | Ingestion | Inhalation | Dermal | Exposure Routes Total |
| Groundwater | Air | Water Vapors from Showerhead | Chloroform | | 1E-05 | | | 1E-05 | Liver | | 5 | | 5 |
| | | | Chemical Total | | 1E-05 | | | 1E-05 | | | 5 | | 5 |
| | | | Radionuclide Total | | | | | | | | | | |
| | | Exposure Point Total | · | | | • | | 1E-05 | | • | • | • | 5 |
| Soil | Soil | Soil at Site 1 | 4,4'-DDE | 1E-06 | | 1E-06 | | 2E-06 | | | | | |
| | | | 4,4'-DDT | 5E-06 | | 5E-006 | | 1E-005 | | | | | |
| | | | Chemical Total | 6E-06 | | 6E-06 | | 1E-05 | | | | | |
| | | | | | | | | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | |
| | Exposure Point Total | | | | • | • | | 1E-05 | | • | • | | |

Total Risk Across All Media

Total Hazard Across All Media = 5

Cancer risks presented are those greater than 1E-06; Non-cancer risks presented are those greater than 1.

Total Liver HI Across All Media = 5

EXAMPLE SCENARIO 8 Option 1

TABLE 1

SELECTION OF EXPOSURE PATHWAYS

The Dean Company

| Scenario Timeframe | Medium | Exposure Medium | Exposure Point | Receptor Population | Receptor Age | Exposure Route | Type of Analysis | Rationale for Selection or Exclusion of Exposure Pathway |
|-----------------------|--------|--------------------|-------------------|------------------------|-----------------|-------------------|---------------------|--|
| Future | Soil | Soil | Soil at Site 1 | Resident | Adult | Dermal | Quant | Future onsite residents may come into contact with soil. |
| | | | | | | Ingestion | Quant | Future onsite residents may ingest soil. |
| | | | | | Child | Dermal | Quant | Future onsite residents may come into contact with soil. |
| | | | | | | Ingestion | Quant | Future onsite residents may ingest soil. |
| | | | | | Child/Adult | Dermal | Quant | Future onsite residents may come into contact with soil. |
| | | | | | | Ingestion | Quant | Future onsite residents may ingest soil. |

EXAMPLE SCENARIO 8 Option 2

TABLE 1

SELECTION OF EXPOSURE PATHWAYS

The Dean Company

| Scenario Timeframe | Medium | Exposure Medium | Exposure Point | Receptor Population | Receptor Age | Exposure Route | Type of Analysis | Rationale for Selection or Exclusion of Exposure Pathway |
|-----------------------|--------|--------------------|-------------------|------------------------|-----------------|-------------------|---------------------|--|
| Future | Soil | Soil | Soil at Site 1 | Resident | Adult | Dermal | Quant | Future onsite residents may come into contact with soil. |
| | | | | | | Ingestion | Quant | Future onsite residents may ingest soil. |
| | | | | | Child | Dermal | Quant | Future onsite residents may come into contact with soil. |
| | | | | | | Ingestion | Quant | Future onsite residents may ingest soil. |
| | | | | | Child/Adult | Dermal | Quant | Future onsite residents may come into contact with soil. |
| | | | | | | Ingestion | Quant | Future onsite residents may ingest soil. |

TABLE 4.1.RME

VALUES USED FOR DAILY INTAKE CALCULATIONS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future

Medium: Soil

Exposure Medium: Soil

| | | | | 1 | | ı | 1 | | |
|----------------|---------------------|--------------|----------------|-------------------|--------------------------------|---------------|-----------|-------------------------|---|
| Exposure Route | Receptor Population | Receptor Age | Exposure Point | Parameter Code | Parameter Definition | Value | Units | Rationale/ Reference | Intake Equation/ Model Name |
| Ingestion | Resident | Adult | Soil at Site 1 | CS | Chemical Concentration in Soil | See Table 3.3 | mg/kg | See Table 3.3 | Chronic Daily Intake (CDI) (mg/kg-day) = |
| | | | | IR | Ingestion Rate of Soil | 100 | mg/day | EPA, 1991 | CS x IR x FI x EF x ED x CF1 x 1/BW x 1/AT |
| | | | | FI | Fraction Ingested | 1 | | Professional Judgment | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 1991 | |
| | | | | ED | Exposure Duration | 24 | years | EPA, 1991 | |
| | | | | CF1 | Conversion Factor | 1E-06 | kg/mg | | |
| | | | | BW | Body Weight | 70 | kg | EPA, 1991 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 1989 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 8,760 | days | EPA, 1989 | |
| | | Child | Soil at Site 1 | cs | Chemical Concentration in Soil | See Table 3.3 | mg/kg | See Table 3.3 | CDI (mg/kg-day) = |
| | | | | IR | Ingestion Rate of Soil | 200 | mg/day | EPA, 1991 | CS x IR x FI x EF x ED x CF1 x 1/BW x 1/AT |
| | | | | FI | Fraction Ingested | 1 | | Professional Judgment | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 1991 | |
| | | | | ED | Exposure Duration | 6 | years | EPA, 1991 | |
| | | | | CF1 | Conversion Factor | 1E-06 | kg/mg | | |
| | | | | BW | Body Weight | 15 | kg | EPA, 1991 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 1989 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 2,190 | days | EPA, 1989 | |
| | | Child/Adult | Soil at Site 1 | | | | | | Child/Adult cancer risks will be calculated as the sum of the Child cancer risk and the Adult cancer risk. |

TABLE 4.1.RME

VALUES USED FOR DAILY INTAKE CALCULATIONS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future

Medium: Soil

Exposure Medium: Soil

| | 1 | | | 1 | | ı | | ı | |
|----------------|---------------------|--------------|----------------|-------------------|---|-------------------|-----------|-------------------------|---|
| Exposure Route | Receptor Population | Receptor Age | Exposure Point | Parameter Code | Parameter Definition | Value | Units | Rationale/ Reference | Intake Equation/ Model Name |
| Dermal | Resident | Adult | Soil at Site 1 | cs | Chemical Concentration in Soil | See Table 3.3 | mg/kg | See Table 3.3 | CDI (mg/kg-day) = |
| | | | | CF1 | Conversion Factor | 1E-06 | kg/mg | | CS x CF1 x SA x AF x AB x EF x ED x 1/BW x 1/AT |
| | | | | SA | Skin Surface Area Available for Contact | 5,000 | cm2 | EPA, 1997 | |
| | | | | AF | Soil to Skin Adherence Factor | 0.19 | mg/cm2 | EPA, 1997 | |
| | | | | AB | Absorption Factor | chemical-specific | unitless | EPA, 1995 | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 1991 | |
| | | | | ED | Exposure Duration | 24 | years | EPA, 1991 | |
| | | | | BW | Body Weight | 70 | kg | EPA, 1991 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 1989 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 8,760 | days | EPA, 1989 | |
| | | Child | Soil at Site 1 | cs | Chemical Concentration in Soil | See Table 3.3 | mg/kg | See Table 3.3 | CDI (mg/kg-day) = |
| | | | | CF1 | Conversion Factor | 1E-06 | kg/mg | | CS x CF1 x SA x AF x AB x EF x ED x 1/BW x 1/AT |
| | | | | SA | Skin Surface Area Available for Contact | 3,600 | cm2 | EPA, 1997 | |
| | | | | AF | Soil to Skin Adherence Factor | 0.11 | mg/cm2 | EPA, 1997 | |
| | | | | AB | Absorption Factor | chemical-specific | unitless | EPA, 1995 | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 1991 | |
| | | | | ED | Exposure Duration | 6 | years | EPA, 1991 | |
| | | | | BW | Body Weight | 15 | kg | EPA, 1991 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 1989 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 2,190 | days | EPA, 1989 | |
| | | Child/Adult | Soil at Site 1 | | | | | | Child/Adult cancer risks will be calculated as the sum of the Child cancer risk and the Adult cancer risk. |

EPA 1989: Risk Assessment Guidance for Superfund. Volume 1: Human Health Evaluation Manual, Part A. OERR EPA/540/1-89/002.

EPA 1991: Risk Assessment Guidance for Superfund. Volume 1: Human Health Evaluation Manual - Supplemental Guidance, Standard Default Exposure Factors. Interim Final. OSWER 9285.6-03.

EPA 1995: Assessing Dermal Exposure from Soil, Technical Guidance Manual, Region III, EPA/903-K-95-003.

EPA 1997: Exposure Factors Handbook, Volume 1. EPA/600/P-95/002Fa.

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TABLE 4.1.RME

VALUES USED FOR DAILY INTAKE CALCULATIONS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future

Medium: Soil

| edium: Soil | | | | | | | | | |
|---------------------|---------------------|--------------|----------------|-----------|--------------------------------|---------------|----------------|-----------------------|---|
| posure Medium: Soil | | | | | | | | | |
| Exposure Route | Receptor Population | Receptor Age | Exposure Point | Parameter | Parameter Definition | Value | Units | Rationale/ | Intake Equation/ |
| | | | | Code | | | | Reference | Model Name |
| Ingestion | Resident | Adult | Soil at Site 1 | CS | Chemical Concentration in Soil | See Table 3.3 | mg/kg | See Table 3.3 | Chronic Daily Intake (CDI) (mg/kg-day) = |
| | | | | IR | Ingestion Rate of Soil | 100 | mg/day | EPA, 1991 | CS x IR x FI x EF x ED x CF1 x 1/BW x 1/AT |
| | | | | FI | Fraction Ingested | 1 | | Professional Judgment | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 1991 | |
| | | | | ED | Exposure Duration | 24 | years | EPA, 1991 | |
| | | | | CF1 | Conversion Factor | 1.0E-06 | kg/mg | | |
| | | | | BW | Body Weight | 70 | kg | EPA, 1991 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 1989 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 8,760 | days | EPA, 1989 | |
| | | Child | Soil at Site 1 | CS | Chemical Concentration in Soil | See Table 3.3 | mg/kg | See Table 3.3 | CDI (mg/kg-day) = |
| | | | | IR | Ingestion Rate of Soil | 200 | mg/day | EPA, 1991 | CS x IR x FI x EF x ED x CF1 x 1/BW x 1/AT |
| | | | | FI | Fraction Ingested | 1 | | Professional Judgment | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 1991 | |
| | | | | ED | Exposure Duration | 6 | years | EPA, 1991 | |
| | | | | CF1 | Conversion Factor | 1.0E-06 | kg/mg | | |
| | | | | BW | Body Weight | 15 | kg | EPA, 1991 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 1989 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 2,190 | days | EPA, 1989 | |
| | | Child/Adult | Soil at Site 1 | CS | Chemical Concentration in Soil | See Table 3.3 | mg/kg | See Table 3.3 | CDI (mg/kg/day) = |
| | | | | IF | Ingestion Factor | 114 | mg-year/kg-day | EPA 1991b | CS x IF x CF x FI x EF x 1/AT |
| | | | | BW-C | Body Weight, Child | 15 | kg | EPA, 1991a | where |
| | | | | BW-A | Body Weight, Adult | 70 | kg | EPA, 1991a | IF = (ED-C x IR-C / BW-C) + (ED-TOT - ED-C) > |
| | | | | IR-C | Ingestion Rate, Child | 200 | mg/day | EPA, 1991a | (IR-A / BW-A) |
| | | | | IR-A | Ingestion Rate, Adult | 100 | mg/day | EPA, 1991a | |
| | | | | ED-C | Exposure Duration, Child | 6 | years | EPA, 1991a | |
| | | | | ED-TOT | Exposure Duration, Total | 30 | years | EPA, 1991a | |
| | | | | CF | Conversion Factor | 1.0E-06 | kg/mg | | |
| | | | | FI | Fraction Ingested | 1 | unitless | Professional Judgment | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 1991a | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 1989 | |

TABLE 4.1.RME

VALUES USED FOR DAILY INTAKE CALCULATIONS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future

Medium: Soil

| nealann. Con | | | | | | | | | |
|-----------------------|---------------------|--------------|----------------|-------------------|---|-------------------|-----------|---------------|---|
| Exposure Medium: Soil | | | | | | | | | |
| Exposure Route | Receptor Population | Receptor Age | Exposure Point | Parameter Code | Parameter Definition | Value | Units | Rationale/ | Intake Equation/ Model Name |
| Dermal | Resident | Adult | Soil at Site 1 | CS | Chemical Concentration in Soil | See Table 3.3 | mg/kg | See Table 3.3 | CDI (mg/kg-day) = |
| | | | | CF1 | Conversion Factor | 1.0E-06 | kg/mg | | CS x CF1 x SA x AF x AB x EF x ED x 1/BW x 1/AT |
| | | | | SA | Skin Surface Area Available for Contact | 5,000 | cm2 | EPA, 1997 | |
| | | | | AF | Soil to Skin Adherence Factor | 0.19 | mg/cm2 | EPA, 1997 | |
| | | | | AB | Absorption Factor | chemical-specific | unitless | EPA, 1995 | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 1991 | |
| | | | | ED | Exposure Duration | 24 | years | EPA, 1991 | |
| | | | | BW | Body Weight | 70 | kg | EPA, 1991 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 1989 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 8,760 | days | EPA, 1989 | |
| | | Child | Soil at Site 1 | CS | Chemical Concentration in Soil | See Table 3.3 | mg/kg | See Table 3.3 | CDI (mg/kg-day) = |
| | | | | CF1 | Conversion Factor | 1.0E-06 | kg/mg | | CS x CF1 x SA x AF x AB x EF x ED x 1/BW x 1/AT |
| | | | | SA | Skin Surface Area Available for Contact | 3,600 | cm2 | EPA, 1997 | |
| | | | | AF | Soil to Skin Adherence Factor | 0.11 | mg/cm2 | EPA, 1997 | |
| | | | | AB | Absorption Factor | chemical-specific | unitless | EPA, 1995 | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 1991 | |
| | | | | ED | Exposure Duration | 6 | years | EPA, 1991 | |
| | | | | BW | Body Weight | 15 | kg | EPA, 1991 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 1989 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 2,190 | days | EPA, 1989 | |

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TABLE 4.1.RME

VALUES USED FOR DAILY INTAKE CALCULATIONS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future

Medium: Soil

Exposure Medium: Soil

| Expodure mediam. Con | | | | | | | | | |
|----------------------|----------------------|--------------|----------------|-------------------|--------------------------------|-------------------|-----------------|-----------------------|---|
| Exposure Route | Receptor Population | Receptor Age | Exposure Point | Parameter Code | Parameter Definition | Value | Units | Rationale/ | Intake Equation/ Model Name |
| Dermal (continued) | Resident (continued) | Child/Adult | Soil at Site 1 | CS | Chemical Concentration in Soil | See Table 3.3 | mg/kg | See Table 3.3 | CDI (mg/kg-day) = |
| | | | | DF | Dermal Factor | 3,154 | cm2-year/kg-day | EPA 1991b | CS x CF1 x DF x AF x AB x EF x 1/AT |
| | | | | BW-C | Body Weight, Child | 15 | kg | EPA, 1991a | where |
| | | | | BW-A | Body Weight, Adult | 70 | kg | EPA, 1991a | DF = (ED-C x SA-C / BW-C) + (ED-TOT - ED-C) x |
| | | | | SA-C | Surface Area, Child | 3,600 | cm2 | EPA, 1997 | (SA-A / BW-A) |
| | | | | SA-A | Surface Area, Adult | 5,000 | cm2 | EPA, 1997 | |
| | | | | ED-C | Exposure Duration, Child | 6 | years | EPA, 1991a | |
| | | | | ED-TOT | Exposure Duration, Total | 30 | years | EPA, 1991a | |
| | | | | AF | Soil to Skin Adherence Factor | 0.15 | mg/cm2 | Professional Judgment | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA 1991a | |
| | | | | AB | Absorption Factor | chemical-specific | unitless | EPA, 1995 | |
| | | | | CF1 | Conversion Factor | 1.0E-06 | kg/mg | | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 1989 | |

EPA 1989: Risk Assessment Guidance for Superfund. Volume 1: Human Health Evaluation Manual, Part A. OERR EPA/540/1-89/002.

EPA 1997: Exposure Factors Handbook, Volume 1. EPA/600/P-95/002Fa.

EPA 1991a: Risk Assessment Guidance for Superfund. Volume 1: Human Health Evaluation Manual - Supplemental Guidance, Standard Default Exposure Factors. Interim Final. OSWER 9285.6-03.

EPA 1991b: Human Health Evaluation Manual, Part B: Development of Risk-Based Preliminary Remediation Goals. OSWER Directive 9285.7-01B

EPA 1995: Assessing Dermal Exposure from Soil, Technical Guidance Manual, Region III, EPA/903-K-95-003.

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TABLE 7.1.RME

CALCULATION OF CHEMICAL CANCER RISKS AND NON-CANCER HAZARDS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future
Receptor Population: Resident
Receptor Ane: Adult

| Medium | Exposure Medium | Exposure Point | Exposure Route | Chemical of | Е | PC | | Ca | ncer Risk Calculati | ons | | | Non- | Cancer Hazard Cal | culations | |
|------------|-----------------------|----------------------|------------------|-------------------|-------|-------|----------------|-----------------|---------------------|------------------|-------------|----------------|-----------------|-------------------|------------------|-----------------|
| | | | | Potential Concern | Value | Units | Intake/Exposur | e Concentration | CSF/L | nit Risk | Cancer Risk | Intake/Exposur | e Concentration | RfD |)/RfC | Hazard Quotient |
| | | | | | | | Value | Units | Value | Units | | Value | Units | Value | Units | |
| Soil | Soil | Soil at Site 1 | Ingestion | 4,4'-DDD | 0.452 | mg/kg | 2.1E-07 | mg/kg/day | 2.4E-01 | 1/mg/kg/day | 5E-08 | 6.2E-07 | mg/kg/day | NA | NA | NA |
| | | | İ | 4,4'-DDE | 6.8 | mg/kg | 3.2E-06 | mg/kg/day | 3.4E-01 | 1/mg/kg/day | 1E-06 | 9.3E-06 | mg/kg/day | NA | NA | NA |
| | | | | 4,4'-DDT | 28.6 | mg/kg | 1.3E-005 | mg/kg/day | 3.4E-01 | 1/mg/kg/day | 5E-06 | 3.9E-05 | mg/kg/day | 5.0E-04 | mg/kg/day | 0.08 |
| | | | | Aluminum | 9964 | mg/kg | 4.7E-003 | mg/kg/day | NA | NA | NA | 1.4E-02 | mg/kg/day | 1.0E+00 | mg/kg/day | 0.01 |
| | | | | Manganese | 201 | mg/kg | 9.5E-005 | mg/kg/day | NA | NA | NA | 2.8E-04 | mg/kg/day | 1.4E-01 | mg/kg/day | 0.002 |
| | | | | Thallium | 1.2 | mg/kg | 5.6E-007 | mg/kg/day | NA | NA | NA | 1.6E-06 | mg/kg/day | NA | NA | NA |
| | | | Exp. Route Total | | | | | | | | 6E-06 | | | | | 0.09 |
| | | | Dermal | 4,4'-DDD | 0.452 | mg/kg | 2.0E-007 | mg/kg/day | 2.7E-01 | 1/mg/kg/day | 5E-08 | 5.9E-07 | mg/kg/day | NA | NA | NA |
| | | | | 4,4'-DDE | 6.8 | mg/kg | 3.0E-06 | mg/kg/day | 3.8E-01 | 1/mg/kg/day | 1E-06 | 8.8E-06 | mg/kg/day | NA | NA | NA |
| | | | | 4,4'-DDT | 28.6 | mg/kg | 1.3E-005 | mg/kg/day | 3.8E-01 | 1/mg/kg/day | 5E-06 | 3.7E-005 | mg/kg/day | 4.5E-004 | mg/kg/day | 0.08 |
| | | | | Aluminum | 9964 | mg/kg | 4.5E-004 | mg/kg/day | NA | NA | NA | 1.3E-003 | mg/kg/day | 2.7E-001 | mg/kg/day | 0.005 |
| | | | | Manganese | 201 | mg/kg | 9.0E-006 | mg/kg/day | NA | NA | NA | 2.6E-005 | mg/kg/day | 7.0E-03 | mg/kg/day | 0.004 |
| | | | | Thallium | 1.2 | mg/kg | 5.3E-008 | mg/kg/day | NA | NA | NA | 1.5E-007 | mg/kg/day | NA | NA | NA |
| | | | Exp. Route Total | | | | | | | | 6E-06 | | | | | 0.09 |
| | | Exposure Point Total | | • | | | | | | | 1E-05 | | | | | 0.2 |
| | Expsoure Medium Total | | • | | | | | | | | 1E-05 | | 0.2 | | | |
| Soil Total | | • | · | | | | | • | | | 1E-05 | | | | · | 0.2 |
| | | | | | | | | Total | of Receptor Risks | Across All Media | 1E-05 | | Total of | Receptor Hazards | Across All Media | 0.2 |

TABLE 7.2.RME

CALCULATION OF CHEMICAL CANCER RISKS AND NON-CANCER HAZARDS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future Receptor Population: Resident Receptor Age: Child

| Medium | Exposure Medium | Exposure Point | Exposure Route | Chemical of | Е | PC | | Ca | ancer Risk Calculati | ions | | | Non- | Cancer Hazard Cal | Iculations | |
|--------|-----------------------|----------------------|------------------|-------------------|-------|-------|----------------|-----------------|----------------------|------------------|-------------|----------------|------------------|-------------------|------------------|-----------------|
| | | | | Potential Concern | Value | Units | Intake/Exposur | e Concentration | CSF/L | Init Risk | Cancer Risk | Intake/Exposur | re Concentration | RfD | D/RfC | Hazard Quotient |
| | | | | | | | Value | Units | Value | Units | | Value | Units | Value | Units | |
| Soil | Soil | Soil at Site 1 | Ingestion | 4,4'-DDD | 0.452 | mg/kg | 5.0E-07 | mg/kg/day | 2.4E-01 | 1/mg/kg/day | 1E-07 | 5.8E-06 | mg/kg/day | NA | NA | NA |
| | | | | 4,4'-DDE | 6.8 | mg/kg | 7.4E-06 | mg/kg/day | 3.4E-01 | 1/mg/kg/day | 3E-06 | 8.7E-05 | mg/kg/day | NA | NA | NA |
| | | | | 4,4'-DDT | 28.6 | mg/kg | 3.1E-005 | mg/kg/day | 3.4E-01 | 1/mg/kg/day | 1E-05 | 3.7E-004 | mg/kg/day | 5.0E-04 | mg/kg/day | 0.7 |
| | | | | Aluminum | 9964 | mg/kg | 1.1E-002 | mg/kg/day | NA | NA | NA | 1.3E-001 | mg/kg/day | 1.0E+00 | mg/kg/day | 0.1 |
| | | | | Manganese | 201 | mg/kg | 2.2E-004 | mg/kg/day | NA | NA | NA | 2.6E-003 | mg/kg/day | 1.4E-01 | mg/kg/day | 0.02 |
| | | | | Thallium | 1.2 | mg/kg | 1.3E-006 | mg/kg/day | NA | NA | NA | 1.5E-005 | mg/kg/day | NA | NA | NA |
| | | | Exp. Route Total | | | | | | | | 1E-05 | | | | | 0.8 |
| | | | Dermal | 4,4'-DDD | 0.452 | mg/kg | 9.8E-08 | mg/kg/day | 2.7E-01 | 1/mg/kg/day | 3E-08 | 1.1E-06 | mg/kg/day | NA | NA | NA |
| | | | | 4,4'-DDE | 6.8 | mg/kg | 1.5E-06 | mg/kg/day | 3.8E-01 | 1/mg/kg/day | 6E-07 | 1.7E-05 | mg/kg/day | NA | NA | NA |
| | | | | 4,4'-DDT | 28.6 | mg/kg | 6.2E-006 | mg/kg/day | 3.8E-01 | 1/mg/kg/day | 2E-06 | 7.2E-005 | mg/kg/day | 4.5E-004 | mg/kg/day | 0.2 |
| | | | | Aluminum | 9964 | mg/kg | 2.2E-004 | mg/kg/day | NA | NA | NA | 2.5E-003 | mg/kg/day | 2.7E-001 | mg/kg/day | 0.009 |
| | | | | Manganese | 201 | mg/kg | 4.4E-006 | mg/kg/day | NA | NA | NA | 5.1E-005 | mg/kg/day | 7.0E-003 | mg/kg/day | 0.007 |
| | | | | Thallium | 1.2 | mg/kg | 2.6E-008 | mg/kg/day | NA | NA | NA | 3.0E-007 | mg/kg/day | NA | NA | NA |
| | | | Exp. Route Total | | | | | | | | 3E-06 | | | | | 0.2 |
| | | Exposure Point Total | | 1 | | • | | • | • | • | 1F-05 | | • | | • | 1 |
| | Exposure Medium Total | | | | | | | · | · | | 1E-05 | | | · | · | 1 |
| Medium | | | | | | | | · | · | | 1E-05 | | | · | · | 1 |
| | | | | | | | | Total | of Receptor Risks | Across All Media | 1E-05 | | Total of | Receptor Hazards | Across All Media | 1 |

TABLE 7.3.RME

CALCULATION OF CHEMICAL CANCER RISKS AND NON-CANCER HAZARDS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future
Receptor Population: Resident
Receptor Age: Child/Adult

| Medium | Exposure Medium | Exposure Point | Exposure Route | Chemical of | EF | PC PC | | Ca | ncer Risk Calculati | ions | | | Non- | Cancer Hazard Cald | culations | |
|--------|-----------------------|----------------------|------------------|-------------------|-------|-------|---------------|------------------|---------------------|------------------|-------------|----------------|------------------|--------------------|------------------|-----------------|
| | | | | Potential Concern | Value | Units | Intake/Exposu | re Concentration | CSF/U | Init Risk | Cancer Risk | Intake/Exposur | re Concentration | RfD | /RfC | Hazard Quotient |
| | | | | | | | Value | Units | Value | Units | | Value | Units | Value | Units | |
| Soil | Soil | Soil at Site 1 | Ingestion | 4,4'-DDD | 0.452 | mg/kg | 7.1E-07 | mg/kg/day | 2.4E-01 | 1/mg/kg/day | 2E-07 | | | | | |
| | | | | 4,4'-DDE | 6.8 | mg/kg | 1.1E-05 | mg/kg/day | 3.4E-01 | 1/mg/kg/day | 4E-06 | | | | | ' |
| | | | | 4,4'-DDT | 28.6 | mg/kg | 4.4E-05 | mg/kg/day | 3.4E-01 | 1/mg/kg/day | 2E-05 | | | | | |
| | | | | Aluminum | 9964 | mg/kg | 1.6E-02 | mg/kg/day | NA | NA | NA | | | | | |
| | | | | Manganese | 201 | mg/kg | 3.2E-05 | mg/kg/day | NA | NA | NA | | | | | |
| | | | | Thallium | 1.2 | mg/kg | 1.9E-06 | mg/kg/day | NA | NA | NA | | | | | |
| | | | Exp. Route Total | | | | | | | | 2E-05 | | | | | |
| | | | Dermal | 4,4'-DDD | 0.452 | mg/kg | 3.0E-07 | mg/kg/day | 2.7E-01 | 1/mg/kg/day | 8E-08 | | | | | |
| | | | | 4,4'-DDE | 6.8 | mg/kg | 4.5E-06 | mg/kg/day | 3.8E-01 | 1/mg/kg/day | 2E-06 | | | | | 1 |
| | | | | 4,4'-DDT | 28.6 | mg/kg | 1.9E-05 | mg/kg/day | 3.8E-01 | 1/mg/kg/day | 7E-06 | | | | | |
| | | | | Aluminum | 9964 | mg/kg | 6.7E-04 | mg/kg/day | NA | NA | NA | | | | | |
| | | | | Manganese | 201 | mg/kg | 1.3E-05 | mg/kg/day | NA | NA | NA | | | | | |
| | | | | Thallium | 1.2 | mg/kg | 7.9E-08 | mg/kg/day | NA | NA | NA | | | | | |
| | | | Exp. Route Total | | | | | | | | 9E-06 | | | | | |
| | | Exposure Point Total | | | | | | | | | 3E-05 | | | | | |
| | Exposure Medium Total | • | | | | | | | | | 3E-05 | | | | | |
| Medium | | | | | | | | | | | 3E-05 | | | | | |
| | | | | - | | | | Total | of Receptor Risks | Across All Media | 3E-05 | | Total o | f Receptor Hazards | Across All Media | |

Note: Child/Adult cancer risk was calculated as the sum of the Child cancer risk (Table 7.2.RME) and the Adult cancer risk (Table 7.1.RME).

TABLE 7.1.RME

CALCULATION OF CHEMICAL CANCER RISKS AND NON-CANCER HAZARDS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future
Receptor Population: Resident
Receptor Age: Adult

| Medium | Exposure Medium | Exposure Point | Exposure Route | Chemical of | EF | PC O | | Can | cer Risk Calculatio | ins | | | Non-0 | Cancer Hazard Cal | culations | |
|------------|-----------------------|----------------------|------------------|-------------------|-------|-------|---------------|------------------|---------------------|------------------|-------------|-----------------|-----------------|-------------------|------------------|-----------------|
| | | | | Potential Concern | Value | Units | Intake/Exposu | re Concentration | CSF/U | nit Risk | Cancer Risk | Intake/Exposure | e Concentration | RfD | /RfC | Hazard Quotient |
| | | | | | | | Value | Units | Value | Units | | Value | Units | Value | Units | |
| Soil | Soil | Soil at Site 1 | Ingestion | 4,4'-DDD | 0.452 | mg/kg | 2.1E-07 | mg/kg/day | 2.4E-01 | 1/mg/kg/day | 5E-08 | 6.2E-07 | mg/kg/day | NA | NA | NA |
| | | | | 4,4'-DDE | 6.8 | mg/kg | 3.2E-06 | mg/kg/day | 3.4E-01 | 1/mg/kg/day | 1E-06 | 9.3E-06 | mg/kg/day | NA | NA | NA |
| | | | | 4,4'-DDT | 28.6 | mg/kg | 1.3E-005 | mg/kg/day | 3.4E-01 | 1/mg/kg/day | 5E-06 | 3.9E-05 | mg/kg/day | 5.0E-04 | mg/kg/day | 0.08 |
| | | | | Aluminum | 9964 | mg/kg | 4.7E-003 | mg/kg/day | NA | NA | NA | 1.4E-02 | mg/kg/day | 1.0E+00 | mg/kg/day | 0.01 |
| | | | | Manganese | 201 | mg/kg | 9.5E-005 | mg/kg/day | NA | NA | NA | 2.8E-04 | mg/kg/day | 1.4E-01 | mg/kg/day | 0.002 |
| | | | | Thallium | 1.2 | mg/kg | 5.6E-007 | mg/kg/day | NA | NA | NA | 1.6E-06 | mg/kg/day | NA | NA | NA |
| | | | Exp. Route Total | | | | | | | | 6E-06 | | | | | 0.09 |
| | | | Demal | 4,4'-DDD | 0.452 | mg/kg | 2.0E-007 | mg/kg/day | 2.7E-01 | 1/mg/kg/day | 5E-08 | 5.9E-07 | mg/kg/day | NA | NA | NA |
| | | | | 4,4'-DDE | 6.8 | mg/kg | 3.0E-06 | mg/kg/day | 3.8E-01 | 1/mg/kg/day | 1E-06 | 8.8E-06 | mg/kg/day | NA | NA | NA |
| | | | | 4,4'-DDT | 28.6 | mg/kg | 1.3E-005 | mg/kg/day | 3.8E-01 | 1/mg/kg/day | 5E-06 | 3.7E-005 | mg/kg/day | 4.5E-004 | mg/kg/day | 0.08 |
| | | | | Aluminum | 9964 | mg/kg | 4.5E-004 | mg/kg/day | NA | NA | NA | 1.3E-003 | mg/kg/day | 2.7E-001 | mg/kg/day | 0.005 |
| | | | | Manganese | 201 | mg/kg | 9.0E-006 | mg/kg/day | NA | NA | NA | 2.6E-005 | mg/kg/day | 7.0E-03 | mg/kg/day | 0.004 |
| | | | | Thallium | 1.2 | mg/kg | 5.3E-008 | mg/kg/day | NA | NA | NA | 1.5E-007 | mg/kg/day | NA | NA | NA |
| | | | Exp. Route Total | 1 | | | | | | | 6E-06 | | | | | 0.09 |
| | | Exposure Point Total | | | | | | | | • | 1E-05 | | | | • | 0.2 |
| | Exposure Medium Total | | | | | | | · | | | 1E-05 | | | | | 0.2 |
| Soil Total | • | | | | · | | | 1E-05 | | | | | | | | 0.2 |
| | | • | | <u> </u> | | | | Total o | of Receptor Risks | Across All Media | 1E-05 | | Total of F | Receptor Hazards | Across All Media | 0.2 |

TABLE 7.2.RME

CALCULATION OF CHEMICAL CANCER RISKS AND NON-CANCER HAZARDS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future Receptor Population: Resident Receptor Age: Child

| Medium | Exposure Medium | Exposure Point | Exposure Route | Chemical of | EI | PC | | Car | cer Risk Calculatio | ns | | | Non-0 | Cancer Hazard Cal | culations | |
|------------------|-----------------------|----------------------|------------------|-------------------|-------|-------|---------------|-------------------|---------------------|------------------|-------------|-----------------|-----------------|-------------------|------------------|-----------------|
| | | | | Potential Concern | Value | Units | Intake/Exposu | ure Concentration | CSF/U | nit Risk | Cancer Risk | Intake/Exposure | e Concentration | RfD | /RfC | Hazard Quotient |
| | | | | | | | Value | Units | Value | Units | | Value | Units | Value | Units | |
| Soil | Soil | Soil at Site 1 | Ingestion | 4,4'-DDD | 0.452 | mg/kg | 5.0E-07 | mg/kg/day | 2.4E-01 | 1/mg/kg/day | 1E-07 | 5.8E-06 | mg/kg/day | NA | NA | NA |
| | | | | 4,4'-DDE | 6.8 | mg/kg | 7.4E-06 | mg/kg/day | 3.4E-01 | 1/mg/kg/day | 3E-06 | 8.7E-05 | mg/kg/day | NA | NA | NA |
| | | | | 4,4'-DDT | 28.6 | mg/kg | 3.1E-005 | mg/kg/day | 3.4E-01 | 1/mg/kg/day | 1E-05 | 3.7E-004 | mg/kg/day | 5.0E-04 | mg/kg/day | 0.7 |
| | | | | Aluminum | 9964 | mg/kg | 1.1E-002 | mg/kg/day | NA | NA | NA | 1.3E-001 | mg/kg/day | 1.0E+00 | mg/kg/day | 0.1 |
| | | | | Manganese | 201 | mg/kg | 2.2E-004 | mg/kg/day | NA | NA | NA | 2.6E-003 | mg/kg/day | 1.4E-01 | mg/kg/day | 0.02 |
| | | | | Thallium | 1.2 | mg/kg | 1.3E-006 | mg/kg/day | NA | NA | NA | 1.5E-005 | mg/kg/day | NA | NA | NA |
| | | | Exp. Route Total | ĺ | | | | | | | 1E-05 | | | | | 0.8 |
| | | | Dermal | 4,4'-DDD | 0.452 | mg/kg | 9.8E-08 | mg/kg/day | 2.7E-01 | 1/mg/kg/day | 3E-08 | 1.1E-06 | mg/kg/day | NA | NA | NA |
| | | | | 4,4'-DDE | 6.8 | mg/kg | 1.5E-06 | mg/kg/day | 3.8E-01 | 1/mg/kg/day | 6E-07 | 1.7E-05 | mg/kg/day | NA | NA | NA |
| | | | | 4,4'-DDT | 28.6 | mg/kg | 6.2E-006 | mg/kg/day | 3.8E-01 | 1/mg/kg/day | 2E-06 | 7.2E-005 | mg/kg/day | 4.5E-004 | mg/kg/day | 0.2 |
| | | | | Aluminum | 9964 | mg/kg | 2.2E-004 | mg/kg/day | NA | NA | NA | 2.5E-003 | mg/kg/day | 2.7E-001 | mg/kg/day | 0.009 |
| | | | | Manganese | 201 | mg/kg | 4.4E-006 | mg/kg/day | NA | NA | NA | 5.1E-005 | mg/kg/day | 7.0E-003 | mg/kg/day | 0.007 |
| | | | | Thallium | 1.2 | mg/kg | 2.6E-008 | mg/kg/day | NA | NA | NA | 3.0E-007 | mg/kg/day | NA | NA | NA |
| | | | Exp. Route Total | | | | | | | | 3E-06 | | | | | 0.2 |
| | | Exposure Point Total | | H | • | | | | | | 1E-05 | | | | | 1 |
| | Exposure Medium Total | | • | • | | | | | | | 1E-05 | | | | | 1 |
| Soil Total 1E-05 | | | | | | | | | | | • | • | | 1 | | |
| | | | | | | | ! | Total | of Receptor Risks | Across All Media | 1E-05 | | Total of F | Receptor Hazards | Across All Media | 1 |
| | | | | | | | | | | | | | | | | |

TABLE 9.1.RME
SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCS
REASONABLE MAXIMUM EXPOSURE
The Dean Company

Scenario Timeframe: Future Receptor Population: Resident Receptor Age: Adult

| Medium | Exposure Medium | Exposure Point | Chemical of Potential | | | Carcinogeni | c Risk | | | Non-Carcinoge | nic Hazard Quo | tient | |
|----------------|--------------------|----------------------|-----------------------|-----------|------------|-------------|-------------|--------------|------------------------|---------------|----------------|--------|--------------|
| | | | Concern | Ingestion | Inhalation | Dermal | External | Exposure | Primary | Ingestion | Inhalation | Dermal | Exposure |
| | | | | | | | (Radiation) | Routes Total | Target Organ(s) | | | | Routes Total |
| Soil | Soil | Soil at Site 1 | 4,4'-DDD | 5E-08 | | 5E-08 | | 1E-07 | | | | | |
| | | | 4,4'-DDE | 1E-06 | | 1E-06 | | 2E-06 | | | | | |
| | | | 4,4'-DDT | 5E-06 | | 5E-06 | | 1E-05 | Liver | 0.08 | | 0.08 | 0.2 |
| | | | Aluminum | | | | | | Central Nervous System | 0.01 | | 0.005 | 0.02 |
| | | | Manganese | | | | | | Central Nervous System | 0.002 | | 0.004 | 0.006 |
| | | | Thallium | | | | | | | | | | |
| | | | Chemical Total | 6E-06 | | 6E-06 | | 1E-05 | | 0.09 | | 0.09 | 0.2 |
| | | | | | | | | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | |
| | | Exposure Point Total | | | • | | | 1E-05 | | • | | • | 0.2 |
| | Exposure Medium To | tal | | | • | | | 1E-05 | | • | | • | 0.2 |
| Soil Total | | | | | • | • | | 1E-05 | | • | | • | 0.2 |
| Receptor Total | | | | | | | | 1E-05 | | | | | 0.2 |

Total Risk Across All Media 1E-05 Total Hazard Across All Media 0.2

TABLE 9.1.RME
SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCS
REASONABLE MAXIMUM EXPOSURE
The Dean Company

Scenario Timeframe: Future Receptor Population: Resident Receptor Age: Adult

| Medium | Exposure Medium | Exposure Point | Chemical of Potential | | | Carcinogen | ic Risk | | | Non-Carcinoge | enic Hazard Quo | otient | |
|----------------|---------------------|----------------------|-----------------------|-----------|------------|------------|-------------|--------------|------------------------|---------------|-----------------|--------|--------------|
| | | | Concern | Ingestion | Inhalation | Dermal | External | Exposure | Primary | Ingestion | Inhalation | Dermal | Exposure |
| | | | | | | | (Radiation) | Routes Total | Target Organ(s) | | | | Routes Total |
| Soil | Soil | Soil at Site 1 | 4,4'-DDD | 5E-08 | | 5E-08 | | 1E-07 | | | | | |
| | | | 4,4'-DDE | 1E-06 | | 1E-06 | | 2E-06 | | | | | |
| | | | 4,4'-DDT | 5E-06 | | 5E-06 | | 1E-05 | Liver | 0.08 | | 0.08 | 0.2 |
| | | | Aluminum | | | | | | Central Nervous System | 0.01 | | 0.005 | 0.02 |
| | | | Manganese | | | | | | Central Nervous System | 0.002 | | 0.004 | 0.006 |
| | | | Thallium | | | | | | | | | | |
| | | | Chemical Total | 6E-06 | | 6E-06 | | 1E-05 | | 0.09 | | 0.09 | 0.2 |
| | | | | | | | | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | |
| | | Exposure Point Total | | | | | | 1E-05 | | • | • | | 0.2 |
| | Exposure Medium Tot | tal | | | | | | 1E-05 | | • | • | | 0.2 |
| Soil Total | · | <u> </u> | | | | | | 1E-05 | | | | | 0.2 |
| Receptor Total | · | · | _ | | | | | 1E-05 | | | | | 0.2 |

Total Risk Across All Media 1E-05 Total Hazard Across All Media 0.2

TABLE 9.2.RME
SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCS
REASONABLE MAXIMUM EXPOSURE
The Dean Company

Scenario Timeframe: Future Receptor Population: Resident Receptor Age: Child

| Medium | Exposure Medium | Exposure Point | Chemical of Potential | | | Carcinogen | ic Risk | | | Non-Carcinoge | enic Hazard Quo | otient | |
|----------------|-----------------------|----------------------|-----------------------|-----------|------------|------------|-------------|--------------|------------------------|---------------|-----------------|--------|--------------|
| | | | Concern | Ingestion | Inhalation | Dermal | External | Exposure | Primary | Ingestion | Inhalation | Dermal | Exposure |
| | | | | | | | (Radiation) | Routes Total | Target Organ(s) | | | | Routes Total |
| Soil | Soil | Soil at Site 1 | 4,4'-DDD | 1E-07 | | 3E-08 | | 1E-07 | | | | | |
| | | | 4,4'-DDE | 3E-06 | | 6E-07 | | 3E-06 | | | | | |
| | | | 4,4'-DDT | 1E-05 | | 2E-06 | | 1E-05 | Liver | 0.7 | | 0.2 | 0.9 |
| | | | Aluminum | | | | | | Central Nervous System | 0.1 | | 0.009 | 0.1 |
| | | | Manganese | | | | | | Central Nervous System | 0.02 | | 0.007 | 0.03 |
| | | | Thallium | | | | | | | | | | |
| | | | Chemical Total | 1E-05 | | 3E-06 | | 1E-05 | | 0.8 | | 0.2 | 1 |
| | | | | | | | | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | |
| | | Exposure Point Total | | | | • | | 1E-05 | | • | • | | 1 |
| | Exposure Medium Total | | | | • | | 1E-05 | | • | • | | 1 | |
| Soil Total | Soil Total | | | 1E-05 | | | 1E-05 | | | | | 1 | |
| Receptor Total | Receptor Total | | | | | | | 1E-05 | | | | | 1 |

Total Risk Across All Media 1E-05 Total Hazard Across All Media 1

TABLE 9.2.RME
SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCS
REASONABLE MAXIMUM EXPOSURE
The Dean Company

Scenario Timeframe: Future Receptor Population: Resident Receptor Age: Child

| Medium | Exposure Medium | Exposure Point | Chemical of Potential | | | Carcinogen | ic Risk | | Non-Carcinogenic Hazard Quotient | | | | |
|----------------|-----------------------|----------------------|-----------------------|-----------|------------|------------|-------------|--------------|----------------------------------|-----------|------------|--------|--------------|
| | | | Concern | Ingestion | Inhalation | Dermal | External | Exposure | Primary | Ingestion | Inhalation | Dermal | Exposure |
| | | | | | | | (Radiation) | Routes Total | Target Organ(s) | | | | Routes Total |
| Soil | Soil | Soil at Site 1 | 4,4'-DDD | 1E-07 | | 3E-08 | | 1E-07 | | | | | |
| | | | 4,4'-DDE | 3E-06 | | 6E-07 | | 3E-06 | | | | | |
| | | | 4,4'-DDT | 1E-05 | | 2E-06 | | 1E-05 | Liver | 0.7 | | 0.2 | 0.9 |
| | | | Aluminum | | | | | | Central Nervous System | 0.1 | | 0.009 | 0.1 |
| | | | Manganese | | | | | | Central Nervous System | 0.02 | | 0.007 | 0.03 |
| | | | Thallium | | | | | | | | | | |
| | | | Chemical Total | 1E-05 | | 3E-06 | | 1E-05 | | 0.8 | | 0.2 | 1 |
| | | | | | | | | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | |
| | | Exposure Point Total | • | | | | | 1E-05 | | • | | | 1 |
| | Exposure Medium Total | | | | | | 1E-05 | | | | | 1 | |
| Soil Total | Soil Total | | | | | 1E-05 | | | | | 1 | | |
| Receptor Total | | | _ | | | | | 1E-05 | | | | | 1 |

Total Risk Across All Media 1E-05 Total Hazard Across All Media 1

TABLE 9.3.RME
SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCS
REASONABLE MAXIMUM EXPOSURE
The Dean Company

Scenario Timeframe: Future
Receptor Population: Resident
Receptor Age: Child/Adult

| Medium | Exposure Medium | Exposure Point | Chemical of Potential | | | Carcinogen | ic Risk | | Non-Carcinogenic Hazard Quotient | | | | |
|----------------|-----------------------|-------------------|-----------------------|-----------|------------|------------|-------------|--------------|----------------------------------|-----------|------------|--------|--------------|
| | | | Concern | Ingestion | Inhalation | Dermal | External | Exposure | Primary | Ingestion | Inhalation | Dermal | Exposure |
| | | | | | | | (Radiation) | Routes Total | Target Organ(s) | | | | Routes Total |
| Soil | Soil | Soil at Site 1 | 4,4'-DDD | 2E-07 | | 8E-08 | | 3E-07 | | | | | |
| | | | 4,4'-DDE | 4E-06 | | 2E-06 | | 6E-06 | | | | | |
| | | | 4,4'-DDT | 2E-05 | | 7E-06 | | 3E-05 | | | | | |
| | | | Aluminum | | | | | | | | | | |
| | | | Manganese | | | | | | | | | | |
| | | | Thallium | | | | | | | | | | |
| | | | Chemical Total | 2E-05 | | 9E-06 | | 3E-05 | | | | | |
| | | | | | | | | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | |
| | Exposure Point Total | | | • | • | | 3E-05 | | • | • | • | | |
| | Exposure Medium Total | | | | | | 3E-05 | | | | | | |
| Soil Total | Soil Total | | | 3E | | | 3E-05 | | | | | | |
| Receptor Total | | | | | | | | 3E-05 | | | | | |

Total Risk Across All Media 3E-05 Total Hazard Across All Media --

Note: This table represents the residential lifetime cancer risk and was derived by combining the adult residential risks and the child residential risks.

TABLE 9.3.RME SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCS REASONABLE MAXIMUM EXPOSURE The Dean Company

Scenario Timeframe: Future
Receptor Population: Resident
Receptor Age: Child/Adult

| Medium | Exposure Medium | Exposure Point | Chemical of Potential | | | Carcinogen | ic Risk | | Non-Carcinogenic Hazard Quotient | | | | |
|----------------|-----------------------|-------------------|-----------------------|-----------|------------|------------|-------------|--------------|----------------------------------|-----------|------------|--------|--------------|
| | | | Concern | Ingestion | Inhalation | Dermal | External | Exposure | Primary | Ingestion | Inhalation | Dermal | Exposure |
| | | | | | | | (Radiation) | Routes Total | Target Organ(s) | | | | Routes Total |
| Soil | Soil | Soil at Site 1 | 4,4'-DDD | 2E-07 | | 8E-08 | | 3E-07 | | | | | |
| | | | 4,4'-DDE | 4E-06 | | 2E-06 | | 6E-06 | | | | | |
| | | | 4,4'-DDT | 2E-05 | | 7E-06 | | 3E-05 | | | | | |
| | | | Aluminum | | | | | | | | | | |
| | | | Manganese | | | | | | | | | | |
| | | | Thallium | | | | | | | | | | |
| | | | Chemical Total | 2E-05 | | 9E-06 | | 3E-05 | | | | | |
| | | | | | | | | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | |
| | Exposure Point Total | | | • | • | • | 3E-05 | | • | • | • | | |
| | Exposure Medium Total | | | | | | 3E-05 | | | | | | |
| Soil Total | Soil Total | | | | | 3E-05 | | | | | | | |
| Receptor Total | | | | | | | | 3E-05 | | | | | |

Total Risk Across All Media 3E-05 Total Hazard Across All Media --

Note: Child/Adult cancer risk was calculated using age-adjusted exposure factor values.

TABLE 1 SELECTION OF EXPOSURE PATHWAYS

The Dean Company

| Scenario Timeframe | Medium | Exposure Medium | Exposure Point | Receptor Population | Receptor Age | Exposure Route | Type of Analysis | Rationale for Selection or Exclusion of Exposure Pathway |
|-----------------------|--------|--------------------|-----------------------------------|------------------------|-----------------|-------------------|---------------------|--|
| Timeframe | Soil | Animal Tissue (1) | Beef from cattle grazing in field | Population 1 | Age 1 | Route 1 | Quant | Rationale |
| | | | | | Age 2 | Route 1 | Quant | Rationale |
| | | | | Population 2 | Age 1 | Route 1 | Quant | Rationale |
| | | | | | Age 2 | Route 1 | Quant | Rationale |

⁽¹⁾ Modeled via plant uptake from soil and beef cattle ingestion of plants. See Appendix x for full details of modeling.

TABLE Y (RAGS D ADULT LEAD WORKSHEET)

Site Name: Example Site, Slag Pile 2

Receptor: Adult Worker, Exposure to Media as Described

1. Lead Screening Questions

| Medium | Lead Concentration used in Model Run | | Basis for Lead Concentration Used | Lead Sc Concent | | Basis for Lead Screening Level |
|--------|--------------------------------------|-------|--------------------------------------|--------------------|-------|----------------------------------|
| | Value | Units | For Model Run | Value | Units | Ü |
| Soil | 2000 | mg/kg | Average Detected Value | 750 | mg/kg | Recommended Soil Screening Level |

2. Lead Model Questions

| Question | Response |
|---|---|
| What lead model was used? Provide reference and version | EPA Interim Adult Lead Model (1996) |
| If the EPA Adult Lead Model (ALM) was not used provide rationale for model selected. | n/a |
| Where are the input values located in the risk assessment report? | Located in Appendix 5 |
| What statistics were used to represent the exposure concentration terms and where are the data on concentrations in the risk assessment that support use of these statistics? | Mean soil concentration. Data are Located in Appendix 2 |
| What was the point of exposure and location? | OU 3 Slag pile area |
| Where are the output values located in the risk assessment report? | Located in Appendix 5 |
| What GSD value was used? If this is outside the recommended range of 1.8-2.1, provide rationale in Appendix <y>.</y> | 1.8 |
| What baseline blood lead concentration (PbB $_0$) value was used? If this is outside the default range of 1.7 to 2.2 provide rationale in Appendix $<$ Y $>$. | 2.0 |
| Was the default exposure frequency (EF; 219 days/year) used? | Yes |
| Was the default BKSF used (0.4 ug/dL per ug/day) used? | Yes |
| Was the default absorption fraction (AF; 0.12) used? | Yes |
| Was the default soil ingestion rate (IR; 50 mg/day) used? | Yes |
| If non-default values were used for any of the parameters listed above, where are the rationale for the values located in the risk assessment report? | Located in Appendix 5 |

3. Final Result

| Medium | Result | Comment/RBRG 1 |
|--------|--|----------------|
| Soil | 2000 ppm lead in soil results in >5% of receptors above a blood lead level of 10 ug/d and geometric mean blood lead = 11.6 ug/dL. This exceeds the blood lead goal as described in the 1994 OSWER Directive of no more than 5% of children (fetuses of exposed women) exceeding 10 ug/dL blood lead. | 1500 ppm |

^{1.} Attach the ALM spreadsheet output file upon which the Risk Based Remediation Goal (RBRG) was based and description of rationale for parameters used. For additional information, see www.epa.gov/superfund/programs/lead

TABLE X (RAGS D IEUBK LEAD WORKSHEET)

Site Name: Example Site, Neighborhood 2
Receptor: Future Residential Child (Age 0 to 84 Months) Exposure to Media as Described

1. Lead Screening Questions

| Medium | Lead Concentration used in Model Run | | Basis for Lead Concentration Used | Lead Screening Concentration | | Basis for Lead Screening Level |
|--------|---|-------|--------------------------------------|---------------------------------|-------|--|
| | Value | Units | for Model Run | Value | Units | C |
| Soil | 1000 | mg/kg | Average Detected Value | 400 | mg/kg | Recommended Soil Screening Level |
| Water | 4 | ug/L | Average Detected Value | 15 | ug/L | Recommended Drinking Water Action Level |

2. Lead Model Questions

| Question | Response for Residential Lead Model |
|---|--|
| What lead model (version and date) was used? | IEUBK version 0.99d, 1994 |
| Where are the input values located in the risk assessment report? | Located in Appendix 3 |
| What range of media concentrations were used for the model? | Refer to sampling data table 2 |
| What statistics were used to represent the exposure concentration terms and where are the data on concentrations in the risk assessment that support use of these statistics? | Mean value of backyard and side yard. Data presented in Appendix 3. |
| Was soil sample taken from top 2 cm? If not, why? | Yes |
| Was soil sample sieved? What size screen was used? If not sieved, provide rationale. | Yes, 250 um |
| What was the point of exposure/location? | Residential yard in Neighborhood 2: back yard and side yard composite. |
| Where are the output values located in the risk assessment report? | Located in Appendix 3 |
| Was the model run using default values only? | Yes, except for soil and dust concentration data. |
| Was the default soil bioavailability used? | Yes. Default is 30% |
| Was the default soil ingestion rate used? | Yes. Default values for 7 age groups are 85, 135, 135, 100, 090, and 85 mg/day |
| If non-default values were used, where are the rationale for the values located in the risk assessment report? | Located in Appendix 3 |

3. Final Result

| Medium | Result | Comment/PRG ¹ |
|--------|---|--|
| Soil | Input value of 1000 ppm in soil (and MSA derived dust of 710 ppm) results in 42.7% of children 0-84 months above a blood lead level of 10 ug/dL. Geometric mean blood lead = 9.5 ug/dL. This exceeds the blood lead goal as described in the 1994 OSWER Directive of no more than 5% of children exceeding 10 ug/dL blood lead. | Based on site conditions, a PRG of 354 ppm in soil is indicated. This PRG is typically rounded to 400 ppm. |

^{1.} Attach the IEUBK text output file and graph upon which the PRG was based as an appendix. For additional information, see www.epa.gov/superfund/programs/lead

TABLE 1 SELECTION OF EXPOSURE PATHWAYS

The Dean Company

| Scenario Timeframe | Medium | Exposure Medium | Exposure Point | Receptor Population | Receptor Age | Exposure Route | Type of Analysis | Rationale for Selection or Exclusion of Exposure Pathway |
|-----------------------|-------------|--------------------|--------------------|------------------------|-----------------|----------------------|---------------------|--|
| Future | Groundwater | Groundwater | Aquifer 1Tap Water | Resident | Adult | Dermal | Quant | Future onsite residents may rely on domestic wells drawing from Aquifer 1. |
| | | | | | | Ingestion | Quant | Future onsite residents may rely on domestic wells drawing from Aquifer 1. |
| | | | | | Child | Dermal | Quant | Future onsite residents may rely on domestic wells drawing from Aquifer 1. |
| | | | | | | Ingestion | Quant | Future onsite residents may rely on domestic wells drawing from Aquifer 1. |
| | | Air | Water Vapors from | Resident | Adult | Inhalation | Quant | Future onsite residents may rely on domestic wells drawing from Aquifer 1. |
| | | | Showerhead | | Child | Inhalation | None | Children are assumed not to shower. |
| | Soil | Soil | Soil at Site 1 | Resident | Adult | Dermal | Quant | Future onsite residents may come into contact with soil. |
| | | | | | | Ingestion | Quant | Future onsite residents may ingest soil. |
| | | | | | | External (Radiation) | Quant | Future onsite residents may come into contact with soil. |
| | | | | | Child | Dermal | Quant | Future onsite residents may come into contact with soil. |
| | | | | | | Ingestion | Quant | Future onsite residents may ingest soil. |
| | | | | | | External (Radiation) | Quant | Future onsite residents may come into contact with soil. |

TABLE 2.1

OCCURRENCE, DISTRIBUTION, AND SELECTION OF CHEMICALS OF POTENTIAL CONCERN

The Dean Company

Scenario Timeframe: Future Medium: Groundwater

Exposure Medium: Groundwater

| Exposure Point | CAS Number | Chemical | Minimum Concentration (Qualifier) | Maximum Concentration (Qualifier) | Units | Location of Maximum Concentration | Detection Frequency | Range of Detection Limits | Concentration Used for Screening (1) | Background Value (2) | Screening Toxicity Value (3) (N/C) | Potential ARAR/TBC Value | Potential ARAR/TBC Source | COPC Flag (Y/N) | Rationale for Selection or Deletion (4) |
|-------------------|---------------|----------------------------|-----------------------------------|-----------------------------------|-------|---|------------------------|---------------------------|--------------------------------------|-------------------------|------------------------------------|--------------------------------|---------------------------------|-----------------------|---|
| Aquifer 1 - | 117817 | Bis(2-ethylhexyl)phthalate | 2 J | 5 J | ug/l | GW3D | 4/12 | 3 - 4 | 5 | NA | 4.8 C | 6 | MCL | Y | ASL |
| Tap Water | 67663 | Chloroform | 0.6 J | 9 | ug/l | GW3D | 3/12 | 1 - 1 | 9 | NA | 0.063 C | 100 | MCL | Y | ASL |
| | 75150 | Carbon Disulfide | 0.3 J | 4.5 | ug/l | GW3D | 3/12 | 1 - 1 | 4.5 | NA | 100 N | NA | NA | N | BSL |
| | 76448 | Heptachlor | 2 J | 33 J | ug/l | GW4D | 6/12 | 0.01 - 0.01 | 33 | NA | 0.015 C | 0.4 | MCL | Υ | ASL |
| | 108883 | Toluene | 0.1 J | 0.2 J | ug/l | GW3D | 3/12 | 1 - 1 | 0.2 | NA | 75 N | 1000 | MCL | N | BSL |
| | 7429905 | Aluminum | 134 J | 1340 | ug/l | GW3D | 2/12 | 29 - 38.2 | 1340 | NA | 3700 N | 50 - 200 | SMCL | N | BSL |
| | 7440393 | Barium | 65 J | 489 | ug/l | GW1D | 6/12 | 0.2 - 1 | 489 | NA | 260 N | 2000 | MCL | Υ | ASL |
| | 7440417 | Beryllium | 0.2 K | 1.5 K | ug/l | GW2D | 3/12 | 0.1 - 1 | 1.5 | NA | 7.3 N | 4 | MCL | N | BSL |
| | 7439921 | Lead | 6 J | 35 J | ug/l | GW3D | 4/12 | 0.1 - 1 | 35 | NA | 15 | 15 | MCL | Υ | ASL |
| | 7439965 | Manganese | 1900 | 12500 | ug/l | GW1D | 6/12 | 0.3 - 1 | 12500 | NA | 73 N | 50 | SMCL | Υ | ASL |
| | 7440020 | Nickel | 0.9 J | 1.5 J | ug/l | GW4D | 3/12 | 0.9 - 7 | 1.5 | NA | 73 N | NA | NA | N | BSL |
| | 7440611 | Uranium | 50 | 500 | ug/l | GW1D | 12/12 | 1 - 2 | 500 | NA | 11 N | NA | NA | Υ | ASL |
| | 7440611 | Uranium 238 | 0.23 | 80 | pCi/l | GW1D | 12/12 | NA | NA | NA | NA | NA | NA | Υ | DET |
| | 13982-63-3 | Radium 226 | 0.2 | 11 | pCi/l | GW1D | 12 / 12 | NA | NA | NA | NA | 5 | MCL | Y | DET |

 Maximum concentration used for screening chemicals. No screening was conducted for radionuclides; all radionuclides detected are selected as COPCs.

(2) To date, no background study has been completed.

(3) All compounds were screened against the Risk-Based Concentration (RBC) Table, U.S. EPA Region III, May 8, 2001 for tap water (cancer benchmark = 1E-06; HQ = 0.1). Lead was screened against the action level of 15 ug/l.

(4) Rationale Codes:

Selection Reason: Above Screening Level (ASL)

Detected at Site (DET)

Deletion Reason: Below Screening Level (BSL)

Definitions: NA = Not Applicable

MCL = Maximum Contaminant Level

SMCL = Secondary Maximum Contaminant Level

J = Estimated Value

K = Estimated Value - Biased High

C = Carcinogen

N = Noncarcinogen

TABLE 2.2

OCCURRENCE, DISTRIBUTION, AND SELECTION OF CHEMICALS OF POTENTIAL CONCERN

The Dean Company

Scenario Timeframe: Future

Medium: Groundwater

Exposure Medium: Air

| Exposure Point | CAS Number | Chemical | Minimum Concentration (Qualifier) | Maximum Concentration (Qualifier) | Units | Location of Maximum Concentration | Detection Frequency | Range of Detection Limits | Concentration Used for Screening (1) | Background Value (2) | Screening Toxicity Value (3) (N/C) | Potential ARAR/TBC Value | Potential ARAR/TBC Source | COPC Flag (Y/N) | Rationale for Selection or Deletion (4) |
|-------------------|---------------|----------------------------|-----------------------------------|-----------------------------------|-------|-----------------------------------|------------------------|---------------------------|--------------------------------------|-------------------------|--|--------------------------------|---------------------------------|-----------------------|---|
| Water Vapors | 117817 | Bis(2-ethylhexyl)phthalate | 2 J | 5 J | ug/l | GW3D | 4/12 | 3-4 | 5 | NA | 4.8 C | 6 | MCL | Υ | ASL |
| from SHowerhead | 67663 | Chloroform | 0.6 J | 9 | ug/l | GW3D | 3/12 | 1 - 1 | 9 | NA | 0.063 C | 100 | MCL | Υ | ASL |
| | 75150 | Carbon Disulfide | 0.3 J | 4.5 | ug/l | GW3D | 3/12 | 1 - 1 | 4.5 | NA | 100 N | NA | NA | N | BSL |
| | 76448 | Heptachlor | 2 J | 33 J | ug/l | GW4D | 6/12 | 0.01 - 0.01 | 33 | NA | 0.015 C | 0.4 | MCL | Υ | ASL |
| | 108883 | Toluene | 0.1 J | 0.2 J | ug/l | GW3D | 3/12 | 1 - 1 | 0.2 | NA | 75 N | 1000 | MCL | N | BSL |
| | 7429905 | Aluminum | 134 J | 1340 | ug/l | GW3D | 2/12 | 29 - 38.2 | 1340 | NA | 3700 N | 50 - 200 | SMCL | N | BSL |
| | 7440393 | Barium | 65 J | 489 | ug/l | GW1D | 6/12 | 0.2 - 1 | 489 | NA | 260 N | 2000 | MCL | Υ | ASL |
| | 7440417 | Beryllium | 0.2 K | 1.5 K | ug/l | GW2D | 3/12 | 0.1 - 1 | 1.5 | NA | 7.3 N | 4 | MCL | N | BSL |
| | 7439921 | Lead | 6 J | 35 J | ug/l | GW3D | 4/12 | 0.1 - 1 | 35 | NA | 15 | 15 | MCL | Υ | ASL |
| | 7439965 | Manganese | 1900 | 12500 | ug/l | GW1D | 6/12 | 0.3 - 1 | 12500 | NA | 73 N | 50 | SMCL | Υ | ASL |
| | 7440020 | Nickel | 0.9 J | 1.5 J | ug/l | GW4D | 3/12 | 0.9 - 7 | 1.5 | NA | 73 N | NA | NA | N | BSL |
| | 7440611 | Uranium | 50 | 500 | ug/l | GW1D | 12/12 | 1 - 2 | 500 | NA | 11 N | NA | NA | Υ | ASL |
| | 7440611 | Uranium 238 | 0.23 | 80 | pCi/l | GW1D | 12/12 | NA | NA | NA | NA | NA | NA | Y | DET |
| | 13982-63-3 | Radium 226 | 0.2 | 11 | pCi/l | GW1D | 12/12 | NA | NA | NA | NA | 5 | MCL | Y | DET |

 Maximum concentration used for screening chemicals. No screening was conducted for radionuclides; all radionuclides detected are selected as COPCs.

(2) To date, no background study has been completed.

(3) All compounds were screened against the Risk-Based Concentration (RBC) Table, U.S. EPA Region III, May 8, 2001 for tap water (cancer benchmark = 1E-06; HQ = 0.1). Lead was screened against the action level of 15 ug/l.

(4) Rationale Codes:

Selection Reason: Above Screening Level (ASL)

Detected at Site (DET)

Deletion Reason: Below Screening Level (BSL)

Definitions: NA = Not Applicable

MCL = Maximum Contaminant Level

SMCL = Secondary Maximum Contaminant Level

J = Estimated Value

K = Estimated Value - Biased High

C = Carcinogen

N = Noncarcinogen

TABLE 2.3

OCCURRENCE, DISTRIBUTION, AND SELECTION OF CHEMICALS OF POTENTIAL CONCERN

The Dean Company

Scenario Timeframe: Future

Medium: Soil

Exposure Medium: Soil

| Exposure Point | CAS Number | Chemical | Minimum Concentration (Qualifier) | Maximum Concentration (Qualifier) | Units | Location of Maximum Concentration | Detection Frequency | Range of Detection Limits | Concentration Used for Screening (1) | Background Value (2) | Screening Toxicity Value (3) (N/C) | Potential ARAR/TBC Value | Potential ARAR/TBC Source | COPC Flag (Y/N) | Rationale for Selection or Deletion (4) |
|-------------------|---------------|--------------------|-----------------------------------|-----------------------------------|-------|---|------------------------|---------------------------------|--------------------------------------|-------------------------|--|--------------------------------|---------------------------------|-----------------------|---|
| Soil at Site 1 | 11096825 | Aroclor-1260 | 15 J | 110 J | ug/kg | SS03 | 6/29 | 33 - 300 | 110 | NA | 320 C | NA | NA | N | BSL |
| | 56553 | Benzo(a)anthracene | 120 J | 230 J | ug/kg | SS03 | 16/29 | 330 - 700 | 230 | NA | 870 C | NA | NA | N | BSL |
| | 50328 | Benzo(a)pyrene | 48 J | 70 J | ug/kg | SS03 | 17/29 | 30 - 70 | 70 | NA | 87 C | NA | NA | N | BSL |
| | 75150 | Carbon Disulfide | 2 J | 33 | ug/kg | SB07 | 4/29 | 10 - 16 | 33 | NA | 780000 N | NA | NA | N | BSL |
| | 72548 | 4,4'-DDD | 1 J | 4200 | ug/kg | SS09 | 22 / 29 | 3.3 - 1900 | 4200 | NA | 2700 C | NA | NA | Υ | ASL |
| | 72559 | 4,4'-DDE | 0.44 J | 7200 J | ug/kg | SS09 | 28 / 29 | 2.2 - 700 | 7200 | NA | 1900 C | NA | NA | Υ | ASL |
| | 50293 | 4,4'-DDT | 0.69 J | 290000 J | ug/kg | SB08 | 29 / 29 | 3.3 - 700 | 290000 | NA | 1900 C | NA | NA | Υ | ASL |
| | 108883 | Toluene | 1 J | 2 J | ug/kg | SS08 | 2/29 | 10 - 16 | 2 | NA | 1600000 N | NA | NA | N | BSL |
| | 7429905 | Aluminum | 1960 | 21700 | mg/kg | SB07 | 29 / 29 | 6.3 - 11 | 21700 | NA | 7800 N | NA | NA | Υ | ASL |
| | 7440417 | Beryllium | 0.1 J | 13.4 | mg/kg | SS06 | 23 / 29 | 0.02 - 0.21 | 13.4 | NA | 16 N | NA | NA | N | BSL |
| | 7439921 | Lead | 56 J | 750 J | mg/kg | SS03 | 16/29 | 10 - 16 | 750 | NA | 400 | NA | NA | Υ | ASL |
| | 7439965 | Manganese | 5.9 | 688 | mg/kg | SS03 | 29 / 29 | 0.05 - 0.5 | 688 | NA | 160 N | NA | NA | Υ | ASL |
| | 7782492 | Selenium | 0.53 J | 1 | mg/kg | SS02 | 9/29 | 0.43 - 0.75 | 1 | NA | 39 N | NA | NA | N | BSL |
| | 7440611 | Uranium | 50 | 700 | mg/kg | SS03 | 17 / 29 | 1 - 2 | 700 | NA | 610 N | NA | NA | Υ | ASL |
| | 7440611 | Uranium 238 | 0.3 | 110 | pCi/g | SS03 | 29 / 29 | 0.2 - 0.3 | NA | NA | NA | NA | NA | Υ | DET |
| | 13982-63-3 | Radium 226 | 0.36 | 41 | pCi/g | SS02 | 29 / 29 | 0.2 - 0.3 | NA | NA | NA | NA | NA | Υ | DET |

(1) Maximum concentration used for screening chemicals. No screening was conducted for radionuclides; all radionuclides detected are selected as COPCs.

(2) To date, no background study has been completed.

(3) All compounds were screened against the Risk-Based Concentration (RBC) Table, U.S. EPA Region III, May 8, 2001 for residential soil (cancer benchmark = 1E-06; HQ = 0.1). Lead was screened against the U.S. EPA screening value of 400 mg/kg.

(4) Rationale Codes:

Selection Reason: Above Screening Level (ASL)

Detected at Site (DET)

Deletion Reason: Below Screening Level (BSL)

Definitions: NA = Not Applicable

J = Estimated Value

C = Carcinogen

N = Noncarcinogen

TABLE 3.1.RME

EXPOSURE POINT CONCENTRATION SUMMARY

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future

Medium: Groundwater

Exposure Medium: Groundwater

| Exposure Point | Chemical of | Units | Arithmetic | 95% UCL | Maximum Concentration | | Exposure | Point Concentration | |
|-----------------------|----------------------------|-------|------------|-----------|--------------------------|-------|----------|---------------------|--------------|
| | Potential Concern | | Mean | (N/T) | (Qualifier) | Value | Units | Statistic | Rationale |
| Aquifer 1 - Tap Water | Bis(2-ethylhexyl)phthalate | ug/l | 4 | 5.5 (T) | 5 J | 5 | ug/l | Max | W-Test (1) |
| | Chloroform | ug/l | 1.9 | 14.9 (T) | 9 | 9 | ug/l | Max | W-Test (1) |
| | Heptachlor | ug/l | 27 | 30 (T) | 33 J | 30 | ug/l | 95% UCL - T | W - Test (2) |
| | Barium | ug/l | 224 | 2835 (T) | 489 | 489 | ug/l | Max | W-Test (1) |
| | Lead | ug/l | 21 | 32 (T) | 35 J | 32 | ug/l | 95% UCL - T | W - Test (2) |
| | Manganese | ug/l | 6052 | 33449 (T) | 12500 | 12500 | ug/l | Max | W-Test (1) |
| | Uranium | ug/l | 62 | 375 (T) | 500 | 375 | ug/l | 95% UCL - T | W - Test (2) |
| | Uranium 238 | pCi/l | 3.2 | 8.3 (T) | 80 | 8.3 | pCi/l | 95% UCL - T | W - Test (2) |
| | Radium 226 | pCi/l | 3.5 | 4 (T) | 11 | 4 | pCi/l | 95% UCL - T | W - Test (2) |

Statistics: Maximum Detected Value (Max); 95% UCL of Transformed Data (95% UCL - T)

T = Transformed

(1) 95% UCL exceeds maximum detected concentration. Therefore, maximum concentration used for EPC.

J = Estimated Value

(2) Shapiro-Wilk W Test indicates data are lognormally transformed.

TABLE 3.2.RME

EXPOSURE POINT CONCENTRATION SUMMARY

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future

Medium: Groundwater

Exposure Medium: Air

| Exposure Point | Chemical of | Units | Arithmetic | 95% UCL | Maximum Concentration | | Exposure | Point Concentration | |
|-------------------|----------------------------|-------|------------|----------------|--------------------------|-------|----------|---------------------|--------------|
| | Potential Concern | | Mean | (Distribution) | (Qualifier) | Value | Units | Statistic | Rationale |
| Water Vapors from | Bis(2-ethylhexyl)phthalate | ug/l | 4 | 5.5 (T) | 5 J | 5 | ug/l | Max | W-Test (1) |
| Showerhead | Chloroform | ug/l | 1.9 | 14.9 (T) | 9 | 9 | ug/l | Max | W-Test (1) |
| | Heptachlor | ug/l | 27 | 30 (T) | 33 J | 30 | ug/l | 95% UCL - T | W - Test (2) |

Statistics: Maximum Detected Value (Max); 95% UCL of Transformed Data (95% UCL - T)

T = Transformed

(1) 95% UCL exceeds maximum detected concentration. Therefore, maximum concentration used for EPC.

J = Estimated Value

(2) Shapiro-Wilk W Test indicates data are log-normally distributed.

TABLE 3.3.RME

EXPOSURE POINT CONCENTRATION SUMMARY

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future

Medium: Soil

Exposure Medium: Soil

| Exposure Point | Chemical of | Units | Arithmetic | 95% UCL | Maximum Concentration | | Exposure | e Point Concentration | |
|----------------|-------------------|-------|------------|-----------|--------------------------|-------|----------|-----------------------|--------------|
| | Potential Concern | | Mean | (N/T) | (Qualifier) | Value | Units | Statistic | Rationale |
| Soil at Site 1 | 4,4'-DDD | ug/kg | 239 | 452 (T) | 4200 | 452 | ug/kg | 95 % UCL -T | W - Test (2) |
| | 4,4'-DDE | ug/kg | 596 | 6793 (T) | 7200 J | 6793 | ug/kg | 95% UCL - T | W - Test (2) |
| | 4,4'-DDT | ug/kg | 11007 | 28619 (N) | 290000 J | 28619 | ug/kg | 95% UCL - N | W - Test (1) |
| | Aluminum | mg/kg | 7450 | 9964 (T) | 21700 | 9964 | mg/kg | 95% UCL - T | W - Test (2) |
| | Lead | mg/kg | 210 | 345 (T) | 750 J | 345 | mg/kg | 95% UCL - T | W - Test (2) |
| | Manganese | mg/kg | 116 | 201 (T) | 688 | 201 | mg/kg | 95% UCL - T | W - Test (2) |
| | Uranium | mg/kg | 125 | 675 (T) | 700 | 675 | mg/kg | 95% UCL - T | W - Test (2) |
| | Uranium 238 | pCi/g | 2.5 | 3.4 (T) | 110 | 3.4 | pCi/g | 95% UCL - T | W - Test (2) |
| | Radium 226 | pCi/g | 3.1 | 3.9 (T) | 41 | 3.9 | pCi/g | 95 % UCL - T | W- Test (2) |

Statistics: 95% UCL of Normal Data (95% UCL - N); 95% UCL of Transformed Data (95% UCL - T)

(1) Shapiro-Wilk W Test indicates data are normally distributed.

(2) Shapiro-Wilk W Test indicates data are lognormally transformed.

N = Normal

T = Transformed

J = Estimated Value

TABLE 4.1.RME

VALUES USED FOR DAILY INTAKE CALCULATIONS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future Medium: Groundwater

Exposure Medium: Groundwater

| Exposure Route | Receptor Population | Receptor Age | Exposure Point | Parameter Code | Parameter Definition | Value | Units | Rationale/ | Intake Equation/ Model Name |
|----------------|---------------------|--------------|-----------------------|------------------------------|--|---------------------------------|---|---|--|
| Ingestion | Resident | Adult | Aquifer 1 - Tap Water | CW IR-W EF ED BW | Chemical Concentration in Water Ingestion Rate of Water Exposure frequency Exposure Duration Body Weight | See Table 3.1 2 350 24 70 | mg/l l/day days/year years kg | See Table 3.1 EPA, 1991 EPA, 1991 EPA, 1991 EPA, 1991 | Chronic Daily Intake (CDI) (mg/kg/day) = CW x IR-W x EF x ED x 1/BW x 1/AT |
| | | | | AT-C AT-N CWR | Averaging Time - Cancer Averaging Time - Non-Cancer Radionuclide Concentration in Water | 25,550 8,760 | days days | EPA, 1989a EPA, 1989a | |
| | | | | IR-W EF ED | Ingestion Rate of Water Exposure Frequency Exposure Duration | See Table 3.1 2 350 24 | pCi/l l/day days/year | See Table 3.1 EPA, 1991 EPA, 1991 EPA, 1991 | Intake (pCi) = CWR x IR x EF x ED |
| | | Child | Aquifer 1 - Tap Water | CW IR-W EF | Chemical Concentration in Water Ingestion Rate of Water Exposure frequency | See Table 3.1 | years mg/l l/day | See Table 3.1 EPA, 1989b | CDI (mg/kg/day) = CW x IR-W x EF x ED x 1/BW x 1/AT |
| | | | | ED BW AT-C | Exposure Duration Body Weight Averaging Time - Cancer | 350 6 15 | days/year years kg | EPA, 1991 EPA, 1991 EPA, 1991 | |
| | | | | AT-N | Averaging Time - Non-Cancer Radionuclide Concentration in Water | 25,550 2,190 | days days | EPA, 1989a EPA, 1989a | |
| | | | | IR-W | Ingestion Rate of Water | See Table 3.1 | pCi/l l/day | See Table 3.1 EPA, 1991 | Intake (pCi) = CWR x IR x EF x ED |
| | | | | EF ED | Exposure Frequency Exposure Duration | 350 6 | days/year years | EPA, 1991 EPA, 1991 | |

TABLE 4.1.RME

VALUES USED FOR DAILY INTAKE CALCULATIONS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future Medium: Groundwater

Exposure Medium: Groundwater

| Exposure Route | Receptor Population | Receptor Age | Exposure Point | Parameter Code | Parameter Definition | Value | Units | Rationale/ Reference | Intake Equation/ Model Name |
|----------------|---------------------|--------------|-----------------------|-------------------|--|-------------------|-------------|-------------------------|--|
| Dermal | Resident | Adult | Aquifer 1 - Tap Water | CW | Chemical Concentration in Water | See Table 3.1 | mg/l | See Table 3.1 | Dermally Absorbed Dose (DAD) (mg/kg-day) = |
| | | | | FA | Fraction Absorbed Water | Chemical Specific | | EPA, 2001 | DA-event x EV x ED x EF x SA x 1/BW x 1/AT |
| | | | | Кр | Permeability Constant | Chemical Specific | cm/hr | EPA, 2001 | where for organic compounds, |
| | | | | SA | Skin Surface Area | 18,000 | cm2 | EPA, 2001 | Absorbed Dose per Event (DA-event) (mg/cm2-event) = |
| | | | | tau-event | Lag time per event | Chemical Specific | hours/event | EPA, 2001 | 2 FA x Kp x CW x CF x SQRT{(6 x tau-event x t-event)/pi} |
| | | | | t-event | Event Duration | 0.58 | hours/event | EPA, 2001 | or |
| | | | | В | Ratio of permeability coefficient of a | Chemical Specific | | EPA, 2001 | DA-event = FA x Kp x CW x {(t-event/(1 + B)) + |
| | | | | | compound through the stratum | | | | 2 x tau-event x ((1 + (3 x B) + (3 x B x B))/(1 + B)2)} |
| | | | | | corneum relative to its permeability | | | | and where for inorganic compounds, |
| | | | | | coefficient across the viable | | | | DA-event = Kp x CW x CF x t-event |
| | | | | | epidermis | | | | |
| | | | | EV | Event Frequency | 1 | events/day | EPA, 2001 | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 2001 | |
| | | | | ED | Exposure Duration | 24 | years | EPA, 1991 | |
| | | | | CF | Volumetric Conversion Factor for Water | 0.001 | I/cm3 | | |
| | | | | BW | Body Weight | 70 | kg | EPA, 2001 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 2001 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 8,760 | days | EPA, 2001 | |

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TABLE 4.1.RME

VALUES USED FOR DAILY INTAKE CALCULATIONS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future Medium: Groundwater

Exposure Medium: Groundwater

| Exposure Route | Receptor Population | Receptor Age | Exposure Point | Parameter Code | Parameter Definition | Value | Units | Rationale/ Reference | Intake Equation/ Model Name |
|--------------------|----------------------|--------------|-----------------------|-------------------|---|------------------------|-------------|-------------------------|--|
| Dermal (continued) | Resident (continued) | Child | Aquifer 1 - Tap Water | CW | Chemical Concentration in Water | See Table 3.1 | mg/l | See Table 3.1 | DAD (mg/kg-day) = |
| | | | | FA Kp | Fraction Absorbed Water Permeability Constant | Chemical Specific | cm/hr | EPA, 2001 EPA, 2001 | DA-event x EV x ED x EF x SA x 1/BW x 1/AT where for organic compounds, |
| | | | | SA | Skin Surface Area | 6.600 | cm2 | EPA, 2001 EPA. 2001 | DA-event (mg/cm2-event) = |
| | | | | - | Lag time per event | Chemical Specific | | EPA, 2001 | 2 FA x Kp x CW x CF x SQRT{(6 x tau-event x t-event)/pi} |
| | | | | t-event B | Event Duration Ratio of permeability coefficient of a | 1 Chemical Specific | hours/event | EPA, 2001 EPA, 2001 | or DA-event = FA x Kp x CW x {(t-event/(1 + B)) + |
| | | | | | compound through the stratum corneum relative to its permeability | | | | 2 x tau-event x ((1 + (3 x B) + (3 x B x B))/(1 + B)2)} and where for inorganic compounds, |
| | | | | | coefficient across the viable | | | | DA-event = Kp x CW x CF x t-event |
| | | | | | epidermis | | | | |
| | | | | EV | Event Frequency | 1 | events/day | EPA, 2001 | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 2001 | |
| | | | | ED | Exposure Duration | 6 | years | EPA, 2001 | |
| | | | | CF | Volumetric Conversion Factor for Water | 0.001 | I/cm3 | | |
| | | | | BW | Body Weight | 15 | kg | EPA, 2001 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 2001 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 2,190 | days | EPA, 2001 | |

EPA 1989a: Risk Assessment Guidance for Superfund. Volume 1: Human Health Evaluation Manual, Part A. OERR EPA/540/1-89/002.

EPA 1989b: Exposure Factors Handbook, July 1989, EPA/600/8-89/043.

EPA 1991: Risk Assessment Guidance for Superfund. Volume 1: Human Health Evaluation Manual - Supplemental Guidance, Standard Default Exposure Factors. Interim Final. OSWER 9285.6-03.

EPA 1992: Dermal Exposure Assessment: Principles and Applications. EPA/600/8-91/011B.

EPA 1997: Exposure Factors Handbook, Volume 1. EPA/600/P-95/002Fa.

EPA 2001: Risk Assessment Guidance for Superfund. Volume 1: Human Health Evaluation Manual (Part E, Supplemental Guidance for Dermal Risk Assessment) Interim.

TABLE 4.2.RME

VALUES USED FOR DAILY INTAKE CALCULATIONS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future Medium: Groundwater

Exposure Medium: Air

| Exposure Route | Receptor Population | Receptor Age | Exposure Point | Parameter | Parameter Definition | Value | Units | Rationale/ | Intake Equation/ |
|----------------|---------------------|--------------|-------------------|-----------|----------------------|-------|-------|------------|------------------------------|
| | | | | Code | | | | Reference | Model Name |
| Inhalation (1) | Resident | Adult | Water Vapors from | (1) | (1) | (1) | (1) | (1) | Foster and Chrostowski Model |
| | | | Showerhead | | | | | | |

⁽¹⁾ Refer to the Risk Assessment text for details on the modeled intake methodology and parameters used to calculate modeled intake values for the Foster and Chrostowski Shower Model.

TABLE 4.3.RME

VALUES USED FOR DAILY INTAKE CALCULATIONS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future

Medium: Soil

Exposure Medium: Soil

| Exposure Route | Receptor Population | Receptor Age | Exposure Point | Parameter | Parameter Definition | Value | Units | Rationale/ | Intake Equation/ |
|----------------|---------------------|--------------|----------------|-----------|------------------------------------|---------------|-----------|-----------------------|--|
| | | | | Code | | | | Reference | Model Name |
| Ingestion | Resident | Adult | Soil at Site 1 | cs | Chemical Concentration in Soil | See Table 3.3 | mg/kg | See Table 3.3 | Chronic Daily Intake (CDI) (mg/kg-day) = |
| | | | | IR-S | Ingestion Rate of Soil | 100 | mg/day | EPA, 1991 | CS x IR x FI x EF x ED x CF1 x 1/BW x 1/AT |
| | | | | FI | Fraction Ingested | 1 | | Professional Judgment | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 1991 | |
| | | | | ED | Exposure Duration | 24 | years | EPA, 1991 | |
| | | | | CF1 | Conversion Factor | 1E-06 | kg/mg | | |
| | | | | BW | Body Weight | 70 | kg | EPA, 1991 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 1989 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 8,760 | days | EPA, 1989 | |
| | | | | CSR | Radionuclide Concentration in Soil | See Table 3.3 | pCi/g | See Table 3.3 | Intake (pCi) = CSR x IR x CF x EF X ED |
| | | | | IR-S | Ingestion Rate of Soil | 100 | mg/day | EPA, 1991 | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 1991 | |
| | | | | ED | Exposure Duration | 24 | years | EPA, 1991 | |
| | | | | CF1 | Conversion Factor | 1.00E-03 | g/mg | | |
| | | Child | Soil at Site 1 | cs | Chemical Concentration in Soil | See Table 3.3 | mg/kg | See Table 3.3 | CDI (mg/kg-day) = |
| | | | | IR-S | Ingestion Rate of Soil | 200 | mg/day | EPA, 1991 | CS x IR x FI x EF x ED x CF1 x 1/BW x 1/AT |
| | | | | FI | Fraction Ingested | 1 | | Professional Judgment | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 1991 | |
| | | | | ED | Exposure Duration | 6 | years | EPA, 1991 | |
| | | | | CF1 | Conversion Factor | 1E-06 | kg/mg | | |
| | | | | BW | Body Weight | 15 | kg | EPA, 1991 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 1989 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 2,190 | days | EPA, 1989 | |

TABLE 4.3.RME

VALUES USED FOR DAILY INTAKE CALCULATIONS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future

Medium: Soil

Exposure Medium: Soil

| | | | | | T . | | | | Т |
|-----------------------|----------------------|-------------------|----------------------------|-----------|---|-------------------|--------------|---------------|---|
| | | | · · | | | | | 5 | |
| Exposure Route | Receptor Population | Receptor Age | Exposure Point | Parameter | Parameter Definition | Value | Units | Rationale/ | Intake Equation/ |
| | | | | Code | | | | Reference | Model Name |
| Ingestion (continued) | Resident (continued) | Child (continued) | Soil at Site 1 (continued) | CSR | Radionuclide Concentration in Soil | See Table 3.3 | pCi/g | See Table 3.3 | Intake (pCi) = CSR x IR x CF x EF X ED |
| | | | | IR-S | Ingestion Rate of Soil | 200 | mg/day | EPA, 1991 | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 1991 | |
| | | | | ED | Exposure Duration | 6 | years | EPA, 1991 | |
| | | | | CF1 | Conversion Factor | 1.00E-03 | g/mg | | |
| Dermal | Resident | Adult | Soil at Site 1 | CS | Chemical Concentration in Soil | See Table 3.3 | mg/kg | See Table 3.3 | Dermal Absorbed Dose (DAD) (mg/kg-day) = |
| | | | | CF | Conversion Factor | 1E-06 | kg/mg | | DA-event x EF x ED x EV x SA X 1/BW x 1/AT |
| | | | | SA | Skin Surface Area Available for Contact | 5,700 | cm2 | EPA, 2001 | where |
| | | | | AF | Soil to Skin Adherence Factor | 0.07 | mg/cm2-event | EPA, 2001 | Absorbed Dose per Event (DA-event) (mg/cm2-event) = |
| | | | | ABS-d | Dermal Absorption Factor | chemical-specific | unitless | EPA, 2001 | CS x CF x AF x ABS-d |
| | | | | EV | Event Frequency | 1 | events/day | EPA, 2001 | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 2001 | |
| | | | | ED | Exposure Duration | 24 | years | EPA, 1991 | |
| | | | | BW | Body Weight | 70 | kg | EPA, 2001 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 2001 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 8,760 | days | EPA, 2001 | |
| | | Child | Soil at Site 1 | CS | Chemical Concentration in Soil | See Table 3.3 | mg/kg | See Table 3.3 | DAD (mg/kg-day) = |
| | | | | CF | Conversion Factor | 1E-06 | kg/mg | | DA-event x EF x ED x EV x SA X 1/BW x 1/AT |
| | | | | SA | Skin Surface Area Available for Contact | 2,800 | cm2 | EPA, 2001 | where |
| | | | | AF | Soil to Skin Adherence Factor | 0.2 | mg/cm2-event | EPA, 2001 | DA-event (mg/cm2-event) = |
| | | | | ABS-d | Dermal Absorption Factor | chemical-specific | unitless | EPA, 2001 | CS x CF x AF x ABS-d |
| | | | | EV | Event Frequency | 1 | events/day | EPA, 2001 | |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 2001 | |
| | | | | ED | Exposure Duration | 6 | years | EPA, 2001 | |
| | | | | BW | Body Weight | 15 | kg | EPA, 2001 | |
| | | | | AT-C | Averaging Time - Cancer | 25,550 | days | EPA, 2001 | |
| | | | | AT-N | Averaging Time - Non-Cancer | 2,190 | days | EPA, 2001 | |

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TABLE 4.3.RME

VALUES USED FOR DAILY INTAKE CALCULATIONS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future

Medium: Soil
Exposure Medium: Soil

| Exposure Route | Receptor Population | Receptor Age | Exposure Point | Parameter Code | Parameter Definition | Value | Units | Rationale/ Reference | Intake Equation/ Model Name |
|----------------------|---------------------|--------------|----------------|-------------------|------------------------------------|---------------|-----------|-------------------------|---|
| External (Radiation) | Resident | Adult | Soil at Site 1 | CSR | Radionuclide Concentration in Soil | See Table 3.3 | pCi/g | See Table 3.3 | External Exposure (pCi-year/g) = |
| | | | | ET | Exposure Time | 17 | hrs/day | | CSR x ET x EF x {(Fi x GSFi) + (Fo x GSFo)] x ED x CF |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 1991 | |
| | | | | Fi | Time Fraction Indoors | 0.75 | | | |
| | | | | Fo | Time Fraction Outdoors | 0.25 | | | |
| | | | | GSFi | Gamma Shielding Factor Indoors | 0.8 | | | |
| | | | | GSFo | Gamma Shielding Factor Outdoors | 1 | | | |
| | | | | ED | Exposure Duration | 24 | years | EPA, 1991 | |
| | | | | CF | Conversion Factor | 0.000114 | years/hr | | |
| | | Child | Soil at Site 1 | CSR | Radionuclide Concentration in Soil | See Table 3.3 | pCi/g | See Table 3.3 | External Exposure (pCi-year/g) = |
| | | | | ET | Exposure Time | 17 | hrs/day | | CSR x ET x EF x {(Fi x GSFi) + (Fo x GSFo)] x ED x CF |
| | | | | EF | Exposure Frequency | 350 | days/year | EPA, 1991 | |
| | | | | Fi | Time Fraction Indoors | 0.875 | | | |
| | | | | Fo | Time Fraction Outdoors | 0.125 | | | |
| | | | | GSFi | Gamma Shielding Factor Indoors | 0.8 | | | |
| | | | | GSFo | Gamma Shielding Factor Outdoors | 1 | | | |
| | | | | ED | Exposure Duration | 6 | years | EPA, 1991 | |
| | | | | CF | Conversion Factor | 0.000114 | years/hr | | |

EPA 1989: Risk Assessment Guidance for Superfund. Volume 1: Human Health Evaluation Manual, Part A. OERR EPA/540/1-89/002.

EPA 1991: Risk Assessment Guidance for Superfund. Volume 1: Human Health Evaluation Manual - Supplemental Guidance, Standard Default Exposure Factors. Interim Final. OSWER 9285.6-03.

EPA 1995: Assessing Dermal Exposure from Soil, Technical Guidance Manual, Region III, EPA/903-K-95-003.

EPA 1997: Exposure Factors Handbook, Volume 1. EPA/600/P-95/002Fa.

EPA 2001: Risk Assessment Guidance for Superfund. Volume 1: Human Health Evaluation Manual (Part E, Supplemental Guidance for Dermal Risk Assessment) Interim.

NA = Not Available

TABLE 5.1 NON-CANCER TOXICITY DATA -- ORAL/DERMAL The Dean Company

| Chemical of Potential | Chronic/ Subchronic | Oral RfD | | Oral Absoprtion Efficiency for Dermal (1) | Absorbed RfD for Dermal (2) | | Primary Target | Combined Uncertainty/Modifying | RfD:Target Organ(s) | |
|----------------------------|------------------------|----------|-----------|---|-----------------------------|-----------|------------------------|--------------------------------|---------------------|--------------|
| Concern | | Value | Units | | Value | Units | Organ(s) | Factors | Source(s) | Date(s) |
| | | | | | | | | | | (MM/DD/YYYY) |
| 4,4'-DDD | NA | NA | NA | 1 | NA | NA | NA | NA | NA | NA |
| 4,4'-DDE | NA | NA | NA | 1 | NA | NA | NA | NA | NA | NA |
| 4,4'-DDT | Chronic | 5.0E-004 | mg/kg/day | 1 | 5.0E-004 | mg/kg/day | Liver | 100 | IRIS | 06/21/2001 |
| 4,4'-DDT | Subchronic | 5.0E-004 | mg/kg/day | 1 | 5.0E-004 | mg/kg/day | Liver | 100 | HEAST | 07/01/1997 |
| Bis(2-ethylhexyl)phthalate | Chronic | 2.0E-02 | mg/kg/day | 1 | 2.0E-02 | mg/kg/day | Liver | 1000 | IRIS | 06/21/2001 |
| Bis(2-ethylhexyl)phthalate | Subchronic | 2.0E-02 | mg/kg/day | 1 | 2.0E-02 | mg/kg/day | Liver | 1000 | HEAST | 07/01/1997 |
| Chloroform | Chronic | 1.0E-02 | mg/kg/day | 1 | 1.0E-02 | mg/kg/day | Liver | 1000 | IRIS | 06/21/2001 |
| Chloroform | Subchronic | 1.0E-02 | mg/kg/day | 1 | 1.0E-02 | mg/kg/day | Liver | 1000 | HEAST | 07/01/1997 |
| Heptachlor | Chronic | 5.0E-04 | mg/kg/day | 1 | 5.0E-04 | mg/kg/day | Liver | 300 | IRIS | 06/21/2001 |
| Heptachlor | Subchronic | 5.0E-04 | mg/kg/day | 1 | 5.0E-04 | mg/kg/day | Liver | 300 | HEAST | 07/01/1997 |
| Aluminum | Chronic | 1.0E+00 | mg/kg/day | 1 | 1.0E+00 | mg/kg/day | Central Nervous System | 100 | NCEA | 06/21/2001 |
| Barium | Chronic | 7.0E-02 | mg/kg/day | 0.07 | 4.9E-03 | mg/kg/day | Heart | 3 | IRIS | 02/02/2001 |
| Barium | Subchronic | 7.0E-02 | mg/kg/day | 0.07 | 4.9E-03 | mg/kg/day | Heart | 3 | HEAST | 07/01/1997 |
| Copper | Chronic | 3.7E-02 | mg/kg/day | 1 | 3.7E-02 | mg/kg/day | Gastrointestinal | NA | HEAST | 07/01/1997 |
| Copper | Subchronic | 3.7E-02 | mg/kg/day | 1 | 3.7E-02 | mg/kg/day | Gastrointestinal | NA | HEAST | 07/01/1997 |
| Iron | Chronic | 3.0E-01 | mg/kg/day | 1 | 3.0E-01 | mg/kg/day | Gastrointestinal | 1 | NCEA | 06/21/2001 |
| Lead | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| Manganese (nonfood) | Chronic | 2.0E-02 | mg/kg/day | 0.04 | 8.0E-04 | mg/kg/day | Central Nervous System | 1 | IRIS | 06/21/2001 |
| Uranium | Chronic | 3.0E-03 | mg/kg/day | 1 | 3E-003 | mg/kg/day | Kidney | 1000 | IRIS | 06/21/2001 |

(1) Source: Risk Assessment Guidance for Superfund. Volume 1: Human Health Evaluation Manual (Part E, Supplemental Guidance for Dermal Risk Assessment) Interim. Section 4.2 and Exhibit 4-1.

(2) See Risk Assessment text for the derivation of the "Absorbed RfD for Dermal".

Definitions: NA = Not Available

IRIS = Integrated Risk Information System

HEAST = Health Effects Assessment Summary Table, July 1997

NCEA = National Center for Environmental Assessment

TABLE 5.2

NON-CANCER TOXICITY DATA -- INHALATION

The Dean Company

| Chemical of Potential | Chronic/ Subchronic | Inhalation RfC | | Extrapolated RfD (1) | | Primary Target | Combined Uncertainty/Modifying | RfC : Target Organ | |
|----------------------------|------------------------|----------------|-------|----------------------|-----------|------------------------|--------------------------------|--------------------|--------------|
| Concern | | Value | Units | Value | Units | Organ(s) | Factors | Source(s) | Date(s) |
| | | | | | | | | | (MM/DD/YYYY) |
| 4,4'-DDD | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| 4,4'-DDE | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| 4,4'-DDT | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| Bis(2-ethylhexyl)phthalate | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| Chloroform | Chronic | 3.0E-04 | mg/m3 | 8.6E-05 | mg/kg/day | Nasal | 1000 | NCEA | 06/21/2001 |
| Chloroform | Subchronic | 3.0E-03 | mg/m3 | 8.6E-4 | mg/kg/day | Nasal | 100 | NCEA | 06/21/2001 |
| Heptachlor | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| Aluminum | Chronic | 5.0E-03 | mg/m3 | 1.4E-03 | mg/kg/day | Central Nervous System | 300 | NCEA | 06/21/2001 |
| Barium | Chronic | 5.0E-04 | mg/m3 | 1.4E-04 | mg/kg/day | Fetus | 1000 | HEAST | 07/01/1997 |
| Barium | Subchronic | 5.0E-03 | mg/m3 | 1.4E-03 | mg/kg/day | Fetus | 100 | HEAST | 07/01/1997 |
| Copper | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| Iron | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| Lead | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| Manganese (nonfood) | Chronic | 5.0E-05 | mg/m3 | 1.4E-05 | mg/kg/day | Central Nervous System | 1000 | IRIS | 06/21/2001 |
| Uranium | NA | NA | NA | NA | NA | NA | NA | NA | NA |

(1) See Risk Assessment text for the derivation of the "Extrapolated RfD".

Definitions: NA = Not Available

IRIS = Integrated Risk Information System

HEAST = Health Effects Assessment Summary Table, July 1997

NCEA = National Center for Environmental Assessment

$\label{eq:table 0} \textbf{SITE RISK ASSESSMENT IDENTIFICATION INFORMATION}$

The Dean Company

Site Name/OU: The Dean Company

Region: III

EPA ID Number: PAD999999999

State: PA

Status: Fund Lead Remedial Investigation

Federal Facility (Y/N): N

EPA Project Manager: John Smith

EPA Risk Assessor: Jane Doe

Document Author: Mary Smith-Johnson

Document Title: Human Health Risk Assessment for the Dean Company Site

Document Date: August 8, 2001

Comments: This site is contaminated with both chemical and radioactive compounds.

TABLE 5.3

NON-CANCER TOXICITY DATA -- SPECIAL CASE CHEMICALS

The Dean Company

| Chemical of Potential | Chronic/ Subchronic | | Parameter | | Primary Target Organ(s) | Combined Uncertainty/Modifying | Parameter:T | arget Organ(s) |
|-----------------------|------------------------|------|-----------|-------|-------------------------|--------------------------------|-------------|----------------|
| Concern | | Name | Value | Units | | Factors | Source(s) | Date(s) |
| | | | | | | | | (MM/DD/YYYY) |
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There are no special case chemicals in this risk assessment. As a result, the table is blank.

TABLE 6.1

CANCER TOXICITY DATA -- ORAL/DERMAL

The Dean Company

| Chemical of Potential | Oral Cancer | Slope Factor | Oral Absorption Efficiency for Dermal (1) | | er Slope Factor | Weight of Evidence/ | C | Oral CSF |
|----------------------------|-------------|--------------|---|----------|-----------------|---------------------|-----------|--------------|
| Concern | Value | Units | | Value | Units | Description | Source(s) | Date(s) |
| | | | | | | | | (MM/DD/YYYY) |
| 4,4'-DDD | 2.4E-01 | 1/mg/kg/day | 1 | 2.4E-01 | 1/mg/kg/day | B2 | IRIS | 06/21/2001 |
| 4,4'-DDE | 3.4E-01 | 1/mg/kg/day | 1 | 3.4E-01 | 1/mg/kg/day | B2 | IRIS | 06/21/2001 |
| 4,4'-DDT | 3.4E-001 | 1/mg/kg/day | 1 | 3.4E-001 | 1/mg/kg/day | B2 | IRIS | 06/21/2001 |
| Bis(2-ethylhexyl)phthalate | 1.4E-02 | 1/mg/kg/day | 1 | 1.4E-02 | 1/mg/kg/day | B2 | IRIS | 06/21/2001 |
| Chloroform | 6.1E-03 | 1/mg/kg/day | 1 | 6.1E-03 | 1/mg/kg/day | B2 | IRIS | 06/21/2001 |
| Heptachlor | 4.5E+00 | 1/mg/kg/day | 1 | 4.5E+00 | 1/mg/kg/day | B2 | IRIS | 06/21/2001 |
| Aluminum | NA | NA | 1 | NA | NA | NA | NA | NA |
| Barium | NA | NA | 0.07 | NA | NA | NA | NA | NA |
| Copper | NA | NA | 1 | NA | NA | NA | NA | NA |
| Iron | NA | NA | 1 | NA | NA | NA | NA | NA |
| Lead | NA | NA | NA | NA | NA | NA | NA | NA |
| Manganese (nonfood) | NA | NA | 0.04 | NA | NA | NA | NA | NA |
| Uranium | NA | NA | NA | NA | NA | NA | NA | NA |

(1) Source: Risk Assessment Guidance for Superfund. Volume 1: Human Health Evaluation Manual (Part E, Supplemental Guidance for Dermal Risk Assessment) Interim. Section 4.2 and Exhibit 4-1.

(2) See Risk Assessment text for the derivation of the "Absorbed Cancer Slope Factor for Dermal".

Definitions: NA = Not Available

IRIS = Integrated Risk Information System

B2 = Probable Human Carcinogen - indicates sufficient evidence

in animals and inadequate or no evidence in humans

TABLE 6.3 CANCER TOXICITY DATA -- SPECIAL CASE CHEMICALS

The Dean Company

| Chemical of Potential Concern | Name | Parameters Value | Units | Source(s) | Date(s) (MM/DD/YYYY) |
|-------------------------------|------|---------------------|-------|-----------|-------------------------|
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There are no special case chemicals in this risk assessment. As a result, this table is blank.

TABLE 6.2 CANCER TOXICITY DATA -- INHALATION

The Dean Company

| Chemical of Potential | Unit | Risk | Inhalation Cand | er Slope Factor | Weight of Evidence/ Cancer Guideline | Unit Risk : I | nhalation CSF |
|----------------------------|----------|---------|-----------------|-----------------|--------------------------------------|---------------|---------------|
| Concern | Value | Units | Value | Units | Description | Source(s) | Date(s) |
| | | | | | | | (MM/DD/YYYY) |
| 4,4'-DDD | NA | NA | NA | NA | NA | NA | NA |
| 4,4-DDE | NA | NA | NA | NA | NA | NA | NA |
| 4,4'-DDT | 9.7E-005 | 1/ug/m3 | 3.4E-001 | 1/mg/kg/day | B2 | IRIS | 06/21/2001 |
| Bis(2-ethylhexyl)phthalate | NA | NA | NA | NA | NA | NA | NA |
| Chloroform | 2.3E-05 | 1/ug/m3 | 8.1E-02 | 1/mg/kg/day | B2 | IRIS | 06/21/2001 |
| Heptachlor | 1.3E-03 | 1/ug/m3 | 4.5E+00 | 1/mg/kg/day | B2 | IRIS | 06/21/2001 |
| Aluminum | NA | NA | NA | NA | NA | NA | NA |
| Barium | NA | NA | NA | NA | NA | NA | NA |
| Copper | NA | NA | NA | NA | NA | NA | NA |
| Iron | NA | NA | NA | NA | NA | NA | NA |
| Lead | NA | NA | NA | NA | NA | NA | NA |
| Manganese (nonfood) | NA | NA | NA | NA | NA | NA | NA |
| Uranium | NA | NA | NA | NA | NA | NA | NA |

Definitions: NA = Not Available

IRIS = Integrated Risk Information System

B2 = Probable Human Carcinogen - indicates sufficient evidence

in animals and inadequate or no evidence in humans

TABLE 6.4 CANCER TOXICITY DATA -- EXTERNAL (RADIATION)

The Dean Company

| Chemical of Potential | Cancer SI | ope Factor | Source(s) | Date(s) (MM/DD/YYYY) |
|--------------------------|-----------|--------------------------|-----------|-------------------------|
| Concern | Value | Units | | |
| Uranium 238 | 6.2E-011 | Risk/pCi | HEAST | 07/01/1997 |
| | 5.3E-008 | Risk/year per pCi/g soil | HEAST | 07/01/1997 |
| Radium 226 | 3.0E-010 | Risk/pCi | HEAST | 07/01/1997 |
| | 6.7E-006 | Risk/year per pCi/g soil | HEAST | 07/01/1997 |
| | | | | |

HEAST = Health Effects Assessment Summary Table, July 1997

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TABLE 7.1.RME

CALCULATION OF CHEMICAL CANCER RISKS AND NON-CANCER HAZARDS REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future

Receptor Population: Resident

Receptor Age: Adult

| Medium | Exposure Medium | Exposure Point | Exposure Route | Chemical of | E | PC | | Ca | ncer Risk Calcula | tions | | | Non-0 | Cancer Hazard Cal | lculations | |
|----------------|-----------------------|-----------------------|------------------|----------------------------|-------|-------|----------------|-----------------|-------------------|-------------|--------------|---------------|------------------|-------------------|------------|----------------|
| | | | | Potential Concern | Value | Units | Intake/Exposur | e Concentration | CSF/L | Init Risk | Cancer Risk | Intake/Exposu | re Concentration | RfD | D/RfC | Hazard Quotier |
| | | | | | | | Value | Units | Value | Units | | Value | Units | Value | Units | |
| iroundwater | Groundwater | Aquifer 1 - Tap Water | Ingestion | Bis(2-ethylhexyl)phthalate | 0.005 | mg/l | 4.7E-005 | mg/kg/day | 1.4E-002 | 1/mg/kg/day | 7E-007 | 1.4E-004 | mg/kg/day | 2.0E-002 | mg/kg/day | 0.007 |
| | | | | Chloroform | 0.009 | mg/l | 8.5E-005 | mg/kg/day | 6.1E-003 | 1/mg/kg/day | 5E-007 | 2.5E-004 | mg/kg/day | 1.0E-002 | mg/kg/day | 0.03 |
| | | | | Heptachlor | 0.03 | mg/l | 2.8E-004 | mg/kg/day | 4.5E+000 | 1/mg/kg/day | 1E-003 | 8.1E-004 | mg/kg/day | 5.0E-004 | mg/kg/day | 2 |
| | | | | Barium | 0.489 | mg/l | 4.6E-003 | mg/kg/day | NA | NA | NA | 1.3E-002 | mg/kg/day | 7.0E-002 | mg/kg/day | 0.2 |
| | | | | Lead (1) | | | | | | | | | | | | |
| | | | | Manganese | 12.5 | mg/l | 1.2E-001 | mg/kg/day | NA | NA | NA | 3.4E-001 | mg/kg/day | 2.0E-002 | mg/kg/day | 17 |
| | | | | Uranium | 0.375 | mg/l | 3.8E-05 | mg/kg/day | NA | NA | NA | 1.0E-02 | mg/kg/day | 3.0E-03 | mg/kg/day | 3 |
| | | | Exp. Route Total | | • | • | | | | • | 1E-003 | | • | | • | 22 |
| | | | Dermal | Bis(2-ethylhexyl)phthalate | 0.005 | mg/l | 7.2E-005 | mg/kg/day | 1.4E-002 | 1/mg/kg/day | 1E-006 | 2.1E-004 | mg/kg/day | 2.2E-002 | mg/kg/day | 0.01 |
| | | | | Chloroform | 0.009 | mg/l | 1.7E-004 | mg/kg/day | 6.1E-003 | 1/mg/kg/day | 1E-006 | 4.9E-004 | mg/kg/day | 1.0E-002 | mg/kg/day | 0.05 |
| | | | | Heptachlor | 0.03 | mg/l | 1.3E-004 | mg/kg/day | 4.5E+000 | 1/mg/kg/day | 6E-004 | 3.9E-004 | mg/kg/day | 5.0E-004 | mg/kg/day | 0.8 |
| | | | | Barium | 0.489 | mg/l | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | | Lead (1) | | | | | | | | | | | | |
| | | | | Manganese | 12.5 | mg/l | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | For Posts Total | Uranium | 0.375 | mg/l | NA | NA | NA | NA | NA 6E-004 | NA | NA | NA | NA | NA 0.0 |
| | | | Exp. Route Total | | | | | | | | | | | | | 0.9 |
| | | Exposure Point Total | | | | | | | | | 2E-003 | | | | | 23 |
| | Exposure Medium Total | | | | | | | | | | 2E-003 | | | | | 23 |
| | Air | Water Vapors from | Inhalation | Bis(2-ethylhexyl)phthalate | 0.005 | mg/l | 2.3E-006 | mg/kg/day | NA | NA | NA | 3.6E-006 | mg/kg/day | NA | NA | NA |
| | | Showerhead | | Chloroform | 0.009 | mg/l | 1.3E-004 | mg/kg/day | 8.1E-002 | 1/mg/kg/day | 1E-005 | 3.9E-004 | mg/kg/day | 8.6E-005 | mg/kg/day | 5 |
| | | | | Heptachlor | 0.03 | mg/l | 2.6E-004 | mg/kg/day | 4.5E+000 | 1/mg/kg/day | 1E-003 | 7.7E-004 | mg/kg/day | NA | NA | NA |
| | | | Exp. Route Total | | | | | - | - | | 1E-003 | | | | | 5 |
| | | Exposure Point Total | | | | | | | | | 1E-003 | | | | | 5 |
| | Exposure Medium Total | | | | | | | | | | 1E-003 | | | | | 5 |
| undwater Total | | | | | | | | | | | 3E-003 | ì | | | | 28 |

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TABLE 7.1.RME

CALCULATION OF CHEMICAL CANCER RISKS AND NON-CANCER HAZARDS REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future

Receptor Population: Resident

Receptor Age: Adult

| | _ | | | | | | | | | | | | | | | |
|------------|-----------------------|----------------------|------------------|-------------------|-------|-------|----------------|-----------------|--------------------|------------------|-------------|-----------------|-----------------|-------------------|------------------|-----------------|
| Medium | Exposure Medium | Exposure Point | Exposure Route | Chemical of | E | PC | | Ca | ncer Risk Calculat | ions | | | Non-0 | Cancer Hazard Cal | culations | |
| | | | | Potential Concern | Value | Units | Intake/Exposur | e Concentration | CSF/U | Init Risk | Cancer Risk | Intake/Exposure | e Concentration | RfD | /RfC | Hazard Quotient |
| | | | | | | | Value | Units | Value | Units | | Value | Units | Value | Units | |
| Soil | Soil | Soil at Site 1 | Ingestion | 4,4'-DDD | 0.452 | mg/kg | 2.1E-07 | mg/kg/day | 2.4E-01 | 1/mg/kg/day | 5E-08 | 6.2E-07 | mg/kg/day | NA | NA | NA |
| | | | | 4,4'-DDE | 6.8 | mg/kg | 3.2E-06 | mg/kg/day | 3.4E-001 | 1/mg/kg/day | 1E-06 | 9.3E-06 | mg/kg/day | NA | NA | NA |
| | | | | 4,4'-DDT | 28.6 | mg/kg | 1.3E-005 | mg/kg/day | 3.4E-001 | 1/mg/kg/day | 5E-06 | 3.9E-05 | mg/kg/day | 5.0E-04 | mg/kg/day | 0.08 |
| | | | | Aluminum | 9964 | mg/kg | 4.7E-003 | mg/kg/day | NA | NA | NA | 1.4E-02 | mg/kg/day | 1.0E+00 | mg/kg/day | 0.01 |
| | | | | Lead (1) | | | | | | | | | | | | |
| | | | | Manganese | 201 | mg/kg | 9.5E-005 | mg/kg/day | NA | NA | NA | 2.8E-04 | mg/kg/day | 1.4E-01 | mg/kg/day | 0.002 |
| | | | | Uranium | 675 | mg/kg | 3.2E-004 | mg/kg/day | NA | NA | NA | 9.2E-04 | mg/kg/day | 3.0E-03 | mg/kg/day | 0.3 |
| | | | Exp. Route Total | | | • | | • | | • | 6E-06 | | • | | • | 0.4 |
| | | | Dermal | 4,4'-DDD | 0.452 | mg/kg | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | | 4,4'-DDE | 6.8 | mg/kg | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | | 4,4'-DDT | 28.6 | mg/kg | 1.6E-006 | mg/kg/day | 3.4E-001 | 1/mg/kg/day | 5E-007 | 4.7E-06 | mg/kg/day | 5.0E-04 | mg/kg/day | 0.009 |
| | | | | Aluminum | 9964 | mg/kg | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | | Lead (1) | | | | | | | | | | | | |
| | | | | Manganese | 201 | mg/kg | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | | Uranium | 675 | mg/kg | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | Exp. Route Total | | | | | | | | 5E-07 | | • | • | | 0.009 |
| | | Exposure Point Total | , | | | | | | | | 7E-006 | | | | | 0.4 |
| | Exposure Medium Total | • | | | | | | | | | 7E-006 | | | | | 0.4 |
| Soil Total | | | | | | | | | | | 7E-006 | | | | | 0.4 |
| | | | | | | | | | of Receptor Risks | Across All Media | 3E-003 | | Total of I | Receptor Hazards | Across All Media | 28 |

⁽¹⁾ Lead is evaluated for the resident using the IEUBK model. See Risk Assessment text for discussion of results and appendix for the lead modeling run results.

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TABLE 7.2.RME

CALCULATION OF CHEMICAL CANCER RISKS AND NON-CANCER HAZARDS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future Receptor Population: Resident Receptor Age: Child

| Medium | Exposure Medium | Exposure Point | Exposure Route | Chemical of | Е | PC | | Ca | ancer Risk Calculat | ions | | | Non- | Cancer Hazard Ca | Iculations | |
|-------------------|-----------------------|-----------------------|------------------|----------------------------|-------|-------|---------------|------------------|---------------------|-------------|-------------|----------------|-----------------|------------------|------------|-----------------|
| | | | | Potential Concern | Value | Units | Intake/Exposu | re Concentration | CSF/L | Init Risk | Cancer Risk | Intake/Exposur | e Concentration | Rff | D/RfC | Hazard Quotient |
| | | | | | | | Value | Units | Value | Units | | Value | Units | Value | Units | |
| Groundwater | Groundwater | Aquifer 1 - Tap Water | Ingestion | Bis(2-ethylhexyl)phthalate | 0.005 | mg/l | 2.7E-005 | mg/kg/day | 1.4E-002 | 1/mg/kg/day | 4E-007 | 3.2E-004 | mg/kg/day | 2.0E-002 | mg/kg/day | 0.02 |
| | | | | Chloroform | 0.009 | mg/l | 4.9E-005 | mg/kg/day | 6.1E-003 | 1/mg/kg/day | 3E-007 | 5.8E-004 | mg/kg/day | 1.0E-002 | mg/kg/day | 0.06 |
| | | | | Heptachlor | 0.03 | mg/l | 1.6E-004 | mg/kg/day | 4.5E+000 | 1/mg/kg/day | 7E-004 | 1.9E-003 | mg/kg/day | 5.0E-004 | mg/kg/day | 4 |
| | | | | Barium | 0.489 | mg/l | 2.7E-003 | mg/kg/day | NA | NA | NA | 3.1E-002 | mg/kg/day | 7.0E-002 | mg/kg/day | 0.4 |
| | | | | Lead (1) | | | | | | | | | | | | |
| | | | | Manganese | 12.5 | mg/l | 6.8E-002 | mg/kg/day | NA | NA | NA | 8.0E-001 | mg/kg/day | 2.0E-002 | mg/kg/day | 40 |
| | | | | Uranium | | mg/l | 2.1E-003 | mg/kg/day | NA | NA | NA | 2.4E-002 | mg/kg/day | 3.0E-003 | mg/kg/day | 8 |
| | | | Exp. Route Total | | | | | • | • | • | 7E-004 | Ī | • | • | | 52 |
| | | | Dermal | Bis(2-ethylhexyl)phthalate | 0.005 | mg/l | 3.1E-005 | mg/kg/day | 1.4E-002 | 1/mg/kg/day | 4E-007 | 3.6E-004 | mg/kg/day | 2.2E-002 | mg/kg/day | 0.02 |
| | | | | Chloroform | 0.009 | mg/l | 7.2E-005 | mg/kg/day | 6.1E-003 | 1/mg/kg/day | 4E-007 | 8.4E-004 | mg/kg/day | 1.0E-002 | mg/kg/day | 80.0 |
| | | | | Heptachlor | 0.03 | mg/l | 5.7E-005 | mg/kg/day | 4.5E+000 | 1/mg/kg/day | 3E-004 | 6.7E-004 | mg/kg/day | 5.0E-004 | mg/kg/day | 1 |
| | | | | Barium | 0.489 | mg/l | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | | Lead (1) | | | | | | | | | | | | |
| | | | | Manganese | 12.5 | mg/l | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | | Uranium | | mg/l | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | Exp. Route Total | | | | | • | • | • | 3E-004 | Ī | • | • | | 1 |
| | | Exposure Point Total | | | | | | | | | 1E-003 | | | | | 1 |
| | Exposure Medium Total | | | | | | Ì | | | | 1E-003 | Ì | | | | 53 |
| Groundwater Total | !! | | | | | | i | | | | 1E-003 | | | | | 53 |
| Soil | Soil | Soil at Site 1 | Ingestion | 4.4'-DDD | 0.452 | mg/kg | 5.0E-07 | mg/kg/day | 2.4E-01 | 1/mg/kg/day | 1E-07 | 5.8E-06 | mg/kg/day | NA NA | NA | NA |
| | | | - | 4,4'-DDE | 6.8 | mg/kg | 7.4E-06 | mg/kg/day | 3.4E-001 | 1/mg/kg/day | 3E-06 | 8.7E-05 | mg/kg/day | NA | NA | NA |
| | | | | 4.4'-DDT | 28.6 | mg/kg | 3.1E-005 | mg/kg/day | 3.4E-001 | 1/mg/kg/day | 1E-005 | 3.7E-004 | mg/kg/day | 5.0E-04 | mg/kg/day | 0.7 |
| | | | | Aluminum | 9964 | mg/kg | 1.1E-002 | mg/kg/day | NA | NA | NA | 1.3E-001 | mg/kg/day | 1.0E+00 | mg/kg/day | 0.1 |
| | | | | Lead (1) | | | | | | | | | | | | |
| | | | | Manganese | 201 | mg/kg | 2.2E-004 | mg/kg/day | NA | NA | NA | 2.6E-003 | mg/kg/day | 1.4E-01 | mg/kg/day | 0.02 |
| | | | | Uranium | | mg/kg | 7.4E-004 | mg/kg/day | NA | NA | NA | 8.6E-003 | mg/kg/day | 3.0E-003 | mg/kg/day | 3 |
| | | | Exp. Route Total | | | | l | | | 1 | 1E-005 | İ | | | 1 | 4 |

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TABLE 7.2.RME

CALCULATION OF CHEMICAL CANCER RISKS AND NON-CANCER HAZARDS

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future
Receptor Population: Resident
Receptor Age: Child

| Medium | Exposure Medium | Exposure Point | Exposure Route | Chemical of | EI | S | | Ca | ncer Risk Calculat | ions | | | Non-0 | Cancer Hazard Cal | culations | |
|------------------|-----------------------|----------------------------|------------------|-------------------|-------|-------|----------------|-----------------|--------------------|------------------|-------------|-----------------|-----------------|-------------------|------------------|-----------------|
| | | | | Potential Concern | Value | Units | Intake/Exposur | e Concentration | CSF/U | nit Risk | Cancer Risk | Intake/Exposure | e Concentration | RfD | /RfC | Hazard Quotient |
| Soil (continued) | Soil (continued) | Soil at Site 1 (continued) | Demal | 4,4'-DDD | 0.452 | mg/kg | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | | 4,4'-DDE | 6.8 | mg/kg | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | | 4,4'-DDT | 28.6 | mg/kg | 2.6E-006 | mg/kg/day | 3.4E-001 | 1/mg/kg/day | 9E-007 | 3.1E-005 | mg/kg/day | 5.0E-004 | mg/kg/day | 0.06 |
| | | | | Aluminum | 9964 | mg/kg | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | | Lead (1) | | | | | | | | | | | | |
| | | | | Manganese | 201 | mg/kg | NA | NA | NA | NA | NA | NA | NA | MA | NA | NA |
| | | | | Uranium | | mg/kg | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | Exp. Route Total | | • | | | | | • | 9E-07 | | • | | • | 0.06 |
| | | Exposure Point Total | | | | | | | | | 1E-005 | | | | | 4 |
| | Exposure Medium Total | | | | | | | | | | 1E-005 | | | | | 4 |
| Soil Total | Soil Total | | | | | | | | · | | 1E-005 | | | | | 4 |
| | | | | | | | • | Total | of Receptor Risks | Across All Media | 1E-03 | | Total of I | Receptor Hazards | Across All Media | 57 |

⁽¹⁾ Lead is evaluated for the resident using the IEUBK model. See Risk Assessment text for discussion of results and appendix for the lead modeling run results.

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TABLE 8.2 CALCULATION OF RADIATION CANCER RISKS The Smith Company

Scenario Timeframe: Future
Receptor Population: Resident
Receptor Age: Child

| Medium | Exposure Medium | Exposure Point | Exposure Route | Radionuclide of Potential Concern | Е | PC | Risk Calculation | | С | ancer Risk Calc | ulations | |
|-------------------|-----------------------|-----------------------|----------------------|-----------------------------------|----------|-------|------------------|------------------|------------------|-----------------|----------------------------|-------------|
| | | | | | Value | Units | Approach | Intake/Ext | ernal Dose | CSF/Conv | ersion Factor | Cancer Risk |
| | | | | | | | | Value | Units | Value | Units | |
| Groundwater | Groundwater | Aquifer 1 - Tap Water | Ingestion | Uranium 238 | 8.3E+000 | pCi/I | USEPA RAGS | 1.7E+004 | pCi | 6.2E-011 | Risk/pCi | 1E-006 |
| | | | | Radium 226 | 4.0E+000 | pCi/I | USEPA RAGS | 8.4E+003 | pCi | 3.0E-010 | Risk/pCi | 3E-006 |
| | | | Exp. Route Total | 1 | | | | | | | | 4E-006 |
| | | Exposure Point Total | | | • | • | | • | | • | | 4E-006 |
| | Exposure Medium Total | | | | | | | | | | | 4E-006 |
| Groundwater Total | | | | | | | | | | | | 4E-006 |
| Soil | Soil | Soil at Site 1 | Ingestion | Uranium 238 | 3.4E+000 | pCi/g | USEPA RAGS | 1.4E+003 | pCi | 6.2E-011 | Risk/pCi | 9E-008 |
| | | | | Radium 226 | 3.9E+000 | pCi/g | USEPA RAGS | 1.6E+003 | pCi | 3.0E-010 | Risk/pCi | 5E-007 |
| | | | Exp. Route Total | 1 | | | | | | | | 6E-007 |
| | | | External (Radiation) | Uranium 238 | 3.4E+000 | pCi/g | USEPA RAGS | 1.1E+001 | pCi-yr/g | 5.3E-008 | Risk/yr per pCi/ g soil | 6E-007 |
| | | | | Radium 226 | 3.9E+000 | pCi/g | USEPA RAGS | 1.3E+001 | pCi-yr/g | 6.7E-006 | Risk/yr per pCi/ g soil | 9E-005 |
| | | | Exp. Route Total | | | | | | | | 1 | 9E-005 |
| | | Exposure Point Total | | | • | • | | | | • | | 9E-005 |
| | Exposure Medium Total | - | | | | • | | · | · | | | 9E-005 |
| Soil Total | | _ | | _ | | • | | • | • | | | 9E-005 |
| | | _ | _ | _ | | | Total of R | eceptor Risks Ac | ross All Media = | | | 9E-005 |

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RADIATION DOSE ASSESSMENT WORKSHEET The Dean Company

Scenario Timeframe: Future Receptor Population: Resident Receptor Age: Adult

| Medium | Exposure Medium | Exposure Point | Exposure Route | Radionuclide of | EPC | | Dose | Internal/Exter | nal Dose | Standard for | Co | nversion Fa | ctor | Risk |
|-------------|-----------------|----------------------|----------------------|-------------------|----------|-------|----------|----------------|----------|---------------|-------|-------------|--------|------|
| | | | | Potential Concern | Value | Units | Approach | Value | Units | Comparison(1) | Value | Units | Source | |
| Groundwater | Groundwater | Aquifer 1 Tap Water | Ingestion | Uranium 238 | 8.3E+000 | pCi/I | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | | Radium 226 | 4.0E+000 | pCi/I | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | Exp. Route Total | ĺ | | | | NA | NA | | | | | NA |
| | | Exposure Point Total | | + | • | • | | NA | NA | | | , | | NA |
| Soil | Soil | Soil at Site 1 | Ingestion | Uranium 238 | 3.4E+000 | pCi/g | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | | Radium 226 | 3.9E+000 | pCi/g | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | Exp. Route Total | | | | | | | | | | | |
| | | | External (Radiation) | Uranium 238 | 3.4E+000 | pCi/g | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | | Radium 226 | 3.9E+000 | pCi/g | NA | NA | NA | NA | NA | NA | NA | NA |
| | | | Exp. Route Total | | | | | NA | NA | | | | | NA |
| | | Exposure Point Total | | • | • | • | | NA | NA | | • | | | NA |

NA = Not Applicable

Total of Receptor Dose Across All Media

NA

NA

Total of Receptor Risks Across All Media

NA

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TABLE 9.1.RME

SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCs REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future Receptor Population: Resident Receptor Age: Adult

| Medium | Exposure Medium | Exposure Point | Chemical of Potential | | | Carcinogen | ic Risk | | | Non-Carcinoge | enic Hazard Quo | otient | |
|-------------|---------------------|-----------------------|----------------------------|-----------|------------|------------|----------------------|--------------------------|----------------------------|---------------|-----------------|--------|--------------------------|
| | | | Concern | Ingestion | Inhalation | Dermal | External (Radiation) | Exposure Routes Total | Primary Target Organ(s) | Ingestion | Inhalation | Dermal | Exposure Routes Total |
| Groundwater | Groundwater | Aquifer 1 - Tap Water | Bis(2-ethylhexyl)phthalate | 7E-07 | | 1E-06 | | 2E-06 | Liver | 0.007 | | 0.01 | 0.02 |
| | | | Chloroform | 5E-07 | | 1E-06 | | 2E-06 | Liver | 0.03 | | 0.05 | 0.08 |
| | | | Heptachlor | 1E-03 | | 6E-04 | | 2E-03 | Liver | 2 | | 0.8 | 3 |
| | | | Barium | | | | | | Heart | 0.2 | | | 0.2 |
| | | | Lead (1) | | | | | | | | | | |
| | | | Manganese | | | | | | Central Nervous System | 17 | | | 17 |
| | | | Uranium | | | | | | Kidneys | 3 | | | 3 |
| | | | Chemical Total | 1E-03 | | 6E-04 | | 2E-03 | | 22 | | 0.9 | 23 |
| | | | Uranium 238 | 9E-06 | | | | 9E-06 | | | | | |
| | | | Radium 226 | 2E-05 | | | | 2E-05 | | | | | |
| | | | Radionuclide Total | 3E-05 | | | | 3E-05 | | | | | |
| | | Exposure Point Total | | | | | | 2E-03 | | | | | 23 |
| | Exposure Medium Tot | al | | | | | | 2E-03 | | | | | 23 |
| | Air | Water Vapors from | Bis(2-ethylhexyl)phthalate | | | | | | | | | | |
| | | Showerhead | Chloroform | | 1E-05 | | | 1E-05 | Liver | | 5 | | 5 |
| | | | Heptachlor | | 1E-03 | | | 1E-03 | | | | | |
| | | | Barium | | | | | | | | | | |
| | | | Lead (1) | | | | | | | | | | |
| | | | Manganese | | | | | | | | | | |
| | | | Uranium | | | | | | | | | | |
| | | | Chemical Total | | 1E-03 | | | 1E-03 | | | 5 | | 5 |
| | | | | <u> </u> | | | <u> </u> | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | |
| | | Exposure Point Total | | | | | | 1E-03 | | | | | 5 |
| | Exposure Medium Tot | tal | | | | | | 1E-03 | | | | | 5 |

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TABLE 9.1.RME

SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCs $\,$

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future
Receptor Population: Resident
Receptor Age: Adult

| Medium | Exposure Medium | Exposure Point | Chemical of Potential | Carcinogenic Risk | | | ic Risk | | Non-Carcinogenic Hazard Quotient | | | | |
|------------------|--------------------|-------------------|-----------------------|-------------------|------------|--------|-------------|--------------|----------------------------------|-----------|------------|--------|--------------|
| | | | Concern | Ingestion | Inhalation | Dermal | External | Exposure | Primary | Ingestion | Inhalation | Dermal | Exposure |
| | | | | | | | (Radiation) | Routes Total | Target Organ(s) | | | | Routes Total |
| Groundwater Tota | Groundwater Total | | | 3E-03 | | | | | 28 | | | | |

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TABLE 9.1.RME

SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCs $\,$

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future
Receptor Population: Resident
Receptor Age: Adult

| Medium | Exposure Medium | Exposure Point | Chemical of Potential | | Carcinogenic Risk Non-Carcinogenic Hazard Quotient | | | | | | otient | | |
|----------------|-----------------------|----------------------|--|-------------------------|--|--------|----------------------|--------------------------|---|------------------------------|------------|--------|--------------------------|
| | | | Concern | Ingestion | Inhalation | Dermal | External (Radiation) | Exposure Routes Total | Primary Target Organ(s) | Ingestion | Inhalation | Dermal | Exposure Routes Total |
| Soil | Soil | Soil at Site 1 | 4,4'-DDD 4,4'-DDE 4,4'-DDT Aluminum Lead (1) Manganese Uranium | 5E-08 1E-06 5E-06 | | 5E-07 | (NadiauUI) | 5E-08 1E-06 6E-06 | Liver Central Nervous System Central Nervous System Kidney | 0.08 0.01 0.002 0.3 | | 0.009 | 0.09 0.01 0.002 |
| | | | Chemical Total | 6E-06 | | 5E-07 | | 7E-06 | - tuano) | 0.4 | | 0.009 | 0.4 |
| | | | Uranium 238 Radium 226 | 2E-07 1E-006 | | | 2E-06 4E-04 | 2E-06 4E-04 | | | | | |
| | | | Radionuclide Total | 1E-06 | | | 4E-04 | 4E-04 | | | | | |
| | | Exposure Point Total | | | | | | 4E-04 | | | | | 0.4 |
| | Exposure Medium Total | | | | | | | 4E-04 | | | | | 0.4 |
| Soil Total | pil Total | | | <u> </u> | | | | 4E-04 | | | | | 0.4 |
| Receptor Total | Receptor Total | | | | | | | 3E-03 | | | | | 28 |

Total Risk Across All Media 3E-03 Total Hazard Across All Media 28

(1) Lead is evaluated for the resident using the IEUBK model. See Risk Assessment text for discussion of results
and appendix for the lead modeling run results.

Total Central Nervous System HI Across All Media = 3

Total Central Nervous System HI Across All Media = 17

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TABLE 9.2.RME SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCS REASONABLE MAXIMUM EXPOSURE The Dean Company

Scenario Timeframe: Future
Receptor Population: Resident
Receptor Age: Child

| Medium | Exposure Medium | Exposure Point | Chemical of Potential | Carcinogenic Risk | | | | Non-Carcinogenic Hazard Quotient | | | | | |
|-----------------|----------------------|-----------------------|----------------------------|-------------------|------------|--------|-------------|----------------------------------|------------------------|-----------|------------|--------|--------------|
| | | | Concern | Ingestion | Inhalation | Dermal | External | Exposure | Primary | Ingestion | Inhalation | Dermal | Exposure |
| | | | | | | | (Radiation) | Routes Total | Target Organ(s) | | | | Routes Total |
| Groundwater | Groundwater | Aquifer 1 - Tap Water | Bis(2-ethylhexyl)phthalate | 4E-07 | | 4E-07 | | 8E-07 | Liver | 0.02 | | 0.02 | 0.04 |
| | | | Chloroform | 3E-07 | | 4E-07 | | 7E-07 | Liver | 0.06 | | 0.08 | 0.1 |
| | | | Heptachlor | 7E-04 | | 3E-04 | | 1E-03 | Liver | 4 | | 1 | 5 |
| | | | Barium | | | | | | Heart | 0.4 | | | 0.4 |
| | | | Lead (1) | | | | | | | | | | |
| | | | Manganese | | | | | | Central Nervous System | 40 | | | 40 |
| | | | Uranium | | | | | | Kidney | 8 | | | 8 |
| | | | Chemical Total | 7E-04 | | 3E-04 | | 1E-03 | | 52 | | 1 | 53 |
| | | | Uranium 238 | 1E-06 | | | | 1E-06 | | | | | |
| | | | Radium 226 | 3E-06 | | | | 3E-06 | | | | | |
| | | | Radionuclide Total | 4E-06 | | | | 4E-06 | | | | | |
| | Exposure Point Total | | | | | | 1E-03 | | | | | 53 | |
| | Exposure Medium To | otal | | | • | | | 1E-03 | | | | | 53 |
| Groundwater Tot | Groundwater Total | | | 1E-03 | | | | | | | | 53 | |

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TABLE 9.2.RME SUMMARY OF RECEPTOR RISKS AND HAZARDS FOR COPCS REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future
Receptor Population: Resident
Receptor Age: Child

| Medium | Exposure Medium | Exposure Point | Chemical of Potential | | Carcinogenic Risk | | | | Non-Carcinogenic Hazard Quotient | | | | |
|----------------|--------------------|----------------------|-----------------------|-----------|-------------------|--------|-------------|--------------|----------------------------------|-----------|------------|--------|--------------|
| | | | Concern | Ingestion | Inhalation | Dermal | External | Exposure | Primary | Ingestion | Inhalation | Dermal | Exposure |
| | | | | | | | (Radiation) | Routes Total | Target Organ(s) | | | | Routes Total |
| Soil | Soil | Soil at Site 1 | 4,4'-DDD | 1E-07 | | | | 1E-07 | | | | | |
| | | | 4,4'-DDE | 3E-06 | | | | 3E-06 | | | | | |
| | | | 4,4'-DDT | 1E-05 | | 9E-07 | | 1E-05 | Liver | 0.7 | | 0.06 | 0.8 |
| | | | Aluminum | | | | | | Central Nervous System | 0.1 | | | 0.1 |
| | | | Lead (1) | | | | | | | | | | |
| | | | Manganese | | | | | | Central Nervous System | 0.02 | | | 0.02 |
| | | | Uranium | | | | | | Kidney | 3 | | | 3 |
| | | | Chemical Total | 1E-05 | | 9E-07 | | 1E-05 | | 4 | | 0.06 | 4 |
| | | | Uranium 238 | 9E-08 | | | 6E-07 | 7E-07 | | | | | |
| | | | Radium 226 | 5E-07 | | | 9E-05 | 9E-05 | | | | | |
| | | | Radionuclide Total | 6E-07 | | | 9E-05 | 9E-05 | | | | | |
| | | Exposure Point Total | | | • | • | | 1E-04 | | | | | 4 |
| | Exposure Medium To | tal | | | • | • | | 1E-04 | | | | | 4 |
| Soil Total | | | | | • | • | | 1E-04 | | | • | | 4 |
| Receptor Total | | | | | | | | 1E-03 | | | | | 57 |

(1) Lead is evaluated for the resident using the IEUBK model. See Risk Assessment text for discussion of results

Total Liver HI Across All Media =

and appendix for the lead modeling run results.

Total Kidney HI Across All Media =

11

Total Central Nervous System HI Across All Media =

40

1E-03

Total Risk Across All Media

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57

Total Hazard Across All Media

TABLE 10.1.RME RISK ASSESSMENT SUMMARY REASONABLE MAXIMUM EXPOSURE The Dean Company

Scenario Timeframe: Future Receptor Population: Resident Receptor Age: Adult

| Medium | Exposure Medium | Exposure Point | Chemical of Potential | | Carcinogenic Risk | | | | Non-Carcinogenic Hazard Quotient | | | | | |
|------------------|-----------------------|---------------------------------|----------------------------|-----------|-------------------|--------|-------------|--------------|----------------------------------|-----------|------------|--------|--------------|--|
| | | | Concern | Ingestion | Inhalation | Dermal | External | Exposure | Primary | Ingestion | Inhalation | Dermal | Exposure | |
| | | | | | | | (Radiation) | Routes Total | Target Organ(s) | | | | Routes Total | |
| Groundwater | Groundwater | Aquifer 1 - Tap Water | Bis(2-ethylhexyl)phthalate | 7E-07 | | 1E-06 | | 2E-06 | | | | | | |
| | | | Chloroform | 5E-07 | | 1E-06 | | 2E-06 | | | | | , | |
| | | | Heptachlor | 1E-03 | | 6E-04 | | 2E-03 | Liver | 2 | | 0.8 | 3 | |
| | | | Manganese | | | | | | Central Nervous System | 17 | | | 17 | |
| | | | Uranium | | | | | | Kidney | 3 | | | 3 | |
| | | | Chemical Total | 1E-03 | | 6E-04 | | 2E-03 | | 22 | | 0.8 | 23 | |
| | | | Uranium 238 | 9E-06 | | | - | 9E-06 | | | | | | |
| | | | Radium 226 | 2E-05 | | | | 2E-05 | | | | | | |
| | | | Radionuclide Total | 3E-05 | | | | 3E-05 | | | | | | |
| | | Exposure Point Total | | | | | | 2E-03 | | | | | 23 | |
| | Exposure Medium To | | | | | | | 2E-03 | | | | | 23 | |
| | Air | Water Vapors from Showerhead | Chloroform | | 1E-05 | | | 1E-05 | Liver | | 5 | | 5 | |
| | | | Heptachlor | | 1E-03 | | | 1E-03 | | | | | | |
| | | | Chemical Total | | 1E-03 | | | 1E-03 | | | 5 | | 5 | |
| | | | | | | | | | | | | | | |
| | | | Radionuclide Total | | | | | | | | | | | |
| | | Exposure Point Total | | | | | | 1E-03 | | | | | 5 | |
| | Exposure Medium Total | | | | | | | 1E-03 | | | | | 5 | |
| Groundwater Tota | oundwater Total | | | | | | | 3E-03 | | | | | 28 | |

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TABLE 10.1.RME RISK ASSESSMENT SUMMARY

REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future

Receptor Population: Resident

Receptor Age: Adult

| Medium | Exposure Medium | Exposure Point | Chemical of Potential | | Carcinogenic Risk Non-Ca | | | | Non-Carcinoge | arcinogenic Hazard Quotient | | | | |
|----------------|-----------------------|----------------------|-----------------------|-----------|--------------------------|--------|----------|----------|---------------|-----------------------------|------------|--------|----------|--|
| | | | Concern | Ingestion | Inhalation | Dermal | External | Exposure | Primary | Ingestion | Inhalation | Dermal | Exposure | |
| Soil | Soil | Soil at Site 1 | 4,4'-DDE | 1E-06 | | | | 1E-06 | | | | | | |
| | | | 4,4'-DDT | 5E-06 | | 5E-007 | | 6E-06 | | | | | | |
| | | | Chemical Total | 6E-06 | | 5E-07 | | 7E-06 | | | | | | |
| | | | Uranium 238 | 2E-07 | | | 2E-06 | 2E-06 | | | | | | |
| | | | Radium 226 | 1E-006 | | | 4E-04 | 4E-04 | | | | | | |
| | | | Radionuclide Total | 1E-06 | | | 4E-04 | 4E-04 | | | | | | |
| | | Exposure Point Total | | | | | | 4E-04 | | | | | | |
| | Exposure Medium Total | | | • | | | 4E-04 | | | | | | | |
| Soil Total | Soil Total | | | | | | | 4E-04 | | | | | | |
| Receptor Total | Receptor Total | | | | • | | | 3E-03 | | | | · | 28 | |

Total Risk Across All Media

Total Liver HI Across All Media

Total Liver HI Across All Media = 8

Total Kidney HI Across All Media = 3

Cancer risks presented are those greater than 1E-06; Non-cancer risks presented are those greater than 1.

Total Central Nervous System HI Across All Media = 17

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TABLE 10.2.RME RISK ASSESSMENT SUMMARY REASONABLE MAXIMUM EXPOSURE

The Dean Company

Scenario Timeframe: Future
Receptor Population: Resident
Receptor Age: Child

| Medium | Exposure Medium | Exposure Point | Chemical of Potential | Carcinogenic Risk | | | | | Non-Carcinogenic Hazard Quotient | | | | | |
|-------------------|-----------------------|-----------------------|-----------------------|-------------------|------------|--------|----------------------|--------------------------|----------------------------------|-----------|------------|--------|--------------------------|--|
| | | | Concern | Ingestion | Inhalation | Dermal | External (Radiation) | Exposure Routes Total | Primary Target Organ(s) | Ingestion | Inhalation | Dermal | Exposure Routes Total | |
| Groundwater | Groundwater | Aquifer 1 - Tap Water | Heptachlor | 7E-04 | | 3E-04 | | 1E-03 | Liver | 4 | | 1 | 5 | |
| | | | Manganese | | | | | | Central Nervous System | 40 | | | 40 | |
| | | | Uranium | | | | | | Kidney | 8 | | | 8 | |
| | | | Chemical Total | 7E-04 | | 3E-04 | | 1E-03 | | 52 | | 1 | 53 | |
| | | | Uranium 238 | 1E-06 | | | [| 1E-06 | | | | | | |
| | | | Radium 226 | 3E-06 | | | | 3E-06 | | | | | | |
| | | | Radionuclide Total | 4E-06 | | | | 4E-06 | | | | | | |
| | | Exposure Point Total | | | | | | 1E-03 | | | | | 53 | |
| | Exposure Medium To | otal | | | | | | 1E-03 | | | | | 53 | |
| Froundwater Total | 1 | | | | | | | 1E-03 | | | | | 53 | |
| Soil | Soil | Soil at Site 1 | 4,4'-DDE | 3E-006 | | | | 3E-06 | | | | | | |
| | | | 4,4'-DDT | 1E-05 | | 9E-07 | | 1E-05 | | | | | | |
| | | | Uranium | | | | | | Kidney | 3 | | | 3 | |
| | | | Chemical Total | 1E-05 | | 9E-07 | | 1E-05 | | 3 | | | 3 | |
| | | | Radium 226 | 5E-07 | | | 9E-05 | 9E-05 | | | | | | |
| | | | Radionuclide Total | 6E-07 | | | 9E-05 | 9E-05 | | | | | | |
| | Exposure Point Total | | | | | | | 1E-04 | | • | • | | 3 | |
| | Exposure Medium Total | | | | | | | 1E-04 | | | | | 3 | |
| Soil Total | Soil Total | | | | | | 1E-04 | | | | | 3 | | |
| Receptor Total | Receptor Total | | | | | | | 1E-03 | | | | | 56 | |

Total Risk Across All Media

Total Liver HI Across All Media =

Total Liver HI Across All Media =

Total Kidney HI Across All Media =

Total Kidney HI Across All Media =

Total Kidney HI Across All Media =

Total Central Nervous System HI Across All Media =

40

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EXHIBIT 1- 2 GUIDELINES FOR PART D APPLICABILITY

| SITE LEAD | PART D APPLICABLE |
|--|-------------------|
| Fund Lead | ✓ |
| Federal Facility Lead | √ |
| PRP Lead | √ |
| State Lead | ✓ |
| SITE TYPE ¹ | |
| Remedial: Scoping, RI/FS, Risk Assessment, Proposed Plan, ROD, RD/RA, Presumptive Remedy | ✓ |
| Post-Remedial: ESD, Amended ROD, Five-Year Review | ✓ |
| Removal: Non-time Critical, Time-Critical, Streamlined | 2 |
| SACM ³ | ✓ |
| RCRA Corrective Action ⁴ | 2 |

Notes:

- 1 The RAGS Part D Workgroup also suggests that RAGS Part D could be a useful tool for quantitative risk assessment for non-NPL, BRAC, and Brownfields sites and encourages its use.
- 2 RAGS Part D use is encouraged as appropriate.
- 3 Superfund Accelerated Cleanup Model.
- 4 As described in the September 1996 EPA memorandum on Coordination Between Resource Conservation and Recovery Act (RCRA) Corrective Action and Closure and CERCLA Site Activities, EPA is "...committed to the principle of parity between the RCRA corrective action and CERCLA programs...".

INTERIM DELIVERABLES FOR EACH SITE

| Interim Deliverable | Scope of Deliverable |
|--|--|
| INTERIM DELIVERABLES ASSOC | IATED WITH PLANNING TABLE 0 |
| TARA Schedule Worksheet | One Worksheet for each Risk Assessment. |
| Planning Table 0 - Site Risk Assessment Identification Information | One Planning Table for each Risk Assessment. |
| INTERIM DELIVERABLES ASSOC | IATED WITH PLANNING TABLE 1 |
| Planning Table 1 - Selection of Exposure Pathways | One Planning Table for each Risk Assessment. |
| INTERIM DELIVERABLES ASSOC | IATED WITH PLANNING TABLE 2 |
| Data Useability Worksheet | One Worksheet for each Medium. |
| Supporting Information on Background Values | Information for all Chemicals listed in Planning Table 2. |
| Planning Table 2 - Occurrence, Distribution, and Selection of Chemicals of Potential Concern (COPCs) | One Planning Table for each unique combination of Scenario Timeframe, Medium, and Exposure Medium. |
| INTERIM DELIVERABLES ASSOC | IATED WITH PLANNING TABLE 3 |
| Supporting Information on EPCs | Information for all EPCs presented in Planning Table 3. |
| Planning Table 3 - Exposure Point Concentration (EPC) Summary | One Planning Table for each unique combination of Scenario Timeframe, Medium, and Exposure Medium. |
| INTERIM DELIVERABLES ASSOC | IATED WITH PLANNING TABLE 4 |
| Supporting Information on Modeled Intake Methodology and Parameters | Information for all Modeled Intake calculations that are not presented in Planning Table 4. |
| Supporting Information on Chemical-Specific Parameters | Information for all Chemical-Specific Parameters used. |
| Dermal Worksheet | Information for calculation of DA(event). |
| Planning Table 4 - Values Used for Daily Intake Calculations | One Planning Table for each unique combination of Scenario Timeframe, Medium, and Exposure Medium. |
| INTERIM DELIVERABLES ASSOCIATI | ED WITH PLANNING TABLES 5 AND 6 |
| Supporting Information on Toxicity Data for Special Case Chemicals | Information for each Special Case Chemical. |
| Planning Table 5 - Non-Cancer Toxicity Data | Three Planning Tables - 5.1 for Oral/Dermal, 5.2 for Inhalation, and 5.3 for Special Case Chemicals. |

INTERIM DELIVERABLES FOR EACH SITE (continued)

| Interim Deliverable | Scope of Deliverable | | | | | | |
|---|---|--|--|--|--|--|--|
| Planning Table 6 - Cancer Toxicity Data | Four Planning Tables - 6.1 for Oral/Dermal, 6.2 for Inhalation, 6.3 for Special Case Chemicals, and 6.4 for External (Radiation). | | | | | | |
| INTERIM DELIVERABLES ASSOCIAT | ED WITH PLANNING TABLES 7 AND 8 | | | | | | |
| Supporting Information on Special Chemical Risk and Hazard Calculations | Information for each Special Case Chemical. | | | | | | |
| Planning Table 7 - Calculation of Chemical Cancer Risks and Non-Cancer Hazards | One Planning Table for each unique combination of Scenario Timeframe, Receptor Population, and Receptor Age, for RME and for CT. | | | | | | |
| Radiation Dose Assessment Worksheet | One Worksheet for each unique combination of Scenario Timeframe, Receptor Population, and Receptor Age (as appropriate). | | | | | | |
| Planning Table 8 - Calculation of Radiation Cancer Risks | One Planning Table for each unique combination of Scenario Timeframe, Receptor Population and Receptor Age. | | | | | | |
| INTERIM DELIVERABLES ASSOCIATE | ED WITH PLANNING TABLES 9 AND 10 | | | | | | |
| Planning Table 9 - Summary of Receptor Risks and Hazards for COPCs | One Planning Table for each unique combination of Scenario Timeframe, Receptor Population, and Receptor Age, for RME and CT. | | | | | | |
| Planning Table 10 - Risk Summary | One Planning Table for each unique combination of Scenario Timeframe, Receptor Population, and Receptor Age, for RME and CT. | | | | | | |
| INTERIM DELIVERABLES | ASSOCIATED WITH LEAD | | | | | | |
| Lead Worksheets (if applicable) | Separate Worksheets for Residential and Non-Residential Scenarios for each unique combination of Scenario Timeframe, Receptor Population, and Receptor Age. | | | | | | |
| INTERIM DELIVERABLES ASSOCIATED WITH UNCERTAINTY ASSESSMENT | | | | | | | |
| Assessment of Confidence and Uncertainty | One Assessment for each Risk Assessment. | | | | | | |
| INTERIM DELIVERABLES ASSOCIAT | ED WITH PROBABILISTIC ANALYSIS | | | | | | |
| Summary of Probabilistic Analysis (if applicable) | One Summary for each Risk Assessment. | | | | | | |

INTERIM DELIVERABLES FOR EACH SITE (continued)

| Interim Deliverable | Scope of Deliverable | | | | | |
|--|--|--|--|--|--|--|
| INTERIM DELIVERABLES ASSOCIATED WITH THE ROD | | | | | | |
| ROD Risk Worksheets | As appropriate to document (in draft form) the need for remedial action. | | | | | |

Notes:

- 1. Each Interim Deliverable should be reviewed and verified by EPA prior to submission of the Draft Baseline Risk Assessment Report.
- 2. Each Interim Deliverable should later be incorporated in the Draft and Final Baseline Risk Assessment Reports.
- 3. The Interim Deliverables are needed for each risk assessment to achieve standardization in risk assessment reporting.

EXHIBIT 3-2 STANDARDIZED RISK ASSESSMENT REPORTING

| Risk Assessment Activity | Corresponding Planning Table/Worksheet |
|--|--|
| Data C | ollection |
| Provide identification information for the risk assessment | Planning Table 0 - Site Risk Assessment Identification Information |
| Plan the risk assessment review process | TARA Schedule Worksheet |
| Develop a conceptual site model | Planning Table 1 - Selection of Exposure Pathways |
| Gather and report appropriate data | Planning Table 2 - Occurrence, Distribution, and Selection of Chemicals of Potential Concern |
| Data Ev | aluation |
| Evaluate detection frequency, background data, and | Data Useability Worksheet |
| site data | Planning Table 2 - Occurrence, Distribution, and Selection of Chemicals of Potential Concern |
| Identify chemicals of potential concern and provide rationale for selection and deletion | Planning Table 2 - Occurrence, Distribution, and Selection of Chemicals of Potential Concern |
| Exposure . | Assessment |
| Characterize physical setting, identify potential pathways and exposed population | Planning Table 1 - Selection of Exposure Pathways |
| Identify exposure assumptions | Planning Table 4 - Values Used for Daily Intake Calculations |
| | Dermal Worksheet |
| Estimate exposure point concentrations | Planning Table 3 - Exposure Point Concentration Summary |
| Estimate exposure intakes | Planning Table 7 - Calculation of Chemical Cancer Risks and Non-Cancer Hazards |
| | Planning Table 8 - Calculation of Radiation Cancer Risks |
| Toxicity A | Assessment |
| Determine toxicity values for carcinogenic and non- | Planning Table 5 - Non-Cancer Toxicity Data |
| carcinogenic effects and provide source information | Planning Table 6 - Cancer Toxicity Data |
| | |

STANDARDIZED RISK ASSESSMENT REPORTING (continued)

| Risk Assessment Activity | Corresponding Planning Table/Worksheet | | | | | |
|--|---|--|--|--|--|--|
| Risk Characterization | | | | | | |
| Quantify cancer and non-cancer risk by pathway | Planning Table 7 - Calculation of Chemical Cancer Risks and Non-Cancer Hazards Planning Table 8 - Calculation of Radiation Cancer Risks Radiation Dose Assessment Worksheet | | | | | |
| Combine risks by media for different receptors | Planning Table 9 - Summary of Receptor Risks and | | | | | |
| Combine risks by media for different receptors | Hazards for COPCs | | | | | |
| Summarize risk drivers for different receptors | Planning Table 10 - Risk Summary | | | | | |
| Prepare draft risk documentation for ROD | ROD Risk Worksheets | | | | | |

SUMMARY OF RAGS PART D REVISION 1 CHANGES

| PLANNING TABLE/WORKSHEET | REVISION 1 CHANGES | | | | |
|---------------------------------------|--|--|--|--|--|
| Planning Table 0 | This is a new Planning Table. | | | | |
| TARA Schedule Worksheet | This is a new Worksheet. | | | | |
| Planning Table 1 | Revision 1 does not include the On-Site/Off-Site field from Revision 0. | | | | |
| Data Useability Worksheet | The Revision 1 Worksheet is the same as the Revision 0 Worksheet. | | | | |
| Planning Table 2 | Exposure Point was moved from the last row of the Summary Box (Revision 0) to the first column of the table (Revision 1). This may reduce the number of versions of Planning Table 2 needed for some sites. The Qualifier information for Minimum and Maximum Concentrations has been moved to the corresponding Concentration fields. | | | | |
| Planning Table 3 | In Revision 1, separate versions of this table should be prepared for RME and CT. Exposure Point was moved from the last row of the Summary Box (Revision 0) to the first column of the table (Revision 1). This may reduce the number of versions of Planning Table 3 needed for some sites. The Qualifier information has been moved to the corresponding Maximum Concentration field. | | | | |
| Planning Table 4 | In Revision 1, separate versions of this table should be prepared for RME and CT. Receptor Population, Receptor Age, and Exposure Point were moved from the Summary Box (Revision 0) to columns in Revision 1. This may reduce the number of versions of Planning Table 4 needed for some sites. | | | | |
| Planning Tables 5.1, 5.2, and 5.3 | The Revision 1 Planning Tables are essentially the same as Revision 0. Some column headings have been slightly reworded, but the data needs are the same. | | | | |
| Planning Table 6.1, 6.2, 6.3, and 6.4 | The Revision 1 Planning Tables 6.1, 6.2, and 6.3 are essentially the same as Revision 0. Some column headings have been slightly reworded, but the data needs are the same. Revision 1 Planning Table 6.4 for radionuclides was not included in Revision 0. | | | | |

SUMMARY OF RAGS PART D REVISION 1 CHANGES (continued)

| PLANNING TABLE/WORKSHEET | REVISION 1 CHANGES | | | |
|--|---|--|--|--|
| Planning Table 7 | Medium, Exposure Medium, and Exposure Point were moved from the Summary Box (Revision 0) to columns in the table (Revision 1). This may reduce the number of versions of Planning Table 7 needed for some sites. Planning Table 7, which previously contained only non-cancer information (Revision 0), now presents cancer and non-cancer information for chemicals. | | | |
| Planning Table 8 | Planning Table 8 (Revision 1) focuses exclusively on the calculation of radiation cancer risks. Planning Table 8 (Revision 0) focused on cancer risk calculations for all chemicals. Medium, Exposure Medium, and Exposure Point were moved from the Summary Box (Revision 0) to columns in the table (Revision 1). This may reduce the number of versions of Planning Table 8 needed for some sites. Medium EPC and Route EPC information (Revision 0) was replaced by EPC information (Revision 1). | | | |
| Radiation Dose Assessment Worksheet | This is a new Worksheet. | | | |
| Planning Tables 9 and 10 | A column for Exposure Route External (Radiation) has been added to the cancer calculations in Revision 1. The second COPC (Planning Table 9) or Chemical (Planning Table 10) column from Revision 0 has been deleted in Revision 1. Accommodations have been made for summing risks and hazards at the Exposure Point, Exposure Medium, Medium, and Receptor Levels. | | | |
| Lead Worksheets | These are new Worksheets. | | | |
| ROD Risk Worksheets (ROD Risk Highlights) | These are new Worksheets that copy the ROD Guidance (U.S. EPA, 1999a) Risk Highlights. | | | |

EXHIBIT 4-1 EXAMPLE TABLES TO STANDARDIZE REPORTING OF FS RISK EVALUATIONS

Example Table 1 REMEDIAL ACTION OBJECTIVES

| | | | | • | | | | | |
|---------------------------------|-----------------------------|---------------------------------------|-------------------|-----------------------|---------------------------|------------------------------|----------------------------------|------------------|----------------------------|
| Medium: | | | | | | | | | |
| Exposure Point Chemical | | | Exposure Route | | Receptor Population | | Remedial Action Objectives | | |
| | | | | | | | | | |
| | | | VALU | Example UES CONSI | e Table 2 DERED AS 1 | PRGs | | | |
| Medium: Receptor Pop | oulation: | | | | | | | | |
| Chemical | Most Restrictive ARAR | Most Restrictive ARAR Source | | isk/Hazard at ARAR | Risk-Bas PRG Cancer | | Risk-Based PRG Non-Cancer* | Other Value** | Other Value** Source |
| | | | | | | | | | |
| **(e.g., detection | | | | Example | e Table 3 SSOCIATEI | o wij | TH PRGs | | |
| Medium: Receptor Pop | oulation: | | | | | | | | |
| Chemical Site PRG Concentration | | | Basis for PRG* | Risk at PF Cancer | | Hazard at PRG: Nor Cancer | n- Target | Target Endpoint | |
| | | | | | | | | | |
| | | | | | | | | | |
| 1 | <u> </u> | <u>I</u> | | Totals | | | | | |

*TBC (Federal ARARs, State ARARs), Risk-based. Background Concentrations, method detection limits