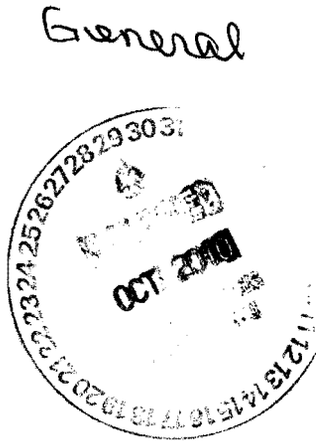




**Environmental Programs**  
P.O. Box 1663, MS M991  
Los Alamos, New Mexico 87545  
(505) 606-2337/FAX (505) 665-1812



**National Nuclear Security Administration**  
Los Alamos Site Office, MS A316  
Environmental Restoration Program  
Los Alamos, New Mexico 87544  
(505) 667-4255/FAX (505) 606-2132

Date: **OCT 29 2010**  
Refer To: EP2010-0496

James Bearzi, Bureau Chief  
Hazardous Waste Bureau  
New Mexico Environment Department  
2905 Rodeo Park Drive East, Building 1  
Santa Fe, NM 87505-6303

**Subject: Submittal of the Ecorisk Database, Release 2.5**

Dear Mr. Bearzi:

Enclosed please find two compact discs (CDs) containing the updates to the Los Alamos National Laboratory's (the Laboratory's) ECORISK Database. The Environmental Programs Directorate maintains and updates the database to ensure the ecological screening levels (ESLs) used to assess potential ecological risk at sites are representative and current. It should be noted that the database and associated files are now available online and can be downloaded from the Laboratory's external website at <http://www.lanl.gov/environment/cleanup/ecorisk.shtml>. The reports submitted to the New Mexico Environment Department-Hazardous Waste Bureau will use the ESLs presented in this release starting October 2010.

If you have any questions, please contact Richard Mirenda at (505) 665-6953 ([rmirenda@lanl.gov](mailto:rmirenda@lanl.gov)) or Hai Shen at (505) 665-5046 ([hshen@doeal.gov](mailto:hshen@doeal.gov)).

Sincerely,

Michael J. Graham, Associate Director  
Environmental Programs  
Los Alamos National Laboratory

Sincerely,

George J. Rael, Manager  
Environmental Projects Office  
Los Alamos Site Office



MG/GR/DN/RM:sm

Enclosure: Two CDs containing ECORISK Database, Release 2.5 (LA-UR-10-6898)

Cy: (w/o enc.)

Laurie King, EPA Region 6, Dallas, TX

Tom Skibitski, NMED-OB, Santa Fe, NM

Steve Yanicak, NMED-DOE-OB, MS M894

Neil Weber, San Ildefonso Pueblo

Hai Shen, DOE-LASO, MS A316

Annette Russell, DOE-LASO (date-stamped letter emailed)

Michael J. Graham, ADEP, MS M991 (date-stamped letter emailed)

Pat Nakagawa, EP-ET, MS M992 (date-stamped letter emailed)

Rich Mirenda, EP-ET, MS M992

William Alexander, EP-BPS, MS M992

Public Reading Room, MS M992

RPF, MS M707

***TO VIEW THE DATABASE  
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HAZARDOUS WASTE  
BUREAU AT 505-476-6000  
TO MAKE AN  
APPOINTMENT***



## **Environmental Programs**

*Engineering and Technology Division*

P.O. Box 1663, MS M992

Los Alamos, New Mexico 87545

505-605-0607/Fax 505-606-0503

October 13, 2010

To Whom It May Concern:

Enclosed is a CD-ROM that contains files for the Los Alamos National Laboratory (LANL) Ecorisk Database Release 2.5 (October 2010).

### **The CD-ROM contains the following folder and files:**

- **ECORISK\_R2.5\_101310.MDB:** A MS Office Access XP/2003 file that is the Ecorisk Database Release 2.5.
- **CoverLetterR2.5\_101310.pdf:** The cover letter you are currently reading.
- **ESLHistorySummary101310.pdf:** A document describing all LANL Ecological Screening Level (ESL) changes since the beta release of the Ecorisk Database to the latest release. This file can be accessed either directly from the CD or from within the database on the 'Main Menu' screen under the 'Supplemental Reports' section.
- **ESLs\_R2.5.xlsx:** A MS Office Excel (XP) file that contains all ESLs from the Ecorisk Database Release 2.5. This file can be accessed either directly from the CD or from within the database on the 'Main Menu' screen under the 'Supplemental Reports' section.
- **GMMTRVDerivationMethods090104b.pdf:** Explanations of the content of Toxicity Reference Value (TRV) Summary Reports associated with Geometric Mean (GMM) TRVs derived by LANL based on reviews of primary toxicity studies. This file can be accessed either directly from the CD or from within the database on the 'Main Menu' screen under the 'Supplemental Reports' section.
- **TRVs\_Methods\_LANL&EcoSSLData.pdf:** A document that explains the methods used to derive TRVs for PAHs and DDT and metabolites using both US Environmental Protection Agency (EPA) Ecological Soil Screening Level (EcoSSL) and LANL data. This file can be accessed either directly from the CD or from within the database on the 'Main Menu' screen under the 'Supplemental Reports' section.
- **TRV\_Dev\_Methods\_091710a.pdf:** A document that explains the methods used to identify/derive TRVs at LANL. This file can be accessed directly from the CD.
- **AppA\_TRV\_Dev\_Methods\_091710a.pdf:** A document that is the appendix to TRV\_Dev\_Methods\_090910a.pdf. This file can be accessed directly from the CD.
- **Interim\_SoilESLs\_R2.5\_101310.xls:** A MS Office Excel (XP) file that contains the interim ESLs and surrogate ESLs that accompany the Ecorisk Database Release 2.5. This file can be accessed either directly from the CD or from within the database on the 'Main Menu' screen under the 'Supplemental Reports' section.
- **EcoriskDbR2.5\_ToxicityData\_ResourceSummary\_SoilESLs\_101310.xlsx** A MS Office Excel (XP) file that contains the search results for toxicity data, as well as the TRVs identified for Ecorisk Database Release 2.5. This file can be accessed either directly from the CD or from within the database on the 'Main Menu' screen under the 'Supplemental Reports' section.

**Installation of Program:**

A directory folder of C:/EcoriskDb must be created in the user's hard drive, and ALL of the files in the CD-ROM must be saved to this location. This is necessary in order to ensure functionality of links to outside files from within the database. If you have files for a previous version of the Ecorisk Database already in this folder, you must delete or move the old files prior to installation. The database file cannot be opened directly from the CD-ROM due to the user-level security component of the database structure. Once the file has been copied to C:/EcoriskDb, please make sure that the read-only file property is NOT checked for the database (.mdb) file.

**Data Issues:**

In this release of the database, ESLs/TRVs were added for chemicals for which no toxicity data was previously available. Online toxicity databases were searched for relevant existing TRVs or for primary toxicity data and/or references from which TRVs could be derived for these chemicals (see EcoriskDbR2.5\_ToxicityData\_ResourceSummary\_SoilESLs\_101310.xls for details of search results). New TRVs have been incorporated into the database. Interim ESLs are reported in a separate file (Interim\_SoilESLs\_R2.5\_101310.xls).

Please refer to the ESL History Summary Report (ESLHistorySummary101310.pdf) for a synopsis of the changes made to the data in the Ecorisk Database since the last release. This file can be accessed either directly from the CD or from within the database on the 'Main Menu' screen under the 'Supplemental Reports' section. Please refer to the 'What's New' screen in the database for specific details on value changes.

**Interface Issues:**

Note, when using the report option in the database, you will receive a blank report if there is no data for the report criteria you selected. You will also receive a blank report if you do not provide all the report criteria.

**Other Issues:**

This database is a work in progress and although we have reviewed the data within it extensively, we still recommend that you verify the data before use by referring to the actual references cited. The project may be able to assist you in obtaining copies of some of the harder to find documents cited in the database.

**Contact Information:**

Please contact Rich Mirinda at [rmirenda@lanl.gov](mailto:rmirinda@lanl.gov) if you have any trouble with your copy of the database or if you have any questions and/or comments about the database.

Thank you for your interest in the database.

The ECORISK Database Team

**Los Alamos National Laboratory**  
**Environmental Stewardship Division**  
**Environmental Remediation and Surveillance Program**  
**Ecorisk Database Release 2.4 (December 2009)**  
**ESL History Summary by Ecorisk Database Release**  
**(October 2010)\***

*\* If you have a specific question(s) that this document does not address adequately, you may contact the database manager for additional help answering your question(s).*

**[Table 1. ESL Changes by Ecorisk Database Release](#)**

**[October 1998 – Beta Release](#)**

**[June 1999 – Release 1.0](#)**

**[April 2000 – Release 1.1](#)**

**[September 2000 – Release 1.2](#)**

**[September 2001 – Release 1.3](#)**

**[March 2002 – Release 1.4](#)**

**[September 2002 – Release 1.5](#)**

**[November 2003 – Release 2.0](#)**

**[September 2004 – Release 2.1](#)**

**[September 2005 – Release 2.2](#)**

**[October 2008 – Release 2.3](#)**

**[December 2009 – Release 2.4](#)**

**[October 2010 – Release 2.5](#)**

**[Table 2. Beta Release \(October 1998\) List of Soil ESLs for Bird Receptors](#)**

**[Table 3. Beta Release \(October 1998\) List of Soil ESLs for Mammalian Receptors](#)**

**[Table 4. Beta Release \(October 1998\) List of Soil ESLs for Earthworm Receptor](#)**

**[Table 5. Beta Release \(October 1998\) List of Soil ESLs for Generic Plant Receptor](#)**

**[Table 6. Beta Release \(October 1998\) List of Sediment and Water ESLs for Aquatic Community Organism Receptors](#)**

**[References](#)**

# Tables

**Table 1. ESL Changes by Ecorisk Database Release**

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<b>Ecorisk Database Release</b>	<b>ESL Changes</b>
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**Table 1. ESL Changes by Ecorisk Database Release**

<b>Ecorisk Database Release</b>	<b>ESL Changes</b>
<b>October 1998 – Beta Release</b>	<p>Original ESL models were as follows:</p> <p>Soil ESLs for Bird Receptors: American kestrel (Avian intermediate carnivore), American kestrel (Avian top carnivore), American robin (Avian insectivore) for 46 non-radionuclides and 18 radionuclides (<a href="#">See Table 2</a>).</p> <p>Soil ESLs for Mammalian Receptors: Deer mouse (Mammalian omnivore), Desert cottontail (Mammalian herbivore), Red fox (Mammalian top carnivore), Vagrant shrew (Mammalian insectivore) for 102 non-radionuclides and 18 radionuclides (<a href="#">See Table 3</a>).</p> <p>Soil ESLs for Invertebrate Receptor: Earthworm (Soil-dwelling invertebrate) for 37 non-radionuclides and 18 radionuclides (<a href="#">See Table 4</a>).</p> <p>Soil ESLs for Plant Receptor: Generic plant (Terrestrial autotroph - producer) for 41 non-radionuclides and 18 radionuclides (<a href="#">See Table 5</a>).</p> <p>Sediment and Water ESLs for 12 radionuclides for Aquatic Community Organism Receptors: Aquatic snails (Aquatic herbivore - grazer), Daphnids (Aquatic omnivore/ herbivore), Fish (Aquatic intermediate carnivore), and Algae (Aquatic autotroph – producer). (<a href="#">See Table 6</a>).</p> <p><a href="#">BACK TO TOP</a></p>
<b>June 1999 – Release 1.0</b>	<p>Addition of sediment ESLs for 19 radionuclides and or 49 non-radionuclides for the new bird receptor, Violet-green Swallow (Avian aerial insectivore).</p> <p>Addition of sediment ESLs for 19 radionuclides and or 106 non-radionuclides for the new Mammal receptor, Occult little brown myotis bat (Mammalian aerial insectivore).</p> <p>Addition of 85 sediment ESLs for non-radionuclides ESLs for the new aquatic community organism receptor.</p> <p>Addition of 7 radionuclides (Cesium-134, Cobalt-60, Europium-152, Radium-228, Sodium-22, Thorium-228, Thorium-230) for sediment and water for aquatic community organism receptors.</p> <p>Addition of non-radionuclide and radionuclide ESLs (19 rad, 48 non-rad) for soil for the new Bird receptors, American robin (Avian omnivore) and American robin</p>

**Table 1. ESL Changes by Ecorisk Database Release**

**Ecorisk Database Release**      **ESL Changes**

(Avian herbivore).

Addition of non-radionuclide and radionuclide ESLs for water for all bird (19 rad, 48 non-rad) and mammal (19 rad, 106 non-rad) receptors.

Addition of 3 ESLs for soil for Boron, Fluoride and Radium-228 for all applicable bird receptors.

Addition of 3 ESLs for soil for Boron, Fluoride, Strontium (stable), Dichlorobenzene[1,4-], and Radium-228 for all applicable mammal receptors.

Addition of 2 ESLs for soil for Trinitrotoluene[2,4,6-], and Radium-228 for the earthworm receptor.

Addition of 3 ESLs for soil for Amino-2, 6-dinitrotoluene[4-], Boron, and Radium-228 for the generic plant receptor.

Numerous ESL updates. Documentation of specific reasons for updates not available at this time. General documentation of reasons for ESL updates indicated that the radionuclide ESL models underwent extensive revisions and the non-radionuclide ESLs were multiplied by a factor of 0.3 per the recommendation of NMED.

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**Table 1. ESL Changes by Ecorisk Database Release**

<b>Ecorisk Database Release</b>	<b>ESL Changes</b>
<b>April 2000 – Release 1.1</b>	<p data-bbox="363 411 1476 520">Addition of 5 ESLs for water for Tetrachlorodibenzodioxin[2,3,7,8-], Dinitrotoluene[2,6-], Fluoride, Pentachloronitrobenzene, and Dichloroethene[1,1-] for the aquatic community organism receptor.</p> <p data-bbox="363 558 1382 625">Addition of soil and water ESLs for Dinitrobenzene[1,3-] for all applicable bird receptors.</p> <p data-bbox="363 663 1305 701">Addition of a soil ESL for Dibenzofuran for the desert cottontail receptor.</p> <p data-bbox="363 739 1476 919">Deletion of sediment ESLs for Butanone[2-], Chloroform, Dichloroethane[1,2-], Dichloroethene[cis-1,2-], Dinitrotoluene[2,6-], and Nitrobenzene for the aquatic community organism receptor. The Chloroform ESL was deleted because the toxicity data it was based on was deemed unsuitable. Reasons for other deletions not available at this time.</p> <p data-bbox="363 957 1365 1024">Deletion of water ESL for Dichloroethene[cis-1,2-] for the aquatic community organism. Reason for deletion not available at this time.</p> <p data-bbox="363 1062 1438 1129">Numerous ESL updates. Documentation of specific reasons for ESL updates is not available at his time. General reasons for ESL updates are described below.</p> <p data-bbox="363 1167 1476 1394">Some ESLs were updated based on reasons documented in the December 1999 Interim ESLs memorandum (<a href="#">Ref ID 1484</a>) and included: 1) the 0.3 factor was removed from the non-radionuclide ESL equations, 2) a correction to the water ESLs to account for a units conversion problem was made (values were multiplied by 1000), 3) all ESL values were rounded down to two significant figures and 4) the aquatic community organism receptor ESL for chlordane was revised.</p> <p data-bbox="363 1432 1476 1722">Some ESLs were updated due to the availability of new PTSE derived CS TRVs to replace secondary data source TRVs in ESL calculations. PTSE CS TRVs derived included Amino-2,6-dinitrotoluene[4-]/ Plant, Amino-4,6-dinitrotoluene[2-]/ Plant, Boron/ Bird, /Mammal and /Plant; Chromium (total)/ Bird and /Mammal, Fluoride/ Bird and / Mammal, Manganese/ Bird, / Mammal and / Plant; Nitroglycerine/ Mammal, Strontium (stable)/ Mammal, Trinitrotoluene[2,4,6-]/ Earthworm, /Mammal and /Plant; Uranium/ Bird, / Mammal and / Plant; and Vanadium/ Bird and / Mammal.</p> <p data-bbox="363 1759 1398 1833">Some ESLs were updated due to quality assurance issues including correction of errors in ESL calculations/parameters, rounding of values or reporting of data.</p>

**Table 1. ESL Changes by Ecorisk Database Release**

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**Ecorisk Database Release**      **ESL Changes**

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**Table 1. ESL Changes by Ecorisk Database Release**

<b>Ecorisk Database Release</b>	<b>ESL Changes</b>
<b>September 2000 – Release 1.2</b>	<p>Addition of soil, sediment and water ESLs for Dichloroethane[1,2-] for all applicable bird and mammal receptors because new PTSE derived TRVs were available.</p> <p>Addition of soil, sediment and water ESLs for Lead-210, Neptunium-237, Thorium-229, Uranium-233, and Uranium-236 for all applicable bird, mammal, earthworm, generic plant and aquatic community organism receptors.</p> <p>Addition of soil ESLs for HMX and RDX for the earthworm receptor. Reason for addition not available at this time.</p> <p>Addition of a water ESL for Dinitrobenzene[1,3-] for the aquatic community organism receptor. Reason for addition not available at this time.</p> <p>Deletion of soil, sediment and water ESLs for Chloro-3-methylphenol[4-] for all applicable bird, mammal, and aquatic community organism receptors. Reasons for deletions not available at this time.</p> <p>Deletion of soil, sediment and water ESLs for Tetrachloroethane[1,1,2,2-] for all applicable mammal, and aquatic community organism receptors. Reasons for deletions not available at this time.</p> <p>Deletion of sediment ESLs for Dinitrobenzene[1,3-], Iron, Polychlorinated Biphenyls, Dimethyl Phthalate, and Phenol for the aquatic community organism receptor.</p> <p>Deletion of water ESLs for Calcium, Nitrate (expressed as NO<sub>3</sub>), and Dichloroethene[1,1-] for the aquatic community organism receptor. Reasons for deletions not available at this time.</p> <p>Deletion of the soil ESL or Dibenzofuran for the desert cottontail receptor. Reason for deletion not available at this time.</p> <p>Numerous ESL updates.</p> <p>Some ESLs were updated because new PTSE derived CS TRVs were available to replace secondary data source TRVs. PTSE CS TRVs available included Acetone/Bird, Barium Bird, Barium/Mammal, Barium/Plant, HMX/Invertebrate, HMX/Mammal, Lead/Mammal, Lead/Bird, Lead/Invertebrate, Lead/Plant, RDX/Invertebrate, RDX/Mammal, Silver/Bird, Silver/Plant, 1,3,5-Trinitrobenzene/Mammal, Thallium/Plant, Zinc/Bird, Zinc Invertebrate.</p>

## Table 1. ESL Changes by Ecorisk Database Release

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**Ecorisk Database Release**    **ESL Changes**

Other ESLs were updated for quality assurance issues including correction of errors in ESL calculations/parameters, rounding of values or reporting of data.

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**Table 1. ESL Changes by Ecorisk Database Release**

<b>Ecorisk Database Release</b>	<b>ESL Changes</b>
<b>September 2001 – Release 1.3</b>	<p>Addition of soil ESL for Chromium (total) for the earthworm receptor due to the availability of a new internally approved secondary data source TRV.</p> <p>Addition of soil ESL for DDT[4,4'-] for the generic plant receptor due to the availability of a new internally approved secondary data source TRV.</p> <p>Addition of water ESL for Dichloroethene[1,1-] for the aquatic community organism receptor due to the availability of a new internally approved secondary data source TRV.</p> <p>Numerous ESL updates.</p> <p>Some ESLs were updated because new PTSE derived CS TRVs were available to replace secondary data source TRVs. PTSE CS TRVs available included DDE[4,4'-]/Bird, DDE[4,4'-]/Mammal, DDT[4,4'-]/Bird, DDT[4,4'-]/Mammal, DDT[4,4'-]/Plant, Aroclor-1016, Aroclor-1242, Aroclor-1248, Aroclor-1254 and Aroclor-1260/Mammal; Aroclor-1242, Aroclor-1248, Aroclor-1254 and Aroclor-1260/Bird; and Aroclor-1254/Plant.</p> <p>Other ESLs were updated for quality assurance issues including correction of errors in ESL calculations/parameters, rounding of values or reporting of data.</p> <p><a href="#">BACK TO TOP</a></p>
<b>March 2002 – Release 1.4</b>	<p>Numerous ESL updates.</p> <p>Radionuclide ESLs, except Tritium, were updated due to revision of TF_plant and TF_invert from a dry weight basis to a fresh weight basis assuming 85% and 61% moisture content of plant and invertebrate diets, respectively (<a href="#">Ref ID 0561</a>). This revision was required for units to cancel correctly in the ESL model equations.</p> <p>Radionuclide ESLs for Tritium were updated due to revision of TF_plant and TF_invert to assume equilibrium between the tritium in soil moisture and tissue waters. The value is calculated by dividing the moisture in tissues by the moisture in soil where 61% moisture content of invertebrates is based on beetles (<a href="#">Ref ID 0561</a>, Table 4-1, p. 4-13) and 85% moisture content of plant material is based on leaves (<a href="#">Ref ID 0561</a>, Table 4-2, p.4-14) and soil moisture of 10% is based on an average soil moisture found in the Los Alamos area. This revision was required for units to cancel correctly in the ESL model equations.</p>

**Table 1. ESL Changes by Ecorisk Database Release**

**Ecorisk Database Release**      **ESL Changes**

Radionuclide ESLs were also updated due to the revision of TF\_flesh, which was revised because it is calculated from TF\_plant and TF\_invert, which were revised as explained above. This revision was required for units to cancel correctly in the ESL model equations.

Radionuclide ESLs were also update due to the revision of all receptor intake rates from a dry weight basis to a fresh weight basis where the moisture content of invertebrates is assumed to be 61% (beetles ([Ref ID 0561](#), Table 4-1, p. 4-13)), of plant materials is assumed to be 85% (leaves ([Ref ID 0561](#), Table 4-2, p.4-14)), and flesh is assumed to be 68% (mammals - mice, voles, rabbits ([Ref ID 0561](#), Table 4-1, p. 4-13)). This revision was required for units to cancel correctly in the ESL model equations.

Radionuclide ESLs were also updated due to the replacement of TF\_beef with TF\_blood in ESL models. TF(blood) is calculated by multiplying TF(beef) by I(food) or in the case of water intake, I(water). TF(blood) is required in all radionuclide ESL models for wildlife, and TF(beef) was used as a surrogate measure to estimate body burdens for internal dose calculations. TF(beef) has been replaced by TF(blood) in all these models so that the units in these models cancel properly. Internal dose calculations require a TF that models the transfer of radionuclides from food to blood.

Other reasons for ESL updates include the rounding of ESL model parameters to 3 significant digits for reporting consistency as well addressing quality assurance issues.

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**Table 1. ESL Changes by Ecorisk Database Release**

<b>Ecorisk Database Release</b>	<b>ESL Changes</b>
<b>September 2002 – Release 1.5</b>	<p>Addition of soil, sediment and water ESLs for Trinitrotoluene[2,4,6-] for all applicable bird receptors due to the availability of a new PTSE derived CS TRV.</p> <p>Addition of soil ESL for Tetrachloroethene for the generic plant receptor due to the availability of a new PTSE derived CS TRV.</p> <p>Numerous ESL updates.</p> <p>Some ESLs were updated due to the availability of new PTSE derived CS TRVs to replace secondary data source TRVs in ESL calculations. Applicable PTSE TRVs derived included Tetrachlorodibenzodioxin[2,3,7,8-]/Bird, Mammal, and Plant; Antimony/Mammal, Cadmium/Bird, Mammal and Invertebrate; Copper/Bird and Mammal; Mercury (inorganic) /Bird, Mammal and Invertebrate; Nickel /Bird, Mammal and Invertebrate; Selenim/Invertebrate, Zinc/Mammal and Plant; Tetrachloroethene/Mammal, Trichloroethane[1,1,1-]/Mammal, Trichloroethene/Mammal, and Xylene (total)/Bird.</p> <p>Some ESLs were updated due to quality assurance issues for TRVs. Specific details of issues are not available at this time.</p> <p><a href="#">BACK TO TOP</a></p>
<b>November 2003 – Release 2.0</b>	<p>Addition of soil ESLs for Antimony, Barium, and Beryllium for the earthworm receptor due to the availability of EPA Eco-SSL TRVs.</p> <p>Deletion of the soil ESL for Trinitrotoluene[2,4,6-] for the earthworm receptor because the toxicity data it was based on was deemed unsuitable.</p> <p>Deletion of soil ESLs for Aluminum for all applicable bird, mammal and generic plant receptors because EPA Eco-SSL uses a soil pH of less than 5.5 as an indicator of toxicity instead of an Aluminum soil concentration.</p> <p>Numerous ESL updates.</p> <p>Some ESLs were updated due to the availability of new PTSE derived GMM TRVs to replace PTSE derived CS TRVs or secondary data source TRVs in ESL calculations. Applicable PTSE GMM TRVs included, Aroclor-1016, Aroclor-1242, Aroclor-1254, Aroclor-1260, DDT[4,4'-], Di-n-Butyl Phthalate, Nickel, RDX, and Tetrachlorodibenzodioxin[2,3,7,8-] for food exposure for Mammals; Antimony, Cadmium, and Lead for drinking water exposure for Mammals; Aroclor-1260,</p>

**Table 1. ESL Changes by Ecorisk Database Release**

**Ecorisk Database Release**      **ESL Changes**

Barium, Boron, Copper, DDE[4,4'-], Nickel, and Zinc for food exposure for Birds; Aroclor-1254, Boron, and Di-n-Butyl Phthalate for soil exposure for Plants; and Zinc for soil exposure for Invertebrates.

Some ESLs were updated due to the availability of EPA Eco-SSL TRVs to replace PTSE or secondary data source TRVs in ESL calculations. Applicable EPA Eco-SSL TRVs available included Antimony, Barium, Beryllium, Cadmium, Cobalt, Lead, and Dieldrin for food exposure for Mammals; Cadmium, Cobalt, Lead, and Dieldrin for food exposures for Birds; Antimony, Barium, Beryllium, Cadmium, and Lead for soil exposure for Invertebrates; and Cadmium, Cobalt, and Lead for soil exposure for Plants.

Some ESLs were updated due to the availability of EPA NRWQC CCC TRVs to replace other secondary data source TRVs. Applicable EPA NRWQC CCC TRVs available included Selenium and Mercury (inorganic) for water exposure for the aquatic community organism receptor.

Other ESLs were updated due to addressing data quality assurance issues or because the previously used toxicity data the ESLs were based on was deemed unsuitable and was revised appropriately to make it suitable. Specific details of issues are not available at this time.

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**Table 1. ESL Changes by Ecorisk Database Release**

<b>Ecorisk Database Release</b>	<b>ESL Changes</b>
<b>September 2004 – Release 2.1</b>	<p>A mammalian screening receptor used in soil and water ESL models for a mammalian insectivore in the database has changed. The vagrant shrew (<i>Sorex vagrans</i>) in New Mexico has been reclassified as the montane shrew, also known as the dusky shrew, (<i>Sorex monticolus</i>) by Eastern New Mexico University (see <a href="http://fwie.fw.vt.edu/states/nmex_main/species/050725.htm">http://fwie.fw.vt.edu/states/nmex_main/species/050725.htm</a> for more information). However, this the ESLs for the vagrant shrew are applicable to the montane shrew because the short-tailed shrew data that was used as surrogates for parameters in the vagrant shrew ESL models are applicable for the montane shrew as a mammalian insectivore. As a result, only the ESL screening receptor common and scientific name has changed.</p> <p>Addition of soil ESL for HMX for the generic plant receptor due to the availability of a new Tier 2 TRV (PTSE GMM TRV).</p> <p>Addition of soil ESL for Trinitrotoluene[2,4,6-] for the earthworm receptor due to the availability of a new Tier 3 TRV (PTSE CS TRV).</p> <p>Addition of sediment and soil ESLs for RDX for all applicable bird receptors due to the availability of a new Tier 2 TRV (PTSE GMM TRV).</p> <p>Addition of sediment, soil and water ESLs for Thallium for all applicable bird receptors due to the availability of a newly approved Tier 4 TRV (secondary data source CS TRV).</p> <p>Addition of 16 air ESLs for Acetone, Benzene, Carbon, Tetrachloride, Chloroform, Chloromethane, Dichlorodifluoromethane, Dichloroethane[1,1-], Dichloroethane[1,2-], Dichloroethene[1,1-], Methylene Chloride, Tetrachloroethene, Toluene, Trichloroethane[1,1,1-], Trichloroethene, Trichlorofluoromethane, and Xylene (Total) for the new Mammal receptor, Botta's Pocket Gopher (Burrowing mammal). These ESL were added due to the availability of new Tier 2 TRVs (PTSE GMM TRVs).</p> <p>Deletion of sediment, soil and water ESLs for Tetrachlorodibenzodioxin[2,3,7,8-] for all applicable bird receptors due to discontinued use of previous Tier 3 (CS) TRV that was deemed unsuitable because it was based on an non-oral exposure (i.p. injection).</p> <p>Numerous ESL updates.</p> <p>Naphthalene soil and sediment ESLs for all applicable bird receptors updated due to the previous Tier 4 TRV (secondary data source CS TRV) being replaced by a new Tier 2 TRV (PTSE GMM TRV).</p>

**Table 1. ESL Changes by Ecorisk Database Release**

**Ecorisk Database Release**      **ESL Changes**

Chromium (+6) soil, sediment and water ESLs for all applicable bird receptors updated due to the previous Tier 4 (CS) TRV being replaced by a new Tier 3 TRV (PTSE CS TRV).

Chromium (total) soil, sediment and water ESLs are based on Chromium (+6) toxicity data and because the oral chromium (+6) TRV for birds was updated (see previous paragraph), the corresponding chromium (total) ESLs for birds were updated accordingly based on the new chromium (+6) data.

HMX soil ESL for the earthworm receptor updated due to the previous Tier 3 TRV (PTSE CS TRV) being replaced by a new Tier 2 TRV (PTSE GMM TRV).

RDX soil ESL for the earthworm receptor updated due to the previous Tier 3 TRV (PTSE CS TRV) being replaced by a new Tier 3 TRV (PTSE CS TRV).

Trinitrotoluene[2,4,6-] soil ESL for the generic plant receptor updated due to the Tier 3 TRV (PTSE CS TRV) being replaced by a new Tier 2 TRV (PTSE GMM TRV).

Plutonium-241 water ESL for the vagrant shrew receptor updated due to the revision of the ESL model parameter, TF\_blood, which was corrected for a previous rounding error.

All ESL for radionuclides in sediment for aquatic receptors were revised based on the guidance of DOE-STD-1153-2002 to not include internal dose for aquatic organisms exposed to radionuclides in sediment. The ESL model parameter, DCF\_int\_fw, was set to 0 to incorporate this guidance.

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**Table 1. ESL Changes by Ecorisk Database Release**

<b>Ecorisk Database Release</b>	<b>ESL Changes</b>
<b>September 2005 – Release 2.2</b>	<p><b>New ESLs</b></p> <p>Sediment and water ESLs for iron for aquatic community organisms due to this analyte being added as a new LANL exposure concern.</p> <p>Water ESLs for perchlorate ion for mammalian and avian receptors due to development of a New Tier 2 (GMM) TRV and New Tier 3 (CS) TRV, respectively.</p> <p>Soil and sediment ESLs for mammalian receptors for BHC[alpha-] due to the development of a New Tier 3 (CS) TRV.</p> <p>Soil ESLs for the earthworm for fluoranthene, phenanthrene and pyrene due to the development of New Tier 3 (CS) TRVs.</p> <p>Soil ESL for the generic plant for naphthalene due to the development of a New Tier 3 (CS) TRV.</p> <p><b>ESL Updates</b></p> <p>Revision of various Transfer Factors (TF) for soil-to-plant and soil-to-invertebrate for both inorganic and organic analytes based on the most current EPA EcoSSL bioaccumulation data or models (Ref ID 1401), which resulted in the revision of the calculated soil-to-flesh TF and as well as numerous ESL updates.</p> <p>Inorganic TFs were replaced with more comprehensive empirical values, median values from the empirical data set.</p> <p>Organic TFs for soil-to-invertebrates were revised based on a more appropriate bioaccumulation model (<math>BAF_{ww} = (K_{ww}/K_d)/0.16</math> where <math>\log K_{ww} = 0.87 * \log K_{ow} - 2.0</math> and <math>K_d = f_{oc} * K_{oc}</math> where <math>f_{oc}</math> is 1%, or 0.01.) cited in the 2005 EPA EcoSSL bioaccumulation data report (REF ID1401, Table 5 and dry to fresh weight ratio (0.16) for earthworms from Ref ID 1574), except for Dieldrin, DDT[4,4'-], and DDE[4,4'-], which were based on the median of comprehensive empirical data sets.</p> <p>Organic TFs for soil-to-plants were revised based on a more appropriate bioaccumulation model (<math>BAF = 10^{(-0.4057 \log K_{ow} + 1.781)}</math> <math>r^2 = 0.3226</math>, <math>n = 228</math>, <math>p &lt; 0.0001</math>) cited in the 2005 EPA EcoSSL bioaccumulation data report (REF ID1401).</p> <p>Furthermore, various TRVs were also updated and this contributed to the ESL updates. TRV updates include replacement of:</p> <ul style="list-style-type: none"> <li>Tier 1 TRVs with new Tier 1 TRVs from EPA from EcoSSL 2005 data</li> <li>Tier 3 or 4 TRVs with new Tier 1 TRVs from EPA EcoSSL 2005 data</li> <li>Tier 3 or 4 TRVs with new Tier 2 TRVs</li> <li>Tier 3 TRVs with a more appropriate Tier 3 TRVs</li> </ul> <p>Below is a list of the 99 analytes updated grouped based on type of revisions* A.) TF</p>

**Table 1. ESL Changes by Ecorisk Database Release**

**Ecorisk  
Database  
Release**

revisions only, B.) TF and TRV revisions, and C.) TRV revisions only.

*\*Detailed information on changes available from the "What's New In this Release" screen in the Ecorisk Database - section Change Type, ESLs, Update).*

**A.) TF REVISIONS ONLY**

**HIGH EXPLOSIVES/ Sediment and Soil ESLs**

Amino-2,6-dinitrotoluene[4-]  
Amino-4,6-dinitrotoluene[2-]  
Dinitrobenzene[1,3-]  
Dinitrotoluene[2,4-]  
Dinitrotoluene[2,6-]  
HMX  
Nitroglycerine  
Nitrotoluene[2-]  
Nitrotoluene[3-]  
Nitrotoluene[4-]  
PETN  
RDX  
Tetryl  
Trinitrobenzene[1,3,5-]  
Trinitrotoluene[2,4,6-]

**INORGANICS/ Sediment and Soil ESLs**

Aluminum (sediment)  
Arsenic  
Barium  
Cadmium  
Chromium (total)  
Copper  
Manganese  
Mercury (inorganic)  
Nickel  
Selenium (soil)  
Silver  
Strontium (stable)  
Uranium  
Zinc

**POLYAROMATIC HYDROCARBONS/ Sediment and Soil ESLs**

Acenaphthene  
Acenaphthylene  
Anthracene  
Benzo(a)anthracene (soil)  
Benzo(a)pyrene (soil)  
Benzo(b)fluoranthene (soil)  
Benzo(g,h,i)perylene  
Benzo(k)fluoranthene (soil)  
Chrysene (soil)  
Dibenzo(a,h)anthracene (soil)

**Table 1. ESL Changes by Ecorisk Database Release**

**Ecorisk Database Release**

Fluoranthene  
Fluorene  
Indeno(1,2,3-cd)pyrene (soil)  
Methylnaphthalene[2-]  
Naphthalene  
Phenanthrene (soil)  
Pyrene

**POLYCHLORINATED BIPHENYLS/ Soil ESLs**

Aroclor-1016  
Aroclor-1242  
Aroclor-1248  
Aroclor-1254  
Aroclor-1260

**PESTICIDES/ Sediment and Soil ESLs**

BHC[beta-]  
BHC[gamma-]  
Chlordane[alpha-]  
Chlordane[gamma-]  
DDE[4,4'-]  
DDT[4,4'-]  
Dieldrin  
Endosulfan  
Endrin  
Heptachlor (soil)  
Kepone  
Methoxychlor[4,4'-]  
Toxaphene (Technical Grade)

**SEMI-VOLATILE ORGANIC COMPOUNDS/ Sediment and Soil ESLs**

Benzoic Acid  
Bis(2-ethylhexyl)phthalate  
Butyl Benzyl Phthalate  
Chlorobenzene  
Chlorophenol[2-]  
Dimethyl Phthalate  
Di-n-Butyl Phthalate  
Di-n-octylphthalate  
Nitrobenzene  
Pentachloronitrobenzene  
Phenol

**VOLATILE ORGANIC COMPOUNDS/ Sediment and Soil ESLs**

Acetone  
Benzene  
Butanone[2-]  
Chloroform  
Dichlorobenzene[1,4-]  
Dichloroethane[1,1-]  
Dichloroethane[1,2-]  
Dichloroethene[1,1-]  
Dichloroethene[cis/trans-1,2-]

**Table 1. ESL Changes by Ecorisk Database Release**

**Ecorisk Database Release**      **ESL Changes**

Methylene Chloride  
Tetrachloroethene  
Toluene  
Trichlorobenzene[1,2,4-]  
Trichloroethane[1,1,1-]  
Trichloroethene  
Xylene (Total)

**B.) TF REVISIONS & TRV REVISIONS**

**INORGANICS/ Sediment and Soil ESLs**

Antimony (sediment)  
Barium  
Beryllium  
Cadmium  
Chromium (total)  
Cobalt  
Lead  
Vanadium

**PESTICIDES/ Sediment and Soil ESLs**

DDT[4,4'-]  
Dieldrin

**SEMI-VOLATILE ORGANIC COMPOUNDS/ Sediment and Soil ESLs**

Pentachlorophenol

**C.) TRV REVISIONS ONLY**

**DIOXIN/FURANS/ Soil ESLs**

Tetrachlorodibenzodioxin[2,3,7,8-]

**INORGANICS/ Sediment, Soil and Water ESLs**

Antimony (soil)  
Arsenic (soil)  
Barium (soil)  
Cadmium (soil)  
Chromium (total) (soil and water)  
Chromium(+6)  
Lead (soil)  
Vanadium (soil)

**POLYAROMATIC HYDROCARBONS/ Sediment and Soil ESLs**

Fluorene (soil)

**SEMI-VOLATILE ORGANIC COMPOUNDS/ Sediment and Soil ESLs**

Pentachlorophenol

**Other Changes:**

Documentation and value for DCF\_int\_fw for aquatic receptors (algae, aquatic snail, daphnid and generic fish) for water Rad ESL model. This change did not affect ESLs, it was only a documentation error after from the previous release that was made after ESLs had been calculated.

Added TF\_beef\_fw for BHC[alpha-]. Needed to calculate ESL for this new

## Table 1. ESL Changes by Ecorisk Database Release

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**Ecorisk Database Release**    **ESL Changes**

exposure concern.

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**Table 1. ESL Changes by Ecorisk Database Release**

<b>Ecorisk Database Release</b>	<b>ESL Changes</b>
<b>October 2008 – Release 2.3</b>	<p><b>New ESLs</b></p> <p>Soil and Sediment ESLs for DDD[4,4'-] <sup>a</sup>, Diethyl Phthalate, Methyl-2-pentanone[4-], Methylphenol[2-], and Aldrin due to these analytes being added as a new LANL exposure concerns.</p> <p>Soil ESLs for Manganese and Anthracene <sup>a</sup> for the earthworm due to availability of New Tier 1 TRV and New Tier 2 (GMM) TRV, respectively.</p> <p><b>ESL Updates</b></p> <p>Revision of the equation used to calculate the Transfer Factor (TF) for soil-to-flesh for both inorganic and organic analytes, which resulted in the revision of the calculated soil-to-flesh TF and as well as numerous ESL updates.</p> <p>The equation is now:</p> $TF_{flesh\_dw} \text{ equals } TF_{beef\_fw} * [I_{foodcomposite\_fw} * MAX(TF_{plant\_dw} * \{1 - MC_{plant}\}, TF_{invert\_dw} * \{1 - MC_{invert}\}) + I_{soilcomposite\_dw}] / (1 - MC_{flesh})$ <p>Previous equation:</p> $TF_{flesh\_dw} \text{ equals } TF_{beef\_fw} * [I_{foodcomposite\_fw} * If(TF_{invert\_dw} > TF_{plant\_dw}, TF_{invert\_dw} * \{1 - MC_{invert}\}, TF_{plant\_dw} * (1 - MC_{plant})) + I_{soilcomposite\_dw}] / (1 - MC_{flesh})$ <p>Where:</p> <p><math>I_{soilcomposite\_dw}</math> is the maximum dry weight intake of soil (0.00281 kg-dry soil/d) for prey species (American robin, deer mouse, desert cottontail and shrew) of the red fox and American kestrel</p> <p>MAX is maximum</p> <p><math>MC_{plant}</math> is the moisture content of plant matter, which is assumed to be 85% (leaves (Ref ID 0561, Table 4-2, p.4-14))</p> <p><math>MC_{invert}</math> is the moisture content of invertebrates, which is assumed to be 61% (beetles (Ref ID 0561, Table 4-1, p. 4-13))</p> <p><math>MC_{flesh}</math> is the moisture content of flesh, which is assumed to be 68% (mammals - mice, voles, rabbits and birds – passerines (Ref ID 0561, Table 4-1, p. 4-13))</p> <p><math>TF_{beef\_fw}</math> is the food to beef transfer factor (mg-COPC/kg-fresh beef per mg-COPC/d)</p> <p>Furthermore, various TRVs were also updated and this contributed to the ESL updates. TRV updates include replacement of:</p> <ul style="list-style-type: none"> <li>Tier 1 TRVs with new Tier 1 TRVs from EPA from EcoSSL 2005 data</li> <li>Tier 3 or 4 TRVs with new Tier 1 TRVs from EPA EcoSSL 2005 data</li> <li>Tier 3 or 4 TRVs with new Tier 2 TRVs</li> </ul>

**Table 1. ESL Changes by Ecorisk Database Release**

**Ecorisk Database Release**

Tier 3 TRVs with a more appropriate Tier 3 TRVs

Below is a list of the analytes updated grouped based on type of revisions\* A.) TF revisions only, B.) TF and TRV revisions and C.) TRV revisions only.

*\*Detailed information on changes available from the "What's New In this Release" screen in the Ecorisk Database - section Change Type, ESLs, Update).*

**A.) TF REVISIONS ONLY**

**HIGH EXPLOSIVES**

Nitrotoluene[3-] (soil)  
RDX (soil)

**INORGANICS**

Barium (soil)  
Cyanide (total) (soil)

**POLYAROMATIC HYDROCARBONS**

Benzo(a)anthracene (soil)  
Chrysene (soil)

**SEMI-VOLATILE ORGANIC COMPOUNDS**

Carbazole (soil)

**VOLATILE ORGANIC COMPOUNDS**

Chloroform (soil)  
Dichloroethane[1,1-] (soil)

**B.) TF REVISIONS & TRV REVISIONS**

NONE

**C.) TRV REVISIONS ONLY**

**INORGANICS**

Chromium(+6) (sediment, soil)  
Copper (sediment, soil)  
Manganese (sediment, soil)  
Nickel (sediment, soil)  
Selenium (sediment, soil)  
Silver (sediment, soil)  
Zinc (sediment, soil)

**POLYAROMATIC HYDROCARBONS<sup>a</sup>**

Benzo(a)pyrene (sediment, soil)  
Fluoranthene (soil)  
Fluorene (soil)  
Naphthalene (sediment, soil)  
Phenanthrene soil)  
Pyrene (soil)

**PESTICIDES<sup>a</sup>**

DDE[4,4'-] (sediment, soil)  
DDT[4,4'-] (sediment, soil)

**Table 1. ESL Changes by Ecorisk Database Release**

Ecorisk Database Release	ESL Changes
	<p><sup>a</sup> TRVs developed for PAHs and DDT and metabolites DDE and DDD were done according to the following methods: <a href="#">TRVs Methods LANL&amp;EcoSSLData</a></p> <p>Other Changes:</p> <ul style="list-style-type: none"> <li>Updated documentation for Aluminum ESL for soil by removing an ESL value of &gt; 5 and indicating in notes “pH dependent. Aluminum is identified as a COPC only at sites where the soil pH is less than 5.5.</li> <li>Added TF_plant_dw, TF_invert_dw, TF_beef_fw and TF_flesh_dw for DDD[4,4’-], Diethyl Phthalate, Methyl-2-pentanone[4-], Methylphenol[2-], and Aldrin. Needed to calculate ESLs for these new LANL exposure concerns.</li> <li>Updated interface screens:</li> </ul>
	Brand new Analyte Search menu screen with concise instructions that shows menu for searching for ESLs by analyte and accessed via the updated Main Menu screen. Contains the same buttons that were originally on old Main Menu screen and leads to the same Analyte Search Result screens.
	Brand new Contact Information screen that shows point of contact information for Ecorisk Db. Accessed via a button on the updated Home screen.
	Updated Home screen to reduce clutter of information. Contains button to access contact information, ESL search menus and report menus, what’s new in this release information, and a button to exit the Db.
	Updated Main Menu screen to reduce clutter of information. Contains button to new screens that show ESL search menus (by analyte or by screening receptor), and summary and custom report menus. Also contains buttons to see the existing screens for ESL radionuclide and non-radionuclide model information.
	Updated Custom Report Menu screen that now has a design similar to the other search menus (e.g., Screening Receptor Search menu) and concise instructions. Contains the same buttons that were on Old Main Menu screen. Accessed via the updated Main Menu screen.
	Updated Primary Toxicity Study (PTS) Description screen that now shows vertical scroll bars that were missing in some fields. Recommended update.
	Updated Primary Toxicity Value (PTV) Evaluation screen that now shows more information to aid in understanding better how the PTV confidence ratings are determined. More specifically, this form now shows Maximum weighted scores for the different exposure scenarios (i.e., bird or mammal, oral ingestion; mammal, inhalation; and plant or invertebrate). Recommended update.
	Brand new Screening Receptor Search menu screen with concise instructions that shows menu for searching for ESLs by screening receptor and accessed via the updated Main Menu screen. Contains the same buttons that were originally on old main Menu screen and leads to the same Receptor Search Result screens.
	Brand new Summary Reports Menu screen with concise instructions that shows menu for summary reports and accessed via the updated Main Menu Screen. Contains the same buttons that were originally on old main Menu screen but improved in presentation.

## Table 1. ESL Changes by Ecorisk Database Release

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**Ecorisk Database Release**    **ESL Changes**

Updated Select TRV Summary Report Criteria screen in which redundancy was removed (the same sentence was repeated twice). Recommended update.

Updated Weighting Factor Description screen that now explains in more detail what is done with the weighting factors and why. Recommended update.

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**Table 1. ESL Changes by Ecorisk Database Release**

<b>Ecorisk Database Release</b>	<b>ESL Changes</b>
<b>December 2009 – Release 2.4</b>	<p>In this release of the database, ESLs/TRVs were added for chemicals for which no toxicity data was previously available. Online toxicity databases were searched for relevant existing TRVs or for primary toxicity data and/or references from which TRVs could be derived for these chemicals (see <a href="#">EcoriskDbR2.4 ToxicityData ResourceSummary SoilESLs 112409.xls</a> for details of search results). Of those 40 chemicals of concern, 11 chemicals now have LANL peer reviewed/ approved TRVs/ESLs incorporated into this release of the database, 5 chemicals have interim ESLs/ TRVs because LANL peer reviewed/ approved values could not be obtained in time for this release of the database (see <a href="#">Interim SoilESLs R2.4 111309.xls</a>), 13 chemicals have surrogate ESLs/TRVs (see <a href="#">Interim SoilESLs R2.4 111309.xls</a>) based on chemicals already in the database, and the remaining 12 chemicals still have no ESL at this time. Note – The sum of the numbers adds up to 41 instead of 40 because Hexanone[2-] has both an incorporated ESL (for birds) and an interim ESL (for mammals). Below is a summary of the ESLs/ TRVs incorporated into Release 2.4 of the Ecorisk Database, as well as other relevant data or interface changes.</p> <p><b>New ESLs</b></p> <p>Soil and Sediment ESLs for birds due to availability of new TRVs:</p> <ul style="list-style-type: none"><li>• Molybdenum</li><li>• Hexachlorobenzene</li><li>• Hexanone[2-]</li></ul> <p>Soil and Sediment ESLs for mammals due to availability of new TRVs:</p> <ul style="list-style-type: none"><li>• Lithium</li><li>• Carbon Disulfide</li><li>• Hexachlorobenzene</li><li>• Dichlorobenzene[1,2-]</li><li>• Dichlorobenzene[1,3-]</li><li>• Vinyl Chloride</li></ul> <p>Soil ESL for earthworm due to availability of new TRVs:</p> <ul style="list-style-type: none"><li>• Chloroaniline[4-]</li><li>• Hexachlorobenzene</li><li>• Styrene</li></ul> <p>Soil ESL for plant due to availability of new TRVs:</p> <ul style="list-style-type: none"><li>• Chloroaniline[4-]</li><li>• Hexachlorobenzene</li><li>• Styrene</li></ul> <p>Alternative screening approach for Iron for plant based on EPA EcoSSL’s report</p> <ul style="list-style-type: none"><li>• See <a href="http://www.epa.gov/ecotox/ecossl/pdf/eco-ssl_iron.pdf">http://www.epa.gov/ecotox/ecossl/pdf/eco-ssl_iron.pdf</a></li></ul> <p>Sediement ESL for aquatic community organism due to availability of a new</p>

**Table 1. ESL Changes by Ecorisk Database Release**

**Ecorisk Database Release**

TRVs:

- Molybdenum

**ESL Updates**

Soil ESLs for deer mouse, desert cottontail and red fox due to TF updates:

- Methylphenol[2-]

**New TRVs**

Tier 2 (Geometric Mean) oral diet TRVs from LANL were developed with the PTSE Process for the following chemicals and receptor groups:

- Lithium/ mammal

Tier 3 (Critical Study) oral diet TRVs from LANL were developed with the PTSE Process for the following chemicals and receptor groups:

- Hexanone[2-]/ bird

Tier 4 (based on secondary data) oral diet TRVs from ORNL were identified for the following chemicals and receptor groups:

- Lithium/ plant
- Molybdenum/ plant
- Molybdenum/ bird
- Styrene/ earthworm
- Vinyl Chloride/ mammal

Tier 4 (based on secondary data) oral diet TRVs from EPA ECOTOX were identified for the following chemicals and receptor groups:

- Carbon Disulfide/ mammal
- Chloroaniline[4-]/ earthworm
- Chloroaniline[4-]/ plant
- Dichlorobenzene[1,2-]/ mammal
- Dichlorobenzene[1,3-]/ mammal
- Hexachlorobenzene/ bird
- Hexachlorobenzene/ mammal
- Hexachlorobenzene/ earthworm
- Hexachlorobenzene/ plant
- Styrene/ plant

**TRV Updates**

The use status of various TRVs changed for the following reasons:

- Vinyl chloride/ mammal oral diet TRV records deleted due to availability of updated toxicity information for oral diet TRV from same data source (ORNL).

**Table 1. ESL Changes by Ecorisk Database Release**

**Ecorisk Database Release**      **ESL Changes**

- Vinyl chloride/ mammal drinking water TRV no longer used because primary toxicity data is for oral diet exposure, which is no longer considered an appropriate TRV surrogate for a drinking water exposure.
- Carbon Tetrachloride/ mammal oral TRVs no longer used because currently not an exposure concern for this exposure pathway.
- Molybdenum/ aquatic community organism sediment TRV selected for use because this chemical is now a chemical of concern.

**New TFs**

All New TFs (except where noted otherwise) were acquired for the following chemicals because these chemicals are new exposure concerns:

- Carbon Disulfide
- Chloroaniline[4-]
- Dichlorobenzene[1,2-]
- Dichlorobenzene[1,3-]
- Hexachlorobenzene
- Hexanone[2-]
- Styrene
- Vinyl Chloride
- Lithium (only TF\_invert and TF\_flesh)
- Molybdenum (only TF\_invert and TF\_flesh)

**TF Updates**

TFs for the following chemicals were updated:

- Methylphenol[2-] – all TFs updated due to availability of more appropriate data
- Molybdenum – TF\_beef and TF\_plant updated due to availability of more appropriate data

**Interface Updates**

- Added “Other Reports” links to the “Menu” screen to allow access to other files on the Ecorisk Db from within the database interface including;

**Table 1. ESL Changes by Ecorisk Database Release**

<b>Ecorisk Database Release</b>	<b>ESL Changes</b>
<b>October 2010 – Release 2.5</b>	<p>In this release of the database, ESLs/TRVs were added for chemicals for which no toxicity data was previously available. Online toxicity databases were searched for relevant existing TRVs or for primary toxicity data and/or references from which TRVs could be derived for x chemicals (see EcoriskDbR2.5_ToxicityData_ResourceSummary_SoilESLs_101310.xls for details of search results). In this release of the database, an additional 11 new chemicals now have LANL peer reviewed/ approved TRVs/ESLs incorporated into this release of the database, no chemicals have interim ESLs/ TRVs at this time, 13 chemicals have surrogate ESLs/TRVs (see Interim_SoilESLs_R2.5_101310.xls) based on chemicals already in the database, and the remaining 8 chemicals from the original data gap list still have no ESLs at this time.</p> <p><b>New ESLs</b></p> <p>Soil and Sediment ESLs for birds due to availability of new TRVs:</p> <ul style="list-style-type: none"><li>• Benz(a)anthracene</li><li>• Diphenylamine</li><li>• Iodomethane</li><li>• Pyrene</li></ul> <p>Soil and Sediment ESLs for mammals due to availability of new TRVs:</p> <ul style="list-style-type: none"><li>• Carbazole</li><li>• Nitroaniline[2-]</li><li>• Benzyl alcohol</li><li>• Hexanone[2-]</li><li>• Trichlorofluoromethane</li></ul> <p>Soil ESL for plant due to availability of new TRVs:</p> <ul style="list-style-type: none"><li>• Methylphenol[2-]</li><li>• Methylphenol[3-]</li></ul> <p><b>ESL Updates</b></p> <p>Water ESL for aquatic community organism due to retraction of previous TRV and replacement with available alternative TRV:</p> <ul style="list-style-type: none"><li>• Beryllium</li></ul> <p><b>New TRVs</b></p> <p>Tier 2 (Geometric Mean) oral diet TRVs from LANL were developed with the PTSE Process for the following chemicals and receptor groups:</p> <ul style="list-style-type: none"><li>• Hexanone[2-]/Mammal</li><li>• Trichlorofluoromethane/Mammal</li></ul> <p>Tier 3 (Critical Study) oral diet TRVs from LANL were developed with the PTSE Process for the following chemicals and receptor groups:</p>

**Table 1. ESL Changes by Ecorisk Database Release**

**Ecorisk Database Release**    **ESL Changes**

- Benzyl alcohol/Mammal
- Carbazole/Mammal
- Nitroaniline[2-]/Mammal

Tier 4 (based on secondary data) oral diet TRVs from identified for the following chemicals and receptor groups:

- Diphenylamine/Bird
- Iodomethane/Bird
- Benz(a)anthracene/Bird
- Pyrene/Bird
- Methylphenol[2-]/Plant
- Methylphenol[3-]/Plant

**TRV Updates**

The use status of various TRVs changed for the following reasons:

- Beryllium/Aquatic community organism water TRV deleted due to retraction of value by publishing data source. TRV replaced with available alternative value.

**New TFs**

All New TFs (except where noted otherwise) were acquired for the following chemicals because these chemicals are new exposure concerns:

- Benzyl alcohol
- Diphenylamine
- Iodomethane
- Nitroanilin[2-]

**TF Updates**

TFs for the following chemicals were updated:

- Carbazole – TF\_beef updated due to availability of more appropriate data
- Trichlorofluoromethane – TF\_plant updated due to availability of more appropriate data

**Interface Updates**

None.

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**Table 2. Beta Release (October 1998) List of Soil ESLs for Bird Receptors**

Analyte Class	Analyte Group	Analyte Name	Analyte Code	ESL Medium	Receptor Group
NONRAD	D/F	Tetrachlorodibenzodioxin[2,3,7,8-]	1746-01-6	SOIL	Bird
NONRAD	INORG	Aluminum	AL	SOIL	Bird
NONRAD	INORG	Arsenic	AS	SOIL	Bird
NONRAD	INORG	Barium	BA	SOIL	Bird
NONRAD	INORG	Cadmium	CD	SOIL	Bird
NONRAD	INORG	Chromium (total)	CR	SOIL	Bird
NONRAD	INORG	Chromium(+6)	CR(+6)	SOIL	Bird
NONRAD	INORG	Cobalt	CO	SOIL	Bird
NONRAD	INORG	Copper	CU	SOIL	Bird
NONRAD	INORG	Cyanide (total)	CN(-1)	SOIL	Bird
NONRAD	INORG	Lead	PB	SOIL	Bird
NONRAD	INORG	Manganese	MN	SOIL	Bird
NONRAD	INORG	Mercury (inorganic)	HGI	SOIL	Bird
NONRAD	INORG	Mercury (methyl)	HGM	SOIL	Bird
NONRAD	INORG	Nickel	NI	SOIL	Bird
NONRAD	INORG	Selenium	SE	SOIL	Bird
NONRAD	INORG	Silver	AG	SOIL	Bird
NONRAD	INORG	Uranium	U	SOIL	Bird
NONRAD	INORG	Vanadium	V	SOIL	Bird
NONRAD	INORG	Zinc	ZN	SOIL	Bird
NONRAD	PAH	Naphthalene	91-20-3	SOIL	Bird
NONRAD	PCB	Aroclor-1242	53469-21-9	SOIL	Bird
NONRAD	PCB	Aroclor-1248	12672-29-6	SOIL	Bird
NONRAD	PCB	Aroclor-1254	11097-69-1	SOIL	Bird
NONRAD	PCB	Aroclor-1260	11096-82-5	SOIL	Bird
NONRAD	PEST	BHC[beta-]	319-85-7	SOIL	Bird
NONRAD	PEST	BHC[gamma-]	58-89-9	SOIL	Bird
NONRAD	PEST	Chlordane[alpha-]	5103-71-9	SOIL	Bird
NONRAD	PEST	Chlordane[gamma-]	5103-74-2	SOIL	Bird
NONRAD	PEST	DDE[4,4'-]	72-55-9	SOIL	Bird
NONRAD	PEST	DDT[4,4'-]	50-29-3	SOIL	Bird
NONRAD	PEST	Dieldrin	60-57-1	SOIL	Bird
NONRAD	PEST	Endosulfan	115-29-7	SOIL	Bird
NONRAD	PEST	Endrin	72-20-8	SOIL	Bird
NONRAD	PEST	Heptachlor	76-44-8	SOIL	Bird
NONRAD	PEST	Kepone	143-50-0	SOIL	Bird
NONRAD	PEST	Methoxychlor[4,4'-]	72-43-5	SOIL	Bird
NONRAD	PEST	Toxaphene (Technical Grade)	8001-35-2	SOIL	Bird
NONRAD	SVOC	Bis(2-ethylhexyl)phthalate	117-81-7	SOIL	Bird

**Table 2. Beta Release (October 1998) List of Soil ESLs for Bird Receptors**

Analyte Class	Analyte Group	Analyte Name	Analyte Code	ESL Medium	Receptor Group
NONRAD	SVOC	Chloro-3-methylphenol[4-]	59-50-7	SOIL	Bird
NONRAD	SVOC	Chlorophenol[2-]	95-57-8	SOIL	Bird
NONRAD	SVOC	Di-n-Butyl Phthalate	84-74-2	SOIL	Bird
NONRAD	SVOC	Pentachloronitrobenzene	82-68-8	SOIL	Bird
NONRAD	SVOC	Pentachlorophenol	87-86-5	SOIL	Bird
NONRAD	VOC	Acetone	67-64-1	SOIL	Bird
NONRAD	VOC	Xylene (Total)	1330-20-7	SOIL	Bird
RAD	RAD	Americium-241	AM-241	SOIL	Bird
RAD	RAD	Cesium-134	CS-134	SOIL	Bird
RAD	RAD	Cesium-137 + Barium-137	CS-137/ BA-137	SOIL	Bird
RAD	RAD	Cobalt-60	CO-60	SOIL	Bird
RAD	RAD	Europium-152	EU-152	SOIL	Bird
RAD	RAD	Plutonium-238	PU-238	SOIL	Bird
RAD	RAD	Plutonium-239, 240	PU-239/240	SOIL	Bird
RAD	RAD	Plutonium-241	PU-241	SOIL	Bird
RAD	RAD	Radium-226	RA-226	SOIL	Bird
RAD	RAD	Sodium-22	NA-22	SOIL	Bird
RAD	RAD	Strontium-90 + Yttrium-90	SR-90/ Y-90	SOIL	Bird
RAD	RAD	Thorium-228	TH-228	SOIL	Bird
RAD	RAD	Thorium-230	TH-230	SOIL	Bird
RAD	RAD	Thorium-232	TH-232	SOIL	Bird
RAD	RAD	Tritium	H-3	SOIL	Bird
RAD	RAD	Uranium-234	U-234	SOIL	Bird
RAD	RAD	Uranium-235	U-235	SOIL	Bird
RAD	RAD	Uranium-238	U-238	SOIL	Bird

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**Table 3. Beta Release (October 1998) List of Soil ESLs for Mammalian Receptors**

Analyte Class	Analyte Group	Analyte Name	Analyte Code	ESL Medium	Receptor Group
NONRAD	D/F	Tetrachlorodibenzodioxin[2,3,7,8-]	1746-01-6	SOIL	Mammal
NONRAD	HE	Amino-2,6-dinitrotoluene[4-]	19406-51-0	SOIL	Mammal
NONRAD	HE	Amino-4,6-dinitrotoluene[2-]	35572-78-2	SOIL	Mammal
NONRAD	HE	Dinitrobenzene[1,3-]	99-65-0	SOIL	Mammal
NONRAD	HE	Dinitrotoluene[2,4-]	121-14-2	SOIL	Mammal
NONRAD	HE	Dinitrotoluene[2,6-]	606-20-2	SOIL	Mammal
NONRAD	HE	HMX	2691-41-0	SOIL	Mammal
NONRAD	HE	Nitroglycerine	55-63-0	SOIL	Mammal
NONRAD	HE	Nitrotoluene[2-]	88-72-2	SOIL	Mammal
NONRAD	HE	Nitrotoluene[3-]	99-08-1	SOIL	Mammal

**Table 3. Beta Release (October 1998) List of Soil ESLs for Mammalian Receptors**

Analyte Class	Analyte Group	Analyte Name	Analyte Code	ESL Medium	Receptor Group
NONRAD	HE	Nitrotoluene[4-]	99-99-0	SOIL	Mammal
NONRAD	HE	PETN	78-11-5	SOIL	Mammal
NONRAD	HE	RDX	121-82-4	SOIL	Mammal
NONRAD	HE	Tetryl	479-45-8	SOIL	Mammal
NONRAD	HE	Trinitrobenzene[1,3,5-]	99-35-4	SOIL	Mammal
NONRAD	HE	Trinitrotoluene[2,4,6-]	118-96-7	SOIL	Mammal
NONRAD	INORG	Aluminum	AL	SOIL	Mammal
NONRAD	INORG	Antimony	SB	SOIL	Mammal
NONRAD	INORG	Arsenic	AS	SOIL	Mammal
NONRAD	INORG	Barium	BA	SOIL	Mammal
NONRAD	INORG	Beryllium	BE	SOIL	Mammal
NONRAD	INORG	Cadmium	CD	SOIL	Mammal
NONRAD	INORG	Chromium (total)	CR	SOIL	Mammal
NONRAD	INORG	Chromium(+6)	CR(+6)	SOIL	Mammal
NONRAD	INORG	Cobalt	CO	SOIL	Mammal
NONRAD	INORG	Copper	CU	SOIL	Mammal
NONRAD	INORG	Cyanide (total)	CN(-1)	SOIL	Mammal
NONRAD	INORG	Lead	PB	SOIL	Mammal
NONRAD	INORG	Manganese	MN	SOIL	Mammal
NONRAD	INORG	Mercury (inorganic)	HGI	SOIL	Mammal
NONRAD	INORG	Mercury (methyl)	HGM	SOIL	Mammal
NONRAD	INORG	Nickel	NI	SOIL	Mammal
NONRAD	INORG	Selenium	SE	SOIL	Mammal
NONRAD	INORG	Silver	AG	SOIL	Mammal
NONRAD	INORG	Thallium	TL	SOIL	Mammal
NONRAD	INORG	Titanium	TI	SOIL	Mammal
NONRAD	INORG	Uranium	U	SOIL	Mammal
NONRAD	INORG	Vanadium	V	SOIL	Mammal
NONRAD	INORG	Zinc	ZN	SOIL	Mammal
NONRAD	PAH	Acenaphthene	83-32-9	SOIL	Mammal
NONRAD	PAH	Acenaphthylene	208-96-8	SOIL	Mammal
NONRAD	PAH	Anthracene	120-12-7	SOIL	Mammal
NONRAD	PAH	Benzo(a)anthracene	56-55-3	SOIL	Mammal
NONRAD	PAH	Benzo(a)pyrene	50-32-8	SOIL	Mammal
NONRAD	PAH	Benzo(b)fluoranthene	205-99-2	SOIL	Mammal
NONRAD	PAH	Benzo(g,h,i)perylene	191-24-2	SOIL	Mammal
NONRAD	PAH	Benzo(k)fluoranthene	207-08-9	SOIL	Mammal
NONRAD	PAH	Chrysene	218-01-9	SOIL	Mammal
NONRAD	PAH	Dibenzo(a,h)anthracene	53-70-3	SOIL	Mammal
NONRAD	PAH	Fluoranthene	206-44-0	SOIL	Mammal

**Table 3. Beta Release (October 1998) List of Soil ESLs for Mammalian Receptors**

Analyte Class	Analyte Group	Analyte Name	Analyte Code	ESL Medium	Receptor Group
NONRAD	PAH	Fluorene	86-73-7	SOIL	Mammal
NONRAD	PAH	Indeno(1,2,3-cd)pyrene	193-39-5	SOIL	Mammal
NONRAD	PAH	Methylnaphthalene[2-]	91-57-6	SOIL	Mammal
NONRAD	PAH	Naphthalene	91-20-3	SOIL	Mammal
NONRAD	PAH	Phenanthrene	85-01-8	SOIL	Mammal
NONRAD	PAH	Pyrene	129-00-0	SOIL	Mammal
NONRAD	PCB	Aroclor-1016	12674-11-2	SOIL	Mammal
NONRAD	PCB	Aroclor-1242	53469-21-9	SOIL	Mammal
NONRAD	PCB	Aroclor-1248	12672-29-6	SOIL	Mammal
NONRAD	PCB	Aroclor-1254	11097-69-1	SOIL	Mammal
NONRAD	PCB	Aroclor-1260	11096-82-5	SOIL	Mammal
NONRAD	PEST	BHC[beta-]	319-85-7	SOIL	Mammal
NONRAD	PEST	BHC[gamma-]	58-89-9	SOIL	Mammal
NONRAD	PEST	Chlordane[alpha-]	5103-71-9	SOIL	Mammal
NONRAD	PEST	Chlordane[gamma-]	5103-74-2	SOIL	Mammal
NONRAD	PEST	DDE[4,4'-]	72-55-9	SOIL	Mammal
NONRAD	PEST	DDT[4,4'-]	50-29-3	SOIL	Mammal
NONRAD	PEST	Dieldrin	60-57-1	SOIL	Mammal
NONRAD	PEST	Endosulfan	115-29-7	SOIL	Mammal
NONRAD	PEST	Endrin	72-20-8	SOIL	Mammal
NONRAD	PEST	Heptachlor	76-44-8	SOIL	Mammal
NONRAD	PEST	Kepone	143-50-0	SOIL	Mammal
NONRAD	PEST	Methoxychlor[4,4'-]	72-43-5	SOIL	Mammal
NONRAD	PEST	Toxaphene (Technical Grade)	8001-35-2	SOIL	Mammal
NONRAD	SVOC	Benzoic Acid	65-85-0	SOIL	Mammal
NONRAD	SVOC	Bis(2-ethylhexyl)phthalate	117-81-7	SOIL	Mammal
NONRAD	SVOC	Butyl Benzyl Phthalate	85-68-7	SOIL	Mammal
NONRAD	SVOC	Chloro-3-methylphenol[4-]	59-50-7	SOIL	Mammal
NONRAD	SVOC	Chlorobenzene	108-90-7	SOIL	Mammal
NONRAD	SVOC	Chlorophenol[2-]	95-57-8	SOIL	Mammal
NONRAD	SVOC	Dimethyl Phthalate	131-11-3	SOIL	Mammal
NONRAD	SVOC	Di-n-Butyl Phthalate	84-74-2	SOIL	Mammal
NONRAD	SVOC	Di-n-octylphthalate	117-84-0	SOIL	Mammal
NONRAD	SVOC	Nitrobenzene	98-95-3	SOIL	Mammal
NONRAD	SVOC	Pentachloronitrobenzene	82-68-8	SOIL	Mammal
NONRAD	SVOC	Pentachlorophenol	87-86-5	SOIL	Mammal
NONRAD	SVOC	Phenol	108-95-2	SOIL	Mammal
NONRAD	VOC	Acetone	67-64-1	SOIL	Mammal
NONRAD	VOC	Benzene	71-43-2	SOIL	Mammal
NONRAD	VOC	Butanone[2-]	78-93-3	SOIL	Mammal

**Table 3. Beta Release (October 1998) List of Soil ESLs for Mammalian Receptors**

Analyte Class	Analyte Group	Analyte Name	Analyte Code	ESL Medium	Receptor Group
NONRAD	VOC	Chloroform	67-66-3	SOIL	Mammal
NONRAD	VOC	Dichloroethane[1,1-]	75-34-3	SOIL	Mammal
NONRAD	VOC	Dichloroethene[1,1-]	75-35-4	SOIL	Mammal
NONRAD	VOC	Dichloroethene[cis/trans-1,2-]	540-59-0	SOIL	Mammal
NONRAD	VOC	Methylene Chloride	75-09-2	SOIL	Mammal
NONRAD	VOC	Tetrachloroethane[1,1,2,2-]	79-34-5	SOIL	Mammal
NONRAD	VOC	Tetrachloroethene	127-18-4	SOIL	Mammal
NONRAD	VOC	Toluene	108-88-3	SOIL	Mammal
NONRAD	VOC	Trichlorobenzene[1,2,4-]	120-82-1	SOIL	Mammal
NONRAD	VOC	Trichloroethane[1,1,1-]	71-55-6	SOIL	Mammal
NONRAD	VOC	Trichloroethene	79-01-6	SOIL	Mammal
NONRAD	VOC	Xylene (Total)	1330-20-7	SOIL	Mammal
RAD	RAD	Americium-241	AM-241	SOIL	Mammal
RAD	RAD	Cesium-134	CS-134	SOIL	Mammal
RAD	RAD	Cesium-137 + Barium-137	CS-137/ BA-137	SOIL	Mammal
RAD	RAD	Cobalt-60	CO-60	SOIL	Mammal
RAD	RAD	Europium-152	EU-152	SOIL	Mammal
RAD	RAD	Plutonium-238	PU-238	SOIL	Mammal
RAD	RAD	Plutonium-239, 240	PU-239/240	SOIL	Mammal
RAD	RAD	Plutonium-241	PU-241	SOIL	Mammal
RAD	RAD	Radium-226	RA-226	SOIL	Mammal
RAD	RAD	Sodium-22	NA-22	SOIL	Mammal
RAD	RAD	Strontium-90 + Yttrium-90	SR-90/ Y-90	SOIL	Mammal
RAD	RAD	Thorium-228	TH-228	SOIL	Mammal
RAD	RAD	Thorium-230	TH-230	SOIL	Mammal
RAD	RAD	Thorium-232	TH-232	SOIL	Mammal
RAD	RAD	Tritium	H-3	SOIL	Mammal
RAD	RAD	Uranium-234	U-234	SOIL	Mammal
RAD	RAD	Uranium-235	U-235	SOIL	Mammal
RAD	RAD	Uranium-238	U-238	SOIL	Mammal

[BACK TO TOP](#)**Table 4. Beta Release (October 1998) List of Soil ESLs for Earthworm Receptor**

Analyte Class	Analyte Class	Analyte Class	Analyte Class	Analyte Class	Analyte Class
NONRAD	D/F	Tetrachlorodibenzodioxin[2,3,7,8-]	1746-01-6	SOIL	Invertebrate
NONRAD	INORG	Arsenic	AS	SOIL	Invertebrate
NONRAD	INORG	Cadmium	CD	SOIL	Invertebrate
NONRAD	INORG	Chromium(+6)	CR(+6)	SOIL	Invertebrate
NONRAD	INORG	Copper	CU	SOIL	Invertebrate

**Table 4. Beta Release (October 1998) List of Soil ESLs for Earthworm Receptor**

Analyte Class	Analyte Class	Analyte Class	Analyte Class	Analyte Class	Analyte Class
NONRAD	INORG	Lead	PB	SOIL	Invertebrate
NONRAD	INORG	Mercury (inorganic)	HGI	SOIL	Invertebrate
NONRAD	INORG	Mercury (methyl)	HGM	SOIL	Invertebrate
NONRAD	INORG	Nickel	NI	SOIL	Invertebrate
NONRAD	INORG	Selenium	SE	SOIL	Invertebrate
NONRAD	INORG	Zinc	ZN	SOIL	Invertebrate
NONRAD	PAH	Fluorene	86-73-7	SOIL	Invertebrate
NONRAD	SVOC	Chlorobenzene	108-90-7	SOIL	Invertebrate
NONRAD	SVOC	Dimethyl Phthalate	131-11-3	SOIL	Invertebrate
NONRAD	SVOC	Nitrobenzene	98-95-3	SOIL	Invertebrate
NONRAD	SVOC	Pentachlorophenol	87-86-5	SOIL	Invertebrate
NONRAD	SVOC	Phenol	108-95-2	SOIL	Invertebrate
NONRAD	VOC	Dichlorobenzene[1,4-]	106-46-7	SOIL	Invertebrate
NONRAD	VOC	Trichlorobenzene[1,2,4-]	120-82-1	SOIL	Invertebrate
RAD	RAD	Americium-241	AM-241	SOIL	Invertebrate
RAD	RAD	Cesium-134	CS-134	SOIL	Invertebrate
RAD	RAD	Cesium-137 + Barium-137	CS-137/ BA-137	SOIL	Invertebrate
RAD	RAD	Cobalt-60	CO-60	SOIL	Invertebrate
RAD	RAD	Europium-152	EU-152	SOIL	Invertebrate
RAD	RAD	Plutonium-238	PU-238	SOIL	Invertebrate
RAD	RAD	Plutonium-239, 240	PU-239/240	SOIL	Invertebrate
RAD	RAD	Plutonium-241	PU-241	SOIL	Invertebrate
RAD	RAD	Radium-226	RA-226	SOIL	Invertebrate
RAD	RAD	Sodium-22	NA-22	SOIL	Invertebrate
RAD	RAD	Strontium-90 + Yttrium-90	SR-90/ Y-90	SOIL	Invertebrate
RAD	RAD	Thorium-228	TH-228	SOIL	Invertebrate
RAD	RAD	Thorium-230	TH-230	SOIL	Invertebrate
RAD	RAD	Thorium-232	TH-232	SOIL	Invertebrate
RAD	RAD	Tritium	H-3	SOIL	Invertebrate
RAD	RAD	Uranium-234	U-234	SOIL	Invertebrate
RAD	RAD	Uranium-235	U-235	SOIL	Invertebrate
RAD	RAD	Uranium-238	U-238	SOIL	Invertebrate

[BACK TO TOP](#)**Table 5. Beta Release (October 1998) List of Soil ESLs for Generic Plant Receptor**

Analyte Class	Analyte Class	Analyte Class	Analyte Class	Analyte Class	Analyte Class
NONRAD	HE	Amino-4,6-dinitrotoluene[2-]	35572-78-2	SOIL	Plant
NONRAD	HE	RDX	121-82-4	SOIL	Plant
NONRAD	HE	Tetryl	479-45-8	SOIL	Plant

**Table 5. Beta Release (October 1998) List of Soil ESLs for Generic Plant Receptor**

Analyte Class	Analyte Class	Analyte Class	Analyte Class	Analyte Class	Analyte Class
NONRAD	HE	Trinitrotoluene[2,4,6-]	118-96-7	SOIL	Plant
NONRAD	INORG	Aluminum	AL	SOIL	Plant
NONRAD	INORG	Antimony	SB	SOIL	Plant
NONRAD	INORG	Arsenic	AS	SOIL	Plant
NONRAD	INORG	Barium	BA	SOIL	Plant
NONRAD	INORG	Beryllium	BE	SOIL	Plant
NONRAD	INORG	Cadmium	CD	SOIL	Plant
NONRAD	INORG	Chromium (total)	CR	SOIL	Plant
NONRAD	INORG	Chromium(+6)	CR(+6)	SOIL	Plant
NONRAD	INORG	Cobalt	CO	SOIL	Plant
NONRAD	INORG	Copper	CU	SOIL	Plant
NONRAD	INORG	Lead	PB	SOIL	Plant
NONRAD	INORG	Manganese	MN	SOIL	Plant
NONRAD	INORG	Mercury (inorganic)	HGI	SOIL	Plant
NONRAD	INORG	Nickel	NI	SOIL	Plant
NONRAD	INORG	Selenium	SE	SOIL	Plant
NONRAD	INORG	Silver	AG	SOIL	Plant
NONRAD	INORG	Thallium	TL	SOIL	Plant
NONRAD	INORG	Uranium	U	SOIL	Plant
NONRAD	INORG	Vanadium	V	SOIL	Plant
NONRAD	INORG	Zinc	ZN	SOIL	Plant
NONRAD	PAH	Acenaphthene	83-32-9	SOIL	Plant
NONRAD	PAH	Benzo(a)anthracene	56-55-3	SOIL	Plant
NONRAD	PAH	Benzo(b)fluoranthene	205-99-2	SOIL	Plant
NONRAD	PCB	Aroclor-1254	11097-69-1	SOIL	Plant
NONRAD	PEST	BHC[gamma-]	58-89-9	SOIL	Plant
NONRAD	PEST	Chlordane[alpha-]	5103-71-9	SOIL	Plant
NONRAD	PEST	Chlordane[gamma-]	5103-74-2	SOIL	Plant
NONRAD	PEST	Dieldrin	60-57-1	SOIL	Plant
NONRAD	PEST	Endrin	72-20-8	SOIL	Plant
NONRAD	PEST	Heptachlor	76-44-8	SOIL	Plant
NONRAD	SVOC	Dibenzofuran	132-64-9	SOIL	Plant
NONRAD	SVOC	Di-n-Butyl Phthalate	84-74-2	SOIL	Plant
NONRAD	SVOC	Pentachlorophenol	87-86-5	SOIL	Plant
NONRAD	SVOC	Phenol	108-95-2	SOIL	Plant
NONRAD	VOC	Methylene Chloride	75-09-2	SOIL	Plant
NONRAD	VOC	Toluene	108-88-3	SOIL	Plant
NONRAD	VOC	Xylene (Total)	1330-20-7	SOIL	Plant
RAD	RAD	Americium-241	AM-241	SOIL	Plant
RAD	RAD	Cesium-134	CS-134	SOIL	Plant

**Table 5. Beta Release (October 1998) List of Soil ESLs for Generic Plant Receptor**

Analyte Class	Analyte Class	Analyte Class	Analyte Class	Analyte Class	Analyte Class
RAD	RAD	Cesium-137 + Barium-137	CS-137/ BA-137	SOIL	Plant
RAD	RAD	Cobalt-60	CO-60	SOIL	Plant
RAD	RAD	Europium-152	EU-152	SOIL	Plant
RAD	RAD	Plutonium-238	PU-238	SOIL	Plant
RAD	RAD	Plutonium-239, 240	PU-239/240	SOIL	Plant
RAD	RAD	Plutonium-241	PU-241	SOIL	Plant
RAD	RAD	Radium-226	RA-226	SOIL	Plant
RAD	RAD	Sodium-22	NA-22	SOIL	Plant
RAD	RAD	Strontium-90 + Yttrium-90	SR-90/ Y-90	SOIL	Plant
RAD	RAD	Thorium-228	TH-228	SOIL	Plant
RAD	RAD	Thorium-230	TH-230	SOIL	Plant
RAD	RAD	Thorium-232	TH-232	SOIL	Plant
RAD	RAD	Tritium	H-3	SOIL	Plant
RAD	RAD	Uranium-234	U-234	SOIL	Plant
RAD	RAD	Uranium-235	U-235	SOIL	Plant
RAD	RAD	Uranium-238	U-238	SOIL	Plant

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**Table 6. Beta Release (October 1998) List of Sediment and Water ESLs for Aquatic Community Organism Receptors**

Analyte Class	Analyte Class	Analyte Class	Analyte Class	Analyte Class	Analyte Class
RAD	RAD	Americium-241	AM-241	WATER and SEDIMENT	Aquatic
RAD	RAD	Cesium-137 + Barium-137	CS-137/ BA-137	WATER and SEDIMENT	Aquatic
RAD	RAD	Plutonium-238	PU-238	WATER and SEDIMENT	Aquatic
RAD	RAD	Plutonium-239, 240	PU-239/240	WATER and SEDIMENT	Aquatic
RAD	RAD	Plutonium-241	PU-241	WATER and SEDIMENT	Aquatic
RAD	RAD	Radium-226	RA-226	WATER and SEDIMENT	Aquatic
RAD	RAD	Strontium-90 + Yttrium-90	SR-90/ Y-90	WATER and SEDIMENT	Aquatic
RAD	RAD	Thorium-232	TH-232	WATER and SEDIMENT	Aquatic
RAD	RAD	Tritium	H-3	WATER and SEDIMENT	Aquatic
RAD	RAD	Uranium-234	U-234	WATER and	Aquatic

**Table 6. Beta Release (October 1998) List of Sediment and Water ESLs for Aquatic Community Organism Receptors**

Analyte Class	Analyte Class				
				SEDIMENT	
RAD	RAD	Uranium-235	U-235	WATER and SEDIMENT	Aquatic
RAD	RAD	Uranium-238	U-238	WATER and SEDIMENT	Aquatic

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### **Ecorisk Database REF ID 0561**

United States Environmental Protection Agency (USEPA). 1993h. Wildlife Exposure Factors Handbook, Vol. I and II. EPA/600/R-93/187. United States Environmental Protection Agency.

### **Ecorisk Database REF ID 1484**

Newell, PG. 1999 (Dec.). Revisions to Ecorisk Database R.1 ESLs. Los Alamos National Laboratory, Environmental Restoration Project, Los Alamos National Laboratory, Los Alamos, NM.

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# **Appendix A**

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*Primary Toxicity Study Evaluation Methods Used to Develop  
Los Alamos National Laboratory Toxicity Reference Values*



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## Attachments

- Attachment A-1 GMM TRV Summary Report Example  
Attachment A-2 CS TRV Summary Report Example

## Acronyms and Abbreviations

ATSDR	Agency for Toxic Substances and Disease Registry
C-CL	chronic-critical life stage
Cal/Ecotox	California OHHEA Wildlife Biology, Exposure Factor, and Toxicity Database
CASRN	Chemical Abstracts Service Registry Number
CEC	cation exchange capacity
CL	critical life stage
CS	critical study

DART/ETIC	Development and Reproductive Toxicology/Environmental Teratology Information Center
EC <sub>xx</sub>	effective concentration for xx% of the population
Eco-SSL	ecological soil screening level
ECOTOX	Ecotoxicology (database)
ED <sub>xx</sub>	effective dose for xx% of the population
EP	Environmental Programs (Directorate)
EPA	U.S. Environmental Protection Agency
ERED	Environmental Residue-Effects Database
ESL	ecological screening level
ETWS	equivalent total weighted score
EXTOXNET	Extension Toxicology Network
Fm	female
GMM	geometric mean
GSD	geometric standard deviation
HMX	1,3,5,7-tetranitro-1,3,5,7-tetrazocine
IRIS	Integrated Risk Information System
ITER	International Toxicity Estimates for Risk
LANL	Los Alamos National Laboratory
LC <sub>xx</sub>	lethal concentration for xx% of the population
LD <sub>xx</sub>	lethal dose for xx% of the population
LOAEL	lowest observed adverse effect level
LOEC	lowest observed effect concentration
LOEL	lowest observed effect level
%MTWS	percent of maximum total weighted score
MF	male and female
MI	male
N/A	not applicable
NLM	National Library of Medicine
NOAEL	no observed adverse effect level
NOEC	no observed effect concentration
NOEL	no observed effect level
NR	not reported
O	other

OC	organic carbon
OECD	Organisation for Economic Co-operation and Development
OHHEA	Office of Environmental Health Hazard Assessment (state of California)
OM	organic matter
ORNL	Oak Ridge National Laboratory
PAN	Pesticide Action Network
PTSE	primary toxicity study evaluation
PTV	primary toxicity value
R/D	reproduction/development
RDX	hexahydro-1,3,5-trinitro-1,3,5-triazine
Ref ID	reference identification
RfC	reference concentration
RfD	reference dose
S	survival
SNL	Sandia National Laboratories
SzC	size change (adult)
T&E	threatened and endangered
TOXLINE	Toxicology Literature Online
TOXNET	Toxicology Data Network
TRV	toxicity reference value
UF	uncertainty factor
USACHPPM	U.S. Army Center for Health Promotion and Preventive Medicine
USGS	U.S. Geological Society
WC	weight change (adult)



## **A-1.0 PRIMARY TOXICITY STUDY EVALUATION METHODS**

### **A-1.1 Primary Toxicity Literature Search and Retrieval**

Before a primary toxicity study evaluation (PTSE) can be started, the primary toxicity literature for the organism, exposure pathway, and chemical scenario of concern (e.g., plant root uptake of barium from soil) must be collected.

A literature search consists of the following two components: (1) an online search of databases that contain citations for primary toxicity literature (see Table A-1), and (2) a review of bibliographies of secondary toxicity data literature that has been identified either through online searches or the risk assessment community (see Table A-2). Each piece of literature (reference) identified is assigned a unique Ecorisk Database reference identification (Ref ID) number for identification, tracking, and citation during the literature search, review, and evaluation process. These numbers will be included throughout this document.<sup>1</sup>

Keyword searches are performed. For example, if the title of a reference in a bibliography (or an online literature search result) indicates that the reference contains the sought-after toxicity information, a paper copy of the reference is retrieved. The abstracts are then reviewed to verify that the reference contains applicable toxicity data for the derivation of a toxicity reference value (TRV). Verification of applicable contents requires scanning the reference for relevant measurement endpoints (including reproduction, development, survival, adult weight changes, and adult size changes) that are considered to have a direct link to the fitness of an organism and its contribution toward population health. Focusing on ecologically relevant endpoints ensures that all levels of ecological organization are considered in the screening process (LANL 2004, 087630, Ref ID 1554). If the reference contains ecologically relevant data, then a PTSE can be performed. In cases where ecologically relevant endpoints are not available for certain chemicals and organism groups, a PTSE may be performed on references with endpoints having a less direct link to the fitness of an organism and its contribution toward population health, such as endpoints associated with physiological functions, cancer, histopathology, clinical observations, and behavioral changes. Values based on endpoints other than reproduction/development, survival, or weight or size change are to be used with caution given the uncertainty surrounding their impact on population health (LANL 2004, 087630, Ref ID 1554).

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<sup>1</sup> Initially, the construction of the Ecorisk Database took precedence over performing extensive toxicity data literature retrieval. The initial literature search for bird, mammal, invertebrate (earthworm), and plant toxicity data was limited to reviewing reference lists in secondary references and conducting minimal searches of online literature databases. As the Ecorisk Database underwent further development, literature searches became more comprehensive and included more extensive online literature searches and reviews of related bibliographies.

**Table A-1**  
**Online Databases and Search Engines to Search for Primary Toxicity Data Literature**

Internet Source	Site Contents / Database Name	Web Address
Australian Government, Department of the Environment and Heritage	National Pollutant Inventory database	<a href="http://www.npi.gov.au/index.html">http://www.npi.gov.au/index.html</a>
First Search	Literature search engine	<a href="http://www.oclc.org/firstsearch/">http://www.oclc.org/firstsearch/</a>
Google	Internet search engine	<a href="http://www.google.com">http://www.google.com</a>
Los Alamos National Laboratory (LANL)	External and internal access to library catalogs	<a href="http://lib-www.lanl.gov/">http://lib-www.lanl.gov/</a>
National Library of Medicine (NLM)	MEDLINE/PubMed literature search engine	<a href="http://www.ncbi.nlm.nih.gov/PubMed/">http://www.ncbi.nlm.nih.gov/PubMed/</a>
	Toxicology Data Network (TOXNET) literature search engine (includes Toxicology Literature Online [TOXLINE], Integrated Risk Information System [IRIS], and several other databases)	<a href="http://toxnet.nlm.nih.gov/">http://toxnet.nlm.nih.gov/</a>
TOXNET	TOXNET is a cluster of databases covering toxicology, hazardous chemicals, environmental health, and related areas. It is managed by the Toxicology and Environmental Health Information Program in the Division of Specialized Information Services of the NLM.  International Toxicity Estimates for Risk (ITER) is a database that contains risk information for over 600 chemicals from authoritative groups worldwide.	<a href="http://toxnet.nlm.nih.gov/cgi-bin/sis/htmlgen?iter">http://toxnet.nlm.nih.gov/cgi-bin/sis/htmlgen?iter</a>
	Development and Reproductive Toxicology/Environmental Teratology Information Center (DART/ETIC) is a bibliographic database covering literature on reproductive and developmental toxicology. DART is managed by NLM and funded by the U.S. Environmental Protection Agency (EPA), the National Institute of Environmental Health Sciences and NLM. DART/ETIC contains references to reproductive and developmental toxicology literature published since 1965.	<a href="http://toxnet.nlm.nih.gov/cgi-bin/sis/htmlgen?DARTETIC">http://toxnet.nlm.nih.gov/cgi-bin/sis/htmlgen?DARTETIC</a>
	TOXLINE is a bibliographic database providing comprehensive coverage of the biochemical, pharmacological, physiological, and toxicological effects of drugs and other chemicals from 1965 to the present. TOXLINE contains over 3 million citations, almost all with abstracts and/or index terms and Chemical Abstracts Service Registry Numbers (CASRN).	<a href="http://toxnet.nlm.nih.gov/cgi-bin/sis/htmlgen?TOXLINE">http://toxnet.nlm.nih.gov/cgi-bin/sis/htmlgen?TOXLINE</a>

Table A-1 (continued)

Internet Source	Site Contents / Database Name	Web Address
Integrated Risk Information System	<p>IRIS is an electronic database containing information on human health effects that may result from exposure to various substances in the environment. IRIS is prepared and maintained by the EPA's National Center for Environmental Assessment within the Office of Research and Development.</p> <p><i>Noncancer effects:</i> Oral reference doses (RfDs) and inhalation reference concentrations (RfCs) for effects known or assumed to be produced through a nonlinear (possibly threshold) mode of action. In most instances, RfDs and RfCs are developed for the noncarcinogenic effects of substances.</p> <p><i>Cancer effects:</i> Descriptors that characterize the weight of evidence for human carcinogenicity, oral slope factors, and oral and inhalation unit risks for carcinogenic effects. Where a nonlinear mode of action is established, RfD and RfC values may be used. Primary toxicity study references for mammalian test species are reported and include body weight and survival data.</p>	<a href="http://www.epa.gov/ncea/iris/search_keyword.htm">http://www.epa.gov/ncea/iris/search_keyword.htm</a>
National Technical Information Service	Source of government-funded information	<a href="http://www.ntis.gov/search/index.aspx">http://www.ntis.gov/search/index.aspx</a>
Pacific Northwest National Laboratory	External access to Pacific Northwest National Laboratory publication catalog	<a href="http://www.pnl.gov/main/publications/index.asp">http://www.pnl.gov/main/publications/index.asp</a>
Web of Science	Literature search engine (accessed via Colorado State University)	<a href="http://libguides.colostate.edu/content.php?pid=30095&amp;sid=220274">http://libguides.colostate.edu/content.php?pid=30095&amp;sid=220274</a>
U.S. Geological Society (USGS)	USGS Contaminant Exposure and Effects--Terrestrial Vertebrates database contains contaminant exposure and effects information for terrestrial vertebrates (birds, mammals, amphibians, and reptiles) that reside in estuarine and coastal habitats along the Atlantic, Gulf, and Pacific Coasts, including Alaska and Hawaii, and in the Great Lakes Region.	<a href="http://www.pwrc.usgs.gov/contaminants-online/pages/CEETV/CEETVintro.htm">http://www.pwrc.usgs.gov/contaminants-online/pages/CEETV/CEETVintro.htm</a>
U.S. Environmental Protection Agency (EPA) Office of Pesticide Programs	EPA Office of Pesticide Programs' Aquatic Life Benchmarks.	<a href="http://www.epa.gov/oppefed1/ecorisk_ders/aquatic_life_benchmark.htm">http://www.epa.gov/oppefed1/ecorisk_ders/aquatic_life_benchmark.htm</a>

Table A-1 (continued)

Internet Source	Site Contents / Database Name	Web Address
Pesticide Action Network (PAN)	The PAN Pesticide Database is a one-stop location for toxicity and regulatory information for pesticides. The PAN Pesticide Database brings together a diverse array of information on pesticides from many different sources, providing human toxicity (chronic and acute), ecotoxicity, and regulatory information for about 6400 pesticide active ingredients and their transformation products, as well as adjuvants and solvents used in pesticide products. Only aquatic ecotoxicity data are reported.	<a href="http://www.pesticideinfo.org/Search_Ecotoxicity.jsp">http://www.pesticideinfo.org/Search_Ecotoxicity.jsp</a>
EPA Ecotoxicology (ECOTOX) Database	The ECOTOX database provides single chemical toxicity information for aquatic and terrestrial life. Values reported include the lethal concentration for 50% of the population (LC <sub>50</sub> ), no observed effect concentration (NOEC), lowest observed effect concentration (LOEC), lowest observed effect level (LOEL), no observed effect level (NOEL), effective concentration for 50% of the population (ED <sub>50</sub> ), etc. Toxicity data for available substances are reported in worksheet "ECOTOX." Only terrestrial data for growth, mortality, reproduction, and population queried from database. Searched by CASRN.	<a href="http://cfpub.epa.gov/ecotox/">http://cfpub.epa.gov/ecotox/</a>
The American Bird Conservancy	The American Bird Conservancy Pesticide Toxicity Database contains acute pesticide toxicity data for birds.	<a href="http://www.abcbirds.org/abcprograms/policy/pesticides/aims/aims/toxicity.cfm">http://www.abcbirds.org/abcprograms/policy/pesticides/aims/aims/toxicity.cfm</a>
The California Office of Environmental Health Hazard Assessment (OHHEA) Wildlife Biology, Exposure Factor, and Toxicity Database (Cal/Ecotox)	Cal/Ecotox is a compilation of physiological and ecological parameters and toxicity data for a number of California fish and wildlife. Species, chemical, endpoint type, endpoint description, endpoint value, endpoint range, study description, and reference are reported. Data for chemicals of interest are reported in worksheet "CalEcotox."	<a href="http://www.oehha.org/cal_ecotox/default.htm">http://www.oehha.org/cal_ecotox/default.htm</a>
The U.S. Army Corps of Engineers/EPA Environmental Residue-Effects Database (ERED)	The ERED is a compilation of data, taken from the literature, where biological effects (e.g., reduced survival, growth, etc.) and tissue contaminant concentrations were simultaneously measured in the same organism. Currently, the database is limited to those instances where biological effects observed in an organism are linked to a specific contaminant within its tissues.	<a href="http://el.erdc.usace.army.mil/ered/Index.cfm">http://el.erdc.usace.army.mil/ered/Index.cfm</a>

**Table A-1 (continued)**

Internet Source	Site Contents / Database Name	Web Address
EPA National Information System of the Regional Integrated Pest Management Centers Office of Pesticide Programs Pesticide Ecotoxicity Database	The Ecological Fate and Effects Division of the EPA Office of Pesticide Programs is continuing efforts to update the database with all EPA-reviewed ecotoxicity endpoints for pesticides registered or previously registered in the U.S. Toxicity data on over 800 active ingredients, metabolites, and multi-ingredient formulations are presently included in the database. The toxicity data input into the database are compiled from actual studies reviewed by EPA in conjunction with pesticide registration or reregistration and studies performed by EPA, U.S. Department of Agriculture, and U.S. Fish and Wildlife Service laboratories, which have been reviewed by Agency biologists and judged acceptable for use in the ecological risk assessment process. The database presently contains over 21,000 records for acute and chronic toxicity endpoints on terrestrial and aquatic plants, aquatic invertebrates, terrestrial invertebrates, insects, amphibians, fish, birds, reptiles, and wild mammals. The database is presented in Microsoft Access and contains 35 fields per record entry. Each record entry summarizes one ecotoxicity study for a single species or one toxicity endpoint from a multiple-species study and includes EPA tracking information regarding that study submission.	<a href="http://www.ipmcenters.org/ECotox/DataAccess.cfm">http://www.ipmcenters.org/ECotox/DataAccess.cfm</a>
U.S. Army Center for Health Promotion and Preventive Medicine (USACHPPM)	The USACHPPM Wildlife Toxicity Assessment Program contains complete chemical toxicological assessments/profiles for wildlife with reference lists.	<a href="http://chppm-www.apgea.army.mil/erawq/tox/index.htm">http://chppm-www.apgea.army.mil/erawq/tox/index.htm</a>
Agency for Toxic Substances and Disease Registry (ATSDR)	The ATSDR website contains toxicological profiles for human health. These profiles succinctly characterize the toxicologic and adverse health effects information for a hazardous substance. Each peer-reviewed profile identifies and reviews the key literature that describes a hazardous substance's toxicologic properties. Other pertinent literature is also presented, but is described in less detail than the key studies. The references are generally for mammalian studies for all routes.	<a href="http://www.atsdr.cdc.gov/">http://www.atsdr.cdc.gov/</a>

**Table A-2**  
**Examples of Secondary Toxicity Data Literature**  
**Bibliographies to Review for Primary Toxicity Data Literature Citations**

Source	Author (Year, ER ID)	Description	Ecorisk Database Reference ID
Oak Ridge National Laboratory (ORNL)*	Efroymson et al. (1997, 059231)	Screening toxicity benchmarks for terrestrial plants	Ref ID 0094
	Efroymson et al. (1997, 059231)	Screening toxicity benchmarks for soil and litter invertebrates	Ref ID 0096
	Sample et al. (1996, 059306)	Screening toxicity benchmarks for wildlife	Ref ID 0344
	Maxwell and Opresko (1996, 059275)	Ecological criteria for HMX (1,3,5,7-tetranitro-1,3,5,7-tetrazocine)	Ref ID 0467
	Talmage and Opresko (1995, 059328)	Ecological criteria for 2,4,6-trinitrotoluene	Ref ID 0469
	Talmage and Opresko (1996, 059329)	Ecological criteria for RDX (hexahydro-1,3,5-trinitro-1,3,5-triazine)	Ref ID 0470
	Talmage et al. (1999, 063021)	Screening values for nitroaromatic munition compounds	Ref ID 0480
Sandia National Laboratories (SNL)	IT Corporation (1997, 057136) (Appendix A, Table A.1)	Ecological risk assessment methodology	Ref ID 0092
LANL threatened and endangered (T&E) species	Gallegos et al. (1997, 059790)	Risk assessment of peregrine falcon (includes toxicity benchmarks for avian species)	Ref ID 0427
U.S. Army	Layton et al. (1987, 014703)	Explosives information	Ref ID 0552
USACHPPM	Johnson and McAtee (2001, 110044)	Wildlife toxicity assessment for 2,4,6-trinitrotoluene	Ref ID 1195
	Johnson and Midgley (2001, 089453)	Wildlife toxicity assessment for nitroglycerine	Ref ID 1446
	Salice and Holdsworth (2001, 089452)	Wildlife toxicity assessment for 1,3,5-trinitrobenzene	Ref ID 1447
	Salice and Holdsworth (2001, 089451)	Wildlife toxicity assessment for dinitrobenzene	Ref ID 1448
	Johnson and Holdsworth (2001, 089454)	Wildlife toxicity assessment for 2-amino-4,6-dinitrotoluene and 4-amino-2,6-dinitrotoluene	Ref ID 1449
	Johnson and Holdsworth (2001, 073781)	Wildlife toxicity assessment for HMX	Ref ID 1450
	Johnson and Holdsworth (2001, 089455)	Wildlife toxicity assessment for pentaerythritol tetranitrate	Ref ID 1451
	Salice and Holdsworth (2002, 073780)	Wildlife toxicity assessment for RDX	Ref ID 1452
EPA Region 5 environmental data quality levels	PRC Environmental Management, Inc. (1996, 059989)	Ecological data quality levels	Ref ID 0574

\*Reports are available online at <http://www.esd.ornl.gov/programs/ecorisk/documents/tm85r3.pdf>.

## A-1.2 Overview of PTSEs

Once a set of references is compiled for an organism, exposure pathway, and chemical scenario of concern, each reference is subjected to the PTSE process. This process is broken down into four main parts:

1. data extraction,
2. study evaluation and primary toxicity value (PTV) calculation,
3. TRV development, and
4. TRV approval.

Data-entry databases were created for each of the first three parts of the PTSE process to guide the reviewer in extracting, scoring, and evaluating the necessary information. The database system also assists in maintaining consistency in the way the toxicity information are tabulated and peer reviewed as well as provides a mechanism for documentation of the PTSE process. Users of the Ecorisk Database can review the data reported and gain an understanding of the information supporting the TRV used to calculate a particular ecological screening level (ESL). A brief description of each part of the PTSE process is presented below, followed by a more detailed breakdown of the components of each part.

### A-1.2.1 Part 1, Data Extraction

Data extraction involves reading each primary toxicity reference thoroughly, extracting pertinent pieces of information, and documenting them in the Part 1 PTSE data-entry database.

*“Data” represents toxicity information from the scientific literature such as details of the study design, test organism, or toxicological effects.*

### A-1.2.2 Part 2, Study Evaluation and PTV Calculation

During the study evaluation process, information obtained from the data extraction process is reviewed and scored based on availability and character of information reported. The data are semiquantitatively scored in the Part 2 data-entry database in four areas: study design and documentation, taxonomic relationship of test organism to ESL screening receptors, exposure conditions, and measurements and results. Components of each of these areas are scored based on their relevancy toward deriving scientifically defensible TRVs. The score for each criterion is then weighted according to its ability to influence the development of a TRV with the least uncertainty. Uncertainty is the extent to which the TRV represents a dose rate or concentration in an exposure medium that is associated with no significant risk for adverse ecological effects for the LANL environmental exposure scenario of concern; therefore, uncertainty can be influenced by how well the data approximates the LANL exposure scenario. The last step in this part is to calculate the PTVs: no observed adverse effect levels (NOAELs) for birds and mammals or NOECs for earthworms and plants, lowest observed adverse effect levels (LOAELs) for birds and mammals or LOECs for earthworms and plants, and/or other effect levels (e.g., effective concentrations for xx% of the population [EC<sub>xx</sub>s] or lethal doses for xx% of the population [LD<sub>xx</sub>s]).

### A-1.2.3 Part 3, TRV Development

In Part 3, the number of PTVs available for TRV development for an organism, exposure pathway, and chemical scenario of concern is determined by selecting one PTV per endpoint category (reproduction/development, survival, and adult weight/size changes) represented in an experiment. If

three or more PTVs exist, a geometric mean (GMM) TRV is calculated. If less than three PTVs are available, professional judgment is used to select the PTV associated with the most applicable study, measurement endpoint, and effect level to derive a critical study (CS) TRV. Uncertainty factors (UFs) are applied to achieve a TRV equivalent to a chronic NOAEL or NOEC where necessary. A summary describing the basis for the TRV is written. This discussion describes the importance of the TRV in protection of wildlife, invertebrate, or plant populations; the data set considered for the selection of the TRV; the justification to support this selection; and the aspects of the study or studies that relate it to the environmental concerns for LANL. Also, UF explanations and calculations are noted.

*Professional judgment considers ecological relevance and is peer reviewed for greater consistency in selection of values.*

#### **A-1.2.4 Part 4, TRV Approval**

Once a TRV is derived, whether it is a GMM or CS TRV, the value and its supporting documentation are peer reviewed by LANL's Environmental Programs (EP) Directorate's Risk Assessment Team to gain approval of the TRV for use in calculations of ESLs in the Ecorisk Database.

### **A-2.0 PTSE PART 1, DATA EXTRACTION**

The PTSE Part 1 consists of four separate tables of data entry. Information is entered into these tables by way of Microsoft Access database forms. There are tables for reference, chemical, experiment, and effect detail information; therefore, the data entry follows this order to ensure the connection of the appropriate Ref IDs with the chemical, experiment, and experiment effect IDs. Also, for control purposes (i.e., maintaining the latest versions of object format and data), PTSE reviewer initials are entered more than once throughout the data entry process to ensure that each record in each table is tracked by reviewer and date.

Each specific field entry (e.g., codes selected from a drop-down list) is usually followed by a comments field to allow the reviewer to further elaborate on the selection and any relevant assumptions. The following sections focus on the specific fields, but will also discuss the types and examples of comments that may be entered in the corresponding comments field.

#### **A-2.1 Data Entry**

Data entry is broken down into four parts: reference and reviewer information, chemical information, experiment information, and measurements and results. Each of these parts has its own table in the Part 1 data-entry database where data are recorded. However, the data are typed into or presented in database forms for easier entry and editing of information.

##### **A-2.1.1 Reference and Reviewer Information**

###### **Reference ID**

The PTSE Ref ID is entered here (see section A-1.1 for a description of the Ref ID).

## Reference Summary

A brief description of the reference and its experiments is written here. This description includes the test organism, chemical, route, medium of exposure, and length of chemical administration for each experiment and also summarizes key differences between experiments, if applicable. Also, the basis for not developing a TRV (e.g., the exposure route is injection, or one of multiple chemicals administered in the study is not a chemical of concern) is noted at the end of the reference description. In addition, the reference summary may describe why the focus of the review is placed on a particular experiment or experiments and not on others. See Example A-1.

### Example A-1 Reference Summaries

(a) Barley (*Hordeum vulgare*) was the test organism used to evaluate the toxicity of copper (Cu+2) or chromium (Cr+6) in two types of soil: artificial and natural forest soil. The nominal exposure concentrations used were 0.1, 1, 10, 100, and 1000 µg/g dry soil for copper and chromium experiments. The endpoints evaluated include plant emergence and shoot and root growth (both 5- and 14-d). Additionally, the levels of copper or chromium in the plant tissues were assessed, but this will not be evaluated in this PTSE because there is not a clear connection between tissue burdens and adverse effects to population health. Additionally, only the 14-d plant emergence measurement will be considered in this evaluation because it is a more chronic measurement than 5-d plant emergence, considering it took place at the end of exposure. A reference toxicant, HgCl<sub>2</sub>, also contributed to another exposure group, but it and its effects will not be evaluated because the results do not give any additional information about the toxicity of copper or chromium.

(b) Fischer 344 rats were intermittently exposed to 0-, 150-, 475-, or 1500-ppm chloromethane by way of inhalation. In the first of two experiments, 40 males and 80 females were exposed to chloromethane for 6 h/d, 5 d/wk for 10 wk. After 10 wk, inhalation occurred for 6 h/d, 7 d/wk during the 2-wk mating season where one male was mated to two exposed females. The females were continued on the 6-h/d, 7-d/wk exposure regimen from the start of mating through postnatal day 28, except from gestation day 18 to postnatal day 4, while 10 males from each group were necropsied. Pups from this experiment were not directly exposed to the chemical until after weaning, and then they were put through the same exposure and mating regimen as their parents. In the second experiment, the remaining 30 males from each group in the first experiment were then mated to unexposed females for another 2 wk. Adult body weight, litter parameters (e.g., pup survival, pup weight), gross pathology, and histopathology were observed. The second experiment is not reviewed in this Part 1 in favor of the more chronic exposure period in the multigenerational experiment.

## Reviewer Initials

The initials of the person responsible for completing the PTSE are selected from the drop-down list.

## Review Start and Finish Date

The dates the review is started and finished are reported here. If a change is made in the reference summary, the date of the change supersedes the finish date. Dates are entered for each record in the tables of the data-entry database for purposes of data control and ensuring the latest information is present in the latest release of the Ecorisk Database.

### A-2.1.2 Chemical Information

#### Chemical ID

The analyte code for the chemical of concern is selected from the drop-down list. Analyte codes follow Johnston (1997, 059791, Ref ID 0576). Generally, the Chemical Abstracts Service Registry Numbers are used for organic compounds (e.g., 11097-69-1 for Aroclor-1254) while element abbreviations are used for inorganic chemicals (e.g., CD for cadmium). Further identification occurs for forms of inorganic chemicals, such as hexavalent chromium vs. trivalent chromium, where the analyte code for these forms are CR(+6) and CR(+3), respectively. Also, chemicals with organic and inorganic forms are also coded differently to distinguish between them (e.g., HGI for inorganic mercury and HGM for methyl mercury).

#### Reviewer Initials

The initials of the person responsible for completing the chemical details in the PTSE are selected from the drop-down list.

#### Record Date

The date the chemical record was created is typed into this field.

### A-2.1.3 Experiment Information

#### Experiment ID

The experiment ID consists of the Ecorisk Database Ref ID, chemical ID (analyte code), and experiment number in the format of Ref ID\_analyte code\_experiment number (see Example A-2).

**Example A-2 Experiment IDs**

0025\_SE\_1

0517\_50-29-3\_2

As mentioned previously, the Ref ID is a unique identifier assigned to each reference for tracking during the literature search, review, and evaluation process. The analyte code is a unique identifier assigned by the reviewer following guidelines set forth in Johnston (1997, 059791, Ref ID 0576) for each element and compound. The experiment number is based on the actual number of experiments reported in a reference. For the purposes of the PTSE process, an experiment is defined by a unique set of exposure parameters (i.e., one chemical administration period, one exposure frequency type, one test organism, one chemical, one exposure medium, one exposure route, and one set of exposure concentrations). The reviewer may have to use his or her own judgment in delineating unique experimental scenarios.

#### Experiment Purpose

The purpose(s) of the experiment is noted here. Also, since each experiment has its own record in the Part 1 database, a brief description of the test organism, exposure route and medium, and length of chemical administration is entered in this field in order for the reviewer and user of the database to distinguish between experiments (see Example A-3).

**Example A-3 Experiment Purposes**

(a) The purpose of the study was to see whether selenium levels similar to those found in raptor prey items from selenium-contaminated environments would affect reproduction in captive eastern screech-owls. The screech-owls were fed a diet containing 0, 4.4, or 13.2 ppm wet weight of selenium in the form of selenomethionine. Growth, reproduction, and liver biochemistry effects were studied.

(b) Authors emphasize the importance of earthworms as a biomonitoring tool for assessing the impact of chemicals in soil quality and fauna. In order to use them as a biomonitoring tool successfully, the effects of various chemicals on earthworms needs to be studied. The investigators determined the effect of zinc on the growth and reproduction of earthworms during a 20-wk study.

**Reviewer**

The initials of the person responsible for completing the experiment details in the PTSE are selected from the drop-down list.

**Review Date**

The date the experiment record is created is typed into this field.

**Organism Type ID**

The test organisms are classified into the following categories and coded accordingly:

SLE soil and/or litter earthworm

TB terrestrial bird

TM terrestrial mammal

TP terrestrial plant

The appropriate code for the test organism of concern in the PTSE is selected from the drop-down list.

**Organism Name**

At a minimum, the common name of the test organism is reported in the reference. In cases where the scientific name is not reported, various references are consulted to find it. This is done to later assess the taxonomic relationship of test species to ecological screening receptor species of concern, especially for bird and mammal receptors. The common name of the organism is selected from a drop-down list that is linked to the test species table. If the name is not found on the list, the name can be typed in. However, the information is still added to the test species table so that it appears on the drop-down list in the future.

Examples of sources consulted for scientific names include

- National Geographic Society, 1987. *Field Guide to the Birds of North America*, 2<sup>nd</sup> Ed., Washington, D.C., 464 pp. (Note: Later editions are available and may have more updated records on names as a result of merging or division of species.)
- Burt, W. H., and R. P. Grossenheider, 1980. *A Field Guide to the Mammals: North America North of Mexico*, Houghton Mifflin Company, New York, New York, 289 pp.
- BIOSIS. Index to Organism Names (<http://www.organismnames.com/>)
- New Mexico Game and Fish Biota System Information of New Mexico (BISON-M) <http://www.bison-m.org/databasequery.aspx>

### **Author's Reason for Studying this Particular Test Organism**

If it is explicitly stated why the author(s) chose to use a particular species of test organism (e.g., Oldfield mouse, *Peromyscus polionotus*) in their research, the reasons are paraphrased. If it is not clearly stated, but the purpose can be deduced for the use of the general organism type (e.g., mouse or rodent), the reasons are noted. However, the reviewer clarifies that these reasons noted are assumptions. For example, if in the introduction of a paper, the authors discuss case histories describing the effects of trichloroethylene inhalation exposure in humans, and they also discuss previous studies of exposure of trichloroethylene to laboratory mammals, it can be reasonably assumed that their choice of the test organism is used as an experimental model to gauge potential effects that may occur in humans (see Example A-4).

#### **Example A-4 Author's Reason for Studying this Particular Test Organism**

(a) The investigators wished to use the same standard toxicity test organisms as described in the Organisation for Economic Co-operation and Development (OECD) contact and artificial soil testing procedures (OECD 1984, 109940, Ref ID 1235). This enabled them to focus on determining influences of contact tests and soil characteristics (pH and organic matter content) on toxicity in the earthworms and compare their data with others.

(b) It is unknown why the authors specifically chose mallards over other aquatic birds, but it is assumed they considered them to be representative of aquatic birds in order to study cadmium toxicity in waterfowl.

### **Age or Life Stage**

The age or life stage of a test organism is coded because later in the Part 1 PTSE process, this information is needed to gauge whether or not measurements occurred during a critical life stage (see Focus Measurement Critical Life Stage Category in section A-2.1.4). Coding for age/life stage of the test organism adheres to the conventions presented in Table A-3.

**Table A-3  
Age Categories, Codes, and Definitions**

Age Category ID	Age Category	Definition
BrA_Unk	Bird Adult	Bird is known to be in reproductive condition or is otherwise mature, but it is unknown if it is breeding for the first time or at later stages.
BrA1	Bird Adult 1	Bird reaches sexual maturity and breeds for the first time.
BrA2	Bird Adult 2	Bird survives to breed at older age.
BrE	Bird Embryo	Fertilization occurs and embryo develops inside an eggshell until hatched.
BrG	Bird Gamete	Unfertilized egg and sperm
BrJ_Unk	Bird Juvenile	Bird is said to be a juvenile but exact phase is unknown.
BrJ1	Bird Juvenile 1	Hatchling, chick, or nestling grows until flight feathers are developed.
BrJ2	Bird Juvenile 2	Sexually immature fledgling or poult that undergoes additional development prior to breeding condition
BrLC	Bird Life Cycle	All life stages
EwA_Unk	Earthworm Adult	Earthworm is known to be in reproductive condition or is otherwise mature, but it is unknown if it is breeding for the first time or at later stages.
EwA1	Earthworm Adult 1	Sexually mature worm (with clitellum) breeds for the first time.
EwA2	Earthworm Adult 2	Earthworm survives to breed at older age.
EwE	Earthworm Cocoon or Embryo	External fertilization, cocoon formation, embryo development, and worm emergence from cocoon
EwG	Earthworm Gamete	Unfertilized egg and sperm
EwJ1	Earthworm Juvenile	Small worm grows until it reaches reproductive condition.
EwLC	Earthworm Life Cycle	All life stages
MmA_Unk	Mammal Adult	Mammal is known to be in reproductive condition or is otherwise mature, but it is unknown if it is breeding for the first time or at later stages.
MmA1	Mammal Adult 1	Mammal reaches sexual maturity and breeds for the first time.
MmA2	Mammal Adult 2	Mammal survives to breed at older age.
MmE	Mammal Embryo or Fetus	<i>In utero</i> organism developing from fertilized egg to birth
MmG	Mammal Gamete	Unfertilized egg and sperm
MmJ_Unk	Mammal Juvenile	Mammal is said to be a juvenile but exact phase is unknown.
MmJ1	Mammal Juvenile 1	Newborn mammal obtaining all or most of its nutrition by nursing until weaning
MmJ2	Mammal Juvenile 2	Immature mammal growing from weaning to more or less adult size and appearance. The typical "juvenile" stage.
MmJ3	Mammal Juvenile 3	Period of additional development is required or time must pass until the organism may breed (next season). Often independent from parents, "subadult."

Table A-3 (continued)

Age Category ID	Age Category	Definition
MmLC	Mammal Life Cycle	All life stages
Pa_Unk	Plant (Annual) Unknown Age/Stage	Not enough information was provided or found to determine what life stage this plant age represents.
PaA1	Plant (Annual) Flowering and Seed Set	Plant is fertilized and seeds develop and disperse.
PaE	Plant (Annual) Seed	Embryo inside seed
PaG	Plant (Annual) Gamete	Unfertilized ova and pollen
PaM	Plant (Annual) Mature	Plant is known to be at a mature stage but it is unknown how else to classify this stage.
PaS_Unk	Plant (Annual) Seedling	Plant is a seedling but it is uncertain/unknown with regards to whether seedling is closer to a sprouting stage or closer to reproductive stage.
PaS1	Plant (Annual) Seedling	Seed sprouts, grows to emerge from soil, and leaves open or some minimum size is attained.
PaS2	Plant (Annual) Seedling 2	Plant continues to grow until reproductive stage achieved.
Po_Unk	Plant (Other) Unknown Age/Stage	Not enough information was provided or found to determine what life stage this plant age represents.
PoA_Unk	Plant (Other)	Plant is in mature, reproductive condition but it is unknown if it is fertilized for the first time or if it is a larger individual producing seeds.
PoA1	Plant (Other) Flowering and Seed Set	Plant is fertilized and seeds develop and disperse.
PoA2	Plant (Other) Larger Reproducing Plant	Larger individuals producing seeds
PoE	Plant (Other) Seed	Embryo inside seed
PoG	Plant (Other) Gamete	Unfertilized ova and pollen
PoLC	Plant (Other) Life Cycle	All life stages
PoM	Plant (Other) Mature	Plant is known to be at a mature stage but it is unknown how else to classify this stage.
PoS_Unk	Plant (Other) Seedling/Sapling	Plant is a seedling but it is uncertain/unknown with regard to whether seedling is closer to a sprouting stage or closer to reproductive stage.
PoS1	Plant (Other) Seedling/Sapling 1	Seed sprouts, grows to emerge from soil, and leaves open or some minimum size is attained.
PoS2	Plant (Other) Seedling/Sapling 2	Plant continues to grow until reproductive stage achieved.

If the age or life stage of a bird or mammal test organism is not provided but body weight is, an age or life stage is estimated for the organism based on other reference sources containing similar organisms, body weights, and age information.

The age coding task becomes difficult when placing organisms in categories that are borderline juvenile/adult or seedling/adult. If more information is needed, related information is first sought in the toxicity references currently on hand for the Ecorisk Database. For example, if a primary toxicity reference states the mouse was 6 wk old at the time of exposure, and it is difficult to determine whether to code this

age as a juvenile or an adult, information in the database is reviewed to find similar records containing mice to see if a correlation can be made between ages and life stages. When information such as this cannot be found in the existing references, additional references specific to the test organism species or genera are consulted, and a note summarizing the information is recorded in the age or life stage comment field of the database.

### Organism Sex

The genders of the test organisms that are directly exposed to the chemical are selected from the drop-down list (Ml for male, Fm for female, or MF for male and female). This field is not applicable (N/A) for invertebrates and plants. If the sex is not reported (NR), NR is selected.

If a situation arises where only the females were exposed to the chemical, and they were then bred with untreated males, the code Fm is entered for sex, and a note of this arrangement is made in the associated comment field. Likewise, if only males were exposed, Ml is entered, and any related notes are made in the comment field.

### Organism Source/Origin

The location of where the test organism was obtained, bred, or collected is noted here. Any other relevant information about the organism (such as if the organism was pathogen-free) is also noted in this field.

### Dose Rate Parameters

Dose rate parameters other than exposure concentrations (i.e., body weights and ingestion or inhalation rates) reported in the study are recorded here for later use in calculating the PTV(s) (see section A-3.2). Exposure concentrations are recorded later in the Part 1 experiment details. For the dose rate parameters, the aim is to use values that will lead to the most conservative PTV in units of mg chemical/kg body weight/d for birds and mammals.

Dose rate parameters are not needed for invertebrates or plants because the dose concentration (in mg/kg) is used for the TRV itself. Note: Default values of 999, N/A, and N/A are entered into the value, units, and comment fields, respectively, for invertebrate and plant studies.<sup>2</sup>

*Dose rate parameters are selected to calculate the most reasonably conservative dose rate to represent the TRV; therefore, TRVs and ESLs are conservative, protective values.*

### Author-Reported Daily Dose Rates for Bird and Mammal Studies

If the exposure concentrations presented in the study are already in, or can be easily converted to, units of mg chemical/kg body weight/d, dose parameters and calculations for a daily dose rate are not needed, and this is indicated in the appropriate fields. However, if dose rate parameters are provided in the study, information is still recorded with the expectation that they may be used for other studies where the parameters are not available but are needed for similar test organisms.

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<sup>2</sup> In the early developmental stages of the Ecorisk Database, dose rate parameters may have been considered inapplicable, and the default value of 999 was used. The dose rate parameter may have not been reported if the authors already provided daily dose rates or if the ingestion rates were already normalized to body weight. In these cases, the dose rate parameter was, and still is, not needed for PTV calculation, but it is now reported for possible future use in other areas of the database.

### Author-Reported Exposure Concentrations Other than Daily Dose Rates for Bird and Mammal Studies

If the exposure levels are presented as concentrations of chemical in the exposure medium (such as mg/kg food, mg/mL water, or mg air/m<sup>3</sup>), the body weight (in kg) and food or water ingestion rate (in kg food/d or L/d, respectively) or inhalation rate (in m<sup>3</sup>/d) dose rate parameters are needed to calculate the PTV in mg chemical/kg body weight/d.

### Reporting Dose Rate Parameters

Table A-4 provides scenarios of how dose rate parameters may be reported in the primary toxicity study and how the parameter is reported in the dose rate parameter field in the database.

**Table A-4**  
**Scenarios of Dose Rate Parameter Information Reported in Primary Toxicity Studies and How Body Weight Values are Reported in the PTSE Part 1 Data-Entry Database Field for Body Weight**

Scenario	Report
Dose rate parameter for controls was measured at intervals throughout the study	Average of all values throughout study <sup>a</sup>
	If values are grouped according to male and female organisms, the average of the male or female values that will lead to a more conservative PTV is used. <sup>b</sup>
Dose rate parameter for controls were measured at beginning and at end of study	Average of the two values <sup>a</sup>
	If values are grouped according to male and female organisms, the average of the male or female values that will lead to a more conservative PTV is used. <sup>b</sup>
Dose rate parameter for controls was measured at beginning of study only.	Measured value
Range of dose rate parameters for controls or all organisms at beginning of study	Either end of this range, depending on which value will lead to a more conservative PTV <sup>b</sup>
	If body weights are grouped according to male and female organisms, the average weight that will produce a more conservative PTV is used. <sup>b</sup>
No dose rate parameter information for controls, only treated organisms	The average of the beginning value of treated organisms, before chemical exposure began <sup>c</sup>
No dose rate parameters reported at all	Default value of 999

<sup>a</sup> In situations where dose rate parameters are measured and provided throughout the study, an average is calculated from those measurements to provide an estimate that is representative of the organism at all stages throughout the study.

<sup>b</sup> The general rule is that if there are dose rate parameters reported for male and female groups, or if a range of dose rate parameters is reported, either the lower or higher average value is used because this value, when used in the PTV calculation, will lead to a more conservative PTV. For example, a larger value for the body weight leads to a lower PTV (see Example A-5a), thus the PTV is more protective. On the other hand, a lower value for an oral ingestion rate leads to a higher PTV (see Example A-5b).

<sup>c</sup> The average of the beginning weight of the organisms in a treatment group before exposure begins is used, rather than the average of the weights throughout the study, because the weights throughout the study may be affected by chemical exposure. Therefore, the daily dose calculation may be influenced if the affected body weights are used, and it may not be representative of a daily dose that would affect a healthy individual.

**Example A-5 The Selection of Dose Rate Parameters to Provide the Most Protective PTV**

Note: Explanations of PTV calculations are more detailed in section A-3.0, PTSE Part 2, Study Evaluation and Primary Toxicity Value Calculation.

(a) Higher vs. lower body weight: A higher body weight leads to a lower PTV when used in the denominator. The following calculations demonstrate the difference by holding the concentration (100 mg/kg) and food ingestion rate (0.0055 kg/d) constant and using body weights of 0.03 and 0.09 kg.

Lower weight:

$$\text{PTV (mg/kg/d)} = \frac{100 \text{ mg/kg} * 0.0055 \text{ kg/d}}{0.03 \text{ kg}} = 18.3 \text{ mg/kg/d}$$

Higher weight:

$$\text{PTV (mg/kg/d)} = \frac{100 \text{ mg/kg} * 0.0055 \text{ kg/d}}{0.09 \text{ kg}} = 6.1 \text{ mg/kg/d}$$

(b) Higher vs. lower ingestion rate or inhalation rate: A lower ingestion or inhalation rate leads to a lower PTV. Since these parameters take the same location in the equation and therefore have the same type of influence on the PTV, only the use of water ingestion will be used to demonstrate the difference. The following calculations hold the concentration of 5 mg/L and body weight of 0.03 kg constant, while using the water ingestion rates of 0.0075 and 0.009 L/d.

Lower ingestion rate:

$$\text{PTV (mg/kg/d)} = \frac{5 \text{ mg/L} * 0.0075 \text{ L/d}}{0.03 \text{ kg}} = 1.25 \text{ mg/kg/d}$$

Higher ingestion rate:

$$\text{PTV (mg/kg/d)} = \frac{5 \text{ mg/L} * 0.009 \text{ L/d}}{0.03 \text{ kg}} = 1.5 \text{ mg/kg/d}$$

**Exposure Environment**

If the study was conducted in a laboratory, a greenhouse, or some other controlled environment, it is marked as a laboratory study. Lab is selected from the drop-down list. If the study was a field study conducted under uncontrolled environmental variables, it is noted as a field study and Fld is selected from the drop-down list. Physical descriptions of the laboratory or greenhouse environment, what the test organisms were housed in, controlled variables (such as temperature and humidity), and other relevant information are noted in the corresponding comment field.

**Test Chemical Form (for Inorganic Chemicals Only)**

If the chemical administered is inorganic, the compound as it is administered in the study is selected from a master pull-down list of chemicals maintained in a separate analyte table. If the compound cannot be found, it must be added to the master list of analytes in the Ecorisk Database before this field can be filled. If the chemical is organic, the default value of N/A is left in the field.

### Test Chemical Description/Source

The purity of the chemical and the company it was purchased from are noted in this field. If the chemical was synthesized by the researchers of the study itself, a brief summary of the process is described.

### Exposure Medium

The medium in which the chemical was administered is noted here. A brief description of any relevant information pertaining to the incorporation of the chemical into the medium and properties of the exposure medium is noted in the comment field. In inhalation exposure studies, a brief description of how the vapors were generated is reported in the comment field as well. Exposure medium codes and descriptions are presented in Table A-5.

**Table A-5**  
**Codes and Descriptions for Exposure Media**

Code	Description
AIR	Air. Used in inhalation exposure studies.
AQS	Aqueous solution. Used in plant studies or as an injection vehicle in bird and mammal studies.
CHM	Chemical only. Used if only the chemical is administered. The chemical is not dissolved in solution, oil, or any other media.
DW	Drinking water
DW+F	Drinking water plus food. Drinking water is the primary exposure medium while a background concentration is reported in the food.
F	Food
F+DW	Food plus drinking water. Food is the primary exposure medium while a background concentration is reported in the drinking water.
FLPP	Filter paper. Used in contact tests with earthworms.
MNU	Manure. Used in earthworm studies.
NR	Not reported
NSOLN	Nutrient solution. Used in plant studies.
OIL	Oil. Used if the exposure medium is known to be an oil solution but type is not specified
OIL_ACHS	Arachis oil. Often used as a vehicle in oral gavage or injection studies.
OIL_CORN	Corn oil. Often used as a vehicle in oral gavage or injection studies.
OIL_O	Other oil. Used if the exposure medium is known to be an oil solution but is a mixture of different types or other types not listed.
OIL_PNT	Peanut oil. Often used as a vehicle in oral gavage or injection studies.
OTH	Other. Exposure medium not listed. Specifics are noted in the corresponding comment field.
SAND	Sand
SAND&OM	Sand and organic matter mixture
SAND_CLTR	Sand culture. A solution is washed through silver sand daily.
SOIL	Soil
SOIL&MNU	Soil and manure mixture. Manure is usually used as a food source for earthworms.
SOIL&SAND	1:1 soil and sand mixture

**Table A-5 (continued)**

Code	Description
SOIL&SLDG	Soil and sludge mixture
SOLN	Solution. Exposure medium is assumed to be a solution but type is unknown.
SOLN_AQS	Aqueous solution. Used if the chemical was inorganic, and it was assumed the chemical is dissolved in an aqueous solution.
SOLN_O	Other solution. Used only if the exposure medium is assumed to be a solution of mixed composition or one not listed.
SOLN_OIL	Oil solution. Assumed.
W	Water

### Exposure Medium Background Data

Any background concentrations of chemicals that have the potential to impact the toxicity of the chemical of concern in soil, water, food, or air are noted here. In cases where the authors provide verified concentrations of the chemical in the control medium, this concentration is entered as background data. Compositions of fertilizer added to soil and any other supplemental substances are also noted here.

### Exposure Route ID

The exposure route code is selected from the drop-down list. Any further information relevant to the exposure route is noted in the comment field. For inhalation exposure studies, this comment field describes the inhalation chamber conditions (e.g., temperature, air flow). Exposure route codes and descriptions are presented in Table A-6.

**Table A-6**  
**Codes and Descriptions for Exposure Routes**

Code	Description
ALL	All exposure routes are used for chemical administration.
DC_SED	Direct contact in sediment
DC_W	Direct contact in water
DERM	Dermal contact (filter paper)
INH	Inhalation
INJ_EGG	Injection (egg)
INJ_IP	Injection (intraperitoneal)
INJ_IV	Injection (intravenous)
NR	Not reported
O	Oral
O/D	Oral and dermal
OC	Oral (capsule)
OD	Oral (diet)
OD+W	Oral (diet) plus exposure in drinking water

**Table A-6 (continued)**

Code	Description
OG	Oral (gavage)
OI	Oral (intubation)
OTH	Other
OW	Oral (water)
OW+D	Oral (water) plus exposure in food
U	Uptake (unknown whether through roots, seed coat, or both)
U_R	Uptake via roots
U_SC	Uptake via seed coat
U_SC+R	Uptake via seed coat and roots

### Length of Chemical Administration

The length of the chemical administration is briefly described here. If the exposure was intermittent (e.g., 4 h/d, 5 d/wk, for 7 wk), the total length of time over which the chemical was administered is reported (e.g., 7 wk). The chemical administration period for purposes of the Ecorisk Database is synonymous with the term exposure duration or period. The terms “chemical administration period” or “length of chemical administration” are used to clarify the difference between exposure duration and test period; test period includes both chemical administration and any periods during the study in which the organisms are acclimatized before exposure or further observed after exposure has ceased.

### Chemical Administration ID

The exposure duration code is selected from a drop-down list. The definitions and coding for exposure duration categories are shown in Table A-7. The exposure duration categories follow EPA (1999, 070923, Ref ID 0716).

**Table A-7**  
**Exposure Duration Categories and IDs for Birds, Mammals, Earthworms, and Plants**

Duration	ID	Birds and Mammals	Earthworms and Plants
Chronic	C	91 d or more	7 d or more
Subchronic	SC	14 to 91 d	3 to 6 d
Acute	A	13 d or less	2 d or less
Single dose	SD	One-time administration	One-time administration

### Exposure Frequency

The frequency of the chemical administration is noted here. For food and drinking water studies, it is often a continuous exposure where the exposure medium was provided throughout (also called *ad libitum*) the study. In inhalation exposure studies, the exposure frequency is either continuous or intermittent. In intermittent exposures, test organisms inhaled the chemical vapors for a certain number of hours per day

and number of days per week for a certain study length (e.g., 4 h/d, 5 d/wk, for 10 wk). In continuous exposures, the test organisms are exposed for 24 h/d, 7 d/wk.

### **Control Group Exposure Concentration(s) and Comment**

If a background concentration of the chemical of concern was reported in the primary exposure medium in addition to the administered amount, this concentration and its units are reported here. If no background concentrations were reported, a value of 0 mg/kg for soil or food, 0 mg/L for water, or 0 ppm for air is assumed.

### **Exposure Group Exposure Concentration(s) and Comment**

The concentrations of the treatment groups are noted here along with their units. If a background concentration was present in the primary exposure medium, this concentration is added to the basal concentration. If nominal (target) and empirical (verified or measured) concentrations are both provided, the verified concentrations are reported in the value field, and the target concentrations are noted in the comment field.

### **Nominal (Target) or Empirical (Verified or Measured) Concentration**

If it was not explicitly stated whether the concentration was nominal (target) or empirical (verified or measured), the concentration is assumed to be nominal (Nom). Otherwise, Nom or empirical (Emp) is noted based upon the information provided in the reference. If both nominal and empirical values were present, the empirical values are preferred over the nominal values, and the field is marked with Emp. Empirical values are preferred because they represent concentrations in the exposure medium that were analyzed and thus measured or verified; therefore, the empirical concentrations more accurately represent the concentrations that are available to the test organisms via the exposure medium. The nominal (target) concentrations are noted in the associated comment field. There are two fields for this data entry, one each for control and exposure groups, along with associated comment fields.

### **Dry or Wet Weight**

If the moisture basis of the concentration in the medium is not explicitly stated, NR is entered into the field. If the exposure route is oral by way of inhalation or by drinking water, gavage, intubation, or capsule, N/A is the entry. Otherwise, the moisture basis of the food or soil exposure medium is noted as WW for wet weight or DWt for dry weight. If both dry weights and wet weights are available from a study, dry weights are preferred. Dry weights are preferred because they eliminate variations in the PTV as a result of the wide variation of moisture contents of exposure media; the weights of the media are more easily compared when reported in dry weight. Furthermore, dry weight is the moisture basis of the TRV required for ESL calculations. There are two fields for this data entry, one each for control and exposure groups.<sup>3</sup>

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<sup>3</sup> During the early developmental stages of the Ecorisk Database, studies using exposure media of filter paper, aqueous solutions, and nutrient solutions for invertebrates and plants were evaluated. The moisture basis for these media was N/A. However, as more attention was placed on how well certain types of exposure media approximated the environmental exposure medium of concern (soil), these studies were not considered representative. Now, experiments containing these types of media are not evaluated.

### **Number of Individuals per Group**

The number of test organisms in each control and exposure group is noted. There are two fields for this data entry, one each for control and exposure groups.

### **Number of Sex per Group**

The number of females and/or males in each control and exposure group is noted. There are two fields for this data entry, one each for control and exposure groups.

### **Number of Replicates per Group**

If the number of replicates per control or exposure group was not clearly identified in a study, usually the number of individual organisms or sexual pairs that were caged separately is a suitable substitute. There are two fields for this data entry, one each for control and exposure groups.

### **Soil Characteristics (for Plant Studies Only)**

When the study is not a plant study, N/A is the default entry.

#### ***Soil Type***

The soil type and content are reported. Any other information not presented in the other fields of the soil characteristics section is also noted. See Example A-6.

#### **Example A-6 Soil Characteristics**

(a) Phaeosem, 3.85% sand, 74.90% silt, and 21.25% loam, water-holding capacity of 55.5%

(b) Ap horizon

(c) Sterilized shredded peat moss passed through 2-mm soil sieve and white silica sand. Base saturation of 93.9.

#### ***Soil Organic Matter***

If provided, the percent of organic matter (%OM) content in the soil medium is noted. If percent total organic carbon (OC), particulate OC, or just OC was reported, it is converted to OM as follows:

$$\%OM = 1.72 * \%OC$$

The notes regarding the conversion, including the source reference (EPA 2003, 85643; Ref ID 1400), are placed in the soil %OM field. If the percent of OM was not provided in the study but the percent content of sphagnum peat moss was, the percent content of the moss is considered to be equivalent to the percent of OM and is reported as so.

#### ***Soil Cation Exchange Capacity***

If provided in the study, the cation exchange capacity (CEC) in meq/100 g of soil is reported. If the CEC is not provided, NR is entered.

**Soil pH**

If provided, the soil pH is reported here. If the soil pH is not provided, NR is entered.

**Growth Medium Characteristics (for Invertebrate Studies Only)**

When the study is not an invertebrate study, N/A is the default entry.

**Growth Medium Type**

The soil type and content are reported. Any other information not presented in the remaining soil characteristics section is also noted. See Example A-7.

**Example A-7 Growth Medium Types**

- (a) Petri dish with 30 g (dry mass) of screened soil mixed with aged horse manure (75% moisture)
- (b) Sand (0.2- to 2-mm particle size) from C horizon mixed with well-decomposed cattle dung (1:2, vol:vol)
- (c) Sandy loam soil with 17% clay, 5.5% CaCO<sub>3</sub>

**Growth Medium Organic Carbon**

If provided, the percent of organic carbon (%OC) content in the soil medium is noted. It is converted from %OM using the following equation:

$$\%OC = \frac{\%OM}{1.72}$$

The conversion is noted along with the source reference of EPA (2003, 085643, Ref ID 1400) in the exposure medium field.

**Growth Medium pH**

If provided, the growth medium pH is reported here. If it is not provided, NR is entered.

**Growth Medium Percent Moisture**

If provided, the moisture content of the growth medium is reported. If it is not provided, NR is entered.

**Food**

If food for the earthworm was also provided in the soil, and it was explicitly noted as such or reasonably deduced, it is reported here. Examples are manure and litter.

**Organic Matter ID (for both Plant and Invertebrate Studies)**

If the %OM content in the soil or growth medium was 10% or less, it is coded as low. If the %OM was greater than 10%, it is coded as high. The high and low IDs are based on EPA (2003, 085643,

Ref ID 1400), where studies are rejected if the soil exposure medium contains greater than 10% OM because OM may affect the bioavailability of the test chemical to the organism. If OM is not reported, NR is entered and the study is excluded from the rest of the PTSE process. Otherwise, the entry is N/A for bird and mammal studies.

If %OC was reported, it is converted to %OM for the determination of the OM ID. If both the %OC and the percent content of sphagnum peat moss were reported, the content of the peat moss is used to set the OM ID.

### **All Measurements Reported**

All measurement endpoints in the study are listed, regardless of whether they are ecologically relevant or not. The purpose of this field is to provide a complete listing of the various measurements applied in the experiment so that users of the database know what was measured, and if they feel a measurement is ecologically relevant but is not evaluated in the PTSE, they can obtain the reference and further supplement their information.

### **Measurements Not Evaluated and Why**

The measurement endpoints that are not evaluated in the PTSE are listed here. These include “other” effects, such as physiological functions, histopathology, cancer, and behavior (see Focus Measurement Category in section A-2.1.4), as well as any ecologically relevant measurements that are accounted for within measurements that are evaluated. If a plant study reported measurements of both fresh and dry weight values of leaves, only the dry weight information would be evaluated. The fresh weight information would not be evaluated and the reason why (i.e., dry weight is a more accurate measurement of the true mass of the plant because it eliminates the additional weight that is dependent upon varying moisture content of individual plants) is noted in this field. Another example would be to evaluate the percent mortality of juveniles but not the number of juveniles that died because the number of juveniles that died is incorporated as a percentage of the total number of juveniles in the experiment. The number of juveniles died would be reported in the measurements not evaluated and why field along with the explanation of why it was not evaluated. See Example A-8.

#### **Example A-8 Measurements Not Evaluated and Why**

(a) Food consumption, organ weights, hematocrits, hemoglobin concentrations, gross pathology, and organ, blood, and egg residues will not be evaluated in this Part 1 review because their relationships to adverse effects on population health are not clear.

(b) Food consumption, testes weight, liver weight, liver manganese, serum T, and general locomotor activities are not evaluated in this Part 1 because their relationships to population health are not clear. Body weight is not considered in this Part 1 because it is part of the growth rate measurement, which is already accounted for in this Part 1 review.

### **Author-Reported Effect Levels**

If the authors calculated their own effect levels, these are reported in this field. The LC<sub>50</sub> (or LD<sub>50</sub>) or EC<sub>50</sub> (or median effective dose, or ED<sub>50</sub>) are most often the effect levels reported. NOAELs/NOECs and LOAELs/LOECs are also reported. The endpoints that the reported effect levels represent are also specified.

## Experiment Comments/Author Conclusions

An overall summary of the data is presented for the reference, along with mention of any other factors that may have contributed to or confounded the results of the focus measurements in the experiment (e.g., mortality attributed to an infection outbreak and not the chemical exposure). Also, any further general observations on focus measurements not carried forth to Part 2 reviews may be reported here. Page numbers and table or figure designators from the reference should be included to support the comments.

### A-2.1.4 Measurements and Results

Focus measurements (endpoints) that are evaluated in the Part 1 PTSE are limited to reproduction, development, survival, weight changes of adult or mature organism, and size changes of adult or mature organism. Only these categories are evaluated because they are ecologically relevant. In other words, these types of measurements are more directly linked with population health. Adverse effects observed in “other” endpoints, such as seminiferous tubule diameter, require too much speculation as to the degree of their impact on population health and are thus not evaluated in the PTSE process.

### Focus Measurement Effect ID

Experiment effect IDs are created by simply adding an alphabetic identifier to the end of the experiment ID for each focus measurement (see Example A-9).

#### Example A-9 Experiment Effect IDs

0025\_SE\_1A

0025\_SE\_1B

0025\_SE\_2A

0517\_50-29-3\_2A

### Focus Measurement

A focus measurement label is provided in the focus measurement field. The label should follow the labeling present in the study, but exceptions occur where symbols such as # are replaced with the word “number” or phrase “number of,” where % is replaced with the word “percent” or “percentage,” or where / (slash) is replaced with the word “per” for clarification and for data consistency.

### Focus Measurement Category

The category of the focus measurement is then coded and entered in this field (see Table A-8).

**Table A-8**  
**Category Codes and Descriptions for Focus Measurements**

Code	Description
WC	Weight change (adult)
NR	Not reported
O	Other
R/D	Reproduction/development
S	Survival
SzC	Size change (adult)

### ***Reproduction/Development***

If development or mortality was measured in juvenile organisms or immature plants and they were exposed to the chemical through parental exposure, the measurement is coded as reproduction/development (R/D) because it is considered to be a measurement of the ability of the parents to produce offspring that can develop into reproductive adults, and exposure reflects the reproductive cycle. Growth of a juvenile organism or immature plant that was directly exposed to the chemical is coded as R/D because it reflects the potential for the juvenile or immature plant to develop normally into a reproductive adult.

### ***Adult Weight or Size Changes***

If weight change for mature organisms is measured, it is considered a weight change and not development. Likewise, if a change occurs in size of a mature organism (e.g., height or root length of plants), it is noted as a size change.

### ***Survival***

If a juvenile organism or immature plant was directly exposed to the chemical, mortality is coded as S (survival) because it is considered a measurement of the ability of the organism to survive to reproductive maturity, and the exposure did not occur during the reproductive cycle.

### ***Other***

Other measurements are those that are considered to be less directly linked to effects on populations (e.g., tumors, tissue residues, cholesterol level, and behavioral changes) and are generally not reviewed unless the author(s) provides a clear correlation with the measurement and its effect(s) on population health (e.g., behavioral effects that impact reproduction, such as number of mounts in mice) or the data set is very limited.<sup>4</sup>

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<sup>4</sup> There were cases where the measurement was associated with reproduction, but the relationship of the parameter to effects on population health is not clear; therefore, these types of measurements are also coded as O. Examples include sperm motility, seminiferous tubule diameter, and testicular enzyme activities. During the development of the database, these measurements were evaluated but later excluded from consideration for TRVs. Currently, these measurements are no longer evaluated unless a clear relationship to population numbers is described.

**Focus Measurement Frequency**

The number of times the measurement was recorded is noted here (e.g., once per week, or 4 h/d, 5 d/wk).

**Focus Measurement Duration**

If the observation of the focus measurement lasted more than just an instant (e.g., behavioral observations that may take 10 min of observation), the length is noted in this field.

**Focus Measurement Critical Life Stage Category**

A life stage of an organism is considered to be a critical life stage if exposure to a chemical during this life stage is expected to result in a negative impact on the population health of that organism. For the purpose of deriving TRVs, a critical life stage is defined as a life stage associated with a chemical exposure that occurs during the reproductive cycle of the test organism and/or during the development of the immature test organism. For an endpoint to be considered development, it has to fall into one of two scenarios in which measurements must reflect either the development of immature organisms that were exposed via parents or the development of immature organisms directly exposed to the chemical. Reproduction and development endpoints directly reflect effects on the size and character of the next generation of the population. Note that not all endpoints associated with seemingly reproductive/development functions are coded as R/D (see Focus Measurement Category above).

***Chronic – Critical Life Stage***

If an endpoint reflects a critical life stage, the associated effect level may be considered to be equivalent to a chronic exposure endpoint regardless of the actual chemical exposure duration associated with this endpoint. The reasoning behind this assumption is as follows: a chronic study is preferred over a single-dose, acute, or subchronic chemical exposure study because it is more likely to capture effects that reflect critical life stages that are relevant to population success. Therefore, it is assumed that any duration of chemical exposure that is associated with a critical life stage endpoint captures potential effects on population success as a chronic study does. This effect is then considered to be equivalent to a chronic exposure effect regardless of the actual chemical administration period. Ultimately, if an endpoint is categorized as chronic because of a critical life stage, our certainty of this effect predicting the impact of a particular chemical on population success increases. Such endpoints are categorized as chronic-critical life stage (C-CL).

***Critical Life Stage Only***

If the critical life stage endpoint is a type that does not directly reflect effects on the size or character of the next generation of the population, certainty in predicting the impact of a particular chemical on population success is not increased. There are nonreproductive and nondevelopmental endpoints that reflect critical life stages because chemical administration occurred during the reproductive cycle of adults, during the development phase of juveniles, or during an embryo stage. Examples of such endpoints include survival for juvenile organisms (who are still undergoing development, a critical life stage), body weight measured for adult organisms in the reproductive cycle, clinical signs during reproductive cycle, or egg length. However, a measurement of these types of endpoints is not a direct-measurement of a critical life stage reproductive/development endpoint; thus, less certainty is associated with the effect level assigned to it. The actual exposure length remains (i.e., single dose remains single dose) when determining the application of UFs in the TRV derivation process. Using juvenile mortality as

an example to further illustrate the logic, it is difficult to assess the extent to which the critical life stage of development of juveniles impacts their mortality rate. Therefore, by not classifying this juvenile mortality endpoint as C-CL, the PTV that results will be lower, thus more protective, in cases where the exposure duration is acute or subchronic because of UFs that must be applied to extrapolate to a chronic effect level. Such endpoints are categorized as just critical life stage (CL).

### **Coding**

In application, coding for critical life stage generally follows the guidelines below:

- All reproduction/development endpoints are coded as C-CL, regardless of actual chemical exposure duration.
- Other endpoints (such as adult or juvenile survival, adult weight or size change, or other characteristics [S, WC, SzC, and O, respectively]) in which chemical administration occurred during a critical life stage are coded as CL.
- Endpoints in which chemical administration did not occur during a critical life stage are coded as non-CL.
- Endpoints in which it is unknown whether or not chemical administration or measurements were taken during critical life stages are coded as NR.

Further exceptions occur where professional judgment deems the coding that would follow the guidelines to be inappropriate. Examples include the following:

- A study where chemical administration occurred for a lengthy amount of time, but measurements of the effects occurred only for part of the chemical administration period. See Example A-10a.
- A study where a critical life stage occurred, but organisms of a certain treatment group died before the critical life stage began. See Example A-10b.
- A study where a survival endpoint can be classified as chronic as a result of a critical life stage because the immature organism was directly exposed to the chemical, and chemical exposure encompassed this immature life stage. See Example A-10c.

**Example A-10 Exceptions to Coding for Critical Life Stages**

(a) Ivankovic and Preussman (1975, 059251, Ref ID 0010), Experiment 1: Adult rats were exposed to a chemical 90 d before mating and through reproduction for at least another 30 d, and body weight measurements took place only up until the mating period began. This endpoint would be characterized as non-CL. The body weight measurements had not taken place while the rats were subjected to additional stress of reproduction; therefore, they were not expected to be more susceptible to adverse weight change effects.

(b) Aulerich et al. (1974, 059794, Ref ID 0016): Adult mink were exposed to 5 ppm of methylmercury or 10 ppm of mercuric chloride. Authors wished to obtain information on adult body weights, kit body weights, adult mortality, reproductive measurements such as number of females mating and number of kits born alive vs. dead, and clinical signs. All organisms fed 5 ppm of methylmercury in the diet died before breeding season. Adult body weights and critical life stage codes for the mink in the 10-ppm mercuric chloride group would be WC and CL, respectively. However, for mink in the 5-ppm methylmercury group, the codes would be WC and non-CL, respectively, because the body weight measurements did not continue through reproduction as the mink died before breeding season.

(c) In Brunström et al. (1991, 070812, Ref ID 0666) and Gogal et al. (2002, 089461, Ref ID 1216), bird eggs received injections and embryo mortality was measured. This measurement would receive an endpoint coding of S and a critical life stage coding of C-CL. This scenario is also evident where germination of seeds (considered survival, from seed to seedling) was measured.

When considering the use of PTVs in TRV derivation, an endpoint associated with a C-CL category is preferred over one with a CL or non-CL life stage effect. All critical life stage designations are considered to provide support of PTV eliminations or selections for use in TRV development.

**Test Period Duration and Category**

The chemical administration plus any additional time before and/or after the exposure is noted here. If the test organisms were quarantined and/or acclimatized for a period of time before exposure started, or if measurements continued to be recorded after exposure ceased, this length of time is counted in the test period. Results observed after exposure ceased are not usually considered because they are not considered relevant for predicting effects of continuous chemical exposures (such as those that may be found in the environment).

**Focus Measurement Dose Response**

First, the table and/or page number from which the results were taken is noted. Notes on which exposure levels resulted in adverse effects for the focus measurement follow. General observations on dose-response trends are also reported. If no statistics were used, a summary of the results suffices. Basically, entry in this field provides an insight into the results observed by the researchers of the study at various exposure levels and compares them to results for controls.

**Focus Measurement Statistical Test and Confidence Level**

If provided, the statistical test and/or alpha level used to determine significant adverse effects for the measurement are noted here.

**Focus Measurement Comments/Effect Levels**

The effect level(s) are assigned (if not already provided by the authors) and documented in this field. Discussion of whether they are author-reported or reviewer-assigned effect levels and whether the assignment was based on statistics that were provided or not is also presented here as well as in Example A-11. Furthermore, any evidence of dose-response trends, post-exposure related effects, insufficient data, or other conditions that may affect the assignment of the effect levels is also discussed in detail (see Example A-11).

**Example A-11 Focus Measurement Comments/Effect Levels****(a) Author-reported effect levels**

(i) The authors reported effect levels for 5-day emergence: NOEC = 312 mg/kg, and LOEC = 1040 mg/kg. The EC<sub>10</sub> is 307.5 mg/kg dry soil, the EC<sub>20</sub> is 3112.6 mg/kg dry soil, and the EC<sub>50</sub> is > 3120 mg/kg dry soil.

(ii) The researchers reported an LD<sub>50</sub> of 2690 mg/kg with 95% confidence limits of 1571 to 57,063 mg/kg. The researchers did not provide a NOAEL or LOAEL, and statistics were not provided; however, sufficient mortality data were available, so the Dunnett's multiple comparison test was applied by the reviewer in order to determine statistical significance at  $p = 0.05$ . Based on this, statistical significance was determined at 1350 mg/kg and higher. Therefore, the 810-mg/kg level will be used in the NOAEL calculation while the 1350-mg/kg level will be used in the LOAEL calculation.

**(b) Reviewer-assigned effect levels**

A NOAEL can be inferred. Since no effects were observed at the highest level of 32 ppm of mercury, this is designated as the NOAEL.

No significant differences at  $p < 0.05$  were found; however, the decreases in fertilization at 2 and 8 ppm were approaching significance ( $0.05 < p < 0.10$ ), and differences between the 2- and 8-ppm and 4- and 0-ppm groups were at least 22%. Note that the 4-ppm group had a higher fertility rate than, or similar fertility rate as, the 0-ppm group. The author discusses possible reasons for the enhancement at 4 ppm, including bacteriostatic or fungicidal activity or stimulation. Based on these results and a conservative approach, the 2-ppm level is used for the LOAEL because adverse effects were seen at this lowest dose level (22% reduction in fertility) compared to control.

**(c) Dose-response trends**

There were no clear dose-related trends in any of the three 10-d groups, but there was a pattern of 4-ppm groups having the highest hatchability of the three exposure groups. This effect (hatchability) will not be carried further because it is difficult to determine a NOAEL and LOAEL based on three different age groups and varying responses.

**(d) Post-exposure related trends**

There were significantly lower body weights in the 30-, 100-, and 300-ppm groups compared to controls on day 13 of gestation. However, only the 100- and 300-ppm groups continued to have significantly lower body weights on day 21 of gestation, after exposure ceased on day 15. The possibility exists that the absence of a significant difference at the 30-ppm level was a result of the rats having had time to recover following the cessation of exposure on day 15 of gestation. Therefore, the assignment of effect levels is based on significant effects that occurred during exposure rather than effects that were present after 6 days of recovery in order to be protective. Based on this, the 30-ppm level is used for the LOAEL calculation.

**Example A-11 (continued) Focus Measurement Comments/Effect Levels**

(e) No reported statistics

Because it is not clear in the text or statistics which treatment level showed a significantly lower percentage of hens laying compared to controls, the treatment that shows a decrease of 20% or greater compared to controls will be considered significant (Suter et al. 1995, 089449, Ref ID 1088). Based on this, the 210-ppm wet-weight level (target concentration of 200 ppm) had 25% fewer hens laying and will be used for the LOAEL. The 15.2-ppm wet-weight level (target concentration of 20 ppm) will be used for the NOAEL.

(f) Data insufficient for TRV development

An increase in mean egg production associated with increasing mercury exposure does not appear to be an adverse effect and will not be evaluated further.

As noted, phencyclidine at the highest concentration tested (60 mg/kg) stimulated growth, as opposed to depressing it; thus, this is considered not detrimental to the organism and not suitable for deriving a TRV. This focus measurement will not be evaluated further.

**Author-Reported Effect Levels**

If the authors reported their own effect level(s) for the focus measurement (e.g., NOAEL for average number of live fetuses) or its category (e.g., NOAEL for reproduction), the effect level(s) and what it represents is entered into this comment field. It is then decided if each effect level accurately represents the results of the focus measurement. For example, if the authors reported a NOEC that was interpolated based on reproductive toxicity data for four plant species in a study, this NOEC, while reported in the Part 1 database, may not be considered appropriate for use as a NOEC for one species in particular. If the author-provided effect level is not considered appropriate, the reviewer must further assess the validity of the reported results for use in Part 2 (see Reviewer-Assigned Effect Levels below).

**Reviewer-Assigned Effect Levels**

If there is no author-reported effect level(s) or the level(s) reported is found to not be suitable for use (see Author-Reported Effect Levels above), the reviewer must assign an effect level or effect levels to the focus measurement based on the reported data using best professional judgment. Dose-response trends, post-exposure related effects, and availability of statistics are considered in whether to continue to assign effect levels or to determine that the data are insufficient for TRV development.

**Dose-Response Trends**

If a clear dose-response trend and an exposure concentration can be noted at which no adverse effects and/or at which adverse effects were first observed, the exposure concentration that produced no observed adverse effects is used for the NOAEL/NOEC, while the exposure concentration at which adverse effects were first observed is used for the LOAEL/LOEC. Where statistics were used by the researchers of the study, the first exposure concentration to show a statistical significance compared to controls is considered to produce an adverse effect and is used in the LOAEL/LOEC calculation. The next lower exposure concentration is then considered for the NOAEL/NOEC calculation.

### ***Post-Exposure Related Effects***

If observations continued after exposure ceased, the results for this period are not usually included in the assignment of effect levels because it is assumed the organisms of concern are continuously exposed to contaminants and thus no time for recovery is allowed. That is, the adverse effects that occur during exposure are most relevant for predicting effects of continuous chemical exposure. The assignment of a NOAEL/NOEC to a concentration at which adverse effects were observed during exposure but not afterwards may not be protective enough, so the concentration is considered a LOAEL/LOEC. However, results that occurred after exposure ceased are still noted and considered to lend support to the effect level assignment.

### ***No Reported Statistics***

If statistics were not reported by the author, the reviewer either applies his or her own statistics or, more often, considers the exposure concentration with a difference of 20% or greater effect compared to control groups to be significant. If this guideline for using a difference of 20% or greater effect is followed, Suter et al. (1995, 089449, Ref ID 1088) is cited. The guideline for using a difference of 20% or greater effect is followed by ORNL (Suter et al. 1995, 089449, Ref ID 1088) in its selection of effect levels, and it is based on EPA regulatory practices. This method for determining biological significance comes from the inference that the LOEC derived from studies in which terrestrial birds are exposed to contaminants in the diet usually corresponds to a 20% effect on individual response parameters (Suter et al. 1995, 089449, Ref ID 1088). Any difference of 20% or greater is considered a biological significance rather than a statistical significance. For purposes of assigning effect levels, biological significance is considered to be equivalent to statistical significance.

Statistics are often used when the appropriate amounts and types of data are clearly presented for each treatment group and control group in tables in the paper. Best professional judgment is used to determine which statistical test would be appropriate for the data presented.

### ***Data Insufficient for TRV Development***

If the reviewer determines that the data for the focus measurement being evaluated are insufficient for TRV derivation, it is noted that a Part 2 evaluation will not be completed for this measurement. Also, “\_NoPTSEP2” is attached to the end of the experiment effect ID (e.g., 0025\_CD\_1A\_NoPTSEP2).

Conditions in which the data are not sufficient for TRV derivation:

- Only trends are mentioned in the text by the investigators, and they do not clearly illustrate the point at which exposure level adverse effects began.
- Numerical data are available, and authors only hint at results.
- Results of the study are too varied (no clear dose-response or time-related trend), and no statistics are applied.

## **A-3.0 PTSE PART 2, STUDY EVALUATION AND PRIMARY TOXICITY VALUE CALCULATION**

The Part 2 review process is based on evaluating and then scoring the data obtained from the reference in the Part 1 and then calculating a PTV and assigning it a confidence rating. Section A-3.1, Data Evaluation and Scoring Guidelines, provides instruction for evaluating the study and documenting the evaluation. Section A-3.2, PTV Calculation Guidelines, provides instruction for calculating the PTV and

documenting the derivation. Section A-3.2.8, PTV Confidence Rating Guidelines, provides instruction for assigning a confidence rating to each PTV.

### **A-3.1 Data Evaluation and Scoring Guidelines**

#### **A-3.1.1 General PTSE Information**

The data in the following fields are imported from the Part 1 data-entry database:

- Reference ID
- Chemical ID
- Experiment ID
- Experiment purpose
- Effect ID
- Focus measurement label

#### **Review Date**

The date the review is started is entered here. It can be superseded by the date the record was updated (edited).

#### **Reviewer Initials**

The initials are entered or selected from a drop-down list of current reviewers. Initially, the original reviewer of the record is entered. This can be superseded by the initials of the reviewer who updated (edited) the record.

#### **A-3.1.2 Study Design and Documentation Score**

##### **Control**

Was a suitable control present? Was it a negative (no toxicant applied, but similar to treatments in all other aspects), positive (standard such as dieldrin used for comparisons of relative toxicities), or solvent control? An example of a solvent control is illustrated in an invertebrate toxicity study in which HMX was first dissolved in a solvent (acetonitrile) before application to the soil medium. The solvent control would consist of the invertebrates exposed to a soil medium containing only acetonitrile.

If a control group is not included in the experiment, but effect levels are provided by the authors, the scoring is based on whether or not the absence of the control group affects the ability of the reviewer to verify these effect levels or assign effect levels. If only an effect level of other (e.g., LC<sub>50</sub>, EC<sub>20</sub>) is provided by the authors, the score is not penalized because usually in these situations it is reasonably assumed that multiple concentrations were administered to extrapolate the lethal or effective concentrations. Also, a published method is often used by the authors to determine these effect levels. Therefore, it can be assumed that at least one control group was built into the study design or that control groups were not needed as long as an appropriate dose-response curve was produced to extrapolate the other effect level.

If a NOAEL/NOEC and/or LOAEL/LOEC was provided by the authors, but the absence of controls makes it difficult for the reviewer to verify the effect levels, the score will be penalized. This indicates that while the effect levels are still used, caution should be exercised in the interpretation of these values within the TRV data set because the reviewers could not ascertain that the effect levels were determined appropriately.

There are situations where control groups and effect levels are not reported, but a NOAEL/NOEC and LOAEL/LOEC, and/or NOAEL/NOEC and LOAEL/LOEC pair is assigned by the reviewer nonetheless. The score is not penalized in this scenario. This can happen for mortality endpoints where only one exposure level was administered, and it is reported that 0% mortality was observed at this concentration. This exposure concentration is used for the NOAEL/NOEC. On the other hand, if a reasonable percentage of mortality occurred (e.g., more than 50% for birds or mammals is considered adverse), this exposure concentration is used for the LOAEL/LOEC. Furthermore, two exposure concentrations in a mortality study can also lead toward the assignment of a NOAEL/NOEC and LOAEL/LOEC pair without controls if the lower concentration resulted in no mortalities while the higher concentration resulted in greater than 50% mortality.

Control group score:

- 1 A control group was included, or a control group was not included or reported but was not needed to verify or assign effect levels.
- 0 A control group was not included, and effect levels provided by the authors could not be verified.

### **Exposure Groups**

Was more than one exposure group present? Exposure concentrations are listed. It is also noted whether these concentrations are nominal or measured.

Exposure group score:

- 1 More than one exposure group was used.
- 0 Only one exposure group was used.

### **Test Organism Details**

The test organism name, age or life stage, sex, and origin/source are listed, if provided.

### **Organism Details Score**

Up to four pieces of information can be provided for birds and mammals: name (common and/or scientific), age, sex, and source/origin. Up to three pieces of information are available for invertebrates and plants: name (common and/or scientific), age, and source/origin. Scoring is as follows:

- 4 All information is provided.
- 3 Three pieces of information are provided.
- 2 Two pieces of information are provided.

- 1 One piece of information is provided.
- 0 No information was available.

### **Dose Rate Parameters**

In bird and mammal studies, are the exposure concentrations reported in daily dose rates of mg/kg/d, or are body weight, food ingestion rate, and/or water ingestion rate parameters available to convert the provided dose units to mg/kg/d?

For earthworm and plant studies, the entry is N/A because the concentrations are already normalized to the amount of chemical in soil (e.g., mg chemical/kg soil), which is what the PTV is based on.

### **Dose Rate Parameter Score**

Dose rates can be calculated using two dose rate parameters: body weight and either an ingestion rate (for water or food) or an inhalation rate.

- 2 Both dose rate parameters were provided, the ingestion or inhalation rate was already normalized to body weight, or none of the dose rates are applicable (N/A) because the daily dose rate was reported by the authors.
- 1 One dose rate parameter was provided.
- 0 No dose rate parameters were provided.

### **Exposure Dose Concentration**

Are the exposure concentrations nominal (target) or empirical (i.e., verified or measured) concentrations, and what is their moisture basis? If the exposure medium is not food or soil (e.g., vapors in an inhalation study, oil vehicle used in an oral gavage administration), moisture basis is N/A. If chemical administration was already provided as daily dose rates, moisture basis is canceled out and this aspect becomes N/A as well.

Dose concentration basis score:

- 2 Measured, dry weight or N/A.
- 1.75 Measured, wet (fresh) weight
- 1.5 Nominal, dry weight or N/A
- 1.25 Nominal, wet (fresh) weight
- 1 Measured, unknown
- 0.75 Nominal, unknown
- 0.5 Unknown, dry weight or N/A
- 0.25 Unknown, wet (fresh) weight
- 0 Unknown, unknown

## Statistics

Are statistics provided, and if so, what are the test and p-value or confidence limit? If statistics were not provided, was data presented in tables in such a way that the reviewer was to apply his/her own statistics or analysis? Did the measurement show no effects that could be analyzed by statistics (e.g., zero mortality)?

Statistics score:

- 1 Both the statistical test and confidence level are reported.
- 0.5 The statistical test or the confidence level is missing, or if neither is reported, data are available for reviewer to run analysis.
- 0 Neither the statistical test nor confidence level are reported, and data are not adequate for reviewer to run analysis.

### A-3.1.3 Test Organism Score

#### Taxonomic Relationship of Test Organism

The screening receptor is a species that represents a functional food group and exposure pathway (e.g., intermediate carnivore [50% flesh/50% invertebrate], burrowing mammal [inhalation]) in an area of concern. The screening receptor group (i.e., bird, mammal, invertebrate, or plant) that the test organism best represents is noted. It is followed by a description of how closely the test organism is related to the screening receptor taxonomically.

Taxonomic relationship score:

- 2 The test organism is related to at least one screening receptor at the order, family, genus, or species level. (Not applicable to plant or invertebrate test organisms)
- 1 The test organism is related to at least one screening receptor at the class level. (Not applicable to plant or invertebrate test organisms)
- 0 The test organism is not related to a screening receptor at the class or more specific level or is a plant or invertebrate.

#### Basis for Use of Test Organism

Did the investigators of the study provide a reason for using the test organism?

Test organism basis score:

- 1 The researchers indicated, or it can be reasonably assumed, why the particular test organism was chosen.
- 0 It is not known why the test organism was chosen.

### A-3.1.4 Exposure Conditions Score

#### Test Environment

Was the study conducted in a laboratory or other controlled environment with exposure only to a single chemical?

Exposure environment score:

- 1 The study is based on a field or laboratory study from which a single chemical exposure can be discerned.
- 0 The study is not based on a field or laboratory study from which a single chemical exposure can be discerned.

#### Test Exposure Chemical

The chemical of potential ecological concern (e.g., cadmium), not the chemical form (e.g., cadmium chloride), is noted here. Scoring is not applicable to this field.

#### Test Exposure Medium (to Represent Food and Drinking Water TRVs)

For bird and mammal studies,

- the test exposure medium is noted, and
- the exposure media for TRVs and ESLs are noted as follows:
  - ❖ for food media studies, “TRVs: food; ESLs: sediment and soil,” and
  - ❖ for drinking water media studies, “TRV: drinking water; ESL: water.”

These fields are not applicable for earthworm and plant studies or mammal inhalation studies (i.e., N/A is entered).

Food equivalency score:

- 1 The test exposure medium is equivalent to food.
- 0.5 The test exposure medium is similar to food (capsule, oil, or solid bolus).
- 0 The test exposure medium is not equivalent or similar to food (drinking water or other).

Drinking water equivalency score:

- 1 The test exposure medium is equivalent to drinking water.
- 0.5 The test exposure medium is similar to drinking water (aqueous solution or chemical).
- 0 The test exposure medium is not equivalent or similar to drinking water (food or other).

### **Test Exposure Medium (to Represent Soil TRV)**

For earthworm and plant studies,

- the test exposure medium is noted, and
- the exposure media for the TRV and ESL are noted (e.g., “TRV: soil; ESL: soil”).

This field is not applicable for bird and mammal studies (i.e., N/A is entered).

Soil equivalency score:

- 1 The test exposure medium is equivalent or similar to soil.
- 0 The test exposure medium is not equivalent or similar to soil.

### **Test Exposure Chemical Interactions**

Even if there are chemicals in the exposure medium besides the chemical of concern, they may be naturally occurring and are not considered an interaction. Only when chemical or physical properties change during the course of the experiment are they considered an interaction. If an interaction is not reported by the author, it is noted that none is expected.

Chemical interaction score:

- 1 Chemicals and properties that could potentially affect the toxicological impact of the test exposure chemical on the test organism are not present in the test exposure medium.
- 0 Chemicals and properties are present and could potentially affect the toxicological impact of the test exposure chemical on the test organism.

### **Test Exposure Route**

The test exposure route and whether it is similar to the exposure route of concern are described. For example, uptake via seed coat and/or roots is the exposure route of concern for plants. If in a study, plants were exposed to the chemical through spraying on the leaves, this is not considered similar to the exposure route of concern.

Exposure route score:

- 1 The test exposure route is equivalent to the ESL exposure route of concern (for birds and mammals, food, drinking water, or inhalation; for invertebrates, oral/dermal; and for plants, uptake).
- 0.5 The test exposure route is similar to the ESL exposure route of concern (for birds and mammals only, oral intubation or gavage).
- 0 The test exposure route is not equivalent or similar to the ESL exposure route of concern.

### **Test Period (Including Chemical Administration)**

The test period duration, which includes any period of acclimatization before exposure and the time period for additional observations after exposure, is noted here. The percent of the test period during which chemical administration occurs is also described. For example, “The test period was 90 d, and

chemical administration occurred the entire time (100%),” or “The test period was 120 d, and chemical administration occurred during the first 90 d and composed 75% of the total test period.”

Test and exposure period score (based on chemical administration period):

- 3 Chronic
- 2 Subchronic
- 1 Acute
- 0 Not reported

Exposure durations are defined in Table A-9.

**Table A-9**  
**Exposure Durations**

Test	Bird or Mammal	Invertebrate or Plant
Chronic	>90 d	>6 d
Subchronic	14 to 90 d	3 to 6 d
Acute	<14 d	<3 d

### Critical Life Stage

If the chemical administration occurred during the reproduction or development period of the test organism, it is noted as a critical life stage in this field.

Critical life stage score:

- 1 Chemical administration occurs during a critical life stage.
- 0 Chemical administration does not occur during a critical life stage.

### Test Exposure Frequency

The frequency of exposure to which the test organisms were exposed to the test chemical is noted here (e.g., continuous or intermittent, 7 h/d, 5 d/wk). For bird and mammal oral ingestion studies, an exposure that is at least once daily or *ad libitum* is considered frequent. For mammal intermittent inhalation studies, an exposure that constitutes 70% of the chemical administration period is considered frequent (based on most studies exposing animals 5 d/wk). Earthworm and plant soil studies typically have an exposure regimen where the test organism is exposed continuously to the chemical in soil. If this is not the case, the frequency score follows the guideline for bird and mammal oral ingestion studies.

Exposure frequency score:

- 1 The test exposure frequency is continuous or frequent enough to represent the chemical administration period.
- 0 The test exposure frequency is not continuous or frequent enough to represent the chemical administration period.

### **A-3.1.5 Measurement(s) and Result(s)**

#### **Focus Measurement Effect Category**

The focus measurement label (i.e., the measurement endpoint) as the author(s) reported it (e.g., number of pups per dam, shoot length) is noted. The endpoint category in which the focus measurement belongs is also sometimes noted for clarification (e.g., development [body weight vs. adult body weight change] or survival [juvenile mortality vs. development, juvenile mortality for those organisms exposed to the chemical via parents]).

Endpoint category score:

- 4 Reproduction or development
- 3 Survival
- 2 Adult weight or size change
- 1 Other

#### **Measurement of Focus Measurement**

If measurements took place at appropriate times during and after exposure to reflect effects and trends that can be attributed to exposure, YES is entered.

Focus measurement length score:

- 1 The focus measurement reflects the entire chemical administration period.
- 0 The focus measurement does not reflect the entire chemical administration period.

#### **Focus Measurement Effect Level**

The effect levels are noted here. If a NOAEL/NOEC and LOAEL/LOEC are both available, the magnitude of difference is calculated and reported.

Effect level score:

- 6 NOAEL and LOAEL, NOEL and LOEL, or NOEC, LOEC, and values are within a factor of 3.
- 5 NOAEL and LOAEL, NOEL and LOEL, or NOEC, LOEC, and values are within a factor of 10.
- 4 NOAEL and LOAEL, NOEL and LOEL, or NOEC, LOEC, and values are not within a factor of 10.
- 3 NOAEL, NOEL, or NOEC only
- 2 LOAEL, LOEL, or LOEC only
- 1 Other effect level (e.g., LD<sub>50</sub>, LC<sub>50</sub>, or EC<sub>50</sub>) only

#### **Effect Level ID**

The appropriate code is selected from a drop-down list. Options are the following:

- NLOTH = NOAEL/NOEC, LOAEL/LOEC, and other effect level, such as LC<sub>50</sub>
- NL = NOAEL/NOEC and LOAEL/LOEC

- N = NOAEL/NOEC
- NOTH = NOAEL/NOEC and other effect level
- L = LOAEL/LOEC
- LOTH = LOAEL/LOEC and other effect level
- OTH = Other effect level

Scoring is not applicable in this field.

### **A-3.1.6 PTV Calculation**

Below are brief descriptions of the data entry fields for this section. See section A-3.2 for detailed instructions on how to complete these calculations.

#### **Value, Units**

The calculated or author-reported daily dose rate value (PTV) is recorded here along with its units. The units are mg/kg/d for birds and mammals and mg/kg for invertebrates and plants.

#### **Duration**

The chemical administration period is noted here. However, if the chemical administration period is acute, subchronic, or chronic, and the measurement is categorized as chronic-critical life stage, "Chronic-Critical Life Stage" replaces the chemical administration period.

#### **Calculation**

The daily dose rate and unit conversion calculations are detailed here.

#### **Notes**

Notes about where moisture content is obtained, any assumptions about daily dose rates and other calculations (e.g., moisture conversions, determining amount of individual element from compound), and/or notes about how the PTV calculations are derived (e.g., conversion of mg/m<sup>3</sup> to ppm are based on the ideal gas law, use of fraction of time in intermittent inhalation exposure studies) are described here.

#### **Parameters**

There is one comment and one Ref ID field for each dose rate parameter: body weight, food ingestion rate, water ingestion rate, and inhalation rate. Values, units, and an explanation of each parameter relevant to calculating the PTV (e.g., body weight and food ingestion rate for an oral via food ingestion PTV) are entered in the comment fields. If the appropriate parameters were not provided in the study, the most representative value for each parameter is located (see section A-3.2.3), a short description of what each value represents is provided, and an allometric equation, if applicable, is detailed. The source of the parameter is entered in the Ref ID field that corresponds with this parameter. Otherwise, N/A is the default in the comment field, and 0001 is the default in the Ref ID field.

### A-3.2 PTV Calculation Guidelines

In deriving PTVs, the default is to use the effect levels or critical levels provided by the author(s) of the study. If provided, the information is reported in the author-reported effect levels field of the PTSE Part 1. The use of the author-reported value(s) is based upon the assumption that the authors have accounted for background concentrations of the primary exposure medium and/or concentrations in other exposure media for the chemical of concern (see section A-3.2.1). It is also assumed that the authors took care in measuring food ingestion rates and body weights for the test organisms in their study and applied the appropriate software and/or calculations to interpolate the desired effect level. If the authors did not provide effect levels in mg/kg/d for birds and mammals or mg/kg for invertebrates and plants, adjustments are made before calculating the daily dose rate, if necessary. Adjustments are not made if any of the following occur.

- Primary exposure medium concentration is empirical and in dry weight (background concentration is assumed to be included in the empirical concentration), and additional exposure from other media was not reported.
- PTV calculations are normalized for moisture content of exposure medium, and no background or other media concentrations are reported. For example, if cadmium was administered as a concentration of 30 mg Cd/kg food wet weight, and the food ingestion rate for rats was 0.03 kg food wet weight/d, the units are canceled out (normalized) when determining the amount of chemical ingested per day as follows:

$$30 \frac{\text{mg Cd}}{\text{kg food wet weight}} * 0.03 \frac{\text{kg food wet weight}}{\text{day}} = 0.9 \frac{\text{mg Cd}}{\text{day}}$$

- Primary exposure medium concentration is a nominal concentration, moisture basis is unknown, and background concentration and/or additional exposure from other media was not present or reported. (The moisture basis is assumed to be dry weight in order to produce a conservative PTV. See section A-3.2.2.)
- Primary exposure medium concentration is empirical and the moisture basis is unknown. (The moisture basis is assumed to be dry weight in order to produce a conservative PTV. See section A-3.2.2.)
- Exposure concentration is provided in units of mg/kg for earthworms and plants or mg/kg/d for birds and mammals.

If the reported concentrations do not fill the above criteria, various types of adjustments may be made. They may include

- wet weight to dry weight conversions (for concentrations in the exposure medium and for food ingestion rates for birds and mammals),
- unit conversions,
- additions of verified background concentrations in the exposure medium/diet of the test animals to target (nominal) exposure concentrations,
- additions of background exposure concentrations from a medium other than the primary exposure medium to the primary exposure concentrations, or
- a combination of the above.

### A-3.2.1 Background Concentration Explanation

If it was noted that background concentrations were present, but the exact concentration could not be determined from the data provided in the study without introducing more uncertainty, this is noted in the Part 2 notes field. The PTV is based upon only the concentration of the chemical added to the exposure medium, and it is still more conservative than one based on the supplemental concentration plus the concentration in the basal medium. The basis for this is that in using only the concentration added to the exposure medium, it is assumed the test organisms ingest less chemical and thus, assuming all other parameters (e.g., body weight, food ingestion rate) remain equal, the PTV is lower. If the test organisms had actually ingested a larger amount of chemical because of a background concentration in the exposure medium that was not reported, the lower PTV calculated based on only the supplemental concentration of chemical is still protective of any possible adverse effects that may result from exposure to the larger amount of chemical. Example A-12 illustrates the differences in the PTVs.

#### Example A-12 Background Concentration Calculations

Japanese quail were administered 5000 ppm of manganese via food. Although manganese is often present in the basal diet, the background concentration of the basal diet used in this study is not reported. A PTV is calculated based on just the supplemental concentration of 5000 ppm and a food ingestion rate of 115 g/kg body weight/d for the quail.

$$\text{PTV (mg / kg / d)} = \text{Concentration (mg / kg)} * \text{Food ingestion rate (kg / kg / d)}$$

$$5000 \text{ mg/kg} * 0.115 \text{ kg/kg/d} = 575 \text{ mg/kg/d}$$

If it had been reported that the background concentration of manganese in the basal diet was 56 ppm, this is added to the supplemental concentration of 5000 ppm, and the calculations are carried out as above.

$$5056 \text{ mg/kg} * 0.115 \text{ kg/kg/d} = 581.44 \text{ mg/kg/d}$$

It can be seen in Example A-12 that the PTV for the concentration added to the medium without knowing the background concentration is lower than the supplemental amount plus background concentration. If a background concentration had been assumed to be present, and a concentration was obtained from other sources, it would have provided a higher PTV. The higher PTV may not be protective enough of adverse effects that may occur at concentrations lower than the supplemental concentration plus the background concentration but higher than the supplemental concentration alone. Therefore, it is safe to use just the supplemental amount in PTV calculations if a background concentration is not reported.

### A-3.2.2 Moisture Basis Explanation

If the moisture basis of the concentration in the exposure medium of the food is not reported, it is assumed to be based on dry weight. The reasoning is that if the true moisture basis is indeed wet weight, the PTV calculated based on the assumed dry weight would be lower than if the wet weight concentration of the medium had been converted to dry weight. Example A-13 shows two scenarios: in the first one, moisture basis is unknown and therefore assumed to be dry weight, and in the second, the moisture basis is known to be wet weight.

**Example A-13 Moisture Basis Calculations**

Scenario 1: An experiment reports administering to chicks a concentration of 30 mg/kg of hexavalent chromium via food. The moisture basis of the food is unknown and therefore assumed to be dry weight. The body weight and food ingestion rate of the chicks are 0.0874 kg and 0.0096 kg/d, respectively. The PTV is calculated as follows:

$$\text{PTV (mg / kg / d)} = \frac{30 \text{ mg / kg} * 0.0096 \text{ kg / d}}{0.0874 \text{ kg}} = 3.3 \text{ mg / kg / d}$$

Scenario 2: In the same experiment as above, it is reported that the moisture basis of the concentration is wet weight, and the moisture content of the food is 25%. The wet weight concentration must first be converted to a dry weight concentration before calculating the PTV.

$$30 \frac{\text{mg Cr(VI)}}{\text{kg wet food}} * \frac{1 \text{ kg wet food}}{0.75 \text{ kg dry food}} = 40 \frac{\text{mg Cr(VI)}}{\text{kg dry food}}$$

$$\text{PTV (mg / kg / d)} = \frac{40 \text{ mg / kg} * 0.0096 \text{ kg / d}}{0.0874 \text{ kg}} = 4.4 \text{ mg / kg / d}$$

Scenario 2 in Example A-13 shows that because the dry weight concentration resulting from the conversion of a wet weight concentration to dry weight is always higher, the associated PTV value will be higher as well. Therefore, assuming the concentration is based on dry weight when the moisture basis is unknown, the derived PTV is lower than and protective of the actual PTV that would have been calculated based on wet weight converted to dry weight. In this way, the estimate errs on the conservative side.

**A-3.2.3 Obtaining Dose Rate Parameters for Use in PTV Calculations**

Using dose rate parameters reported in the study leads to a more certain PTV than one that is based on estimated values obtained from another source; reported parameters represent direct measurements of the organisms used in the study and thus give a more accurate dose rate.

If dose rate parameters (i.e., body weight, food or water ingestion rate, and inhalation rate) were not provided in the study, they are obtained from other sources, such as

- Wildlife Exposure Factors Handbook (EPA 1993, 059384, Ref ID 0561) and
- Body Weights of 686 North American Birds (Dunning 1984, 089463, Ref ID 0086).

Often, in cases where dose rate parameters are not provided in the primary toxicity study, the body weight is obtained from another source and then the food or water ingestion rate or inhalation rate is allometrically calculated using equations from the Wildlife Exposure Factors Handbook (EPA 1993, 059384, Ref ID 0561) or Recommendations for and Documentation of Biological Values for Use in Risk Assessment (EPA 1988, 089464, Ref ID 0084). The reverse happens occasionally where the food ingestion rate is provided, and the body weight needs to be allometrically calculated. If the dose rate parameters are not in units of kg body weight, kg food/d, kg water/d, or m<sup>3</sup> air/d, the appropriate conversions are made before using the values in the PTV calculation. See Example A-14.

**Example A-14 Unit Conversions**

For example, converting  $\frac{\mu\text{g chemical}}{\text{mL water}}$  to  $\frac{\text{mg chemical}}{\text{kg water}}$  would be as follows:

$$\frac{\mu\text{g chemical}}{\text{mL water}} * \frac{1000 \text{ mL water}}{1 \text{ L water}} * \frac{1 \text{ mg chemical}}{1000 \mu\text{g chemical}} * \frac{1 \text{ L water}}{1 \text{ kg water}} = \frac{\text{mg chemical}}{\text{kg water}}$$

The following hierarchy for obtaining dose rate parameters is adhered to.

1. Empirical data from the reference being reviewed.
2. Empirical data from *Wildlife Exposure Factors Handbook* (EPA 1993, 059384, Ref ID 0561) or from *Recommendations for and Documentation of Biological Values for Use in Risk Assessment* (EPA 1988, 089464, Ref ID 0084), if available.
3. Empirical data from other references.
4. Allometrically derived values from equations available in the *Wildlife Exposure Factors Handbook* (EPA 1993, 059384, Ref ID 0561) or *Recommendations for and Documentation of Biological Values for Use in Risk Assessment* (EPA 1988, 089464, Ref ID 0084).

**A-3.2.4 PTV Calculation for Oral Ingestion via Food (Birds and Mammals)**

If the body weight was provided or obtained from another source (and converted to kg, if required), the food ingestion rate was provided in kg food/d or similar, and exposure concentrations were provided and converted to mg chemical/kg food, the following equation is used:

$$\text{PTV}_{i,j} = \frac{C_i * F_{Ij}}{\text{BW}_j},$$

Where  $\text{PTV}_{i,j}$  is the primary toxicity value (mg/kg/d) for chemical i in organism j

$C_i$  is the concentration (mg/kg) of chemical i in food

$F_{Ij}$  is the food intake rate (kg food/d) for organism j

$\text{BW}_j$  is the body weight (kg) of organism j

If a body weight was provided and converted to kilograms, and the exposure concentration was provided in terms of mg chemical/organism/d, the following equation is used:

$$\text{PTV}_{i,j} = \frac{C_{ij}}{\text{BW}_j},$$

Where  $\text{PTV}_{i,j}$  is the primary toxicity value (mg/kg/d) for chemical i in organism j

$C_{ij}$  is the concentration (mg/organism/d) of chemical i in food for organism j

$\text{BW}_j$  is the body weight (kg) of organism j

**A-3.2.5 PTV Calculation for Oral Ingestion via Drinking Water (Birds and Mammals)**

If the body weight was provided or obtained from another source (and converted to kg if required), water ingestion rate was provided in L water/d or similar, and exposure concentrations were provided and converted to mg chemical/L water, the following equation is used:

$$PTV_{i,j} = \frac{C_i * W_{ij}}{BW_j},$$

Where  $PTV_{i,j}$  is the primary toxicity value (mg/kg/d) for chemical i in organism j

$C_i$  is the concentration (mg/L) of chemical i in water

$W_{ij}$  is the water intake rate (L water/d) for organism j

$BW_j$  is the body weight (kg) of organism j

If a body weight was provided and converted to kilograms, and the exposure concentration was provided in terms of mg/organism/d, the following equation is used:

$$PTV_{i,j} = \frac{C_{ij}}{BW_j},$$

Where  $PTV_{i,j}$  is the primary toxicity value (mg/kg/d) for chemical i in organism j

$C_{ij}$  is the concentration (mg/organism/d) of chemical i in food for organism j

$BW_j$  is the body weight (kg) of organism j

As explained previously, in the Dose Rate Parameters subsection of section A-2.1.3, Experiment Information, a heavier body weight leads to a more conservative PTV. Assuming the concentration and food ingestion rate remain the same, a heavier body weight leads to a lower PTV, which is more protective of possible effects produced by the exposure concentration to the organism of concern. Likewise, assuming the concentration and body weight remain the same, a lower food or water ingestion rate produces a lower PTV. Therefore, when presented with more than one option for the dose rate parameters, the value that leads to a more conservative PTV is usually chosen in order to be over-conservative rather than under-conservative.

**A-3.2.6 PTV Calculation for Continuous and Intermittent Air Exposure via Inhalation (Mammals)**

A continuous inhalation exposure indicates that the test organism was exposed to air containing chemical vapors for 24 h/d, 7 d/wk, for the duration of the chemical administration period. In an intermittent inhalation exposure study, the organism is exposed to air containing chemical vapors for a set amount of time each day or during a certain number of days per week. Because of the differences in the exposure frequency between continuous and intermittent exposures, and therefore the different amounts of chemical the organisms receive over similar chemical administration periods, the actual amount of time exposed to the chemical over the total length of the study must be determined for intermittent studies to determine the actual dose rate.

For both continuous and intermittent studies, the general equation used to calculate a PTV for continuous or intermittent inhalation exposure is as follows:

$$PTV_{i,j} = \frac{C_i * IR_j}{BW_j} * T_f,$$

Where PTV<sub>i,j</sub> is the primary toxicity value (mg/kg/d) for chemical i in organism j  
 C<sub>i</sub> is the concentration (mg/m<sup>3</sup>) of chemical i in air  
 IR<sub>j</sub> is the inhalation rate (m<sup>3</sup>/d) for organism j  
 BW<sub>j</sub> is the body weight (kg) of organism j  
 T<sub>f</sub> is the fraction of time organism j was exposed

Two parameters in this equation must be converted to the units necessary to derive the PTV before the PTV is calculated. The first is the concentration; it often needs to be converted from ppm to mg/m<sup>3</sup>. The second parameter is the inhalation rate; if it is not provided in the paper, it is obtained from another source or calculated using an allometric equation, usually from EPA (1993, 059384, Ref ID 0561; 1988, 089464, Ref ID 0084), and the body weight, whether it is one reported from the study or obtained from another source. One additional parameter needs to be determined for intermittent studies: the fraction of time. In continuous studies, the fraction of time equals 1.

### Converting Concentration from ppm to mg/m<sup>3</sup>

The conversion of a concentration in ppm to mg/m<sup>3</sup> is conveyed by the following equation:

$$\text{Conc (mg / m}^3\text{)} = \text{ppm(v)} * \frac{\text{MW}}{24.45}$$

Where Conc (mg/m<sup>3</sup>) is the concentration of the chemical in mg/m<sup>3</sup>  
 ppm(v) is the concentration of the chemical administered in the study, by volume  
 MW is the molecular weight of the chemical in grams  
 24.45 is the constant molar volume at standard temperature and pressure

The gram molecular weight for the chemical of concern is obtained from the ChemBioFinder.Com website (<http://chemfinder.cambridgesoft.com>) or any other appropriate source containing chemical property information. The value in grams is then multiplied by 1000 to achieve the amount in milligrams, and this value is then used with the units of mg/m<sup>3</sup> in the PTV calculation along with an inhalation rate either provided in the study or obtained from another source. Often, the inhalation rate is calculated using an allometric equation from EPA (1993, 059384, Ref ID 0561) and a body weight that was provided in the study or obtained elsewhere.

**Determining the Inhalation Rate**

Unless already provided in the paper, the inhalation rate for a mammal is obtained from another source if the information supporting it closely matches the information for the test organism of concern (e.g., similar organism type, body weight of organism, and age/life stage of organism). Otherwise, the inhalation rate is usually derived using allometric equations from EPA (1993, 059384, Ref ID 0561, which cites Stahl 1967, 063119, Ref ID 1522), dependent on whether the body weight is presented in grams or kilograms:

$$IR = 0.002173(BW^{0.80}),$$

Where IR is the inhalation rate in m<sup>3</sup>/d  
BW is body weight in grams

OR

$$IR = 0.5458(BW^{0.80}),$$

Where IR is the inhalation rate in m<sup>3</sup>/d  
BW is body weight in kilograms

**Determining the Fraction of Time for One-Phase Intermittent Inhalation Exposure Scenarios**

After the concentrations are converted from units of ppm to mg/m<sup>3</sup>, the actual exposure period is determined as a percentage of the chemical administration period and used as the fraction of time the test organisms are exposed to vapors. Often, in intermittent inhalation toxicity studies, the chemical administration regimen is presented as a rate of number of hours per day and number of days per week. To determine the fraction of time, these numbers must be converted into one total number, in days, to represent the total amount of time the test organisms were actually exposed to the chemical in air. This total number represents the actual exposure period and is divided by the chemical administration period, which should also be converted to days. The following equation is used:

$$T_f = \frac{H * D * W / 24}{P_d},$$

Where  $T_f$  is the fraction of time (unitless)  
H is the number of hours per day  
D is the number of days per week  
W is the number of weeks in the chemical administration period  
 $P_d$  is the chemical administration period, in days

See Example A-15.

**Example A-15 PTV Calculation for a One-Phase Intermittent Inhalation Exposure**

In Goldberg et al. (1964, 089460, Ref ID 1348), rats were exposed to 300 ppm trichloroethene at a rate of 4 h/d, 5 d/wk, for 5 wk.

Step 1: Converting ppm to mg/m<sup>3</sup>:

$$\text{mg/m}^3 = 300 \text{ ppm} * \frac{131.3824}{24.45} = 1600$$

Step 2: Determining the fraction of time:

$$T_f = \frac{(4 \text{ h/d} * 5 \text{ d/wk} * 5 \text{ wk})/24 \text{ h/d}}{35 \text{ d}} = 0.1190 .$$

Step 3: Determining daily inhalation rate of test organism:

The higher end of the body weight range of the rats at the beginning of the study (450 g) was used in an allometric equation for all mammals (EPA 1993, 059384, Ref ID 0561) to determine the daily inhalation rate for rats (0.29 m<sup>3</sup>/d).

$$\text{IR} = 0.002173(\text{Wt}^{0.80}) = 0.002173(450^{0.80}) = 0.29 \text{ m}^3/\text{d}.$$

Step 4: Calculating the PTV:

$$\text{PTV} = \frac{1600 \text{ mg/m}^3 * 0.29 \text{ m}^3 / \text{d}}{0.45 \text{ kg}} * 0.1190 = 122.7 \text{ mg/kg/d}$$

The PTV is rounded to 120 mg/kg/d.

**Determining the Fraction of Time for Two-Phase Intermittent Exposure Scenarios**

In studies where the same group of organisms is exposed to the same exposure concentration of the same chemical in two different exposure regimens (e.g., 4 h/d, 5 d/wk for the first 2 wk, and then 6 h/d, 7 d/wk in the last 5 wk), the actual exposure period for each exposure scenario is determined separately, and then the exposure periods are added together before determining the fraction of the chemical administration period they represent. See Example A-16.

**Example A-16 PTV Calculation for a Two-Phase Intermittent Inhalation Exposure in which the Exposure Frequency is Different from One Phase to the Next**

In York et al. (1982, 089462, Ref ID 1359), female rats were exposed to 2100 ppm 1,1,1-trichloroethane at a rate of 6 h/d, 5 d/wk during the first 2 wk (including pre-mating and mating periods), and then for 6 h/d, 7 d/wk from day 1 to 20 of gestation.

Step 1: Converting ppm to mg/m<sup>3</sup>:

$$\text{mg/m}^3 = 2100 \text{ ppm} * \frac{133.4033}{24.45} \text{ g} = 11500$$

Step 2: Determining the fraction of time:

$$T_f = \frac{((6 \text{ h/d} * 5 \text{ d/wk} * 2 \text{ wk}) + (6 \text{ h/d} * 7 \text{ d/wk} * 20 \text{ d/7 d/wk}))/24 \text{ h/d}}{34 \text{ d}} = 0.2206$$

Step 3: Determining daily inhalation rate of test organism:

The average body weight range of the control rats and rats in the treatment group before exposure was 252.6 g. This body weight is used in an allometric equation to derive an inhalation rate.

$$\text{IR} = 0.002173(\text{Wt}^{0.80}) = 0.002173(252.6^{0.80}) = 0.18 \text{ m}^3/\text{d}.$$

Step 4: Calculating the PTV:

$$\text{PTV} = \frac{11500 \text{ mg/m}^3 * 0.18 \text{ m}^3/\text{d}}{0.2526 \text{ kg}} * 0.2206 = 8194.77 \text{ mg/kg/d}$$

The PTV is rounded to 8200 mg/kg/d.

In studies where the same group of organisms is exposed to two different exposure concentrations under the same exposure conditions (e.g., inhalation of 2000 ppm for the first week and then 500 ppm in the remaining 3 wk), the steps are as follows:

1. The actual exposure period, in days, for each concentration is determined separately.
2. Each concentration of the chemical is converted from ppm to mg/m<sup>3</sup>, if needed.
3. Each concentration of the chemical in mg/m<sup>3</sup> is multiplied by the daily inhalation rate (obtained from reference or calculated allometrically using body weight) and the actual exposure period associated with that concentration to determine the amount of chemical received by the test organism from each exposure concentration.
4. The amounts of chemical from each exposure concentration are added to determine the total amount of chemical received by the test organism throughout the entire chemical administration period.
5. The PTV is calculated by dividing this total amount of chemical by body weight in kilograms and by the total number of days in the chemical administration period.

See Example A-17 for a two-phase intermittent study in which concentrations differ from one phase to the next.

**Example A-17 PTV Calculation for a Two-Phase Intermittent Inhalation Exposure in which the Exposure Concentration is Different from One Phase to the Next**

In Quast et al. (1986, 109942, Ref ID 1360), male and female rats were exposed to 35.8 ppm 1,1-dichloroethene at a rate of 6 h/d, 5 d/wk, during the first 6 wk, then to 72.6 ppm at the same rate for the remaining 66 wk of the 72-wk exposure period.

Step 1a: Determining actual exposure period (in days) for the 35.8-ppm dose regimen:

$$Pd = \frac{6 \text{ h/d} * 5 \text{ d/wk} * 6 \text{ wk}}{24 \text{ h/d}} = 7.5 \text{ d}$$

Step 1b: Determining actual exposure period (in days) for the 72.6-ppm dose regimen:

$$Pd = \frac{6 \text{ h/d} * 5 \text{ d/wk} * 66 \text{ wk}}{24 \text{ h/d}} = 82.5 \text{ d}$$

Step 2a: Converting ppm to mg/m<sup>3</sup> for the 35.8-ppm dose regimen:

$$\text{mg/m}^3 = 35.8 \text{ ppm} * \frac{96.9427 \text{ g}}{24.45} = 140$$

Step 2b: Converting ppm to mg/m<sup>3</sup> for the 72.6-ppm dose regimen:

$$\text{mg/m}^3 = 72.6 \text{ ppm} * \frac{96.9427 \text{ g}}{24.45} = 290$$

Step 3: Determining daily inhalation rate of test organism:

The average body weight range of 10 male control rats throughout 24 mo of the study was 542.2 g. This average body weight is used in an allometric equation to derive an inhalation rate.

$$IR = 0.002173(Wt^{0.80}) = 0.002173(542.2^{0.80}) = 0.33 \text{ m}^3/\text{d}.$$

Step 4a: Determining the amount of chemical received by the rats during the first 6 wk (35.8-ppm dose regimen) using the concentration in mg/m<sup>3</sup>, daily inhalation rate, and actual exposure period.

$$140 \text{ mg/m}^3 * 0.33 \text{ m}^3/\text{d} * 7.5 \text{ d} = 346.5 \text{ mg}$$

Step 4b: Determining the amount of chemical received by the rats during the last 66 wk (72.6 ppm dose regimen) using the concentration in mg/m<sup>3</sup>, daily inhalation rate, and actual exposure period.

Step 5: Calculating the total amount of chemical received by the rats during the entire exposure period:

$$346.5 \text{ mg} + 7895 \text{ mg} = 8242 \text{ mg}$$

Step 6: Calculating the PTV by dividing the total amount of chemical by body weight (0.5422 kg) and by the total number of days in the chemical administration period (72 wk, or 504 d).

$$PTV = 8242 \text{ mg} / 0.5422 \text{ kg} / 504 \text{ d} = 30 \text{ mg/kg/d}.$$

### A-3.2.7 Significant Digits and Rounding Procedure

The rules for significant digits in computations are generally followed in the PTV calculations. In multiplication and division, the product or quotient contains as many significant digits as the number in the

operation with the least number of significant digits. In addition and subtraction, the sum or difference is no more precise than the least precise number involved in the operation. When it comes to rounding off nonessential digits, if the last reported digit was followed by a number less than 5, the reported digit is kept as is. If it was followed by a number greater than 5, it is rounded up. Finally, if the last reported digit was followed by a 5, and that 5 is in turn followed by no other digits or zeroes, then the last reported digit is kept as is. On the other hand, if the 5 is followed by an odd number, the reported digit is rounded up one, and if the 5 is followed by an even number, the reported digit is left as is. Sometimes, significant digit rules are difficult to apply because although numbers are reported, they are often not reported in scientific format. It is difficult to tell whether a zero is significant or not in a number such as 2500. In such situations where the use of significant digits becomes vague, best professional judgment is used. The number is often rounded to a minimum of two significant digits. For example, 1247 is rounded to 1200 and 1.464 is rounded to 1.5.

In inhalation exposure studies, when the concentration in ppm is used to calculate  $V_{\text{analyte}}$ , all numbers resulting in the  $V_{\text{analyte}}$  value are then used in the conversion of ppm to  $\text{mg}/\text{m}^3$  (e.g., 3800 ppm leads to 3.8 L, which is used in calculation of mg). Furthermore, when rounding grams to milligrams, two integers are used (e.g., 15.37 to 15 or 1.611 to 1.6) so that the  $\text{mg}/\text{m}^3$  value then has two foremost numbers followed by zeroes (e.g., 1600 or 15000). Two decimal places are used for the inhalation rate (e.g.,  $0.29 \text{ m}^3/\text{d}$ ). Four decimal places are usually used in the formula weights (e.g., 131.3842 g/mol) and the fraction of time (e.g., 0.2917). The PTV is then rounded to two significant digits (e.g., 122.7 to 120).

The general guideline is to be consistent in the application of significant digit rules where possible, followed by consistent rounding procedures. After the rules for significant digits and rounding procedures are applied, the number that is entered into the PTV field is automatically rendered to scientific notation with two decimal points. This does not denote three significant digits but is rather a truncated way of reporting the values.

### **A-3.2.8 PTV Confidence Rating Guidelines**

The abundance or lack of information provided by the study associated with a PTV is reflected in the scoring of Part 2, and these scores are then weighted according to the ability of each criterion to influence the magnitude of the TRV and the uncertainty associated with it. The following is a list of multipliers and the situations in which they are applied.

- 1 There is little to no influence on the TRV. Most studies have already been eliminated based on nonfulfillment of these fields (e.g., a bird study is not going to be used for a mammal study).
- 2 There is more influence on the TRV as to deciding whether or not to keep the PTV in the TRV data set, but little influence on the actual TRV.
- 3 There is a medium influence on the TRV. This weighting scheme can also be used for criteria in which TRVs are defined (e.g., oral in diet or drinking water) or it can be used for those areas where if data are not provided, other means by the reviewer can be employed (e.g., statistics).
- 4 There is a medium-high influence on the TRV. If the original score is low, this leads to more uncertainty. This weighting scheme is also used for those criteria defining TRVs (e.g., reproduction/development, chronic, NOAEL or NOEC).
- 5 There is a high influence on the TRV where a low original score leads to the most uncertainty and greatest difference in TRVs compared to those criteria derived from extra detail provided in the study (e.g., chronic vs. acute).

Table A-10 illustrates each criterion, its multiplier, and the justification for use of that multiplier.

**Table A-10**  
**Weighting Schemes for Criteria in Part 2 of the Data-Entry Database**

Field that is Scored	Multiplier	Justification
<b>Study Design and Documentation Score</b>		
Control group included	3	While controls are needed for a stronger assessment of effect levels, unbounded NOAELs/NOECs or LOAELs/LOECs (i.e., NOAELs/NOECs without accompanying LOAELs/LOECs or vice versa) can also be derived. Therefore, the magnitude of the influence on the TRV is medium; that is, the TRV is not solely reliant on controls being available.
Multiple exposure groups	3	While multiple exposure groups are needed for a stronger assessment of effect levels, unbounded NOAELs/NOECs and LOAELs/LOECs can also be derived. Therefore, the magnitude of the influence on the TRV is medium; that is, the TRV is not solely reliant on there being more than one exposure group.
Test organism details	1	There is little influence of test organism details on the TRV. The details help to gauge the rigorousness of the study.
Dose rate parameters	4	Those parameters that are specifically related to the organism and study at hand are best suited for deriving the PTV. Parameters can also be obtained elsewhere but their use increases uncertainty, although the difference in the TRV vs. a TRV that would be derived from the use of study-specific dose rate parameters is small.
Exposure dose concentration	3	Measured concentrations in dry weight are preferred. However, if the information is not reported, nominal concentrations based on dry weight are assumed and can result in overly conservative TRVs. Also, uncertainty may be introduced if the moisture basis is in wet weight and conversion to dry weight is needed. If the moisture basis is not reported in the study, a surrogate value must be used. The TRV is not solely reliant on moisture basis; therefore, a medium degree of influence is given.
Statistics	3	Statistics provided in the study are preferred and lead to determination of dose-response trends and assignment of effect levels. However, if not provided, data may be analyzed by the reviewer. The influence on the TRV receives medium weight because of this and because if no statistics or data are provided, the assignment of an effect level is more difficult.
<b>Test Organism Score</b>		
Taxonomic relationship of test organism	2	Less weight is afforded for the taxonomic relationship of test organisms because studies that are not related to a screening receptor by at least the class level are not evaluated. However, more certainty results when the test organism is more closely related to screening receptor.
Basis for use of test organism	1	There is little influence of the authors' basis for the test organism on the TRV. This detail helps in consideration if the study is more attuned to the test organism itself rather than as a model for human exposure or other types of organisms.

Table A-10 (continued)

Field that is Scored	Multiplier	Justification
<b>Exposure Conditions Score</b>		
Test environment	1	There is little influence of the test environment on the TRV because only those studies with appropriate experimental conditions are evaluated in the PTSE. This detail helps gauge the degree of control in a study (laboratory vs. field). Uncontrolled studies are usually eliminated up front.
Test exposure medium similar to food	3	There is little influence of the test exposure medium similar to food on the value of the TRV because the exposure medium in the studies selected for oral exposures is bound to be similar or related to one of the exposure media present here. However, the test exposure medium is one of the more critical parameters evaluated in the study with respect to determining ecological relevance of the experimental conditions.
Test exposure medium similar to drinking water	3	There is little influence of the test exposure medium similar to drinking water on the value of the TRV because the exposure medium in the studies selected for oral exposures is bound to be similar or related to one of the exposure media present here. However, the test exposure medium is one of the more critical parameters evaluated in the study with respect to determining ecological relevance of the experimental conditions.
Test exposure medium similar to soil	3	There is little influence of the test exposure medium similar to soil on the value of the TRV because the exposure medium in the studies selected for oral uptake and dermal exposures or root and/or seed coat uptake is bound to be similar or related to one of the exposure media present here. However, the test exposure medium is one of the more critical parameters evaluated in the study with respect to determining ecological relevance of the experimental conditions.
Chemical interactions	2	Chemical interactions do not influence the value of the TRV much because any study that has chemical interaction is automatically eliminated from the data set before Part 1 is started. If other influences are present, they are likely to be of natural conditions.
Test exposure route	3	There is little influence of the test exposure route on the value of the TRV. However, the test exposure medium is one of the more critical parameters evaluated in the study with respect to determining ecological relevance of the experimental conditions.
Test period and chemical administration period	5	The influence of the test and chemical administration periods on the TRV is high because the assignment of chronic vs. subchronic vs. acute leads to application of UFs, which are the leading factor in TRV differences.
Critical life stage	4	The influence of the critical life stage on the TRV is high because the assignment of chronic to subchronic or acute studies leads to elimination of the use of UFs, which are the leading factor in TRV differences.
Test exposure frequency	2	The value of the TRV is influenced slightly by accounting for actual exposure time in the daily dose rate in intermittent exposure regimens.
<b>Measurement(s) and Result(s)</b>		
Focus measurement category	4	The focus measurement category may not influence TRVs as much because studies with "other" endpoints are eliminated before TRV consideration. However, the type of endpoint is a strong consideration with reproduction/development being the preferred endpoint, followed by survival, and then growth. High weight is given because a wider spread of the score results in clearer distinction between these endpoints.
Measurement length	1	The TRV is influenced slightly by the consideration of whether or not the measurement actually reflects the entire exposure period.

**Table A-10 (continued)**

Field that is Scored	Multiplier	Justification
Effect level category	5	Effect level category receives the highest weight because assignment of NOAEL/NOEC vs. LOAEL/LOEC vs. other effect level leads to the application of UFs, which are the leading factor in TRV differences.

The percent maximum score is achieved by dividing the weighted score of the study by the maximum weighted score possible for the type of study (bird or mammal oral ingestion study, mammal inhalation study, or earthworm or plant study). Bird and mammal oral ingestion studies will have a higher maximum score because the test exposure medium similar to food or drinking water category is not scored in mammal inhalation studies, whereas only the test exposure medium similar to soil is used in plant and invertebrate studies. The percent maximum score determines whether the PTV is assigned a low, medium, or high confidence according to Table A-11.

**Table A-11**  
**Percent Maximum Scores and Confidence Ratings**

Confidence Rating	Percent of Maximum Total Weighted Score (%MTWS)*
High	%MTWS $\geq$ 76%
Medium	51% $\leq$ %MTWS<76%
Low	26% $\leq$ %MTWS<51%
Unacceptable	%MTWS<26%

\* Percent of maximum total weighted score (%MTWS) = (total score/maximum weighted score for appropriate receptor)\*100.

#### **A-4.0 PTSE PART 3, TOXICITY REFERENCE VALUE DEVELOPMENT**

A PTSE Part 3 is used to develop a TRV following the completion of the PTSE Part 1 and Part 2 for all references in the data set for a particular screening receptor group (i.e., bird, invertebrate, mammal, plant), chemical, and exposure route scenario of concern. Either a GMM or CS TRV can be developed; a GMM TRV is preferred. The determination of which TRV is developed is dependent on the characteristics of the data set under consideration. Furthermore, if a GMM TRV is developed but not deemed to be appropriate for protection of ecologically relevant endpoints in the data set or of sensitive species, a subset GMM TRV can be calculated where a portion of the original GMM TRV is used to calculate a new GMM TRV. If a subset GMM TRV cannot be calculated or is still not considered protective enough, a LANL CS TRV is developed. However, the GMM TRV and subset GMM TRVs that were calculated but not used in ESL models (or were replaced with a more preferred TRV in ESL models) are still kept on record in the Ecorisk Database to allow risk assessors, risk managers, and regulators to assess for themselves the appropriateness of the values, if needed. Furthermore, keeping these unused values in the database also tracks the history of TRV development and why these values were replaced or not used. Details for the Part 3 process for GMM and CS TRVs are presented below, starting with section A-4.1.

#### **A-4.1 Creation of the GMM TRV Data Set**

A geometric mean is used instead of an arithmetic mean because it better represents the central tendency of toxicological data sets that tend to be skewed. Selecting the geometric mean as a representative effect level limits the influence of valid data points that are far removed from the general cluster of data points. The ideal GMM TRV for screening-level ecological risk assessments is one that is based on a data set representing the most ecologically relevant endpoints (i.e., reproduction/development), exposure routes (i.e., oral ingestion via food or drinking water in birds or mammals, inhalation in mammals, uptake via seed coat and/or roots in plants, or oral and dermal contact in invertebrates), exposure media (i.e., food or drinking water in birds and mammals, air for mammals, or soil for plants and invertebrates), exposure period (chronic), and effect levels (NOAEL for birds and mammals or NOEC for plants and invertebrates). A GMM TRV based on these characteristics is protective of wildlife, plant, or invertebrate populations because it represents a central tendency of the no adverse effect levels for ecologically relevant effects (i.e., adverse effects on ability of individuals to develop into viable organisms, search for mates, breed successfully, and produce live and equally viable offspring).

The data set for the GMM TRV is developed by including only ecologically relevant records for the receptor group, chemical, and exposure route scenario of concern (e.g., Aroclor-1260 in mammals for food ingestion). PTVs derived from PTSE Part 2 are included in the data set only if they are associated with exposure conditions similar to that of the exposure environment of concern. To create this data set of ecologically relevant PTVs, the PTVs must be evaluated against a set of exclusion criteria, and if they meet any of the criteria, they are excluded from the data set. The three categories of exclusion criteria are (1) exposure conditions, (2) measured endpoints, and (3) repetitive values. All are described below. After the exclusion criteria have been applied and the final GMM TRV data set has been created, there must be three or more PTVs available for a GMM TRV to be developed. If less than three PTVs exist, a CS TRV is developed instead (see section A-4.2). Before the calculation of the GMM TRV, the PTVs are extrapolated to chronic NOAEL- or NOEC-based effect levels. The GMM TRV and its data set are then graphed, and details are documented in the PTSE Part 3 data-entry database for later incorporation into the most current version of the Ecorisk Database.

##### **A-4.1.1 Exclusion Criteria for Study Exposure Conditions**

The PTVs included in the GMM TRV data set for the receptor group, chemical, and exposure route scenario of concern (e.g., Aroclor-1260 in mammals for food ingestion) are those associated with ecologically relevant studies (experiments). An ecologically relevant study is a study that uses exposure conditions and measured endpoints that are considered to be predictive of population level effects in a real world ecosystem. Table A-12 lists the exclusion criteria for exposure conditions used in a study. First, each study is evaluated against the exposure conditions exclusion criteria, and if one of the exclusion criteria is met, any PTVs associated with this study are excluded from the GMM TRV data set. If the exclusion criteria for exposure conditions are not met, then the endpoints measured in the study are evaluated against the measured endpoint exclusion criteria described in the next section.

**Table A-12**  
**Exclusion Criteria for Exposure Conditions Used in a Study**

Organism Group	TRV Type	Exposure Condition	Exclusion Criteria
Bird or mammal	Food	Exposure medium	Drinking water
			Aqueous solution
			Unknown
		Exposure route	Injections
	Unknown		
	Drinking water	Exposure medium	Food
			Peanut oil
			Corn oil
Other types of oil or oil mixtures			
		Exposure route	Injections
Invertebrate	Soil	Exposure medium	Manure
			Soil and manure
			Unknown
		Exposure route	Filter paper
		Soil property	OM greater than 10% or not reported
Plant	Soil	Exposure medium	Nutrient or aqueous solution
		Exposure route	Filter paper
		Soil property	OM greater than 10% or not reported

#### A-4.1.2 Exclusion Criteria for Endpoints Measured in a Study

For all organism groups, the endpoints excluded are those that do not fall into the reproduction/development, survival, adult weight change, or adult size change categories. Examples of these endpoints are

- tumors,
- histopathology,
- nonreproductive organ toxicity,
- biochemistry,
- hematology,
- serum chemistry, and
- nonreproductive behavior.

If one of the measured endpoint exclusion criteria is met, the PTV associated with the measured endpoint is excluded from the GMM TRV data set. If the exclusion criteria for measured endpoints are not met, then the measured endpoints for each study are evaluated against the repetitive values exclusion criteria described in the next section.

### A-4.1.3 Exclusion Criteria for Repetitive Values

An exclusion procedure is performed to remove repetitive endpoints within a study, which entails making sure that there is only one PTV per ecologically relevant endpoint category (reproduction/development, survival, and adult weight or size changes) per study. Best professional judgment is used to select the most ecologically relevant and/or sensitive PTV per ecologically relevant endpoint category per study. For example, if one experiment had three reproduction/development endpoints, one survival endpoint, and one adult weight change endpoint, the most ecologically relevant and/or sensitive reproduction/development endpoint of the three available would be included in the GMM TRV data set along with the single survival and single weight change endpoints. This exclusion process minimizes the possibility of a GMM TRV being skewed to the results of any particular study as a result of repetitive values for the same endpoint category within a study. Those PTVs whose measured endpoints do not meet the repetitive values exclusion criteria are included in the GMM TRV data set.

### A-4.1.4 Deriving Chronic NOAEL- or NOEC-Based Effect Levels

After the exclusion criteria have been applied, the GMM TRV data set now contains a variety of original effect levels (PTVs) derived from the PTSE process ranging from chronic NOAEL/NOEC or LOAEL/LOEC pairs to acute, other effect levels such as LC<sub>50</sub>s or EC<sub>20</sub>s. Effect levels other than chronic NOAELs/NOECs must first be extrapolated to chronic NOAEL- or NOEC-based effect levels before the calculation of the GMM TRV can take place. If the PTV is an acute or subchronic NOAEL/NOEC, it is extrapolated to a chronic NOAEL- or NOEC-based effect level with the application of a UF. If the PTV is a LOAEL/LOEC or other effect level (LC<sub>50</sub>), it is first extrapolated to a NOAEL with the application of a UF, and then it is extrapolated to chronic exposure duration if needed. See Table A-13 for a description of UFs.

**Table A-13**  
**Uncertainty Factors Applied to Derive**  
**Chronic NOAEL- or NOEC-Based Effect Levels**

Type of Effect Level Available	UF Applied to Derive a TRV that is a Chronic NOAEL- or NOEC-Based Effect Level
C-CL or chronic NOAEL/NOEC	1
C-CL or chronic LOAEL/LOEC	10
C-CL or chronic LD <sub>50</sub> /LC <sub>50</sub>	100
C-CL or chronic ED <sub>50</sub> /EC <sub>50</sub>	100
Subchronic NOAEL/NOEC	10
Subchronic LOAEL/LOEC	100
Subchronic LD <sub>50</sub> /LC <sub>50</sub>	100
Subchronic ED <sub>50</sub> /EC <sub>50</sub>	100
Acute or single-dose NOAEL/NOEC	100
Acute or single-dose LOAEL/LOEC	100
Acute or single-dose LD <sub>50</sub> /LC <sub>50</sub>	100
Acute or single-dose ED <sub>50</sub> /EC <sub>50</sub>	100

#### A-4.1.5 Deriving Chronic LOAEL- or LOEC-Based Effect Levels

If a chronic LOAEL/LOEC effect level does not already exist for an endpoint from a particular study, a LOAEL- or LOEC-based effect level is approximated from an effect level (NOAEL, NOEC, LC<sub>xx</sub>, LD<sub>xx</sub>, EC<sub>xx</sub>, or ED<sub>xx</sub>). If the effect level is an acute or subchronic LOAEL/LOEC, a UF of 100 or 10 is applied to extrapolate to a chronic LOAEL/LOEC. On the other hand, if the effect level is a chronic NOAEL/NOEC or chronic NOAEL- or NOEC-based effect level extrapolated from an acute or subchronic NOAEL/NOEC, a test organism-specific LOAEL/LOEC or NOAEL/NOEC factor must be applied to derive a LOAEL- or LOEC-based effect level. Based on Dourson and Stara (1983, 073474, Ref ID 1379), 96% of the ratios between NOAELs and LOAELs for mammals in oral ingestion experiments have values of 5 or less (Dourson and Stara [1983, 073474, Ref ID 1379, p. 232 and Figure 4]). However, because these data are only applicable to oral ingestion exposure in mammals, ratios for the remaining exposure pathways (oral ingestion in birds, oral ingestion and dermal contact in earthworms, uptake via seed coats and/or roots in plants, and inhalation in mammals) were determined from NOAEL/NOEC or LOAEL/LOEC pairs specific to each of the exposure pathways. The data used to develop the ratios are from the Ecorisk Database. The smallest and largest ratios developed for each exposure pathway were used to approximate a minimum and maximum LOAEL- or LOEC-based effect level to bracket a range of concentrations at which the adverse effects may first be observed. Figure A-1 offers a step-by-step process for determining how to derive the LOAEL- or LOEC-based effect levels.

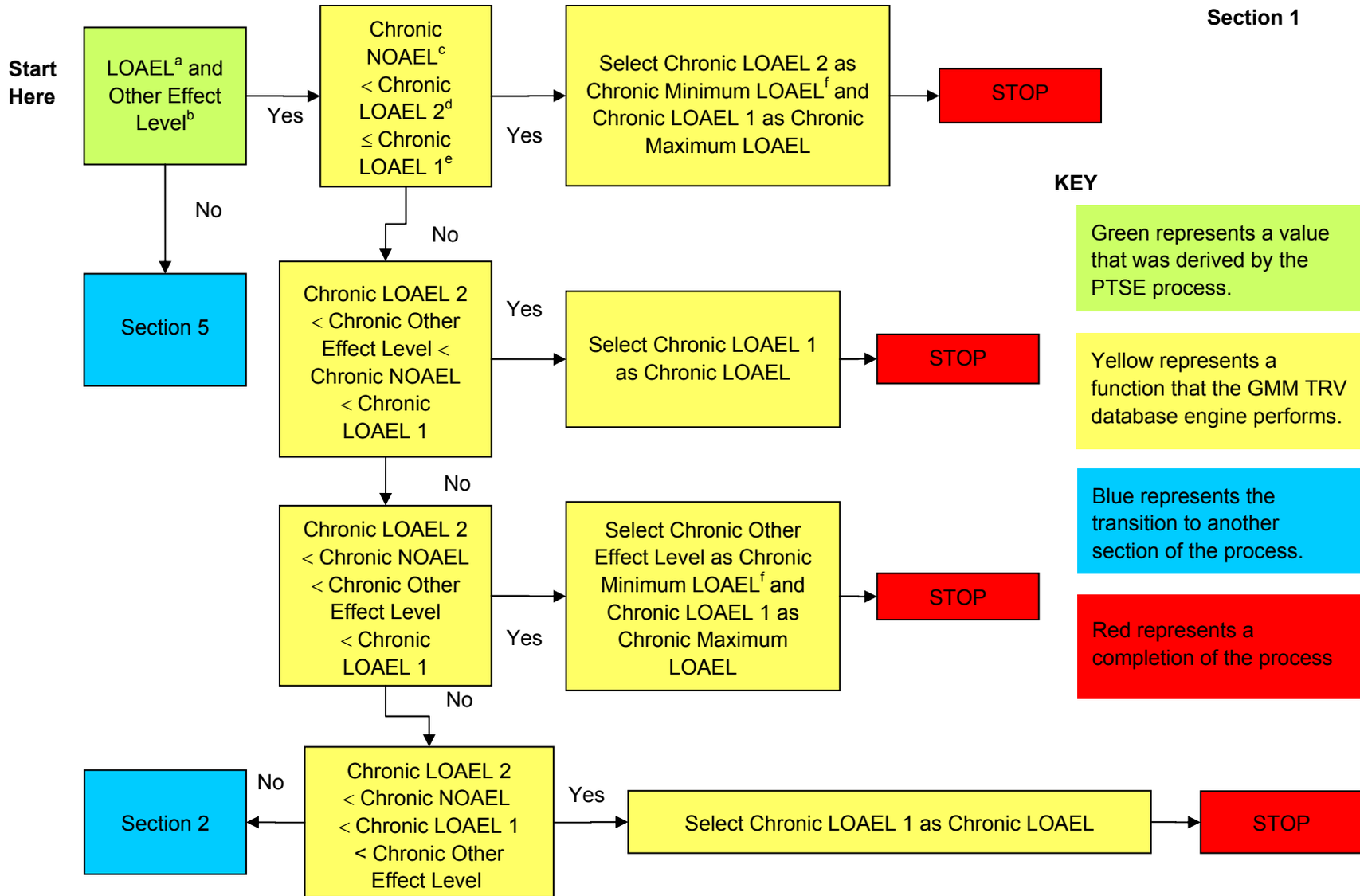


Figure A-1 Process for selecting the chronic LOAEL- or LOEC-based effect level for each endpoint in the GMM TRV data set

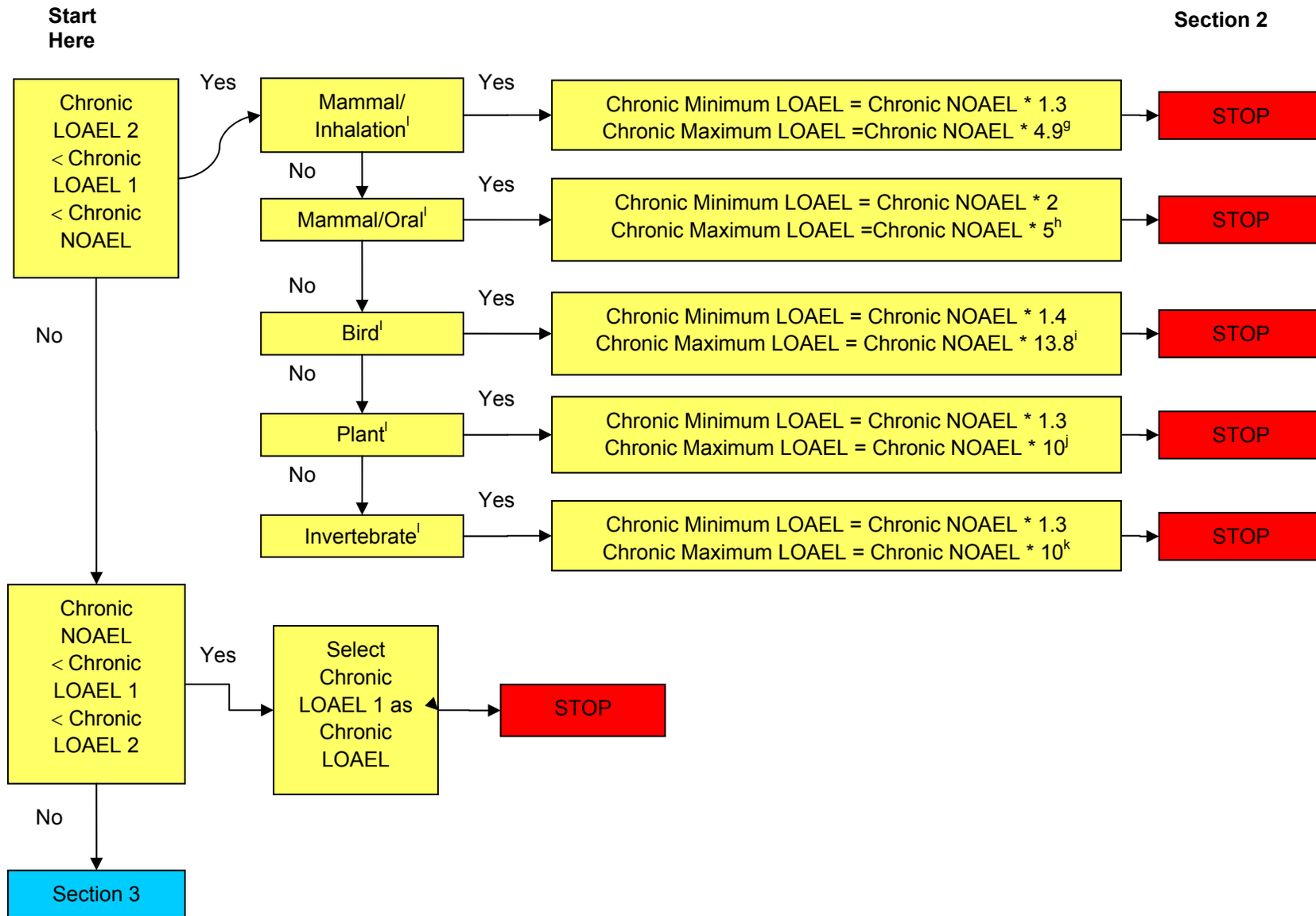
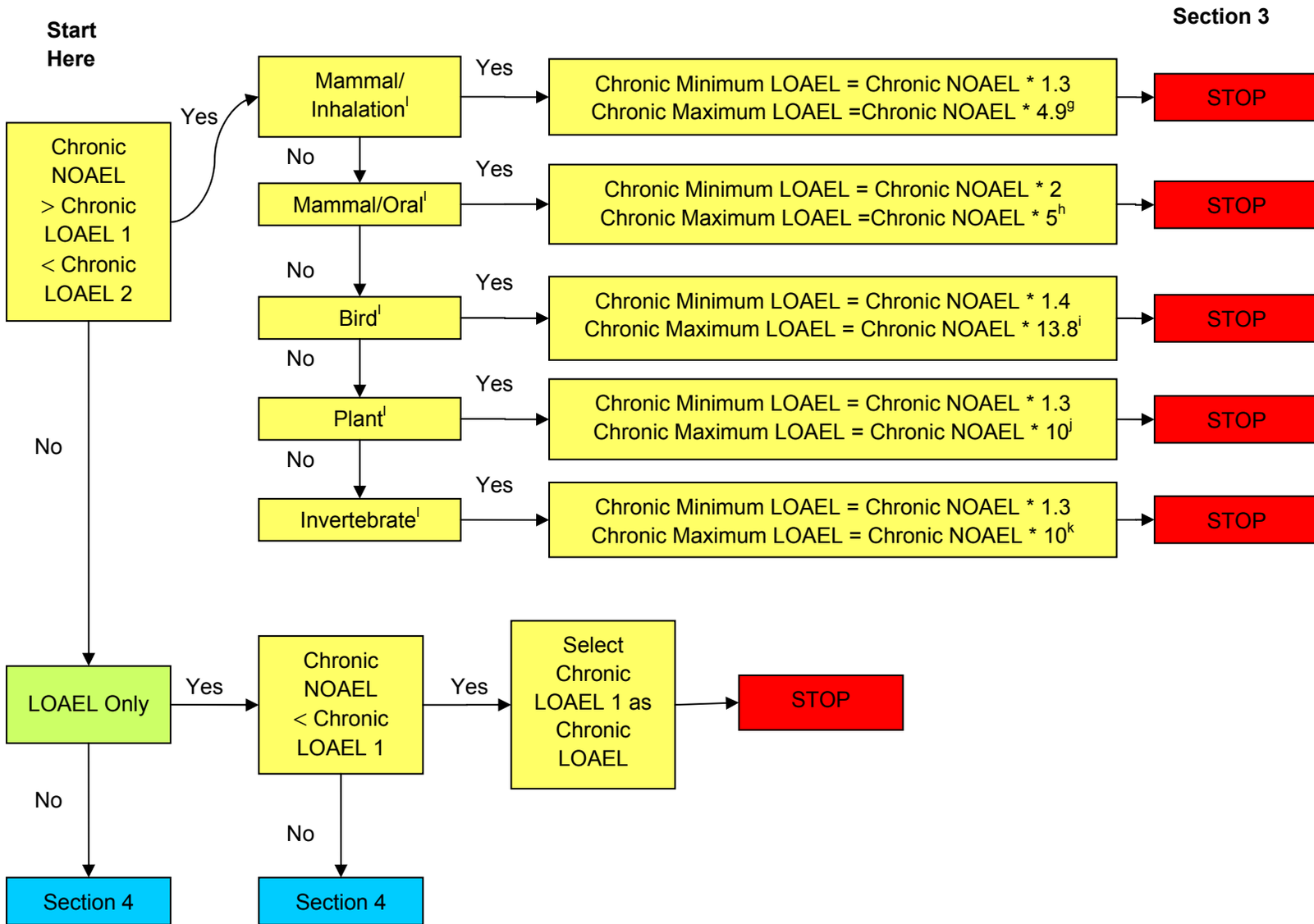


Figure A-1 (continued) Process for selecting the chronic LOAEL- or LOEC-based effect level for each endpoint in the GMM TRV data set



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Figure A-1 (continued) Process for selecting the chronic LOAEL- or LOEC-based effect level for each endpoint in the GMM TRV data set

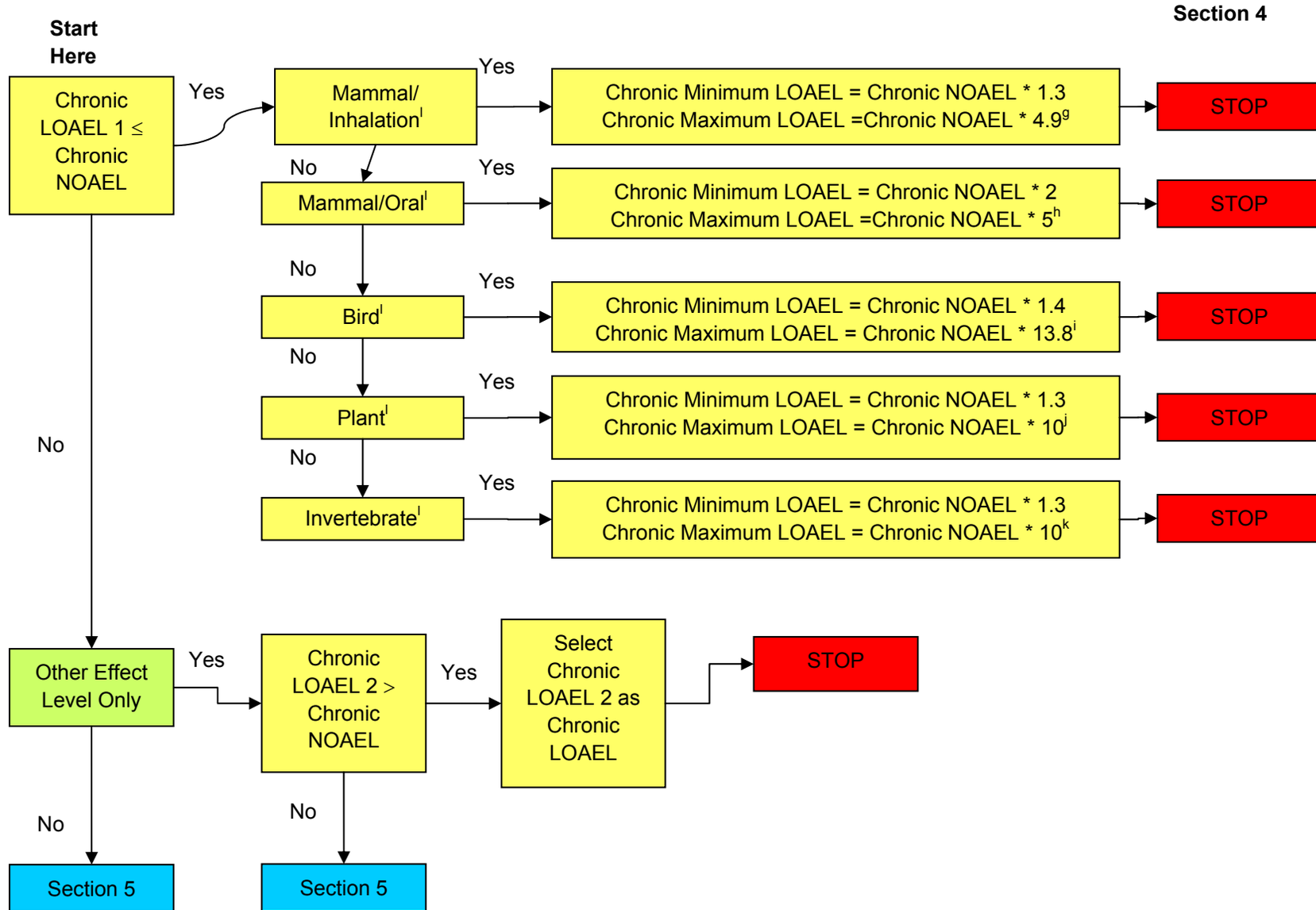
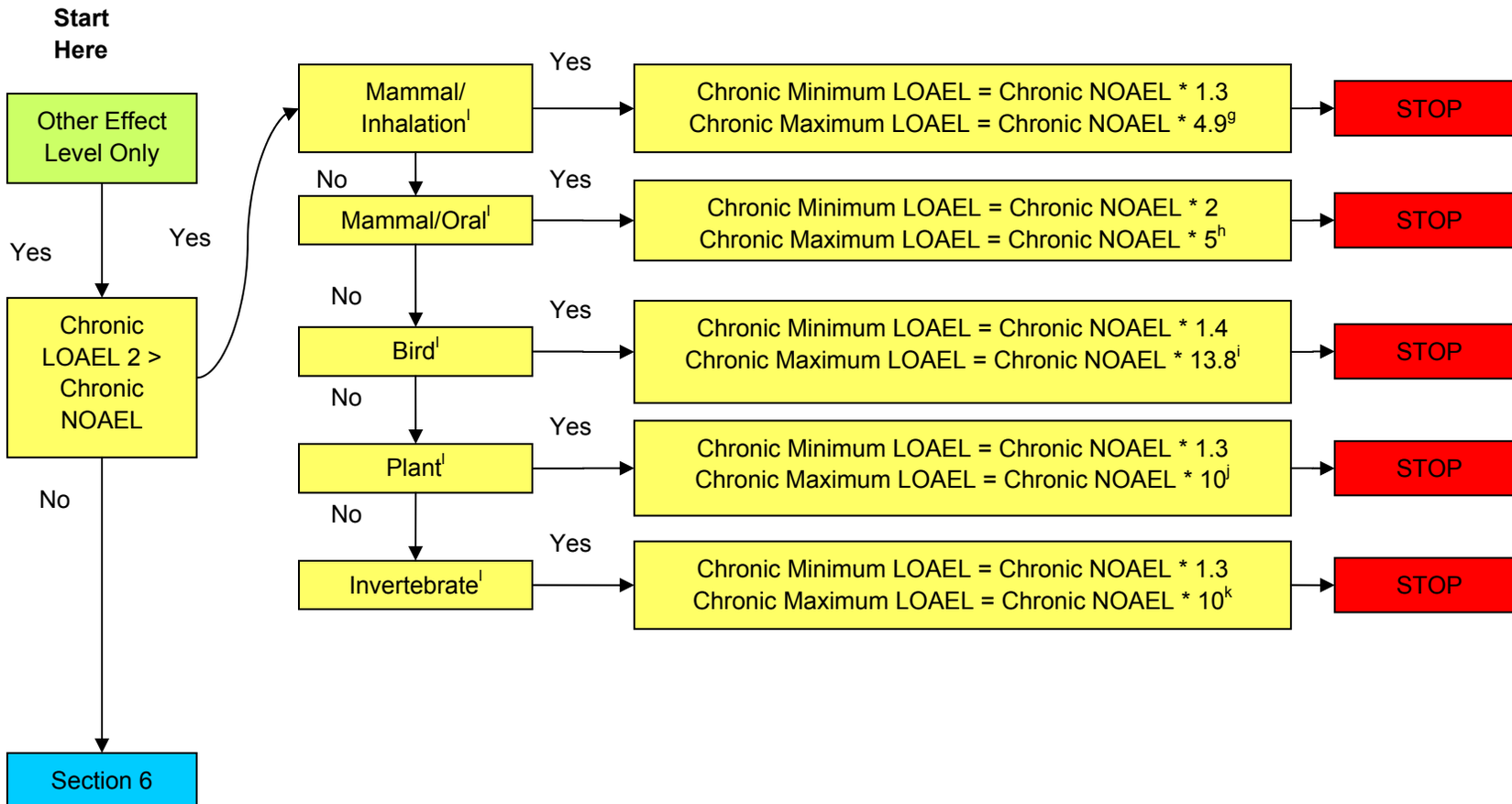


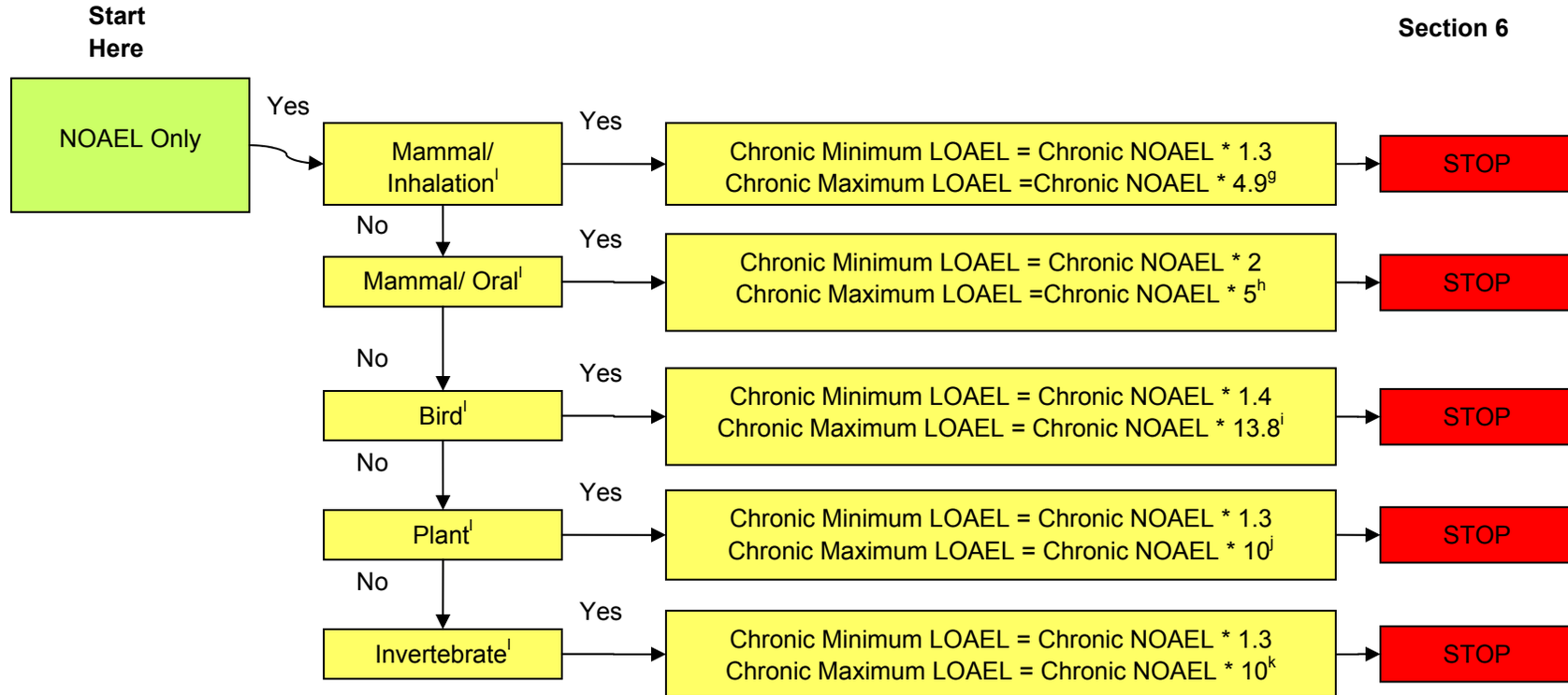
Figure A-1 (continued) Process for selecting the chronic LOAEL- or LOEC-based effect level for each endpoint in the GMM TRV data set

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Figure A-1 (continued) Process for selecting the chronic LOAEL- or LOEC-based effect level for each endpoint in the GMM TRV data set



**Figure A-1 (continued) Process for selecting the chronic LOAEL- or LOEC-based effect level for each endpoint in the GMM TRV data set**

**Notes for Figure A-1:**

<sup>a</sup> Refers to the reported LOAEL/LOEC.

<sup>b</sup> Refers to the reported other effect level (e.g., LD<sub>50</sub>, LC<sub>50</sub>, ED<sub>50</sub>, EC<sub>50</sub>).

<sup>c</sup> Chronic NOAEL/NOEC represents either a reported chronic NOAEL, or it was derived by extrapolating from another reported effect level (e.g., LOAEL, LD<sub>50</sub>) using UFs.

<sup>d</sup> Refers to the chronic LOAEL/LOEC estimated from a reported other effect level.

<sup>e</sup> Refers to the chronic LOAEL/LOEC estimated from a reported LOAEL/LOEC.

<sup>f</sup> Maximum and minimum chronic LOAELs/LOECs are estimated to bound the actual chronic LOAEL/LOEC when the chronic LOAEL/LOEC estimated from a reported LOAEL/LOEC is less than the chronic LOAEL/LOEC estimated from a reported other effect level.

<sup>g</sup> These factors are obtained from the minimum and maximum of a range of ratios determined using NOAEL and LOAEL pairs in the Ecorisk Database (LANL 2004, 087386, Ref ID 1442). These NOAEL and LOAEL pairs represent ecologically relevant data for inhalation of volatile organic compounds by terrestrial mammals.

<sup>h</sup> Factors are obtained from Dourson and Stara (1983, 073474, Ref ID 1379).

**Notes for Figure A-1 (continued):**

<sup>i</sup> Factors are obtained from the minimum and maximum of a range of ratios determined using NOAEL and LOAEL pairs based on ecologically relevant bird data in the Ecorisk Database (LANL 2003, 080117, Ref ID 1294).

<sup>j</sup> Factors are obtained from the minimum and maximum of a range of ratios determined using NOEC and LOEC pairs based on ecologically relevant plant data in the Ecorisk Database (LANL 2003, 080117, Ref ID 1294).

<sup>k</sup> Factors are obtained from the minimum and maximum of a range of ratios determined using NOEC and LOEC pairs based on ecologically relevant invertebrate data in the Ecorisk Database (LANL 2003, 080117, Ref ID 1294).

<sup>l</sup> Maximum and minimum chronic LOAELs/LOECs are estimated to bound the actual chronic LOAEL/LOEC when only a reported NOAEL/NOEC is available. First, the reported NOAEL/NOEC is used to estimate a chronic NOAEL/NOEC from which the maximum and minimum chronic LOAELs/LOECs are sometimes estimated by using extrapolation factors specific to the receptor data set being processed.

#### A-4.1.6 Calculation of the GMM TRV

Next, if three or more ecologically relevant chronic NOAEL- or NOEC-based effect levels are available, the GMM TRV is calculated as follows:

$$\text{GMM TRV} = \sqrt[n]{\text{EL}_1 * \text{EL}_2 * \text{EL}_3 * \dots * \text{EL}_n}$$

Where n is greater than 3 and each effect level (EL) represents a chronic NOAEL- or NOEC-based effect level for an ecologically relevant effect (i.e., reproduction, development, survival, adult weight change, or adult size change). The GMM TRV and effect levels are in units of mg/kg/d for birds and mammals and mg/kg for invertebrates and plants.

#### A-4.2 CS TRVs

If there are two or less ecologically relevant PTVs available in a GMM TRV data set for a chemical, receptor, and exposure medium scenario of concern, a CS TRV is developed instead. However, because there are two or less ecologically relevant PTVs available, the data set becomes limited. As a result, PTVs that were eliminated from the GMM TRV data set because of their lesser ecological relevance are added back into the CS TRV data set for consideration.

The ideal CS TRV for ecological risk screening assessments is one that is conservative in protecting the most sensitive ecologically relevant endpoint (i.e., reproduction/development), exposure route (i.e., oral ingestion via food or drinking water in birds or mammals, inhalation in mammals, uptake via seed coat and/or roots in plants, or oral and dermal contact in invertebrates), exposure medium (i.e., food or drinking water in birds and mammals, air for mammals, or soil for plants and invertebrates), exposure period (chronic), and effect level (NOAEL for birds and mammals or NOEC for plants and invertebrates). Before consideration for the TRV, each PTV is extrapolated to a chronic NOAEL- or NOEC-based effect level, if needed, using UFs (see Table A-13). Next, the information for each PTV is reviewed in detail and then the PTV that best represents the most sensitive ecological exposure scenario of concern (e.g., chronic, low-level exposure via food ingestion) is selected as the CS TRV. Typically, the most chronic, highest NOAEL/NOEC under the lowest LOAEL/LOEC for similar endpoints is selected. If there is a LOAEL/LOEC lower than the lowest NOAEL/NOEC, this effect level is usually selected and extrapolated to a chronic NOAEL- or NOEC-based effect level. Usually, if NOAELs/NOECs and/or LOAELs/LOECs are available, LC<sub>xx</sub>s or LD<sub>xx</sub>s, and EC<sub>xx</sub>s or ED<sub>xx</sub>s are eliminated early in the consideration process. The CS TRV and the data set from which it was selected are graphed and documented in detail in the PTSE Part 3 data-entry database.

#### A-4.3 Organization and Presentation of TRV Data Set Information

##### A-4.3.1 Organization of TRV Data in Tabular Format

Before data entry in the PTSE Part 3 database begins, all information is first organized and documented in Microsoft Word, Excel, and Access applications. This facilitates the gathering of information into organized formats for drafting, reviewing, and editing the TRV summary report before it is entered into numerous fields of the database. First, an output of the TRV data set in Excel is generated and exported from the Access database that runs the exclusion criteria for GMM TRV data sets, or if a GMM TRV cannot be developed, the output includes all values in the data set to be considered for the CS TRV. This output contains basic, crucial information for the PTVs considered in the data set, such as the chemical, test organism name and order, types of original effect levels, chronic NOAEL- or NOEC-based effect levels, chronic LOAEL- or LOEC-based effect levels, and UFs applied. Information from this table is used

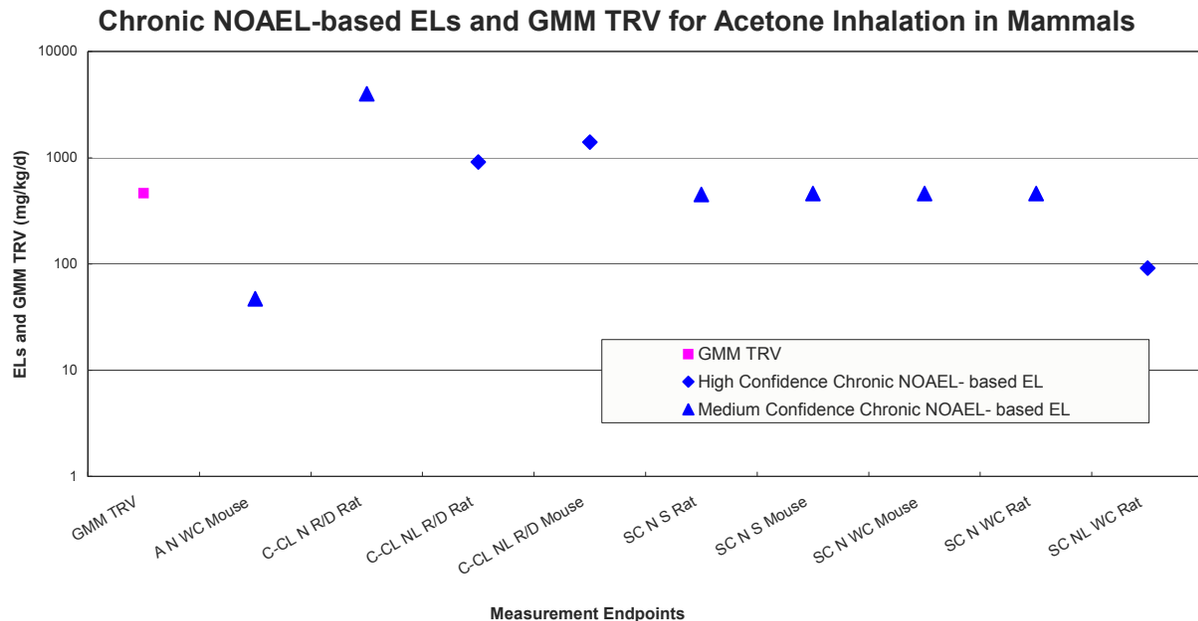
to create two other tables for GMM TRVs: test organism orders and original effect level types. An additional worksheet in the Excel file for the GMM TRV is also created to calculate the geometric standard deviation (GSD) and any outliers (values greater than 2 GSDs from the GMM TRV) that result. The outliers are not eliminated from the data set; therefore, the GMM TRV is not recalculated (see section A-4.3.3, Table A-15 for further explanation of outliers). Finally, the NOAEL- or NOEC-based effect level and LOAEL- or LOEC-based effect level graphs are created. Only graphs for CS TRVs are created from this output.

#### **A-4.3.2 Presentation of TRV Data in Graphs**

Before the TRV summary report is drafted in Word, a graph of the GMM or CS TRV and the chronic NOAEL- or NOEC-based effect levels in its data set is created in Microsoft Excel. The GMM TRV data set is defined as all of the PTVs for a particular receptor group/chemical/exposure route scenario of concern that have passed the exclusion criteria and that have been extrapolated to chronic NOAEL- or NOEC-based effect levels. Similarly, the graph for the CS TRV data set also includes the TRV as well the chronic NOAEL- or NOEC-based effect levels in the data set. However, the graph for CS TRVs can also include other data values that were originally eliminated from the GMM TRV data set.

Regardless of the type of TRV, in larger data sets, the y-axis on the graph is sometimes set to logarithmic scale to show the numerous values clearly. Each NOAEL- or NOEC-based effect level data point on the graph has a shape that represents the PTV confidence rating (diamond, triangle, and circle for high, medium, and low confidence, respectively). Dark blue data points (diamonds, triangles, or circles) represent chronic NOAEL- or NOEC-based effect levels, while the pink data square represents the TRV. An example of a GMM TRV graph is seen in Figure A-2. Graphs presented in the Ecorisk Database will usually not show low confidence PTVs because they will have been eliminated from the data set. They are eliminated at this early stage because insufficient data preclude producing effect levels that can be used in confidently predicting toxicity.

A graph is also created, in a similar manner as the one for NOAEL- or NOEC-based effect levels, for chronic LOAEL- or LOEC-based effect levels in the TRV data set. However, confidence ratings are not highlighted in this graph, and the LOAEL- or LOEC-based effect level data points are represented by dark blue diamonds.



**Figure A-2** Example of a graph illustrating the GMM TRV for the inhalation of acetone in mammals and its corresponding NOAEL-based effect levels

#### A-4.3.3 Assigning Confidence Ratings to TRVs

For GMM TRVs, a second Excel file is created for scoring criteria and confidence ratings. This type of file is not needed for CS TRVs because the confidence rating of the CS TRV is the PTV confidence rating (see section A-3.2.8) for the value upon which the CS TRV is based. The confidence ratings for GMM TRVs are based on a different set of criteria with the purpose of determining how well the GMM TRV represents the ideal GMM TRV, which represents the true TRV. The true TRV is the dose rate or concentration that is equivalent to a no adverse effect level for population level effects (i.e., decreased population size) for a particular receptor under a specific exposure scenario for a particular chemical in the real world. The confidence rating for the GMM TRV is based on how well the GMM TRV meets various criteria within specific evaluation categories. A weighted scoring system based on the degree of influence each evaluation category has on the GMM TRV is used to assess the validity of the GMM TRV for estimating the true TRV. The following sections describe the structure of the confidence rating system for GMM TRVs, including descriptions and justifications for the evaluation processes used to assign the confidence ratings.

#### GMM TRV Confidence Rating System Structure

The first step in assigning a confidence rating to a GMM TRV is to assign a score for each of 11 evaluation categories. Each evaluation category contains individual criteria associated with ranked scores that reflect how well the GMM TRV data set being evaluated represents the characteristics of the ideal GMM TRV. The higher the score, the better the GMM TRV represents the ideal GMM TRV and thus the true TRV.

The second step in assigning a confidence rating to a GMM TRV is to calculate a weighted score for each evaluation category by multiplying the individual scores of each evaluation category by the weighting factor of the evaluation category. The weighted score for each evaluation category is based on the

weighting factor level assigned to the evaluation category. The weighting factor level is based on the degree of influence the evaluation category has on setting the GMM TRV. The higher the weighting factor, the greater the influence the evaluation category has on setting the GMM TRV. The possible weighting factor levels are presented in Table A-14.

**Table A-14**  
**Weighting Factor Levels**

Weighting Factor Level	Definition	Weighting Factor Applied
Critical	A low score for a critical evaluation category triggers reinvestigation of the GMM TRV and possible revision or decision not to use.	2
Noncritical	A high score for a noncritical evaluation category indicates the GMM TRV data set is very robust, highly relevant to the scenario for which the TRV is being developed, or is based primarily on effect levels that were not derived by applying UFs to PTVs. A low score rarely influences revision of the GMM TRV because it is an added benefit if the evaluation category scores high, but not a requirement.	1

The third step in assigning a confidence rating to a GMM TRV is to calculate a total weighted score for the GMM TRV being evaluated. The total weighted score is equal to the sum of weighted scores of all 11 evaluation categories. Table A-15 presents the scores, weighting factors, weighting factor levels, and weighted scores for each evaluation category. The justifications for the scores and weighting factor levels are presented in the Justification for Scoring Criteria and Weighting Factor Levels subsection of section A-4.3.

**Table A-15**  
**Scores, Weighting Factors, and Weighted Scores for each Evaluation Category and Criterion**

Evaluation Category	Evaluation Criterion	Score	Weighting Factor	Weighted Score
Number of experiments	Equal to 10 or more	1.5	1	1.5
	Between 4 and 9	1	1	1
	Less than or equal to 3	0.5	1	0.5
Type of exposure medium	Test exposure medium matches that of concern	1	1	1
	Test exposure medium partially matches that of concern	0.5	1	0.5
Number of test organism orders	Equal to 3 or more	1.5	1	1.5
	Equal to 2	1	1	1
	Equal to 1	0.5	1	0.5
Number of unique measurements (endpoints)	More than 3	1.5	1	1.5
	Equal to 3	1	1	1
	Less than 3	0.5	1	0.5

Table A-15 (continued)

Evaluation Category	Evaluation Criterion	Score	Weighting Factor	Weighted Score
Type of endpoint category	R/D	3.5	1	3.5
	Combination of R/D and S	3	1	3
	Combination of R/D, S, and WC or SzC	2.5	1	2.5
	Combination of R/D and WC or SzC	2	1	2
	S	1.5	1	1.5
	Combination of S and WC or SzC	1	1	1
	WC or SzC	0.5	1	0.5
Number and type of effect levels of PTVs associated with the individual NOAEL- or NOEC-based effect levels in GMM TRV data set	2 or more chronic (or C-CL) NOAELs/NOECs with LOAELs/LOECs	3.5	1	3.5
	1 chronic (or C-CL) NOAELs with LOAELs	3	1	3
	1 or more chronic (or C-CL) NOAELs without LOAELs	2.5	1	2.5
	1 or more chronic (C-CL) LOAELs	2	1	2
	1 or more subchronic NOAEL with LOAEL	1.5	1	1.5
	1 or more subchronic NOAEL without LOAEL	1	1	1
	1 or more subchronic LOAEL or other effect level or acute NOAEL, LOAEL, or other effect level	0.5	1	0.5
Confidence rating of PTVs associated with the individual NOAEL- or NOEC-based effect levels in GMM TRV data set	100% of the effect levels have high confidence ratings	2	1	2
	Effect levels have a mixture of high and medium confidence ratings	1.5	1	1.5
	100% of the effect levels have medium confidence ratings	1	1	1
	Effect levels have a mixture of high, medium, and low confidence ratings	0.5	1	0.5
Outlier(s) in chronic NOAEL- or NOEC-based effect level distribution	100% of data are within a GSD less than or equal to 2	4	2	8
	75%–99% of data are within a GSD less than or equal to 2	3	2	6
	75% or more of data are within a GSD of 6	2	2	4
	75% or more of data are within a GSD of 10	1	2	2
	None of the above	0	2	0
Chronic NOAEL- or NOEC-based effect level distribution is bimodal*	No	2	2	4
	N/A - Evaluation is not possible because data set is too limited	1	2	2
	Yes	0	2	0

Table A-15 (continued)

Evaluation Category	Evaluation Criterion	Score	Weighting Factor	Weighted Score
Relationship of GMM TRV to chronic LOAEL- or LOEC-based effect levels	The GMM TRV is less than the lowest LOAEL- or LOEC-based effect level	3	2	6
	The GMM TRV is higher than the lowest chronic LOAEL- or LOEC-based effect level by a factor of 3 or less and is protective of the majority of R/D endpoints. Furthermore, the lowest chronic LOAEL- or LOEC-based effect level represents a chronic or C-CL LOAEL or other effect level for an R/D endpoint.	2	2	4
	The GMM TRV is higher than the lowest chronic LOAEL- or LOEC-based effect level by a factor of 3 or less, and the lowest chronic LOAEL- or LOEC-based effect level represents a chronic LOAEL or other effect level for an S, WC, or SzC endpoint.	1.5	2	3
	The GMM TRV is higher than the lowest chronic LOAEL- or LOEC-based effect level by a factor of 3 or less, and the lowest chronic LOAEL-based effect level is extrapolated from a subchronic or acute LOAEL or other effect level (e.g., EC <sub>20</sub> , LD <sub>50</sub> ) for an R/D, S, WC, or SzC endpoint.	1	2	2
	The GMM TRV is higher than the lowest chronic LOAEL- or LOEC-based effect level by a factor of 3 or less, and the lowest LOAEL-based effect level is derived from a subchronic or acute NOAEL for an R/D, S, WC, or SzC endpoint.	0.5	2	1
	None of the above	0	2	0
Relationship of GMM TRV to other published TRVs	Acceptable	2	2	4
	No comparison available	1.5	2	3
	Not acceptable	0	2	0

\*Bimodality can only be evaluated for data sets with 10 or more chronic NOAEL- or NOEC-based effect levels.

The fourth step in assigning a confidence rating to a GMM TRV is to determine the percentage the total weighted score is of the maximum total weighted score for the evaluation (i.e., 36.5 points based on summing the highest scores from each evaluation category). The total weighted score percentage of the maximum total weighted score is the ultimate basis for assigning the confidence rating of a GMM TRV. Table A-16 presents the confidence ratings and the corresponding percentage of the maximum total weighted score and the equivalent total weighted score.

**Table A-16**  
**Confidence Ratings for GMM TRVs**

Confidence Rating	Percent of Maximum Total Weighted Score (%MTWS)	Equivalent Total Weighted Score (ETWS)
High	$\%MTWS \geq 75\%$	$27.375 \leq ETWS \leq 36.5$
Medium	$50\% \leq \%MTWS < 75\%$	$18.25 \leq ETWS < 27.375$
Low	$25\% < \%MTWS < 50\%$	$9.125 < ETWS < 18.25$
Unacceptable	$\%MTWS \leq 25\%$	$ETWS \leq 9.125$

### Justification for Scoring Criteria and Weighting Factor Levels

Table A-17 provides the justification for the scoring criteria and weighting factor levels of each evaluation category.

**Table A-17**  
**Justifications for Scoring Criteria and Weighting Factor Levels for Each Evaluation Category**

Evaluation Category	
Justification for Scoring Criteria	Justification for Weighting Factor Level
<b>Number of Experiments</b>	
The preference is to have a high number of experiments because this reduces the potential for the data set to be biased toward a particular study design. Based on best professional judgment, having 10 experiments is considered to provide a more than adequate representation of the toxicity of a chemical for the test organism group of concern. Having 4 to 9 experiments is considered to provide an adequate representation, while having 3 or fewer experiments is considered to provide a minimal representation of the toxicity of a chemical for the test organism group of concern.	This category is given a noncritical weighting factor level. This evaluation category has a strong relationship to the robustness of the data set and its ability to represent the ideal GMM TRV; thus, the true TRV is estimated. The higher the number of experiments, the more robust the data set. This evaluation category is not, however, a primary factor for determining whether or not the GMM TRV should be used because a high number of experiments in the data set is not a requirement, but rather an additional benefit for assessing confidence in the TRV.
<b>Type of Exposure Medium</b>	
The preference is for all the effect levels in the data set to be associated with an exposure medium that is equivalent to the exposure medium of concern. However, if the data set is limited (i.e., less than four effect levels for a particular exposure medium), effect levels that have an appropriate surrogate exposure medium (i.e., exposure medium that has the same exposure route as the exposure route of concern) may be used to supplement the data set so that a GMM TRV can be derived. For example, for an oral ingestion via food TRV, only food effect levels should be used, but if the data set is limited, oral ingestion via drinking water effect levels may be used to supplement the data set so that a GMM TRV may be calculated.	This category is given a noncritical weighting factor level. This evaluation category indicates the degree of relevance the data set has to the TRV that is being developed. The higher the degree of relevance, the more closely the GMM TRV represents the ideal GMM TRV; thus, the true TRV is estimated. This evaluation category is not, however, a primary factor for determining whether or not the GMM TRV should be used because an exact match of the exposure medium for which the TRV is being developed is not a requirement, but rather an additional benefit for assessing confidence in the TRV. Only the exposure route must match the exposure for which the TRV is being developed. However, the toxicity can vary greatly in different exposure media as a result of the differences in bioavailability of the chemical in one compared to the other. Therefore, a complete match of the exposure medium is preferred to more accurately estimate the true TRV.

Table A-17 (continued)

Evaluation Category	
Justification for Scoring Criteria	Justification for Weighting Factor Level
<b>Number of Test Organism Orders</b>	
The preference is to have a high number of test organism orders because this reduces the potential for the data set to be biased toward one order of test organisms. The scoring criteria are based upon the USACHPPM guidance that states that having at least two different taxonomic orders in a TRV data set helps define the quality of the data set (Ryti et al. 2004, 076074, Ref ID 1481).	This category is given a noncritical weighting factor level. This evaluation category has a strong relationship to the robustness of the data set and its ability to represent the ideal GMM TRV; thus, the true TRV is estimated. The higher the number of test organism orders, the more robust the data set. This evaluation category is not, however, a primary factor for determining whether or not the GMM TRV should be used because a high number of test organism orders in the data set is not a requirement, but rather an additional benefit for assessing confidence in the TRV.
<b>Number of Unique Measurements (Endpoints)</b>	
The preference is to have a high number of unique measurements (endpoints) because this helps ensure the robustness of the GMM TRV by including multiple toxicological effects. Unique measurements are those that represent different parameters of measurement for an endpoint category. For example, the endpoints of "mortality" and "LC <sub>50</sub> " may both be categorized as S endpoints because they are both measurements of survival/mortality, but they are each considered a unique measurement because they measure different aspects of survival/mortality. Based on best professional judgment, having more than three unique measurements is considered to provide a more than adequate representation of the toxicity of a chemical for the test organism group of concern. Having three unique measurements is considered to provide an adequate representation while having fewer than three unique measurements is considered to provide a minimal representation of the toxicity of a chemical for the test organism group of concern.	This category is given a noncritical weighting factor level. This evaluation category is related to the robustness of the data set and its ability to represent the ideal GMM TRV; thus, the true TRV is estimated. The higher the number of unique measurements, the more robust the data set. This evaluation category is not, however, a primary factor for determining whether or not the GMM TRV should be used because a high number of unique measurements in the data set is not a requirement, but rather an additional benefit for assessing the validity of the GMM TRV to estimate the true TRV. Furthermore, all the unique measurements that are allowed in the data set are, by definition, relevant to the TRV being developed for population effects. The relevance of the endpoint category of each unique measurement is scored separately under the Type of Endpoint Category evaluation category below.
<b>Type of Endpoint Category</b>	
The preference is to have more reproduction and development endpoints followed by survival endpoints and then by adult body weight or size change endpoints because the first category of endpoints is the most ecologically relevant group for determining long-term effects on populations, followed by the second and third categories.	This category is given a noncritical weighting factor level. This evaluation category indicates the degree of relevance the data set has to the effects of concern, population level effects, for which the GMM TRV is being developed. The higher the degree of relevance, the more closely the GMM TRV represents the ideal GMM TRV; thus, the true TRV is estimated. This evaluation category is not, however, a primary factor for determining whether or not the GMM TRV should be used because all the endpoint categories considered are ecologically relevant by definition. However, reproduction or development endpoints can more closely approximate population level effects, so having more endpoints in this category is an added benefit for assessing the validity of the GMM TRV for estimating the true TRV.

<b>Number and Type of Effect Levels of PTVs Associated with the Individual NOAEL- or NOEC-Based Effect Levels in the GMM TRV Data Set</b>	
<p>The preference is to have chronic NOAELs/NOECs with LOAELs/LOECs, followed by chronic NOAELs/NOECs without LOAELs/LOECs, then by subchronic NOAELs/NOECs with LOAELs/LOECs, then by subchronic NOAELs/NOECs without LOAELs/LOECs and finally by all other effect levels. This hierarchy is based on two factors. One factor is whether or not UFs have to be applied to a PTV to extrapolate to a chronic NOAEL/NOEC. Extrapolated values are less preferred because they may be overly conservative and thus less representative of the actual chronic NOAEL/NOEC. The second factor is whether or not there are any NOAELs/NOECs with accompanying LOAELs/LOECs. NOAELs/NOECs with LOAELs/LOECs are most preferred because these values bracket the range of possible effects better than just a NOAEL/NOEC or just a LOAEL/LOEC alone.</p>	<p>This category is given a noncritical weighting factor level. This evaluation category is directly related to the certainty in the GMM TRV. The more effect levels in the GMM TRV data set that were extrapolated to chronic NOAEL- or NOEC-based effect levels by applying UFs, the greater the level of conservatism that is built into the GMM TRV. Even though being overly conservative is acceptable for screening-level ecological risk assessments, it is preferred that TRVs not be overly conservative if more certain data are available. On the other hand, the higher the number of original effect levels that are chronic NOAELs/NOECs in the GMM TRV data set, the higher the confidence that the GMM TRV represents the ideal GMM TRV and thus estimates the true TRV (chronic NOAEL). A high score in this evaluation category is not required, but is an additional benefit for assessing confidence in the TRV.</p>
<b>Confidence Rating of PTVs Associated with the Individual NOAEL- or NOEC-Based Effect Levels in GMM TRV Data Set</b>	
<p>The preference is to have more effect levels (PTVs) with high confidence ratings, followed by those with medium ratings and then by those with low ratings. A PTV confidence rating indicates to what degree the PTV is ecologically relevant, defensible, and well documented based on the PTSE Part 2 study evaluation criteria. Effect levels associated with a low confidence rating are not included in the data set unless the data set is limited (i.e., less than three effect levels based on PTVs with either a high or medium confidence rating.).</p>	<p>This category is given a noncritical weighting factor level. This evaluation category indicates the degree of relevance the data set has to the TRV that is being developed. The higher the degree of relevance, the more closely the GMM TRV represents the ideal GMM TRV; thus, the true TRV is estimated. The PTV confidence rating is based upon scoring various study elements that are considered to be relevant for developing a scientifically defensible and ecologically relevant TRV. A high PTV confidence rating indicates the value is highly relevant for deriving a TRV and more likely to accurately estimate the true TRV.</p>
<b>Outliers(s) in the Chronic NOAEL- or NOEC-Based Effect Level Distribution</b>	
<p>The data set cannot have invalid outliers (i.e., values associated with error or study designs that do not meet the minimum requirements for deriving a TRV). Invalid outliers must be removed from the data set before calculation of the GMM TRV. An invalid outlier is determined by a low confidence rating of a PTV associated with an effect level in the data set. However, valid outliers, or extreme values, are allowed (e.g., sensitive species) as long as the data set is not bimodal (see the Chronic NOAEL- or NOEC-based Effect Level Distribution is Bimodal evaluation category below). The GSD is used to determine the variance of the GMM TRV. A lower variance (smaller GSD) indicates that the GMM TRV is more likely to represent the ideal GMM TRV and thus more accurately estimate the true TRV while a high variance (higher GSD) indicates that the GMM TRV is less likely to represent the ideal GMM TRV and thus less accurately estimate the true TRV. In most cases of high variance, the GMM TRV may be overly conservative because the large variance in the values is a result of the averaging of effect levels that are based on PTVs other than chronic NOAELs/NOECs and the application of UFs to extrapolate these values to chronic NOAEL- or NOEC-based effect levels. A data set that contains both the smaller, extrapolated values and the nonextrapolated values (i.e., original effect levels that were already chronic NOAELs/NOECs) leads to a high variance.</p>	<p>This category is given a critical weighting factor level. This evaluation category represents the variance of the GMM TRV dataset, which is important because it indicates how well the GMM TRV represents the ideal GMM TRV. Thus, this evaluation category indicates how well the GMM TRV estimates the true TRV, which is directly related to the confidence in the GMM TRV. Low variance equals high confidence. High variance equals low confidence and may require reconsideration of the GMM TRV.</p>

<b>Chronic NOAEL- or NOEC-Based Effect Level Distribution is Bimodal</b>	
<p>The preference is for the GMM TRV data set to not have a bimodal distribution. A bimodal distribution is determined based on two distinct clusters of values associated with different test species, original exposure durations, original effect levels, or endpoint categories of each effect level in the data set. If a data set is bimodal, best professional judgment must be used to determine if a subset GMM TRV(s) (i.e., a TRV calculated from a data set smaller than the original) needs to be calculated or if the GMM TRV can be used as is.</p>	<p>This category is given a critical weighting factor level. This evaluation category has a high influence on whether or not the GMM TRV will be used. If the GMM TRV data set is found to have a bimodal distribution, the GMM TRV may need to be revised to represent the most sensitive and/or ecologically relevant distribution (e.g., one distinct cluster is rodent [omnivore] data while the other is mink [carnivore] data. A TRV calculated from rodent data is more appropriate for the omnivorous deer mouse ESL receptors, while a TRV calculated from the mink is more appropriate for carnivorous red fox ESL receptor.)</p>
<b>Relationship of GMM TRV to Chronic LOAEL- LOEC-Based Effect Levels</b>	
<p>The preference is to have the GMM TRV below the lowest chronic LOAEL- or LOEC-based effect level because that indicates it is protective of the most sensitive adverse effect in the data set. If the GMM TRV is not below the lowest chronic LOAEL- or LOEC-based effect level, the next preference is for it to be no more than 3 times higher than a chronic LOAEL- or LOEC-based effect level based on a chronic or C-CL LOAEL/LOEC for an R/D or less ecologically relevant endpoint. The next preference is to have the GMM TRV at no more than 3 times higher than a chronic LOAEL- or LOEC-based effect level extrapolated from an original effect level other than a LOAEL. Because some of the chronic LOAEL- LOEC-based effect levels are extrapolated from NOAELs/NOECs or other effect levels by applying UFs, they may be overly conservative and not represent the true chronic LOAELs/LOECs for particular endpoints. In such cases, the GMM TRV is considered adequately protective as a result of the conservatism built into the extrapolated chronic LOAEL- or LOEC-based effect levels. Furthermore, the GMM TRV may be considered adequately protective if it is below the chronic LOAEL- or LOEC-based effect levels for the most ecologically relevant endpoints (reproduction and development) even though it may exceed the lowest chronic LOAEL- or LOEC-based effect level for an adult body weight or size change endpoint or for a survival endpoint. Another consideration is to determine, based on best professional judgment, whether or not the GMM TRV is unacceptably higher or lower than the lowest chronic LOAEL- or LOEC-based effect level. If the difference is unacceptable, further investigation is warranted to determine if the GMM TRV is inappropriate (i.e., unacceptably over- or under-conservative). If it is found to be unacceptable, then the GMM TRV may need to be revised.</p>	<p>This category is given a critical weighting factor level. This evaluation category has a high influence on whether or not the GMM TRV will be used. If the difference between the GMM TRV and the lowest chronic LOAEL- or LOEC-based effect level is unacceptable, the GMM TRV is unacceptable and an alternative (e.g., a subset GMM TRV, CS TRV) needs to be considered.</p>
<b>Relationship of GMM TRV to other Published TRVs</b>	
<p>The preference is that any differences between the GMM TRV and other published TRVs be explained based on the experiments, endpoints, test organisms, and test chemical forms, etc., considered. It is also important that the explanation provide support for or against the use of the GMM TRV. It should be verified that the GMM TRV has considered all relevant data. If relevant data have not been considered, the GMM TRV data set may need to be expanded to include the missing data. If no published TRVs are available for comparison, the GMM TRV is considered to be acceptable.</p>	<p>This category is given a critical weighting factor level. This evaluation category has a high influence on whether or not the GMM TRV will be used. If differences between the GMM TRV and other published TRVs are unacceptable (i.e., unexplainable, error based, or lack of data based), the GMM TRV is unacceptable and an alternative (e.g., subset GMM TRV, CS TRV) needs to be considered.</p>

#### A-4.3.4 Drafting the TRV Summary Report and PTSE Part 3 Data Entry

The information organized in the Excel file(s) and presented in the graphs is used as reference and supporting documentation for the TRV summary report as it is drafted in a Microsoft Word format that contains the fields in the PTSE Part 3 data-entry database. The report is created in Word for ease of drafting, peer reviewing, and revising. The final report is then entered into the PTSE Part 3 data-entry database by copying and pasting sections one at a time into Access data fields. The graphs are copied and pasted into fields as well. However, the information in the test organism orders and original effect levels tables and the GSDs worksheet is not entered because these data are automatically generated and presented by the Ecorisk Database. Rather, this information has been created in Excel for reference while working on the TRV summary report.

The PTSE Part 3 data-entry fields are detailed below, and Attachments A-1 and A-2 contain examples of user-printable TRV summary reports for GMM and CS TRVs, respectively. Note that some fields such as reviewer initials and date are not included in the printable reports because they are for quality assurance documentation purposes only.

##### Reviewer Initials

The initials of the person entering the information in the PTSE Part 3 record are entered here. If significant changes are made to a record at a later time, the initials of the new reviewer replace the original reviewer initials.

##### Date

The date the PTSE Part 3 record is created or modified is entered here.

##### Last Updated

If any changes are made to the TRV in the record, the version date of the Ecorisk Database that these changes will appear in is entered in the last updated field.

##### Part 3 TRV Summary ID

A unique ID for the record is entered in this field (see Example A-18). The format, in one continuous string with each parameter separated by an underscore symbol, is as follows:

Analyte Code\_ESL Medium\_ESL Screening Receptor Group ID\_Test Organism Group ID\_Test Organism Common Name\_Test Exposure Medium\_TRV Type\_TRV Ref ID\_Primary Toxicity Study Ref ID

##### Example A-18 Part 3 TRV Summary IDs

107-06-2\_AIR\_M\_TM\_Mammal\_Air\_ChronicGMMNOAEL\_1442\_0001

HGI\_SEDIMENT\_B\_TB\_QuailJapanese\_Diet\_ChronicCSNOAEL\_1230\_0017

### **GMM TRV Record ID**

The ID for GMM TRVs provides the following information in a continuous string with no spaces or underscores: Analyte Code, Test Organism Type, the acronym GMM, and Test Exposure Medium. This field is left blank for CS TRV records. See Example A-19.

#### **Example A-19 GMM TRV Record IDs**

107-06-2MGMMMA  
11096-82-5MGMMF

### **Graph Group ID**

This field helps to identify all graphs belonging to a particular TRV and its data set. The format, in one continuous string with each parameter separated by an underscore symbol, is Analyte Code\_Test Organism Type\_TRV Type (see Example A-20).

#### **Example A-20 Graph Group IDs**

1746-01-6\_TM\_CS  
11096-82-5\_TM\_GMM

### **TRV Type**

The final TRV type is noted here. For birds and mammals, Chronic GMM NOAEL or Chronic CS TRV is entered. TRV type for earthworms and plants is entered as Chronic GMM NOEC or Chronic CS NOEC. In cases where a subset GMM TRV is created (i.e., a TRV calculated from a data set smaller than the original GMM TRV data set), the type is entered as Chronic subset GMM NOAEL or Chronic subset GMM NOEC.

### **TRV Final Value**

The value of the GMM TRV, subset GMM TRV, or CS TRV is entered here. This is the value after all calculations have been completed. Calculations include those for daily dose rates, moisture conversions, and any others from Part 2 records plus any contributions from UFs to be accounted for in this Part 3 record.

### **TRV Units**

For birds and mammals, the GMM or CS TRV is presented in units of mg/kg/d (representing mg chemical/kg body weight/d), while earthworms and plants have units of mg/kg (mg chemical/kg soil).

### **Selected TRV**

In this field, YES or NO is entered for each LANL GMM or CS TRV depending on whether or not it will be used in the ESL models for the Ecorisk Database. According to the tiered TRV development approach for the Ecorisk Database, the most preferred TRV is an EPA ecological soil screening level (Eco-SSL) TRV.

If one does not exist, the LANL GMM TRV is used, followed by the LANL CS TRV, then a secondary source TRV from another published source. Based on this hierarchy, it is likely that if a GMM TRV is developed, an EPA Eco-SSL TRV does not exist; therefore, YES is almost always entered for GMM TRVs. However, if the GMM TRV is not considered suitable, NO will be placed in its corresponding field, and YES will be entered for an alternative TRV (i.e., subset GMM TRV, CS TRV, or secondary source TRV), whichever of the more preferred TRVs is available and most suitable. This field can later be updated should an EPA Eco-SSL become available to replace a GMM TRV or should a GMM TRV or CS TRV be developed to replace a CS TRV or secondary source TRV, respectively.<sup>5</sup>

### **ESL Media**

For birds and mammals, the ESL media are soil, sediment, or water. For plants and earthworms, only one ESL medium of soil is used. If the GMM TRV data set or CS TRV represents food exposure for birds and mammals, two records are created: one each for soil and sediment ESLs. If the GMM TRV data set or CS TRV represents drinking water exposure for birds and mammals, only one record for water is created. Only one record (soil) is needed for each earthworm or plant and chemical combination.

### **Functional Group**

The code A, for all functional groups relevant to the test organism group (bird, invertebrate, mammal, or plant), is entered for GMM or CS TRVs unless it has been determined that the TRV is protective of certain functional groups only. An example is Aroclor-1260, where it was decided that the GMM TRV was not protective enough of the carnivore functional group because according to the data set, the TRV was not protective of mustelids, in which the reproductive effects of polychlorinated biphenyl exposure is well-documented. Instead, the LANL CS TRV for Aroclor-1260 was used. The GMM TRV for Aroclor-1260, however, was used for all other functional groups (all noncarnivores). The coding for the Aroclor-1260 GMM TRV record was N-C for noncarnivores while the coding for the Aroclor-1260 CS TRV was C for carnivores.

### **TRV Confidence Rating**

High, medium, or low is typed in this field for GMM or CS TRVs. Low is rarely, if ever, seen because data receiving a low confidence rating results in the primary toxicity study being rereviewed and eliminated from the data set for GMM or CS TRVs. For CS TRVs, a brief description of the number and type of experiments, confidence ratings, and endpoint categories also follows (e.g., "Medium. Data set consists of 1 experiment, 1 medium confidence PTV, and 1 survival endpoint."). This extra information helps Ecorisk Database users to see the breadth of the data set from which the CS TRV was chosen in addition to the confidence rating of the single value, which is based on the type and degree of detail of information of the study from which it was obtained.

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<sup>5</sup> In the early developmental stages of the Ecorisk Database, before GMM TRVs were developed, CS TRVs representing food exposure were used in soil, sediment, and water ESL models. Likewise, CS TRVs representing drinking water exposures were used in all ESL models as well. Notes regarding bioavailability of the chemical in one medium versus the other were made in the report. Currently, GMM and CS TRVs for food are limited to soil and sediment ESL models only, while TRVs representing drinking water exposures are used only in water ESL models.

### **Primary Toxicity Paper Reference ID**

Because the GMM TRV is usually based on more than one primary toxicity reference, this field is not applicable. Ref ID 0001, which represents not applicable, is entered. For CS TRVs, this field contains the Ref ID of the reference containing the information from which the TRV originated.

### **TRV Reference ID**

The Ref ID for the version of the Ecorisk Database in which this new record (GMM or CS TRV) will appear is entered.

### **Description of TRV Source**

There are various options in the list, but for new Part 3 records that result in the addition of a new LANL GMM or CS TRV to the Ecorisk Database, the selection should be “LANL derived value based on reviewed primary data.”

### **Exposure Medium**

The exposure medium that the GMM or CS TRV represents is selected from the drop-down list.

### **Exposure Route**

The primary exposure route that the GMM or CS TRV represents is selected from the drop-down list.

### **Organism Name**

The organism group representing the organisms in the GMM TRV data set (i.e., bird, mammal, invertebrate, or plant) is selected from the drop-down list. For CS TRVs, the organism name is the common name of the organism represented (e.g., “Rat, Sprague-Dawley”). This is selected from the drop-down list as well.

### **Organism ID**

The code for the organism categories represented by the GMM or CS TRV (as seen in PTSE Part 1, Data Entry) is selected from the drop-down list. The four choices usually selected in new Part 3 records are SLE for earthworms, TB for terrestrial bird, TM for terrestrial mammals, and TP for terrestrial plants. Note that sometimes a bird that is considered an aquatic species is represented in the terrestrial data set (e.g., mallard duck). The TB code is still used for these organisms because they are considered to toxicologically represent a surrogate for terrestrial species. Other aquatic species for mammals, invertebrates, or plants are rejected from the literature set used for review, so they should not be encountered this far into the PTSE process.

### **Screening Receptor Group ID**

The code for the organism group represented by the GMM or CS TRV is selected from the drop-down list. The four choices usually selected in new Part 3 records are B for bird, I for invertebrates (earthworms), M for mammals, and P for plants.

## Chemical ID

The analyte code that the GMM or CS TRV represents is selected from the drop-down list.

## Surrogate Chemical ID

If a surrogate chemical is used, the analyte code for the surrogate chemical is selected. Otherwise, the analyte code the GMM or CS TRV represents is selected from the drop down list; it matches the Chemical ID.

## Discussion

### ***GMM TRVs***

For GMM TRVs, this field holds two paragraphs, the first discusses an overview of the data set used to derive the TRV, and the second is a conclusion summary. The first paragraph includes the following information:

- type of TRV (GMM),
- exposure medium,
- chemical and organism group of concern,
- value of GMM TRV and its units,
- number of chronic NOAEL- and NOEC-based effect levels (PTVs) used to calculate the GMM TRV,
- number of references in the data set,
- number of experiments in the data set,
- number of unique measurements (endpoints) in the data set,
- number of phylogenetic test organism orders,
- endpoint categories represented in the data set,
- number or percent of high, medium, and low PTV confidence ratings,
- exposure routes, and
- relevance or relationship between test exposure route and exposure route of concern for the particular ESL of concern (i.e., sediment, soil, water).

The conclusion paragraph for GMM TRVs summarizes the suitability of the GMM TRV for use in ESL models. The suitability of the GMM TRV is based on further evaluation of the distribution of chronic NOAEL- or NOEC-based effect levels, comparison of the GMM TRV to the lowest chronic LOAEL- or LOEC-based effect level, and comparison of the GMM TRV to other published TRVs. Although this general discussion field is the first of the discussion fields, this field is usually completed last in the data entry process for Part 3. Each of the other discussion fields is explained in detail below. The conclusion paragraph for GMM TRVs includes

- the GMM TRV confidence rating;
- a numbered list of scoring criteria in support of this confidence rating;

- a statement of whether the comparison of the GMM TRV to other published TRVs is acceptable;
- a statement of why bimodality of the data set distribution could not be assessed, if needed;
- another numbered list of criteria, not listed above, that lowered or do not support the confidence rating;
- brief explanation(s) of why criteria did not score well or did not strongly support confidence rating;
- explanation of whether GMM TRV is suitable or not; and
- suggested alternatives for TRVs, if needed.

### **CS TRVs**

The discussion for CS TRVs usually consists of four paragraphs. The first offers a summary of what the ideal TRV represents (i.e., the most protective value that best represents an ecologically relevant endpoint, exposure route and medium, exposure period, and effect level). The second paragraph is titled, "Data Set Considered for Selection of Value," and describes the contents of the data set from which the CS TRV was selected. The following information is presented in the second paragraph:

- number of references,
- number of experiments,
- number of endpoint types,
- types of measurement endpoint categories,
- test organisms represented,
- types of exposure media and routes,
- types of exposure duration categories, and
- types of effect levels.

The third paragraph in the discussion for CS TRVs is "Justification for Selection of Value." The value and effect level type of the PTV selected for use in development of the CS TRV are entered here as well as an explanation of why the PTV was selected over others in the data set. Usually, the highest NOAEL below the lowest LOAEL is selected for use, and this statement is entered. However, if this is not the case, an explanation is needed with further support as to why the TRV is still considered suitable. Some examples of further discussion supporting the selection of the PTV include the following: a comparison of the measurement endpoint the PTV represents to other measurement endpoints available in the data set, an explanation of the sensitivity of certain test organisms over others, and/or a comparison of the exposure conditions (e.g., length of exposure durations, exposures that occurred during critical life stages, *ad libitum* oral ingestion vs. scheduled feedings).

The fourth and final paragraph, "Description of Critical Study," provides more detail of the specific study from which the PTV was selected. The following information is provided:

- exposure length,
- whether exposure occurred during a critical life stage,
- chemical,
- chemical form,

- exposure medium,
- exposure route,
- test organism,
- dose or range of doses and units,
- whether doses were nominal (target) or empirical (verified/measured) concentrations,
- relationship of test exposure route to exposure route of concern,
- whether dose rate parameters (e.g., body weight, ingestion or inhalation rates) were provided or obtained from another source, and
- whether exposure concentrations were in dry or wet weight, and if in wet weight, the moisture basis and an explanation of the conversion to dry weight.

### Uncertainty Factor(s)

This field is left blank for GMM TRVs because UFs should already have been applied to PTVs to approximate chronic NOAEL- or NOEC-based effect levels used in the calculation. Rather, the statement “Prior to the calculation of the GMM TRV, the PTVs in the data set were extrapolated to chronic NOAEL-based effect levels by applying UFs.” is entered, and a table of applied UFs is provided in the Ecorisk Database. For CS TRVs, a brief explanation of whether UFs are needed or not is provided here. If UFs are needed, a brief description outlines the type (e.g., “A UF of 100 for extrapolation from an acute to a chronic exposure duration was applied.”). Table A-13 shows the UFs applied to approximate chronic NOAEL- or NOEC-based effect levels, or TRVs, from PTVs.

### Calculations

Essentially, the calculation for the GMM TRV ( $GMM\ TRV = \sqrt[n]{EL_1 * EL_2 * EL_3 * \dots * EL_n}$ ) should be entered here. However, because this exact equation cannot be entered in an Access field, the following description is entered instead, “GMM TRV = nth root of (EL1 x EL2 x EL3 x ... ELn) where n is greater than or equal to 3, and each effect level represents a chronic NOAEL-based effect level for an oral ingestion exposure for an ecologically relevant effect (i.e., reproduction or development, survival or adult body weight or size changes).”

For CS TRVs, if a UF is applied to the PTV to derive the TRV, this calculation is entered here [e.g., Chronic NOAEL = Chronic LOAEL(0.1)].

### Data Set Distribution

This field is not applicable for CS TRVs; N/A is entered. For GMM TRVs, the data set of chronic NOAEL- or NOEC-based effect levels is evaluated to determine the type of distribution (e.g., normal, positively skewed, negatively skewed, bimodal) and the variance of the distribution based on the number of GSDs from the GMM TRV. Also, any effect levels that may appear to be outliers are discussed (see the Geometric Standard Deviations and Outliers section below). Furthermore, the distribution is also evaluated for patterns or trends based on test organisms, exposure durations, original effect level types, or endpoint categories. Any observed trends are discussed.

### ***Types of Distributions***

If the distribution is negatively skewed, there are a larger number of higher values that most likely represent chronic or C-CL NOAELs/NOECs for ecologically relevant endpoints because no UFs are applied for exposure duration or effect level type; therefore, the GMM TRV is influenced by these higher values and is more likely to approximate a true NOAEL/NOEC. A negatively skewed distribution, in the context of a GMM TRV, is preferred because of this. On the other hand, if the GMM TRV is based on a positively skewed distribution, this means it is usually biased towards the lower values of the distribution and is therefore protective of the higher ones, which are usually associated with chronic or C-CL NOAELs/NOECs. For this reason, a positively skewed distribution is also acceptable because the GMM TRV is overly conservative as a result of the large number of lower values extrapolated from original effect levels other than chronic NOAELs/NOECs. If the distribution shows a bimodal pattern, this indicates there are two clusters of values according to test organisms, original effect levels, exposure durations, and/or endpoint categories. For example, there may be a large group of effect levels associated with acute and subchronic values and another large group of effect levels associated with chronic and C-CL values. It becomes difficult to determine if the GMM TRV is appropriate in this case. Revision of the GMM TRV to a subset GMM TRV may be preferred to represent the group of values that is more ecologically relevant (e.g., the chronic and C-CL values, which are more likely to represent more ecologically relevant endpoints such as reproduction/development effects).

### ***Geometric Standard Deviations and Outliers***

Because the TRV is based on a GMM of a minimum of three NOAEL/NOEC-based effect levels, the spread of data is assessed by calculating the GSD of the GMM TRV. GSDs and outliers are discussed in the assessment of data set distributions in order to (1) describe the variability of the data set, (2) outline any patterns associated with extreme values vs. those within 2 GSDs (e.g., outliers with high values may be associated with chronic durations because no UFs were applied, while values closer to the GMM were extrapolated from exposure durations and/or effect levels other than chronic NOAELs/NOECs with the application of UFs), and (3) provide support to the confidence rating of the GMM TRV where distributions with lower variance have higher confidence (i.e., GMM TRV is a better estimate of the NOAEL) vs. where distributions with higher variance have lower confidence. Some researchers consider any values beyond 2 standard deviations extreme values, or outliers (StatSoft Inc. 2005, 089447, Ref ID 1486). However, while outliers are described to be observations that do not exist within the characteristic distribution of the data, the decision to keep or remove an outlier often relies on professional judgment based on knowledge of the parameter being studied (Samuels 1989, 089450, Ref ID 1485; StatSoft Inc. 2005, 089447, Ref ID 1486). Therefore, in GMM TRV data sets, outliers are usable because they have been evaluated and screened using the same rigorous process as all other values derived using the PTSE process. All effect levels are based on PTVs derived from the PTSE process, and if a PTV was associated with a low confidence based on little or no supporting data, it was eliminated before the formulation of the data set used for the calculation of the GMM TRV. Furthermore, effect levels allowed in the data set that have larger values are often associated with chronic or C-CL PTVs, whereas the lower effect levels allowed in the data set were extrapolated from PTVs that were subchronic or acute NOAELs/NOECs, LOAELs/LOECs, or other effect levels (e.g., LD<sub>50</sub>s) with the use of UFs. The lower, extrapolated values are accepted in the GMM TRV data set because in screening-level ecological risk assessments, the use of a TRV that is conservative, rather than under-protective, is preferred (LANL 2004, 087630, Ref ID 1554). It is important to note that the nature of the data set distribution such as bimodality is evaluated for data sets with 10 or more chronic NOAEL (NOEC)-based effect levels, so for smaller data sets the reasonability of assessing true outliers is less.

**Lowest LOAEL or LOEC**

This field is not applicable for CS TRVs; N/A is entered. The GMM TRV is compared to the lowest chronic LOAEL- or LOEC-based effect level derived from the GMM TRV data set (see section A-4.1.5) to determine whether it is protective of the most sensitive endpoint in the data set. If the GMM TRV is below the lowest LOAEL- or LOEC-based effect level, it is protective of all possible effects in the data set. However, the GMM TRV may be much less than the LOAEL- or LOEC-based effect level, and some consideration must be taken into account to determine whether it is overly protective. On the other hand, if the GMM TRV is greater than the LOAEL- or LOEC-based effect level, further investigation is needed to determine if the GMM TRV may not be protective enough. Examples of information to examine include what endpoint the LOAEL/LOEC or LOAEL- or LOEC-based effect level represents, whether it is more or less ecologically relevant than other endpoints in the data set, if there are other similar endpoints available and how their effect levels compare to the GMM TRV, and what original effect level was used to approximate the LOAEL- or LOEC-based effect level. The application of UFs may have made the chronic LOAEL- or (LOEC)-based effect level overly conservative; therefore, the GMM TRV may still be protective even though it is above the LOAEL (LOEC)-based effect level. This is further strengthened if it can be shown that the GMM TRV includes more ecologically relevant endpoints and chronic exposure durations.

**LANL CS TRV Comparison**

This field is not applicable for CS TRVs; N/A is entered. The GMM TRV is compared to the LANL CS TRV if one is available for the same chemical, organism, and exposure route/medium scenario of concern. It is noted whether it is above or below the LANL CS TRV and by how much. Justification is provided for the continued use of the GMM TRV if it is deemed reasonable. If not, justification is provided for using the LANL CS TRV or an alternative TRV.

**ORNL CS TRV Comparison**

This field is not applicable for CS TRVs; N/A is entered. The GMM TRV is compared to the ORNL CS TRV. It is noted whether it is above or below the ORNL CS TRV and by how much. Justification is provided for the continued use of the GMM TRV if it is deemed reasonable. If not, justification is provided for using the LANL CS TRV or an alternative TRV.

**USEPA R6 CS TRV Comparison**

This field is not applicable for CS TRVs; N/A is entered. The GMM TRV is compared to the USEPA R6 CS TRV. It is noted whether it is above or below the USEPA R6 CS TRV and by how much. Justification is provided for the continued use of the GMM TRV if it is deemed reasonable. If not, justification is provided for using the LANL CS TRV or an alternative TRV.

**SNL CS TRV Comparison**

This field is not applicable for CS TRVs; N/A is entered. The GMM TRV is compared to the SNL CS TRV. It is noted whether it is above or below the SNL CS TRV and by how much. Justification is provided for the continued use of the GMM TRV if it is deemed reasonable. If not, justification is provided for using the LANL CS TRV or an alternative TRV.

### **LANL T&E CS TRV Comparison**

This field is not applicable for CS TRVs; N/A is entered. The GMM TRV is compared to the LANL T&E CS TRV. It is noted whether it is above or below the LANL T&E CS TRV and by how much. Justification is provided for the continued use of the GMM TRV if it is deemed reasonable. If not, justification is provided for using the LANL CS TRV or an alternative TRV.

Note: More comparisons of the LANL TRV to other published TRVs may become necessary if a LANL TRV is developed and there exists a TRV from another organization not mentioned above (e.g., USACHPPM TRVs). Comparison fields will be added should this situation arise.

### **Associated References**

A button is clicked to bring up a pop-up form for entry of Ref IDs cited in any of the fields above. First, the Part 3 Record ID is copied from the main data entry form and pasted into the Part 3 Record ID field of this new pop-up form. If references other than the primary toxicity study noted in the Primary Toxicity Paper Reference ID field are noted in the Discussion, Uncertainty Factor(s), Calculations, Data Set Distribution, Lowest LOAEL (LOEC) Comparison, or Other Published TRV Comparison fields, the Ref IDs for these are listed in the appropriate spaces. If no other references were mentioned, the default Ref ID is 0001.

### **A-5.0 PTSE PART 4, TRV APPROVAL**

After new GMM or CS TRVs are developed, the summary report Excel files containing the tables and graphs are sent to the EP Directorate's Risk Assessment Team for review. Based on their areas of knowledge and expertise, Risk Team members return comments, usually done in tracked-changes mode in the TRV summary report in Word, on TRV derivation methods, approximations of effect levels, chemical bioavailability, biological test organism or screening receptor information, etc. Sometimes their judgment may lead to an exception where a CS TRV may be used in spite of the availability of a GMM TRV. This may be done if the GMM TRV is judged to be under-protective of sensitive organisms to a particular chemical. Other times, Risk Team members may suggest a change from a GMM TRV to a subset GMM TRV, which is based on a subset of the original data set for a particular chemical, receptor group, and exposure scenario of concern, based on their knowledge of the behavior of that chemical with organisms in the wild under certain conditions. The PTSE reviewers consider the Risk Team comments and revise the information as appropriate. Documentation of any deviations is provided in the appropriate places in the PTSE Part 3 process (TRV summary report), especially in the discussion field.

### **A-6.0 REFERENCES**

*The following list includes all documents cited in this appendix. Parenthetical information following each reference provides the author(s), publication date, and ER ID. This information is also included in text citations. ER IDs are assigned by the Environmental Programs Directorate's Records Processing Facility (RPF) and are used to locate the document at the RPF and, where applicable, in the master reference set.*

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# **Attachment A-1**

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*GMM TRV Summary Report Example*



**TRV Summary Report****Ecorisk Database Release 2.1 (September 2004)**

\*TRV Summary ID: 118-96-7\_SOIL\_P\_TP\_Plant\_Soil\_ChronicGMMNOEC\_1442\_0001

GMM TRV ID: 118-96-7PGMM

LANL TRV: YES Data Source: LANL derived value based on reviewed primary data

Analyte Name: Trinitrotoluene[2,4,6-] Analyte Code: 118-96-7 Analyte Group: High Explosive

ESL Receptor Group: P Functional Group: A ESL Media: SOIL

Test Chemical Code: 118-96-7

Test Organism ID: TP Test Organism Common Name: Plant

Final TRV: Chronic GMM NOEC 62.1 mg/kg Exposure Route U\_SC+R

**Derivation Notes:** The GMM TRV for 2, 4, 6- trinitrotoluene in soil of plants is equal to a chronic NOEC of 62.1 mg/kg. This GMM TRV is derived from a data set of 12 PTVs representing 3 references, 12 experiments, 6 unique measurements, and 3 phylogenetic test organism orders. Endpoint categories included in the data set are reproduction and development. Six of 12 PTVs (50%) are associated with high confidence while the rest are associated with medium confidence. Only uptake via seed coat and/or roots exposure route studies were included in the GMM TRV data set; therefore, the test exposure route matches the exposure route of concern for soil ESLs for plants. See the PTVs Considered, Test Organisms, and Original Effect Level Types tables for more details of the data set.

**Conclusion:**

Based on the evaluation of the GMM TRV data set distribution and trends (see Data Set Distribution Comments section) and the comparison of the GMM TRV to the lowest chronic LOEC-based EL (see the Lowest LOEC Comparison section) and other published TRVs (see the Comparison of GMM TRV to other Published TRVs section), the confidence in the GMM TRV is medium because the data set contains: 1) 10 or more experiments, 2) only uptake via seed coat and/or roots which match the exposure route of concern for plant, soil-ESLs, 3) 3 or more test organism orders, 4) more than 3 unique measurements, 5) only R/D endpoints, 6) 2 or more chronic or C-CL NOEC/LOEC pairs, 7) ELs associated with a mixture of high and medium confidence ratings, and 8) no bimodality or other pattern that negatively biases the GMM TRV. Also, the comparison of the GMM TRV to other published TRVs (LANL CS TRV and SNL CS TRV) is acceptable because it is lower than the LANL CS TRV, higher than the SNL CS TRV by only a factor of 2.1, and represents more supporting data than both CS TRVs. The confidence rating was lowered from high to medium because: 1) the GMM TRV is higher than the lowest chronic LOEC-based EL by a factor of 3 or more, and 2) greater than 75% of the ELs were more than 10 GSDs from the GMM TRV, indicating a moderately high variance for the distribution. The lowest chronic-LOEC based EL represents a study in which barley was exposed to TNT in forest soil, which may hold different soil properties than soil exposure media in other studies of this data set (e.g., artificial soil, soil collected from experimental field in Germany). The forest soil has a pH of 7.6, which is within the range of soil values at LANL (5.2 to 8.2; Ref ID 1380). Furthermore, the GMM TRV for different soil properties minimizes the chance that the value can be over or under conservative. Also, the moderately high variance is overridden by the fact that the GMM TRV is protective of the majority (8 of 12) of R/D endpoints in the data set. See the GMM TRV Confidence Rating table for details. In conclusion, the GMM TRV is considered protective of plant populations and the more sensitive individuals of threatened and endangered species because it considers multiple ecologically relevant endpoints and thus provides a more comprehensive TRV than a single CS TRV.

**Uncertainty:** Prior to the calculation of the GMM TRV, the PTVs in the data set were extrapolated to chronic NOEC-based ELs by applying UFs.

**Calculations:**  $GMM\ TRV = \sqrt[n]{(EL1 \times EL2 \times EL3 \times \dots \times ELn)}$  where n is greater than or equal to 3 and each EL represents a chronic NOEC-based EL for a seed coat and/or root uptake via soil exposure for an ecologically relevant effect (i.e., reproduction or development, survival, or mature plant weight or size changes).

Log Kow: KocVu: Foc:

Text Last Updated On: 10-Sep-04 Value Last Updated On: 20-Aug-04

Confidence Rating: Medium NMED Concurrence Date:

\* Further details on the study/ effects/ toxicity values reviewed for this TRV are provided in the PTSE Part 1 (Study Details) and 2s (Study Evaluations) and in the Part 3 (TRV Summary) graph.

**TRV Summary Report****Ecorisk Database Release 2.1 (September 2004)**

\*TRV Summary ID: 118-96-7\_SOIL\_P\_TP\_Plant\_Soil\_ChronicGMMNOEC\_1442\_0001

GMM TRV ID: 118-96-7PGMM

LANL TRV: YES

Data Source: LANL derived value based on reviewed primary data

**Data Set Distribution Comments:**

The distribution of chronic NOEC-based ELs ranging from 5.59 to 355 mg/kg is positively skewed. One of the 12 ELs (8%) is within 2 GSDs, 3 (25%) are between 2 and 6 GSDs, 2 (17%) are between 6 and 10 GSDs, and the rest extend out to 71 GSDs from the GMM TRV, indicating moderately high variance. All but one of the chronic NOEC-based ELs are considered to be outliers (extreme values, or values beyond 2 GSDs), yet they are still usable because the high GSDs indicate a larger spread of data rather than errors in the values. The moderately high variance indicates that the GMM TRV may not as closely approximate the true TRV as one with a lower variability in its data set would. No bimodality was present in the data set distribution. It was observed that the test species in the Order Capparales (cress and turnip) had a narrow range in their chronic NOEC-based values (24-49 mg/kg) compared to Order Cyperales (barley, oat, wheat, yellow nutsedge) values which ranged from 5.59 to 355 mg/kg. Lettuce was the only test species present for Order Asterales. Original effect level types may have also played a role in the 2 lowest chronic NOEC-based values in Order Cyperales (and the data set) because UF<sub>s</sub> of 10 were applied to C-CL LOECs to extrapolate them to NOEC-based ELs. Patterns could not be evaluated for endpoint category or exposure duration because all chronic NOEC-based ELs represent C-CL values for R/D endpoints. The GMM TRV is below 42% of NOEC-based ELs. However, it's below 67% LOEC-based ELs (see the Lowest LOEC Comparison section), so it is still protective of the majority of endpoints.

Based on the evaluation of the distribution of the GMM TRV data set of chronic NOEC-based ELs, the GMM TRV is suitable because 1) it is based on a positively skewed distribution, and 2) and 3) it represents a variety of test species with different sensitivities and is protective of the majority of the data set because it is lower than 67% of the LOEC-based ELs. See the Graph of NOEC-based ELs for details.

**Lowest LOAEL (LOEC) Comparison:**

The range of chronic LOEC-based ELs is 13.66 to 461.5 mg/kg. The GMM TRV is above the lowest chronic LOEC-based EL (13.66 mg/kg) by a factor of 4.5. The lowest chronic LOEC-based EL is based on a C-CL LOEC for an R/D endpoint. The lowest chronic LOEC-based EL is based on barley exposure in forest soil, whereas in the other barley study, barley is exposed to TNT in artificial soil. The chronic LOEC-based EL for the barley exposure in artificial soil is also below the GMM TRV but by a factor of only 1.1, indicating that barley may be less sensitive in artificial soil. The GMM TRV is also above 2 other chronic LOEC-based ELs representing C-CL LOECs for R/D endpoints. These chronic LOEC-based ELs represent exposure to cress and turnip test species via soil collected from an experimental field at a biological station in Berlin, Germany. There are two other studies using the cress and turnip species as well, but they use a different type of soil that was provided by a Germany company. Therefore, the lower sensitivities the cress and turnip in the soil collected from the biological station may be due to the soil properties (e.g., pH, organic matter content). These 2 types of Germany soils were also used in studies for oat and wheat test species, but these plants were less sensitive; 3 of 4 chronic LOEC-based ELs were derived from C-CL NOECs for R/D endpoints, indicating that no adverse effects were observed at the highest concentration administered in the study and that the chronic LOEC-based ELs may be overly conservative due to the application of test organism specific LOEC/NOEC factors (Ref ID 1487) to extrapolate the LOECs from the NOECs. Still, the GMM TRV is protective of these 3 chronic LOEC-based ELs as well as the 4th one which is based on a C-CL LOEC for an R/D endpoint. The GMM TRV is also below the remaining 4 chronic LOEC-based ELs which are based on C-CL NOECs (1) and LOECs (3) for R/D endpoints. Although the GMM TRV is below 4 chronic LOEC-based ELs, it is protective of the majority of the data set (67%) which contains a variety of test species and soil types. See the Graph of LOEC-based ELs for details.

**LANL CS TRV Comparison:**

The GMM TRV is lower than the LANL CS TRV (80 mg/kg) by a factor of 4. This CS TRV is based on a chronic NOEC for a WC endpoint (PTV ID 0379\_118-96-7\_1A) and is included in the data set for the GMM TRV. The LANL CS TRV represents effects on yield (as above-ground plant material) of yellow nutsedge. This endpoint was selected for the CS TRV because at the time, it was the only endpoint available in a data set of 1 reference and 1 experiment. More data was obtained, leading to the derivation of a GMM

**TRV Summary Report**

**Ecorisk Database Release 2.1 (September 2004)**

\*TRV Summary ID: 118-96-7\_SOIL\_P\_TP\_Plant\_Soil\_ChronicGMMNOEC\_1442\_0001

GMM TRV ID: 118-96-7PGMM

LANL TRV: YES

Data Source: LANL derived value based on reviewed primary data

TRV.

ORNL CS TRV Comparison:

ORNL does not have a CS TRV available for comparison.

USEPA R6 CS TRV Comparison:

USEPA R6 does not have a CS TRV available for comparison.

SNL CS TRV Comparison:

The GMM TRV is higher than the SNL CS TRV (30 mg/kg) by a factor of only 2.1. This CS TRV is based on a LOAEL for growth effects on blando brome grass in soil. No UFs were applied. The endpoint that the SNL CS TRV represents is not included in the GMM TRV data set because a hard copy of the reference (Ref ID 0463) could not be located at the time.

**TRV Summary Report**

**Ecorisk Database Release 2.1 (September 2004)**

\*TRV Summary ID: 118-96-7\_SOIL\_P\_TP\_Plant\_Soil\_ChronicGMMNOEC\_1442\_0001

GMM TRV ID: 118-96-7PGMM

LANL TRV: YES

Data Source: LANL derived value based on reviewed primary data

**REFERENCE LIST**

**Ref ID Citation**

Primary Toxicity Study Reference CS TRV: 0001 NOT APPLICABLE

Primary Toxicity Study Reference(s) GMM TRV: (NOT APPLICABLE, if no references are listed in this section)

0379	Pennington, JC. 1988. Soil Sorption and Plant Uptake of 2,4,6-Trinitrotoluene. AD A200 502. Technical Report EL-88-12. US Army Biomedical Research and Development Laboratory, Fort Detrick, Frederick, MD.
1455	Robidoux, PY, G Bardai, L Paquet, G Ampleman, S Thiboutot, J Hawari, and GI Sunahara. 2003. Phytotoxicity of 2,4,6-Trinitrotoluene (TNT) and Octahydro-1,3,5,7-Tetranitro-1,3,5,7-Tetrazocine (HMX) in Spiked Artificial and Natural Forest Soils. Arch. Environ. Contam. Toxicol., 44: 198-209.
1459	Gong, P, B-M Wilke, and S Fleischmann. 1999. Soil-Based Phytotoxicity of 2,4,6-Trinitrotoluene (TNT) to Terrestrial Higher Plants. Arch. Environ. Contam. Toxicol., 36: 152-157.
<b>TRV reference:</b>	1442 Los Alamos National Laboratory (LANL), 2004 (Sept.). ECORISK Database (Release 2.1). LA-UR-04-7304. RRES-R package #186, ER ID 87386. Risk Reduction and Environmental Stewardship Remediation Service Program, Los Alamos National Laboratory, Los Alamos, NM.
<b>Additional References:</b>	0463 Cataldo, DA, SD Harvey, RJ Fellows, et al. 1989. An evaluation of environmental fate behavior of munitions material (TNT, RDX) in soil and plant systems. PNL-7370: AD-A223 5446. US Army Medical Research and Development Command, Fort Detrick, Frederick, MD.
1380	Longmire, PA, SL Reneau, PM Watt, LD McFadden, JN Gardner, CJ Duffy, and RT Rytli. 1996 (May). Natural Background Geochemistry, Geomorphology, and Pedogenesis of Selected Soil Profiles and Bandelier Tuff, Los Alamos, New Mexico. Los Alamos National Laboratory Report LA-12913-MS. Los Alamos, New Mexico. Pages 21-33
1487	Newell, PG, and JS Podolsky. 2004. PTSE Methods (Draft). Risk Reduction and Environmental Stewardship Remediation Services, Los Alamos National Laboratory, Los Alamos, New Mexico.
0001	NOT APPLICABLE
0001	NOT APPLICABLE

\*\* Citations for up to 5 additional references associated with this TRV are listed. If the Ref ID for one or more additional references is 0001 that indicates that there are not any or anymore references associated with the TRV.

# **Attachment B-1**

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*CS TRV Summary Report Example*



**TRV Summary Report****Ecorisk Database Release 2.1 (September 2004)**

\*TRV Summary ID: 11097-69-1\_SOIL\_B\_TB\_ChickenWhiteLeghorn\_Diet\_ChronicNOAEL\_1105\_0756

GMM TRV ID:

LANL TRV: YES

Data Source: LANL derived value based on reviewed primary data

Analyte Name: Aroclor-1254	Analyte Code: 11097-69-1	Analyte Group: Polychlorinated Biphenyl
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ESL Receptor Group: B	Functional Group: A	ESL Media: SOIL
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Test Chemical Code: 11097-69-1

Test Organism ID: TB	Test Organism Common Name: Chicken, White Leghorn
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Final TRV: Chronic CS NOAEL 0.1	mg/kg/d	Exposure Route OD
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**Derivation Notes:** The chronic NOAEL of 0.1 with an accompanying chronic LOAEL of 1 was derived from a primary toxicity value (PTV) selected from a data set of 3 references and 8 effects (4 reproduction/development, 2 survival, and 1 growth). Effects considered in the selection included adult and chick mortality, adult and chick body weight, egg production, and hatchability. The PTV chosen for the derivation of the toxicity reference value (TRV) is from Ref ID 0756 and is based on hatchability (Experiment Effect ID (0756\_11097-69-1\_1A). Mortality in Ref ID 0707 was eliminated from consideration because it was from a study in which only high-dose, relatively short-term (5 day) exposures were evaluated. The other study (Ref ID 0758) reported adverse results (LOAEL) at 2.63 mg/kg/d, and with conversion to NOAEL, this would produce a value of 0.263 mg/kg/d which is close to the value selected for the TRV. The 0.1 mg/kg/d TRV is considered protective of wildlife populations because hatchability is an indicator of the ability of the species to successfully reproduce. Poor reproduction leads to lower success of breeding and less individuals to maintain a viable population. The NOAEL and LOAEL were based on two concentrations (unknown whether they were nominal or empirical) administered.

In this chronic (9 weeks and during a critical life stage) study, Aroclor 1242 was administered orally through food to white leghorn chicken. This test exposure route is related to the exposure route of concern for soil ESLs (food web transfer through consumption of contaminated plants and/or animals and incidental ingestion of soil) because both are oral through the diet. Dose rates were not reported in mg/kg/d, and body weight and food intake data were not available in the primary study; therefore, these parameters had to be obtained from other sources. The moisture basis of the dose is unknown, but will be considered dry weight for conservatism.

**Uncertainty:** Because the exposure was chronic during a critical life stage and the TRV is based on a no observed adverse effects level, the application of an Uncertainty Factor is unnecessary.

**Calculations:** N/A

Log Kow:	KocVu:	Foc:
----------	--------	------

Text Last Updated On: 10-Sep-04	Value Last Updated On: 28-Sep-01
---------------------------------	----------------------------------

Confidence Rating:	NMED Concurrence Date:
--------------------	------------------------

\* Further details on the study/ effects/ toxicity values reviewed for this TRV are provided in the PTSE Part 1 (Study Details) and 2s (Study Evaluations) and in the Part 3 (TRV Summary) graph.

**TRV Summary Report**

***Ecorisk Database Release 2.1 (September 2004)***

\*TRV Summary ID: 11097-69-1\_SOIL\_B\_TB\_ChickenWhiteLeghorn\_Diet\_ChronicNOAEL\_1105\_0756

GMM TRV ID:

LANL TRV: YES

Data Source: LANL derived value based on reviewed primary data

---

Data Set Distribution Comments:

Lowest LOAEL (LOEC) Comparison:

LANL CS TRV Comparison:

ORNL CS TRV Comparison:

USEPA R6 CS TRV Comparison:

SNL CS TRV Comparison:

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**TRV Summary Report**

**Ecorisk Database Release 2.1 (September 2004)**

\*TRV Summary ID: 11097-69-1\_SOIL\_B\_TB\_ChickenWhiteLeghorn\_Diet\_ChronicNOAEL\_1105\_0756

GMM TRV ID:

LANL TRV: YES

Data Source: LANL derived value based on reviewed primary data

**REFERENCE LIST**

<u>Ref ID</u>	<u>Citation</u>
0756	Cecil, HC, J Bitman, RJ Lillie, GF Fries and J Verrett. 1974. Embryotoxic and Teratogenic Effects in Unhatched Fertile Eggs for Hens Fed PCBs. Bull Environ Contam Toxicol 11(6):489-495.

Primary Toxicity Study Reference(s) GMM TRV: (NOT APPLICABLE, if no references are listed in this section)

TRV reference:	1105	Los Alamos National Laboratory (LANL), 2001 (Sept). ECORISK Database (Release 1.3), ER package #186. Environmental Restoration Project, Los Alamos National Laboratory, Los Alamos, NM.
Additional References:	0707	Hill, EF and MB Camardese. 1986. Lethal Dietary Toxicities of Environmental Contaminants and Pesticides to Coturnix. United States Fish And Wildlife Service: Fish and Wildlife Tech Rep 2 (NTIS PB86-176914). Laurel, MD. 154 pp.
	0758	Dahlgren, RB, RL Linder, and CW Carlson. 1972. Polychlorinated Biphenyls: Their Effects on Pened Pheasants. Environ Health Perspec 1:89-101.
	0001	NOT APPLICABLE
	0001	NOT APPLICABLE
	0001	NOT APPLICABLE

\*\* Citations for up to 5 additional references associated with this TRV are listed. If the Ref ID for one or more additional references is 0001 that indicates that there are not any or anymore references associated with the TRV.



**Los Alamos National Laboratory**  
**Environmental Stewardship Division**  
**Environmental Remediation and Surveillance**  
**Program**

**Ecorisk Database Release 2.2 (September 2005)**

**TRV DEVELOPMENT METHODS:  
USING THE GEOMETRIC MEAN**

**DRAFT**

**September 1, 2004**

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## Introduction

The ideal GMM TRV\* for ecological risk screening assessments is one that is based on a data set representing the most ecologically relevant endpoints (reproduction/development), exposure routes (birds and mammals: oral ingestion via food or drinking water; mammals: inhalation; plants: uptake via seed coat and/or roots; or invertebrates: oral and dermal exposure), exposure media (birds and mammals: food, drinking water, air; plants and invertebrates: soil), exposure period (chronic), and effect levels (NOAEL for birds and mammals or NOEC for plants and invertebrates). A GMM TRV based on these characteristics is protective of wildlife, plant, and invertebrate populations and sensitive individuals because it represents an exposure that is not associated with adverse impacts of low-level, long-term chemical effects (i.e., adverse effects on ability of individuals to develop into viable organisms, search for mates, breed successfully, and produce live and equally viable offspring).

The GMM TRV is derived from a data set of chronic NOAEL (NOEC)-based ELs (see the [Deriving chronic NOAEL \(NOEC\)-based ELs](#) section). The suitability of the GMM TRV for the chemical and ecological screening receptor of concern is determined by the following assessments (1) examining the distribution of its data set (see the [Assessing the distribution of the GMM TRV data set](#) section), (2) comparing it to the lowest chronic LOAEL (LOEC)-based EL derived from the GMM TRV data set (see the [Comparing the GMM TRV to the LOAEL \(LOEC\)-based ELs](#) section), and (3) comparing it to TRVs derived by other organizations, if available (see the [Comparing the GMM TRV to other Published TRVs](#) section). The information gathered from the assessments is used to assign the GMM TRV a confidence rating by scoring various aspects of each of the assessments on how well they meet the ideal GMM TRV criteria (see the [Scoring Criteria and Confidence Ratings](#) section). Summary tables and graphs are provided in the specific GMM TRV Summary reports for easier distillation of information contained within the GMM TRV data set (see the [Tables and Graphs for GMM TRV Data Set Information](#) section).

\*See the **Acronyms** table ([Table 5](#)) for definitions of all acronyms and abbreviations.

## Deriving chronic NOAEL (NOEC)-based ELs

The data set used to calculate the GMM TRV (see the **Calculations** section in the specific GMM TRV Summary Report) contains a variety of effect levels (PTVs derived from the PTSE Process as described in Ref ID 1487) ranging from chronic NOAEL (NOEC)/LOAEL (LOEC) pairs to acute, other effect levels such as LC50s or EC20s. The GMM TRV is calculated using chronic NOAEL (NOEC)-based ELs that are either chronic NOAELs (NOECs) or derived from other effect levels. If the effect level (PTV) is an acute or subchronic NOAEL (NOEC), it is extrapolated to a chronic NOAEL (NOEC)-based EL with the application of a UF. If the PTV is a LOAEL (LOEC) or other EL (LC50), it is first extrapolated to a NOAEL with the application of a UF, and then it is extrapolated to a chronic exposure duration if needed. See the **UFs Description** table ([Table 4](#)).

## Assessing the distribution of the GMM TRV data set

The data set of chronic NOAEL (NOEC)-based ELs is evaluated to determine the type of distribution (e.g., normal, positively skewed, negatively skewed, bimodal) and the variance of the distribution based on the number of GSDs from the GMM. Also, any ELs that may appear to be outliers are discussed (see the [Geometric Standard Deviations and Outliers](#) section). The

distribution is also evaluated for patterns or trends based on test organisms, exposure durations, original effect level types, or endpoint categories. The distribution and patterns of the ELs in the GMM TRV data set are discussed in the **Data Set Distribution Comments** section and presented in the **Graph of NOAEL-based ELs**, both which can be found in the specific GMM TRV Summary Report. The experiment details such as exposure duration, test organism, and original effect level are presented in the **PTVs Considered** table in the specific GMM TRV Summary Report.

#### *Types of Distributions*

If the distribution is negatively skewed, there are a larger number of higher values which most likely represent chronic or C-CL NOAELs (NOECs) for ecologically relevant endpoints; therefore, the GMM TRV is influenced by them and more likely to approximate a true NOAEL (NOEC). A negatively skewed distribution, in the purpose of a GMM TRV, is preferred because of this. On the other hand, if the GMM TRV belongs to a positively skewed distribution, this means it is usually biased towards the lower values of the distribution and is therefore protective of the higher ones which are usually associated with chronic or C-CL NOAELs (NOECs). For this reason, a positively skewed distribution is also acceptable because the GMM TRV is overly conservative due to the large number of lower values extrapolated from original effect levels other than chronic NOAELs (NOECs). If the distribution shows a bimodal pattern, this indicates there are two clusters of values according to test organisms, original effect levels, exposure durations, and/or endpoint categories. For example, there may be a large group of ELs associated with acute and subchronic values and another large group of ELs associated with chronic and C-CL values. It becomes difficult to determine if the GMM TRV is appropriate in this case. Revision of the GMM TRV to a subset GMM TRV may be preferred in order to represent the group of values that is more ecologically relevant (e.g., the chronic and C-CL values, which are more likely to represent more ecologically relevant endpoints such as R/D effects).

#### *Geometric Standard Deviations and Outliers*

Because the TRV is based on a geometric mean, the spread of data is assessed by calculating the GSD of the GMM TRV. GSDs and outliers are discussed in the assessment of data set distributions in order to 1) describe the variability of the data set, 2) outline any patterns associated with extreme values vs. those within 2 GSDs (e.g., outliers with high values may be associated with chronic durations while those values closer to the GMM are values extrapolated from original PTVs), and 3) provide support to the confidence rating of the GMM TRV where distributions with lower variance have higher confidence (i.e., GMM TRV is a better estimate of the no observed adverse effect level) vs. distributions having higher variance have lower confidence. Some researchers consider any values beyond 2 standard deviations extreme values, or outliers (Ref ID 1486). However, while outliers are described to be observations that do not exist within the characteristic distribution of the data, the decision to keep or remove an outlier often relies on professional judgment based on knowledge of the parameter being studied (Ref IDs 1485 and 1486). Therefore, in GMM TRV data sets, outliers are usable because they have been evaluated and screened in the same rigorous process as all other values derived via the PTSE process (Ref ID 1487). All ELs are based on PTVs derived from the PTSE Process and if a PTV was associated with a low confidence based on lack of or little supporting data, it was eliminated prior to the formulation of the data set used for the calculation of the GMM TRV.

Furthermore, ELs allowed in the data set that have larger values are often associated with chronic or C-CL PTVs whereas the lower ELs allowed in the data set were extrapolated from PTVs that were subchronic or acute NOAELs (NOECs), LOAELs (LOECs), or Other ELs (e.g., LD50s) with the use of UFs. The lower, extrapolated values are accepted in the GMM TRV data set because in screening level ecological risk assessments the use of a TRV that is overly protective, rather than under protective is preferred.

### **Comparing the GMM TRV to the LOAEL (LOEC)-based ELs**

The GMM TRV is compared to the lowest chronic LOAEL (LOEC)-based EL derived from the GMM TRV data set (see the [Deriving chronic LOAEL \(LOEC\)-based ELs](#) section) in order to determine whether it may or may not be protective of the most sensitive endpoint in the data set. If it is below the lowest LOAEL (LOEC)-based EL, it is protective of all possible effects in the data set. However, it may be too far below the LOAEL (LOEC)-based EL and some consideration must be taken into account to determine whether or not it is overly protective. On the other hand, if the GMM TRV is above the LOAEL (LOEC)-based EL, further investigation is needed to determine if the GMM TRV may not be protective enough. Examples of information to examine include what endpoint the LOAEL (LOEC) or LOAEL (LOEC)-based EL represents, whether it is more or less ecologically relevant than other endpoints in the data set, if there are other similar endpoints available and how their ELs compare to the GMM TRV, and what original effect level was used to approximate the LOAEL (LOEC)-based EL. The application of UFs may have made the chronic LOAEL (LOEC)-based EL overly conservative; therefore, the GMM TRV may still be protective even though it is above the LOAEL (LOEC)-based EL. This is further strengthened if it can be shown that the GMM TRV includes more ecologically relevant endpoints and chronic exposure durations. The comparison is discussed in the **Lowest LOAEL (LOEC) Comparison** section and presented in the **Graph of LOAEL (LOEC)-based ELs**, both which can be found in the specific GMM TRV Summary Report. Experiment details such as exposure duration, test organism, and original effect level are presented in the **PTVs Considered** table in the specific GMM TRV Summary Report.

#### *Deriving chronic LOAEL (LOEC)-based ELs*

The data set may contain a variety of effect levels (PTVs derived from the PTSE Process as described in Ref ID 1487) ranging from chronic NOAEL (NOEC)/LOAEL (LOEC) pairs to acute, other effect levels such as LC50s or EC20s. The extrapolation of NOAELs (NOECs) and other effect levels to chronic LOAEL (LOEC)-based ELs is done to maximize the number of data points that can be compared to the GMM TRV. If the original effect level (PTV) is not a chronic LOAEL (LOEC), it is extrapolated to a chronic LOAEL (LOEC)-based EL by applying standard UFs (see [Table 4](#)) or NOAEL (NOEC) to LOAEL (LOEC) ratios that are specific to the exposure scenario of concern (i.e., exposure route, exposure medium, and ecological receptor) See [Explanation of LOAEL \(LOEC\)/NOAEL \(NOEC\) Ratios](#)

#### *Explanation of LOAEL (LOEC)/NOAEL (NOEC) Ratios*

If only a chronic NOAEL (NOEC) or chronic NOAEL (NOEC)-based EL extrapolated from an acute or subchronic NOAEL (NOEC) is available, a factor must be applied to derive a LOAEL (LOEC)-based EL from this value. Based on Dourson and Stara (1983; Ref ID 1379), 96% of the ratios between NOAELs (NOECs) and LOAELs (LOECs) for mammals in oral ingestion experiments have values of 5 or less (page 232 and figure 4). However, because this data is only

applicable to oral ingestion exposure in mammals, ratios for the remaining exposure pathways (oral ingestion in birds, oral ingestion and dermal contact in earthworms, uptake via seed coats and/or roots, and inhalation in mammals) were determined from NOAEL (NOEC)/LOAEL (LOEC) pairs specific to each of the exposure pathways. The data used to develop the ratios is from the Ecorisk Database. The smallest and largest ratios developed for each exposure pathway were used to approximate a minimum and maximum LOAEL (LOEC)-based EL, in order to bracket a range of concentrations at which the adverse effects may first be observed. The **PTVs Considered** table in the specific GMM TRV Summary Report presents this data.

### **Comparing the GMM TRV to other Published TRVs**

The GMM TRV is compared to TRVs from other organizations in order to present all available data and to ensure the validity of the GMM TRV in light of this other data. See **LANL CS TRV**, **ORNL CS TRV**, **USEPA R6 CS TRV**, and **SNL CS TRV Comparison** sections in the specific GMM TRV Summary Report.

### **Scoring Criteria and Confidence Ratings**

A confidence rating for the GMM TRV indicates how well the GMM TRV represents the ideal GMM TRV, which is representative of the true TRV. The true TRV is the dose rate or concentration that is equivalent to a no adverse effect level for population level effects (i.e., decreased population size) for a particular receptor under a specific exposure scenario to a particular chemical in the real world. The confidence rating for the GMM TRV is based on how well the GMM TRV meets various criteria within specific evaluation categories. A weighted scoring system based on the degree of influence each evaluation category has on the GMM TRV is used to assess the validity of the GMM TRV for estimating the true TRV. The following sections describe the structure of the confidence rating system including descriptions and justifications for the evaluation processes used to assign the confidence ratings.

#### *Confidence Rating System Structure*

The first step in assigning a confidence rating to a GMM TRV is to assign a score for each evaluation category listed below. Each evaluation category contains individual criterion associated with ranked scores that reflect how well the GMM TRV data set being evaluated represents the characteristics of the ideal GMM TRV. The higher the score, the better the GMM TRV represents the ideal GMM TRV and thus the true TRV. The possible scores for each evaluation category are presented in *GMM TRV Scoring Criteria\_R6b.xls* and the justifications for the scores are presented in the [\*Justification for Scoring and Weighting Factor Levels\*](#) section.

#### Data Set Evaluation Categories:

- 1) Number of Experiments
- 2) Type of Exposure Medium
- 3) Number of Test Organism Orders
- 4) Number of Unique Measurements (Endpoints)
- 5) Type of Endpoint Category
- 6) Number and Type of Effect Levels
- 7) Confidence Rating of PTVs Associated with Individual NOAEL (NOEC)-based ELs in GMM TRV Data Set
- 8) Outlier(s) in Chronic NOAEL (NOEC)-based EL Distribution

- 9) Chronic NOAEL (NOEC)-based EL Distribution is Bimodal
- 10) Relationship of GMM TRV to Chronic LOAEL (LOEC)-based ELs
- 11) Relationship of GMM TRV to Other Published TRVs.

The second step in assigning a confidence rating to a GMM TRV is to calculate a weighted score for each evaluation category by multiplying the individual scores of each evaluation category by the weighting factor of the evaluation category. The weighted score for each evaluation category is based on the weighting factor level assigned to the evaluation category. The weighting factor level is based on the degree of influence the evaluation category has on setting the GMM TRV. The higher the weighting factor, the greater the influence the evaluation category has on setting the GMM TRV. The possible weighting factor levels are presented in [Table 1](#).

**Table 1. Weighting Factor Levels.**

<b>Weighting Factor Level</b>	<b>Definition</b>	<b>Weighting Factor Applied</b>
Critical	A low score for a critical evaluation category triggers re-investigation of the GMM TRV and possible revision or decision not to use.	2
Non-critical	A high score for a non-critical evaluation category indicates the GMM TRV data set is very robust, highly relevant to the scenario for which the TRV is being developed, or is based primarily on ELs that were not derived by applying UFs to PTVs. A low score rarely influences revision of GMM TRV because it is an added benefit if the evaluation category scores high, but not a requirement.	1

The third step in assigning a confidence rating to a GMM TRV is to calculate a total weighted score for the GMM TRV being evaluated. The total weighted score is equal to the sum of weighted scores of all 11 evaluation categories. The weighting factor levels assigned to each evaluation category are presented in [Table 2](#) and the justifications for them are presented in the [Justification for Scoring and Weighting Factor Levels](#) section.

**Table 2. Weighting Factor Levels Assigned to Evaluation Categories.**

<b>Evaluation Category</b>	<b>Weighting Factor Level</b>
Number of Experiments	Non-critical
Type of Exposure Medium	Non-critical
Number of Test Organisms Orders	Non-critical
Number of Unique Measurements (Endpoints)	Non-critical
Type of Endpoint Category	Non-critical
Number and Type of Effect Levels	Non-critical

Confidence Rating of PTVs Associated with Individual NOAEL (NOEC)-based ELs in GMM TRV Data Set	Non-critical
Outlier(s) in Chronic NOAEL (NOEC)-based EL Distribution	Critical
Chronic NOAEL (NOEC)-based EL Distribution is Bimodal	Critical
Relationship of GMM TRV to Chronic LOAEL (LOEC)-based ELs	Critical
Relationship of GMM TRV to Other Published TRVs.	Critical

The fourth step in assigning a confidence rating to a GMM TRV is to determine the percentage the total weighted score is of the maximum total weighted score for the evaluation (i.e., 36.5 points based on summing the highest scores from each evaluation category). The percentage the total weighted score is of the maximum total weighted score is the ultimate basis for assigning the confidence rating of a GMM TRV. [Table 3](#) presents the possible confidence ratings and the corresponding percentage of the maximum total weighted score and the equivalent total weighted score.

**Table 3. Confidence Ratings.**

Confidence Rating	% of Maximum Total Weighted Score (%MTWS)	Equivalent Total Weighted Score (ETWS)
High	$\%MTWS \geq 75\%$	$27.375 \leq ETWS \leq 36.5$
Medium	$50\% \leq \%MTWS < 75\%$	$18.25 \leq ETWS < 27.375$
Low	$25\% \leq \%MTWS < 50\%$	$9.125 < ETWS < 18.25$
Unacceptable	$\%MTWS \leq 25\%$	$ETWS \leq 9.125$

#### *Justification for Scoring and Weighting Factor Levels*

The following sections provide the justification for the scoring criteria and weighting factor levels of each evaluation category.

#### Number of Experiments

##### Justification for Scoring:

The preference is to have a high number of experiments because this reduces the potential for the data set to be biased toward a particular study design. Based on best professional judgment, 10 experiments are considered to provide a more than adequate representation of the toxicity of a chemical for the test organism group of concern. Four to 9 experiments are considered to provide an adequate representation while 3 or fewer experiments are considered to provide a minimal representation of the toxicity of a chemical for the test organism group of concern.

##### Justification for Weighting Factor Level:

This evaluation category is given a **Non-critical** weighting factor level. This evaluation category has a strong relationship to the robustness of the data set and its ability to represent the ideal GMM TRV; thus the true TRV is estimated. The higher the number of experiments, the more robust the data set. This evaluation category is not, however, a

primary factor for determining whether or not the GMM TRV should be used because a high number of experiments in the data set is not a requirement, but rather an additional benefit for assessing confidence in the TRV.

#### Type of Exposure Medium

##### Justification for Scoring:

The preference is for all the ELs in the data set to be associated with an exposure medium that is equivalent to the exposure medium of concern. However, if the data set is limited (i.e., less than 3 ELs for a particular exposure medium), ELs that have an appropriate surrogate exposure medium (i.e., exposure medium that has the same exposure route as the exposure route of concern) may be used to supplement the data set so that a GMM TRV can be derived. For example, for an oral ingestion via food TRV only food ELs should be used, but if the data set is limited, oral ingestion via drinking water ELs may be used to supplement the data set so that a GMM TRV may be calculated.

##### Justification for Weighting Factor Level:

This category is given a **Non-critical** weighting factor level. This evaluation category indicates the degree of relevance the data set has to the TRV that is being developed. The higher the degree of relevance, the more closely the GMM TRV represents the ideal GMM TRV; thus the true TRV is estimated. This evaluation category is not, however, a primary factor for determining whether or not the GMM TRV should be used because an exact match on the exposure medium for which the TRV is being developed is not a requirement, but rather an additional benefit for assessing confidence in the TRV. Only the exposure route must match the exposure for which the TRV is being developed. However, the toxicity can vary greatly in different exposure media due to the differences in bioavailability of the chemical in one compared to the other. Therefore, a complete match on the exposure medium is preferred to be able to more accurately estimate the true TRV.

#### Number of Test Organism Orders

##### Justification for Scoring:

The preference is to have a high number of test organism orders because this reduces the potential for the data set to be biased toward one order of test organisms. The scoring criteria are based upon the USACHPPM guidance that states that having at least 2 different taxonomic orders in a TRV data set helps define the quality of the data set (Ref ID 1481).

##### Justification for Weighting Factor Level:

This category is given a **Non-critical** weighting factor level. This evaluation category has a strong relationship to the robustness of the data set and its ability to represent the ideal GMM TRV; thus the true TRV is estimated. The higher the number of test organism orders, the more robust the data set. This evaluation category is not, however, a primary factor for determining whether or not the GMM TRV should be used because a high number of test organism orders in the data set is not a requirement, but rather an additional benefit for assessing confidence in the TRV.

## Number of Unique Measurements (Endpoints)

### Justification for Scoring:

The preference is to have a high number of unique measurements (endpoints) because this reduces the potential for the data set to be biased toward one type of toxicological effect. Unique measurements are those that represent different parameters of measurement for an endpoint category. For example, the endpoints of "Mortality" and "LC50" may both be categorized as S endpoints because they are both measurements of survival/mortality, but they are each considered a unique measurement because they measure different aspects of survival/mortality. Based on best professional judgment, more than 3 unique measurements are considered to provide a more than adequate representation of the toxicity of a chemical for the test organism group of concern. Three unique measurements are considered to provide an adequate representation while fewer than 3 unique measurements are considered to provide a minimal representation of the toxicity of a chemical for the test organism group of concern.

### Justification for Weighting Factor Level:

This category is given a **Non-critical** weighting factor level. This evaluation category is related to the robustness of the data set and its ability to represent the ideal GMM TRV; thus the true TRV is estimated. The higher the number of unique measurements, the more robust the data set. This evaluation category is not, however, a primary factor for determining whether or not the GMM TRV should be used because a high number of unique measurements in the data set is not a requirement, but rather an additional benefit for assessing the validity of the GMM to estimate the true TRV. Furthermore, all the unique measurements that are allowed in the data set are by definition relevant to the TRV being developed for population effects. The relevance of the endpoint category of each unique measurement is scored separately under [\*Type of Endpoint Category\*](#).

## Type of Endpoint Category

### Justification for Scoring:

The preference is to have more reproduction and development endpoints followed by survival endpoints and then by adult body weight or size change endpoints because the first category of endpoints is the most ecologically relevant group for determining long-term effects on populations followed by the second and third category.

### *Justification for Weighting Factor Level*

This category is given a **Non-critical** weighting factor level. This evaluation category indicates the degree of relevance the data set has to the effects of concern, population level effects, for which the GMM TRV is being developed. The higher the degree of relevance, the more closely the GMM TRV represents the ideal GMM TRV; thus the true TRV is estimated. This evaluation category is not, however, a primary factor for determining whether or not the GMM TRV should be used because all the endpoint categories considered are ecologically relevant by definition. However, reproduction or development endpoints can more closely approximate population level effects, so having more endpoints in this

category is an added benefit for assessing the validity of the GMM TRV for estimating the true TRV.

#### Number and Type of Effect Levels of PTVs Associated with the Individual NOAEL (NOEC)-based ELs in the GMM TRV Data Set

##### Justification for Scoring:

The preference is to have chronic NOAELs (NOECs) with LOAELs (LOECs), followed by chronic NOAELs (NOECs) without LOAELs (LOECs), then by subchronic NOAELs (NOECs) with LOAELs (LOECs), then by subchronic NOAELs (NOECs) without LOAELs (LOECs) and finally by all other ELs. This hierarchy is based on two factors. One factor is whether or not UFs have to be applied to a PTV to extrapolate to a chronic NOAEL (NOEC). Extrapolated values are less preferred because they may be overly conservative, thus less representative of the actual chronic NOAEL (NOEC). The second factor is whether or not there are any NOAELs (NOECs) with accompanying LOAELs (LOECs). NOAELs (NOECs) with LOAELs (LOECs) are most preferred because these values bracket the range of possible effects better than just a NOAEL (NOEC) or just a LOAEL (LOEC) alone.

##### Justification for Weighting Factor Level:

This category is given a **Non-critical** weighting factor level. This evaluation category is directly related to the certainty in the GMM TRV. The more ELs in the GMM TRV data set that were extrapolated to chronic NOAEL (NOEC)-based ELs by applying UFs, the greater the level of conservatism that is built into the GMM TRV. Even though being overly conservative is acceptable for screening level ecological risk assessments, it is preferred that TRVs not be overly conservative if more certain data is available. On the other hand, the higher the number of original ELs that are chronic NOAELs (NOECs) in the GMM TRV data set, the higher the confidence that the GMM TRV represents the ideal GMM TRV and thus estimates the true TRV (chronic no observed adverse effect level). A high score in this evaluation category is not required, but is an additional benefit for assessing confidence in the TRV.

#### Confidence Rating of PTVs Associated with the Individual NOAEL (NOEC)-based ELs in GMM TRV Data Set

##### Justification for Scoring:

The preference is to have more effect levels (PTVs) with high confidence ratings, followed by those with medium and then by low. A PTV confidence rating indicates to what degree the PTV is ecologically relevant, defensible and well documented based on the PTSE Part 2 Study Evaluation criteria (See Ref ID 1487). Effect levels associated with a low confidence rating are not included in the data set unless the data set is limited (i.e., less than 3 ELs based on PTVs with either a High or Medium confidence rating.).

##### Justification for Weighting Factor Level:

This category is given a **Non-critical** weighting factor level. This evaluation category indicates the degree of relevance the data set has to the TRV that is being developed. The higher the degree of relevance, the more closely the GMM TRV represents the ideal GMM

TRV; thus the true TRV is estimated. The PTV confidence rating is based upon scoring various study elements that are considered to be relevant for developing a scientifically defensible and ecologically relevant TRV. A high PTV confidence rating indicates the value is highly relevant for deriving a TRV and more likely to accurately estimate the true TRV.

#### Outliers(s) in the Chronic NOAEL (NOEC)-based EL Distribution

##### Justification for Scoring:

The data set cannot have invalid outliers (i.e., values associated with error or study designs that do not meet the minimum requirements for deriving a TRV). Invalid outliers must be removed from the data set prior to calculation of the GMM TRV. An invalid outlier is determined by a low confidence rating of a PTV associated with an EL in the data set. However, valid outliers, or extreme values, are allowed (e.g. sensitive species) as long as the data set is not bimodal (see [the \*Chronic NOAEL \(NOEC\)-based EL Distribution is Bimodal\*](#) section). The GSD is used to determine the variance of the GMM TRV. A lower variance (smaller GSD) indicates that the GMM TRV is more likely to represent the ideal GMM TRV and thus more accurately estimate the true TRV while a high variance (higher GSD) indicates that the GMM TRV is less likely to represent the ideal GMM TRV and thus less accurately estimate the true TRV. In most cases of high variance, the GMM TRV may be overly conservative because the large variance in the values is a result of the averaging of ELs that are based on PTVs other than chronic NOAELs (NOECs) and the application of UFs to extrapolate these values to chronic NOAEL (NOEC)-based ELs. A data set that contains both the smaller, extrapolated values and the non-extrapolated values (i.e., original effect levels that were already chronic NOAELs (NOECs)), leads to a high variance.

##### Justification for Weighting Factor Level:

This category is given a **Critical** weighting factor level. This evaluation category represents the variance of the GMM TRV dataset, which is important because it indicates how well the GMM TRV represents the ideal GMM TRV; thus how well the GMM TRV estimates the true TRV, which is directly related to the confidence in the GMM TRV. Low variance equals high confidence. High variance equals low confidence and may require reconsideration of the GMM TRV.

#### Chronic NOAEL (NOEC)-based EL Distribution is Bimodal

##### Justification for Scoring:

The preference is for the GMM TRV data set not to have a bimodal distribution. A bimodal distribution is determined based on 2 distinct clusters of values associated with different test species, original exposure durations, original effect levels or endpoint categories of each EL in the data set. If a data set is bimodal, best professional judgment must be used to determine if a subset GMM TRV(s) needs to be calculated or if the GMM TRV can be used as is.

##### Justification for Weighting Factor Level:

This category is given a **Critical** weighting factor level. This evaluation category has a large influence on whether or not the GMM TRV will be used. If the GMM TRV data set is found

to have a bimodal distribution, the GMM TRV may need to be revised in order to represent the most sensitive and/or ecologically relevant distribution (E.g., One distinct cluster is rodent (omnivore) data while the other is mink (carnivore) data. A TRV calculated from rodent data is more appropriate for the omnivorous deer mouse ESL receptors, while a TRV calculated from the mink is more appropriate for carnivorous red fox ESL receptor.)

#### Relationship of GMM TRV to chronic LOAEL (LOEC)-based ELs

##### Justification for Scoring:

The preference is that the GMM TRV be below the lowest chronic LOAEL (LOEC)-based EL because that indicates that it is protective of the most sensitive adverse effect in the data set. If the GMM TRV is not below the lowest chronic LOAEL (LOEC)-based EL, the next preference is for it to be no more than 3 times higher than a chronic LOAEL (LOEC)-based EL based on a chronic or C-CL LOAEL (LOEC) for an R/D or less ecologically relevant endpoint. Next preference is that the GMM TRV is not more than 3 times higher than a chronic LOAEL (LOEC)-based EL extrapolated from an original effect level other than a LOAEL. Because some of the chronic LOAEL (LOEC)-based ELs are extrapolated from NOAELs (NOECs) or other effect levels by applying UFs, they may be overly conservative and not represent the true chronic LOAELs (LOECs) for particular endpoints. In such cases, the GMM TRV is considered adequately protective due to the conservatism built into the extrapolated chronic LOAEL (LOEC)-based ELs. Furthermore, the GMM TRV may be considered adequately protective, if it is below the chronic LOAEL (LOEC)-based ELs for the most ecologically relevant endpoints (reproduction and development) even though it may exceed the lowest chronic LOAEL (LOEC)-based EL for an adult body weight or size change endpoint or for a survival endpoint. Another consideration is to determine based on best professional judgment whether or not the GMM TRV is unacceptably higher or lower than the lowest chronic LOAEL (LOEC)-based EL. If the difference is unacceptable, further investigation is warranted to determine if the GMM TRV is inappropriate (i.e., unacceptably over or under conservative). If it is found to be unacceptable, then the GMM TRV may need to be revised.

##### Justification for Weighting Factor Level:

This category is given a **Critical** weighting factor level. This evaluation category has a large influence on whether or not the GMM TRV will be used. If the difference between the GMM TRV and the lowest chronic LOAEL (LOEC)-based EL is unacceptable, the GMM TRV is unacceptable and an alternative (e.g., a subset GMM TRV, CS TRV) must be sought.

#### Relationship of GMM TRV to other published TRVs

##### Justification for Scoring:

It is preferred that any differences between the GMM TRV and other published TRVs be explainable on the basis of the experiments, endpoints, test organisms, and test chemical forms, etc., considered. It is also important that the explanation provide support for or against the use of the GMM TRV. Also to be verified is that the GMM TRV has considered all relevant data. If relevant data has not been considered, the GMM TRV data set may need

to be expanded to include the missing data. If no published TRVs are available for comparison, the GMM TRV is considered to be acceptable.

**Justification for Weighting Factor Level:**

This category is given a **Critical** weighting factor level. This evaluation category has a large influence on whether or not the GMM TRV will be used. If differences between the GMM TRV and other published TRVs are unacceptable (i.e., unexplainable, error based or lack of data based), the GMM TRV is unacceptable and an alternative (e.g., subset GMM TRV, CS TRV) needs to be considered

**Tables and Graphs for GMM TRV Data Set Information**

For easier presentation of the data in support of the GMM TRV, the following tables are included in the specific GMM TRV Summary Report: **PTVs Considered**, **Test Organisms**, and **Original Effect Level Types**. The **Graph of NOAEL-based ELs** and **Graph of LOAEL-based ELs** are also included for illustration of the data. In the graphs, the x-axis labels contain coding that indicates the original exposure durations, effect levels, endpoint categories, and test organisms from which the NOAEL (NOEC)- or LOAEL (LOEC)-based ELs are approximated. For example, if a data point is associated with the label "SC L WC Rat" in a NOAEL-based ELs graph, this label indicates that the value is a NOAEL-based EL approximated from a subchronic (SC) LOAEL (L) for weight change (WC) in the rat. Finally, it should be noted that when minimum and maximum LOAEL-based ELs are approximated, the minimum LOAEL-based ELs are used in the graph for LOAEL (LOEC)-based ELs.

**Table 4. UF Descriptions.**

Type of effect level available	UF applied to derive a TRV that is:	
	Chronic NOAEL-based	Chronic LOAEL-based
C-CL or chronic NOAEL	1	N/A
C-CL or chronic LOAEL	10	1
C-CL or chronic LD50 (or LC50)	100	10
C-CL or chronic ED50 (or EC50)	100	10
Subchronic NOAEL	10	N/A
Subchronic LOAEL	100	10
Subchronic LD50 (or LC50)	100	100
Subchronic ED50 (or EC50)	100	100
Acute or single dose NOAEL	100	N/A
Acute or single dose LOAEL	100	100
Acute or single dose LD50 (or LC50)	100	100
Acute or single dose ED50 (or EC50)	100	100

**Table 5. Acronyms.**

Acronym	Word/Phrase
A	Acute

<b>Acronym</b>	<b>Word/Phrase</b>
AIR	Air
ALL	All
B	Bird
C	Chronic
C-CL	Chronic - Critical Lifestage
CS	Critical Study
D/F	Dioxin/ Furan
DW	Drinking Water exposure medium
DW+F	Drinking water exposure medium plus additional exposure to background in food
EC10	Effective Concentration for 10% of population
EC20	Effective Concentration for 20% of population
EC50	Median Effective Concentration (for 50% of population)
EL	Effect Level
ESL	Ecological Screening Level
F	Food exposure medium
F&DW	Food and drinking water exposure media
F&DW&O	Food and drinking water and other exposure media
F+DW	Food exposure medium plus additional exposure to background in drinking water
GMM	Geometric Mean
GP	Generic plant (Terrestrial autotroph - producer)
GSD	Geometric Standard Deviation
HE	High Explosive
I	Invertebrate
INH	Inhalation exposure route
INORG	Inorganic Compound
L	LOAEL or LOEC
LANL	Los Alamos National Laboratory
LC10	Lethal Concentration for 10% of population
LC20	Lethal Concentration for 20% of population
LC50	Median Lethal Concentration (for 50% of population)
LOAEL	Lowest Observed Adverse Effect Level
LOEC	Lowest Observed Effect Concentration
LOEL	Lowest Observed Effect Level
M	Mammal
N	NOAEL or NOEC
N/A	Not Applicable
NL	NOAEL (or NOEC) and LOAEL (or LOEC)
NLOTH	NOAEL (or NOEC), LOAEL (or LOEC), and other effect level (e.g., LC50, EC50)
NMED	New Mexico Environment Department, inter- and intrastate stream standards
NOAEL	No Observed Adverse Effect Level
NOEC	No Observed Effect Concentration

<b>Acronym</b>	<b>Word/Phrase</b>
NOEL	No Observed Effect Level
NOTH	NOAEL (or NOEC) and other effect level (e.g., LC50, EC50)
NR	Not Reported
OIL	Oil exposure medium
OIL_ACHS	Arachis oil exposure medium
OIL_CORN	Corn oil exposure medium
OIL_O	Other oil exposure medium
OIL_PNT	Peanut oil exposure medium
ORNL	Oak Ridge National Laboratory
OTH	Other
P	Plant
PCB	Polychlorinated Biphenyl
PEST	Pesticide
PTSE	Primary Toxicity Study Evaluation
PTV	Primary Toxicity Value
R/D	Reproduction and Development
R6	Region 6
Ref	Reference
RRES-R	Risk Reduction and Environmental Stewardship - Remediation program
S	Survival
SAND&OM	Sand and Organic Matter mixture exposure medium
SAND_CLTR	Sand culture exposure medium (Solution is washed through silver sand daily)
SC	Subchronic
SD	Single Dose
SLE	Soil and Litter Earthworm
SLERA	Screening Level Ecological Risk Assessment
SNL	Sandia National Laboratory
SOIL&MNU	Soil and Manure mixture exposure medium
SOIL&SAND	Soil and Sand mixture exposure medium
SOIL&SLDG	Soil and Sludge mixture exposure medium
SOLN_O	Other solution exposure medium (assumed)
SOLN_OIL	Oil solution exposure medium (assumed)
SVOC	Semivolatile Organic compound
SzC	Size Change
T&E	Threatened and Endangered (Species)
TB	Terrestrial bird
TM	Terrestrial mammal
TP	Terrestrial plant
TRV	Toxicity Reference Value
UF	Uncertainty Factor
UNK	Unknown
USEPA	United States Environmental Protection Agency
VOC	Volatile Organic Compound
W	Water

<b>Acronym</b>	<b>Word/Phrase</b>
WC	Weight Change

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# **Toxicity Reference Value Development Methods for the Los Alamos National Laboratory**



Prepared by the Environmental Programs Directorate

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### Appendix

Appendix A     Primary Toxicity Study Evaluation Methods Used to Develop Los Alamos National Laboratory Toxicity Reference Values

### Acronyms and Abbreviations

COPC	chemical of potential concern
COPEC	chemical of potential ecological concern
CS	critical study
EP	Environmental Programs (Directorate)
EPA	U.S. Environmental Protection Agency
ESL	ecological screening level
HI	hazard index
HQ	hazard quotient
LANL or the Laboratory	Los Alamos National Laboratory

LC <sub>50</sub>	lethal median concentration
LD <sub>50</sub>	lethal median dose
LOAEL	lowest observed adverse effect level
LOEC	lowest observed effect concentration
NOAEL	no observed adverse effect level
NOEC	no observed effect concentration
PTSE	primary toxicity study evaluation
PTV	primary toxicity value
SLERA	screening-level ecological risk assessment
TRV	toxicity reference value

## 1.0 INTRODUCTION

This document details the process used to develop toxicity reference values (TRVs) for various chemical exposure pathways for selected wildlife at the Los Alamos National Laboratory (LANL or the Laboratory). These TRVs are used in ecological screening level (ESL) models representing the following exposure media for various chemicals to receptors.

- *Air*. Inhalation exposure pathway for burrowing mammals (volatile organic compounds only)
- *Soil and sediment*. Direct and food chain exposure pathways to birds and mammals
- *Water*. Drinking water ingestion to birds and mammals
- *Soil*. Direct exposure pathways to invertebrates (e.g., earthworms) and plants
- *Water and sediment*. Direct exposure pathways to aquatic community organisms

ESLs are used in screening-level ecological risk assessments (SLERAs) at the Laboratory. The TRVs, ESLs, model parameters, and all supporting documentation are archived in the Laboratory's Ecorisk Database (LANL 2005, 090032). The SLERA methodology is documented in "Screening-Level Ecological Risk Assessment Methods, Revision 2" (LANL 2004, 087630).

This document serves as guidance for risk assessors, risk managers, and others who wish to understand the logic behind the literature, evaluations, and documentation that leads to the development of TRVs used to calculate ESLs for SLERAs at the Laboratory.

Section 2 of this document provides a summary of ESL development and use.

Section 3 provides a summary of TRV development. It includes the working definition of a TRV at the Laboratory, an overview of the tiered TRV development process applied by the Laboratory, and a detailed description of each of the four tiers: Tier 1 (national value), Tier 2 (Laboratory-derived geometric mean TRV), Tier 3 (Laboratory-derived critical study TRV), and Tier 4 (non-Laboratory-derived critical study TRV).

Appendix A contains the primary toxicity study evaluation (PTSE) methods used to develop Laboratory TRVs. The PTSE process is used to develop the Laboratory's Tier 2 and Tier 3 TRVs from the primary toxicity literature. Appendix A contains data sources and a detailed step-by-step process for data entry for the PTSE databases created in Microsoft Access for documentation purposes.

Please note that this document best describes the PTSE process for Ecorisk Database Release 2.4 (LANL 2009, 107524). Any updates/revisions to the methods can be obtained from the current Risk Assessment Team Leader for the Laboratory's Environmental Programs (EP) Directorate.

## 2.0 SUMMARY OF ESL DEVELOPMENT AND USE

ESLs are used to evaluate potential hazards associated with chemicals and radionuclides found at the Laboratory. The Laboratory has developed chemical-, media-, and receptor-specific ESLs using a tiered TRV development approach, as described in section 3 of this document. ESLs are developed and maintained by the Laboratory as part of the Ecorisk Database, which archives the ESLs, TRVs, associated exposure parameters, and all supporting documentation. The Ecorisk Database was initially developed in 1998, with the most current release (2.4) provided in December 2009.

The development of an ESL is a two-step process. The first step involves identifying or developing a TRV. In the second step, the TRV and exposure parameters are used to calculate ESLs for chemicals and ecological receptors representative of the ecosystems at the Laboratory. Eleven different receptors were selected to be representative of mammals, birds, plants, and invertebrates inhabiting terrestrial and aquatic ecosystems at the Laboratory. At the time of this publication, 182 analytes, including inorganic chemicals, organic chemicals, and radionuclides, have ESLs documented in the database.

## **2.1 Goals of the Risk Assessment Process at the Laboratory**

The goals of the risk assessment process are two-fold: (1) to quantify hazards to the environment and associated exposure to radioactive and chemical wastes from past treatment, storage, and disposal practices; and (2) to facilitate meeting the environmental cleanup requirements of the Laboratory's permit to operate hazardous waste facilities.

In accordance with these goals, the SLERA is used to determine whether there is a potential ecological risk that needs to be more fully considered in a baseline ecological risk assessment.

## **2.2 The Screening Level Ecological Risk Assessment Process**

The purpose of the screening assessment is to provide information to risk managers so that informed risk management decisions can be made. The SLERA process follows the U.S. Environmental Protection Agency's (EPA's) "Ecological Risk Assessment Guidance for Superfund: Process for Designing and Conducting Ecological Risk Assessments" (EPA 1997, 059370) and the "Guidelines for Ecological Risk Assessment" (EPA 1998, 062809). The SLERA process uses information on the environmental setting, contaminant fate and transport, exposure pathways, and functional food webs to establish a conceptual site model that can be assessed for impacts using assessment endpoints and a select group of screening receptors. The SLERA process then uses ESLs as threshold values to aid in determining whether a chemical is of potential ecological concern and requires further investigation. The ESLs are developed for individual chemicals and are medium and receptor specific. If a site has levels of a chemical above the ESL in any medium, then this site may pose a potential risk to ecological receptors. To evaluate the potential risk for each chemical of potential concern (COPC), the ESL and the representative site concentration are used to calculate the hazard quotient (HQ). If the HQ for a COPC is greater than 1.0 at a site with only a single COPC, or the HQ for a COPC is greater than 0.3 for a site with multiple COPCs, then that chemical is identified as a chemical of potential ecological concern (COPEC) and evaluated further.

ESLs are specific to each medium (air, soil, sediment, and water) and do not account for exposure to multiple media. A method to account for wildlife exposure to multiple media includes a multimedia exposure calculation that results in a hazard index (HI) value for each wildlife receptor. The HI is a sum of HQ values. HQs are calculated for each screening receptor and each contaminant and may be thought of as a ratio of a receptor's exposure at the site to an acceptable effect level. If the HI is greater than 1.0, then the site may pose an ecological risk. An uncertainty analysis follows COPEC identification and can result in adding chemicals to, or removing them from, the list of COPECs. The SLERA process is described in detail in "Screening-Level Ecological Risk Assessment Methods, Revision 2" (LANL 2004, 087630).

## **2.3 Description of Ecological Screening Levels**

ESLs are media- and receptor-specific values. Air, soil, sediment, and water ESLs are calculated for ecological screening receptors in various functional feeding guilds (e.g., carnivores, herbivores,

insectivores). The ESLs are calculated using ecological exposure parameters (e.g., ingestion rate and bioconcentration factors) and the TRV. The ESL calculations are described in detail in "Screening-Level Ecological Risk Assessment Methods, Revision 2" (LANL 2004, 087630), and ESL values are archived in the Ecorisk Database along with the models and model parameter values including the TRVs.

## **2.4 Description of the Ecorisk Database**

The Ecorisk Database was created in 1998 as a user-friendly database application to document and archive information for the ESLs and associated parameters including TRVs. The Ecorisk Database also provides detailed documentation for justifying the type of information collected and used and illustrates how values are calculated. The database can be searched by chemical or screening receptor and generates on-screen and printable reports for all ESLs, TRVs, and exposure parameters. The database is a Microsoft Access file that is distributed to all project risk assessors and is provided upon request to federal and state agencies and other contractors, both nationally and internationally.

## **2.5 Update of ESLs and the Ecorisk Database**

The selection of the specific chemicals for which ESLs are derived is primarily dependent upon project needs. ESLs are updated based on changes to the ESL equations, the calculation or source of ESL parameters, and more recent or updated TRVs. The need for ESLs is reviewed to determine priorities for TRV development. If new ESLs are not needed, then existing TRVs are reviewed to determine priorities for retrieving and reviewing new literature to supplement information in the database.

A new release of the database is provided as necessary. All new/updated ESL parameters and TRVs are recorded in the database, and the new ESLs are calculated. All ESLs, TRVs, and calculations undergo quality assurance checks. Each database release contains an ESL history report that documents any changes made to data or the database interface since the last release.

## **2.6 Interim and Surrogate ESLs/TRVs**

Interim and surrogate ESLs/TRVs are also included along with the most recent release of the Ecorisk Database. Interim values are those that have not been formally peer reviewed by the EP Directorate's Risk Assessment Team. Interim values are provided to risk assessors as needed between database releases.

Surrogate ESLs/TRVs are used for chemicals lacking toxicity data but are structurally similar to, or a degradation product of, chemicals with an ESL/TRV.

## **3.0 SUMMARY OF TRV DEVELOPMENT**

### **3.1 TRV**

A TRV represents an exposure rate associated with an acceptable risk from chronic exposure of an ecological receptor to a specific contaminant via a specific exposure pathway. In other words, exposures exceeding the TRV may pose adverse effects to wildlife species, while exposures below the TRV are not expected to result in adverse effects (EPA 2005, 089448).

TRVs are important parameters in ESL calculations because "they represent the component of the model that determines whether or not a contaminant in a media may present potential harm to ecological receptors in the area" (Podolsky et al. 2001, 072586). For any given chemical, TRV values vary among

government agencies and private sectors because the methods used to develop them vary according to the site-specific concerns of the organization that developed them (i.e., receptor species, chemical, type of exposure pathway, type and magnitude of uncertainty factors applied).

The ideal TRV for ecological risk screening assessments at the Laboratory is one that is based on literature representing the most ecologically relevant effects (reproduction/development, survival and/or adult weight/size change), exposure routes (oral ingestion via food or drinking water for birds and mammals, inhalation for mammals, uptake via seed coat and/or roots for plants, and direct contact exposure for invertebrates and aquatic community organisms), exposure media (food and drinking water for birds and mammals, air for mammals, soil for plants and invertebrates, and water and sediment for aquatic community organisms), exposure period (chronic), and effect levels (no observed adverse effect level [NOAEL] for vertebrates or no observed effect concentration [NOEC] for plants and invertebrates). A TRV based on these characteristics is considered protective of the wildlife; aquatic community organism, plant, and invertebrate populations; and sensitive individuals because it represents an exposure that is not associated with adverse impacts of low-level, long-term chemical effects (i.e., adverse effects on ability of individuals to develop into viable organisms, search for mates, breed successfully, and produce live and equally viable offspring).

## **3.2 Definitions**

### **3.2.1 Ecologically Relevant Effects**

An ecologically relevant toxicity study effect is defined as a measurement that is considered most closely related to population effects, i.e., an effect that directly influences reproductive success and survival. Reproduction, development, survival, and weight/size change measurements are considered to be more ecologically relevant than biochemical, physiological, or cancer measurements because they more closely reflect effects on population health/size (EPA 2005, 089448); thus, the former are selected for use in developing TRVs at the Laboratory.

### **3.2.2 Ecologically Relevant Media and Exposure Routes**

An ecologically relevant toxicity study exposure medium/route is defined as one that is most closely related to that which is found in the natural environment of concern.

Wildlife receptors are exposed to chemicals in their natural environment primarily through their diet, so ingestion of food or food-like substances is considered the most ecologically relevant toxicity study exposure medium/route for developing TRVs at the Laboratory for wildlife. Oral exposure using capsules, gavage, or intubation is considered similar to ingestion of food and thus also ecologically relevant. Wildlife receptors are also exposed, although to a lesser degree, to chemicals through ingestion of drinking water and, under special circumstances, through the inhalation of air (e.g., burrowing mammal), so separate TRVs are developed with toxicity data for chemicals being ingested in drinking water, and separate TRVs are developed for chemicals inhaled in air. Because of differences in bioavailability of chemicals depending on the exposure media/routes, those that do not represent chemical exposure through the digestive system or through the lungs are not considered ecologically relevant, e.g., intraperitoneal, intravenous, or intramuscular. Wildlife receptors are also exposed dermally to chemicals, but this exposure route is not considered for TRV development because the contribution of dermal exposure to the overall exposure is considered minimal compared to the other exposure scenarios mentioned above (i.e., fur and feathers as barrier, dermal exposure less significant than oral exposure [EPA 2005, 089448]).

Terrestrial plants and worms are exposed to chemicals in their natural environment primarily through direct uptake from soil, which is the most ecologically relevant toxicity study exposure medium/route for developing TRVs for plants and worms at the Laboratory. Because of differences in the bioavailability of chemicals in different exposure media, exposure in solution or on filter paper is not considered ecologically relevant. Also, worms ingest chemicals in soil in their natural environment, but this exposure medium/route is not considered separately. The contribution to the overall exposure from ingestion is difficult to discern because the worm's alimentary tract is in contact with soil the majority of the time as well.

Aquatic community organisms are exposed to chemicals in their natural environment primarily through direct contact with water and sediment, which are the most ecologically relevant toxicity study exposure media/routes for developing TRVs at the Laboratory for aquatic community organisms. Also, some aquatic community organisms may ingest chemicals in water and/or sediment in their natural environment, but this exposure medium/route is not considered because the contribution to the overall exposure is considered minimal compared to the direct contact uptake since the organism's body is in complete contact with the water and/or sediment at all times.

### **3.2.3 Ecologically Relevant Test Organisms (species)**

An ecologically relevant toxicity study test organism (species) is defined as one that represents the ecological receptor of concern at least at the taxonomic class level, e.g., mammal, bird, plant, or earthworm class. Although there are species differences within a class, the toxicity data are generally not robust enough to evaluate such differences, except qualitatively.

### **3.2.4 Exposure Duration Categories**

To be ecologically relevant, the toxicity study exposure duration is defined as one with a chemical exposure encompassing the majority of the test organism's lifespan or the critical period / life stage of reproduction. The definition of chronic varies depending on the interpretation of lifespan data, and the definition of chronic critical life stage varies depending on the interpretation of life stage data. The Laboratory uses the definitions stated in EPA's "Screening Level Ecological Risk Assessment Protocol for Hazardous Waste Combustion Facilities" (EPA 1999, 070923).

Because not all toxicity studies are chronic or focused on a critical life stage, less than chronic data are used after the application of appropriate uncertainty factors to extrapolate data to a chronic value. Uncertainty factors for subchronic, acute, and single-dose exposures are described in more detail in section 3.2.7, Uncertainty Factors. Less than chronic data are deemed appropriate for use to increase the size of otherwise limited data sets.

### **3.2.5 Selection of Dose Calculation**

To be ecologically relevant, a dose calculation parameter for wildlife exposure models such as body weight, ingestion, or inhalation rate is defined as one that best matches the age / life stage of the test organism, as well as best reflects the entire chemical administration period of the toxicity study. Furthermore, food ingestion rates in units of dry weight are preferred in order to normalize the rate for moisture content of different dietary items.

### 3.2.6 Dose Calculation

An ecologically relevant dose calculation for wildlife exposure models is defined as one that is continuous/daily because this best represents a chronic exposure, which is generally the exposure of concern in SLERAs. If a datum from an intermittent dosing design is used to develop a toxicity value, it is normalized to a continuous rate before calculating a toxicity value (e.g., normalizing an intermittent inhalation study design to a continuous/daily dose).

### 3.2.7 Uncertainty Factors

In order to best represent an ecologically relevant TRV, uncertainty factors are used to extrapolate toxicity values from studies with less than chronic exposure durations, as well as from toxicity values representing effect levels other than a NOAEL/NOEC, such as a lowest observed adverse effect level / lowest observed effect concentration (LOAEL/LOEC), lethal median dose (LD<sub>50</sub>), or lethal median concentration (LC<sub>50</sub>). Uncertainty factor application allows the use of more data to increase an otherwise limited data set available for developing a TRV. Uncertainty factors are generally based on the relationship identified between no effect and low or lethal effect levels as well as best risk management practices. The Laboratory uses uncertainty factors as defined in EPA's "Screening Level Ecological Risk Assessment Protocol for Hazardous Waste Combustion Facilities" (EPA 1999, 070923).

## 4.0 TIERED TRV APPROACH

TRVs are identified/developed in one of four ways. Depending on how it is developed, the TRV is assigned a tier of 1 to 4. A Tier 1 TRV has the most certainty in the toxicity data used to derive it, and a Tier 4 TRV has the least certainty in its derivation. This tiered process reduces data gaps by allowing for the maximal use of available toxicity data by considering a variety of sources, while at the same time communicating the degree of certainty in the data supporting the value.

Tiers are presented in the order of preference and confidence used to derive the TRVs and are as follows:

- *Tier 1.* A published, nationally accepted TRV such as an EPA ecological soil screening level TRV or International Atomic Energy Agency radionuclide dose limit of 0.1 rad per day for the protection of ecological receptors at the population level.
- *Tier 2.* A TRV equal to the geometric mean of ecologically relevant NOAEL- or NOEC-based effect levels derived from review of the primary toxicity literature by the Laboratory (three or more data points are available) using the PTSE process (see Appendix A).
- *Tier 3.* A critical study (CS) TRV, which is based on an ecologically relevant maximum NOAEL- or NOEC-based effect level that is lower than the lowest reported LOAEL- or LOEC-based effect level derived from review of the primary toxicity literature by the Laboratory using the PTSE process (see Appendix A).
- *Tier 4.* A CS TRV derived using ecologically relevant primary toxicity values (PTVs) or TRVs reported by a secondary data source such as Oak Ridge National Laboratory, Sandia National Laboratories, U.S. Army Center for Health Promotion and Preventive Medicine, or EPA Region 5 environmental data quality levels.

Tier 1 TRVs are considered to have the greatest certainty considering the rigorous national peer review they have undergone before publication. The certainty associated with the Tier 2 and Tier 3 TRVs is

based on the ecological relevance of available toxicity information based on the internal peer review by the Laboratory. Tier 2 TRVs have more certainty than Tier 3 TRVs because they are based on more toxicity information from the literature. Tier 4 TRVs are considered to have the most uncertainty because these secondary compilations of the literature do not provide as much documentation as is available for Tiers 1, 2, or 3.

## 5.0 CONVERSION OF TRVs TO ESLs

ESLs are chemical- and medium-specific screening levels pertaining to a given receptor (e.g., avian omnivore, earthworm) and medium (sediment, soil, water, and/or air). The TRV is used in the receptor-specific ESL calculation, which converts the toxicity value from a dose (mg-contaminant/kg body weight/d) to an environmental concentration (e.g., mg-contaminant/kg-soil) using factors to estimate the transfer of chemical from soil, sediment, or water to dietary media (e.g., soil-to-plant transfer factor) and receptor-specific exposure parameters (e.g., ingestion/inhalation rates and body weight). In the case of plants, earthworms, and aquatic organisms, the TRV is equal to the ESL because the toxicity value is already in environmental concentration units.

## 6.0 OVERVIEW OF APPENDIX A

The Laboratory's PTSE process is used to develop Tier 2 and Tier 3 TRVs. Because this process is detailed and the supporting documentation is contained in a standardized format within the Ecorisk Database, a document that explains the field names, standardized or explanatory data entries, and justification thereof is needed for risk assessors and managers to understand the foundation of the values being used in SLERAs.

Appendix A also provides detailed instructions for performing PTSEs of the literature on the toxicity of chemicals to terrestrial birds, mammals, invertebrates (earthworms), and plants. The data obtained through the PTSE process are used to calculate PTVs. A PTV or group of PTVs is used to derive a Tier 3 CS TRV or Tier 2 geometric mean TRV, respectively, depending on the size of the data set available.

In the case of birds or mammals, a PTV is a daily dose rate (mg chemical/kg body weight/d) derived from the experiment and based on up to three dose rate parameters: (1) the concentration of the chemical administered in the study, (2) the food or water ingestion rate or inhalation rate of the test organism, and (3) the body weight of the test organism. In the case of plants or invertebrates, a PTV is a soil concentration (mg chemical/kg soil) based on the concentration of the chemical administered in the study. A PTV can be designated as a certain effect level (e.g., NOAEL or LC<sub>50</sub>), depending on whether and to what extent the daily dose rate potentially leads to adverse effects in the test organisms.

The PTSE process consists of the following four main steps: (1) data extraction, (2) study evaluation and PTV calculation, (3) TRV development, and (4) TRV peer review and approval. Each of the first three steps has their own data-entry database to facilitate the evaluation and to document the process. The fourth step consists of having the EP Directorate's Risk Assessment Team peer review each TRV derived through the PTSE process. Once a TRV is approved, the new PTSE TRV and all supporting data are incorporated into the Ecorisk Database for calculating appropriate ESLs for specific chemicals, exposure pathways, and screening receptors. These ESLs are ultimately used in SLERAs. Although the TRVs are just one component of the Ecorisk

*“Data” represents toxicity information from the scientific literature such as details of the study design, test organism, or toxicological effects.*

Database, they play a crucial role in the derivation of ESLs. Much consideration of the toxicological data takes place during TRV development to best estimate the exposure concentration in environmental media that will not harm key screening receptors and possibly other organisms in the Laboratory's environment.

In summary, Appendix A includes guidelines for the literature search and collection, data extraction, default value assignment, and exception ruling for various fields of data entry in customized PTSE databases, PTV calculation, and TRV derivation. Before performing a PTSE, the primary toxicity literature for the organism and for the exposure pathway and chemical scenario of concern must be identified and collected. As a result, the appendix begins with guidelines for literature searches and retrieval.

## 7.0 REFERENCES

*The following list includes all documents cited in this report. Parenthetical information following each reference provides the author(s), publication date, and ER ID. This information is also included in text citations. ER IDs are assigned by the Environmental Programs Directorate's Records Processing Facility (RPF) and are used to locate the document at the RPF and, where applicable, in the master reference set.*

EPA (U.S. Environmental Protection Agency), June 5, 1997. "Ecological Risk Assessment Guidance for Superfund: Process for Designing and Conducting Ecological Risk Assessments, Interim Final," Office of Emergency and Remedial Response, Washington, D.C. (EPA 1997, 059370)

EPA (U.S. Environmental Protection Agency), April 1998. "Guidelines for Ecological Risk Assessment," EPA/630/R-95/002F, Risk Assessment Forum, Washington, D.C. (EPA 1998, 062809)

EPA (U.S. Environmental Protection Agency), August 1999. "Screening Level Ecological Risk Assessment Protocol for Hazardous Waste Combustion Facilities," Volume One, peer review draft, EPA530-D-99-001A, Office of Solid Waste and Emergency Response, Washington, D.C. (EPA 1999, 070923)

EPA (U.S. Environmental Protection Agency), February 2005. "Guidance for Developing Ecological Soil Screening Levels," OSWER Directive No. 9285.7-55, Office of Solid Waste and Emergency Response, Washington, D.C. (EPA 2005, 089448)

LANL (Los Alamos National Laboratory), December 2004. "Screening-Level Ecological Risk Assessment Methods, Revision 2," Los Alamos National Laboratory document LA-UR-04-8246, Los Alamos, New Mexico. (LANL 2004, 087630)

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Podolsky, J.S., P.G. Newell, and O.B. Myers, 2001. "A Comparison of Toxicity Reference Values and Their Derivation Methods: Implications for Ecological Risk Assessment," presentation, Los Alamos National Laboratory document LA-UR-01-6503, Los Alamos, New Mexico. (LANL 2001, 072586)

## **Derivation of chemical-specific TRVs for PAHs and DDT & metabolites**

### **Objective**

The objective of this process is to develop toxicity reference values (TRVs) for individual polycyclic aromatic hydrocarbons (PAHs) and DDT and metabolites utilizing the toxicity data published in 2007 by the Environmental Protection Agency's (EPA) Ecological Soil Screening Level (EcoSSL) workgroup. These TRVs are used to calculate Los Alamos National Laboratory (LANL)-specific receptor ecological screening levels (ESLs).

### **Background**

The EPA EcoSSL workgroup reviewed the primary literature to develop TRVs and EcoSSLs for high and low molecular weight PAHs ([Table 1](#)). This class of organic compounds is grouped into two condensed aromatic ring structures with low molecular weight compounds composed of fewer than four rings and high molecular weight compounds composed of four or more rings. They also developed TRVs and EcoSSLs for DDT and metabolites as a group ([Table 2](#)).

In accordance with LANL SLERA methods, LANL generates TRVs for individual chemicals to be used to calculate LANL-specific receptor ecological screening levels (ESLs). Therefore, to remain consistent with the LANL screening level ecological risk assessment (SLERA) methods, the chemical-group TRVs/ ESLs derived by EPA were not adopted. LANL is, however, using the primary toxicity values for birds, mammals, plants and invertebrates (earthworms) for reproduction/ development, growth and survival endpoints that the EPA compiled with EcoSSL methodology to derive LANL TRVs and ESLs per LANL methods.

The EPA generates nationally-accepted EcoSSLs/TRVs through EcoSSL methodology and these toxicity values are considered to have high confidence compared to other sources. Therefore, the EcoSSL dataset is appropriate for use in the LANL Primary Toxicity Study Evaluation (PTSE) method, which is similar in many respects to the EcoSSL method. One notable exception is that LANL uses acute/ subacute and subchronic data by applying exposure duration uncertainty factors to extrapolate to a chronic effect level while EPA excludes these data, even if they have an expectable evaluation score otherwise, in order to focus their efforts on establishing a dose protective of most species from adverse effects associated with long term exposures and sublethal reproductive and growth effects. Another notable exception is that LANL utilizes reproduction/ development, growth and survival endpoints to calculate a TRV while EPA only uses the reproduction/ development and growth endpoints to calculate the TRV and then uses the survival endpoints in a comparative manner to evaluate the protectiveness of the TRV for lethality.

LANL has chosen to include along with chronic studies those of acute, subacute and subchronic duration and to utilize reproduction/ development, growth and survival endpoints to minimize data gaps for toxicological information for chemicals of potential ecological concern (COPECs) in the SLERA process.

The EPA primary toxicity values are used to augment existing LANL primary toxicity values compiled using the LANL PTSE method or to fill data gaps using the LANL PTSE method for LANL COPECs.

## **Methods**

### **Data Acquisition:**

- Primary toxicity values reported in the EPA EcoSSL reports for PAHs (USEPA, 2007a) and DDT and metabolites (USEPA, 2007b) were reviewed.

### **Data Coding – Effect Levels and Endpoints:**

- Selected No effect levels (NOAELs/ NOECs), low effect levels (LOAELs/LOECs), median effect levels (ED50s/ EC50s) and median lethality effect levels (LD50s/LC50s) data for individual PAHs and DDT and metabolites that are LANL COPECs ([Table 3](#)) that represented reproduction/ development, growth, or survival endpoints were selected for use in the LANL TRV data set. See [Table 4](#) for a description of endpoint group coding.

### **Data Coding - Handling of Repetitive Values:**

- In the cases where LANL and EPA derived toxicity values from the same reference, the LANL derived value(s) is used. The exception to this rule is if the LANL value is associated with LANL tier 4 TRV data ([Table 5](#)). Tier 4 TRV data are not included because this type of toxicity data was taken from secondary data sources other than the nationally accepted EPA EcoSSL documents and is not considered appropriate for deriving higher tier LANL TRVs due to differences in the level of detail in documentation of the TRV derivation process compared to the LANL PTSE Method. Only tier 1, 2 and 3 TRV data are included in the LANL TRV data sets. [Table 5](#) defines the LANL TRV tiers and their hierarchy for use in calculating TRVs/ ESLs.
- Only one effect type per reference per receptor/ COPEC pair is included in the data set. Best professional judgment is used to select the most ecologically relevant and/or sensitive value per ecologically relevant endpoint category per study/ reference. For example, if one experiment had 3 reproduction/development endpoints, 1 survival endpoint, and 1 adult growth endpoint, the most ecologically relevant and/or sensitive reproduction/development endpoint of the 3 available would be included in the data set along with the single survival and single growth change endpoints. This exclusion process minimizes the possibility of a TRV being skewed to the results of any particular study as a result of repetitive values for the same endpoint category within a study.

### **Normalization of toxicity values to chronic No Effect Levels:**

- All toxicity values were normalized to chronic no effect levels (NOAELs/NOECs) using uncertainty factors (UFs) for differences in exposure duration ([Table 6](#)) and or effect level per LANL PTSE Methods. [Table 7](#) indicates the UFs applied for various exposure durations and effect level combinations.

- One exposure duration classification that is used that is not necessarily based on the actual chemical administration period is the chronic – critical life stage (C-CL) designation. A C-CL endpoint is equivalent to a chronic exposure endpoint regardless of the actual chemical exposure duration associated with the endpoint because it is more likely to capture effects that reflect critical life stages that are relevant to population success. For the purpose of deriving TRVs, a critical life stage is defined as a life stage associated with a chemical exposure occurring during the reproductive cycle of the test organism and/or during the development of the immature test organism. For an endpoint to be considered development, it has to fall into one of two scenarios in which measurements must reflect either the development of immature organisms that were exposed via parents or the development of immature organisms directly exposed to the chemical.

#### **Calculation of TRV:**

- A Tier 2 geometric mean (GMM) TRV was calculated per LANL PTSE methods (Equation 1) when there were 3 or more primary toxicity values for a particular COPEC and receptor group. A critical study (CS) TRV was derived per LANL PTSE methods when there were less than 3 primary toxicity values for a particular COPEC and receptor group.

#### **Equation 1:**

GMM TRV = nth root of (EL1 x EL2 x EL3 x ... ELn)

where n is greater than or equal to 3 and each EL represents a chronic NOAEL-based EL for an oral ingestion exposure for an ecologically relevant effect (i.e., reproduction or development, survival or adult body weight or size changes)

#### **Results (See individual TRV Summary Reports and supporting PTSE documentation nthe Ecorisk Database)**

[Table 8](#) contains TRVs generated through this process.

#### **Summary**

Based on the primary toxicity data available in the the EPA EcoSSL 2007 reports for PAHs (USEPA, 2007a) and DDT and metabolites (USEPA, 2007b), LANL was able to augment existing PTSE method derived data sets or fill LANL COPEC TRV data gaps for 10 COPEC/ receptor group pairs. GMM TRVs were derived for 2 high MW PAHs [benzo(a)pyrene/ mammal and pyrene/ invertebrate (earthworm)], 2 low MW PAHs [fluorene/ invertebrate (earthworm), naphthalene/ bird, and naphthalene/ mammal), DDD/ bird, DDD/ mammal, DDE/ bird, DDE/ mammal, DDT/ bird and DDT/ mammal.

#### **References**

United States Environmental Protection Agency (USEPA), 2007a (Jun.). Ecological Soil Screening Levels for Polycyclic Aromatic Hydrocarbons (PAHs), Interim Final. OSWER Directive 9285.7-78. US Environmental Protection Agency, Office of Solid Waste and Emergency Response, Washington, D.C.

United States Environmental Protection Agency (USEPA), 2007b (Apr.). Ecological Soil Screening Levels for DDT and Metabolites. OSWER Directive 9285.7-78. US Environmental Protection Agency, Office of Solid Waste and Emergency Response, Washington, D.C.

## TABLES

Table 1. EPA EcoSSL TRVs for PAHs

Receptor	Low Molecular Weight TRV	High Molecular Weight TRV
<b>Soil Invertebrate</b>	29 m/kg soil dry wt.	18 mg/kg soil dry wt.
<b>Mammals</b>	170 mg/kg/day	0.615 mg/kg/day
<b>Birds</b>	Not Available	Not Available
<b>Plants</b>	Not Available	Not Available

Table 2. EPA EcoSSL TRVs for DDT and Metabolites

Receptor	DDT& Metabolite TRV
<b>Birds</b>	0.227 mg/kg/day
<b>Mammals</b>	0.147 mg/kg/d
<b>Soil Invertebrate</b>	Not Available
<b>Plants</b>	Not Available

Table 3. EPA EcoSSL toxicity data for PAHs and DDT and metabolites that are LANL COPECs

MW	COPEC	Receptor Group
LMW	Anthracene	P
HMW	Benzo(a)pyrene	M
LMW	Fluoranthene	I
LMW	Fluorene	I
LMW	Naphthalene	M
LMW	Phenanthrene	I
HMW	Pyrene	I
NA	DDT[4,4'-]	B, M
NA	DDE[4,4'-]	B, M
NA	DDD[4,4'-]	B, M
B = bird, H= high, I = invertebrate, L = Low, M = mammal, MW = molecular weight, P = plant		

Table 4. LANL endpoint groups

Endpoint Group	Description
Reproduction/ development (R/D)	Development or mortality measured in juvenile organisms or immature plants that were exposed to the chemical through

	parental exposure because it is considered to be a measurement of the ability of the parents to produce offspring that can develop into reproductive adults. Also, growth of a juvenile organism or immature plant that was directly exposed to the chemical because it reflects the potential for the juvenile or immature plant to develop normally into a reproductive adult.
Survival (S)	Mortality in an adult organism or in a juvenile organism or immature plant directly exposed to the chemical because it is considered a measurement of the ability of the organism to survive to reproductive maturity.
Growth (G)	Weight change (WC) for mature organisms is measured or a change occurs in size (WSz) of a mature organism (e.g., height or root length of plants).

Table 5. LANL TRV tiers and hierarchy for use calculating ESLs

TRV Tier	Description	Hierarchy for Use
1	Nationally accepted TRV (e.g., EPA EcoSSL TRV)	First
2	Geometric Mean (GMM) TRV derived through the primary toxicity evaluation (PTSE) process	Second
3	Critical Study (CS) TRV derived through the primary toxicity evaluation (PTSE) process	Third
4	Secondary source TRV (e.g., ORNL, SNL)	Fourth

Table 6. Exposure Duration Categories and IDs for Birds, Mammals, Earthworms, and Plants

Duration	Duration ID	Birds and Mammals	Earthworms and Plants
Chronic	C	91 days or more	7 days or more
Chronic- critical life stage	C-CL	All R/D endpoints	
Subchronic	SC	14 to 90 days	3 to 6 days
Acute	A	13 days or less	2 days or less
Single dose	SD	One time administration	One time administration

Duration	Duration ID	Birds and Mammals	Earthworms and Plants
Not Reported	NR	Not applicable	Not applicable
R/D = reproduction/ development			

Table 7. Uncertainty Factors Applied to Derive Chronic NOAEL- or NOEC-based Effect Levels

Type of effect level available	UF applied to derive a TRV that is a Chronic NOAEL (NOEC)-based EL:
C-CL or C NOAEL (NOEC)	1
C-CL or C LOAEL (LOEC)	10
C-CL or C LD50 (LC50), ED50 (EC50)	100
SC NOAEL (NOEC)	10
SC LOAEL (LOEC), LD50 (LC50), ED50 (EC50)	100
A or SD NOAEL (NOEC)	100
A or SD LOAEL (LOEC), LD50 (LC50), ED50 (EC50)	100

A = Acute, C = chronic, C-CL = chronic – critical life stage, EC50 = median effective concentration (for 50% of the population), ED50 = median effective dose (for 50% of the population), LC50 = median lethal concentration (for 50% of the population), ED50 = median lethal dose (for 50% of the population), LOAEL = low observed adverse effect level, LOEC = low observed effect concentration, NOAEL = no observed adverse effect level, NOEC = no observed effect concentration, SC = subchronic, SD = single dose

Table 8. TRVs

MW	COPEC	Receptor Group	GMM TRV*	CS TRV*
LMW	Anthracene	P	6.88	-
HMW	Benzo(a)pyrene	M	5.58	-
LMW	Fluoranthene	I	10.2	-
LMW	Fluorene	I	3.7	-
LMW	Naphthalene	M	14.3	-
LMW	Phenanthrene	I	5.5	-
HMW	Pyrene	I	10.6	-
NA	DDD	B	0.016	-
NA	DDD	M	5.83	-
NA	DDE	B	0.48	-
NA	DDE	M	9.02	-
NA	DDT	B	2.01	-
NA	DDT	M	-	0.139

B = bird, COPEC = chemical of potential ecological concern, CS = critical study, GMM = geometric mean, H= high, I = invertebrate, L = Low, M =

mammal, MW = molecular weight, N = number of toxicity values in data set,  
NA = not applicable, P = plant, TRV = toxicity, \* units are mg/kg for receptor  
groups I and P and mg/kg/d for receptor groups B and M









Year	Month	Day	Time	Location	Activity	Remarks
2000	1	1	08:00	...	...	...
2000	1	2	08:00	...	...	...
2000	1	3	08:00	...	...	...
2000	1	4	08:00	...	...	...
2000	1	5	08:00	...	...	...
2000	1	6	08:00	...	...	...
2000	1	7	08:00	...	...	...
2000	1	8	08:00	...	...	...
2000	1	9	08:00	...	...	...
2000	1	10	08:00	...	...	...
2000	1	11	08:00	...	...	...
2000	1	12	08:00	...	...	...
2000	1	13	08:00	...	...	...
2000	1	14	08:00	...	...	...
2000	1	15	08:00	...	...	...
2000	1	16	08:00	...	...	...
2000	1	17	08:00	...	...	...
2000	1	18	08:00	...	...	...
2000	1	19	08:00	...	...	...
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2000	1	23	08:00	...	...	...
2000	1	24	08:00	...	...	...
2000	1	25	08:00	...	...	...
2000	1	26	08:00	...	...	...
2000	1	27	08:00	...	...	...
2000	1	28	08:00	...	...	...
2000	1	29	08:00	...	...	...
2000	1	30	08:00	...	...	...
2000	1	31	08:00	...	...	...
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2000	2	26	08:00	...	...	...
2000	2	27	08:00	...	...	...
2000	2	28	08:00	...	...	...
2000	2	29	08:00	...	...	...
2000	2	30	08:00	...	...	...
2000	2	31	08:00	...	...	...
2000	3	1	08:00	...	...	...
2000	3	2	08:00	...	...	...
2000	3	3	08:00	...	...	...
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Chemical	Analyte Code	Cas No.	Final Soil ESL (mg/kg)	Receptor	Surrogate
BHC[delta-]	319-86-8	319-86-8	9.40E-03	Montane shrew (Mammalian insectivore)	BHC[gamma-]
Bis(2-chloroethyl)ether	111-44-4	111-44-4			
Bromomethane	74-83-9	74-83-9			
Chloromethane	74-87-3	74-87-3			
Chloronaphthalene[2-]	91-58-7	91-58-7	1.00E+00	Generic plant (Terrestrial autotroph - producer)	Naphthalene
Dinitroaniline[3,5-]	618-87-1	618-87-1			
Endosulfan sulfate	1031-07-8	1031-07-8	6.40E-01	Deer mouse (Mammalian omnivore)	Endosulfan
Endrin aldehyde	7421-93-4	7421-93-4	1.40E-03	American robin (Avian insectivore)	Endrin
Endrin ketone	53494-70-5	53494-70-5	1.40E-03	American robin (Avian insectivore)	Endrin
Ethylbenzene	100-41-4	100-41-4	2.40E+01	Deer mouse (Mammalian omnivore)	Benzene
Heptachlor epoxide	1024-57-3	1024-57-3	5.90E-02	Montane shrew (Mammalian insectivore)	Heptachlor
Isopropylbenzene	98-82-8	98-82-8	2.40E+01	Deer mouse (Mammalian omnivore)	Benzene
Isopropyltoluene[4-]	99-87-6	99-87-6	2.30E+01	Montane shrew (Mammalian insectivore)	Toluene
Methylphenol[4-]	106-44-5	106-44-5	7.90E-01	Generic plant (Terrestrial autotroph - producer)	Phenol
Nitrate (expressed as NO3)	NO3(-1)	14797-55-8			
N-propylbenzene	103-65-1	103-65-1	2.40E+01	Deer mouse (Mammalian omnivore)	Benzene
Perchlorate	ClO4(-1)	14797-73-0			
TATB (triaminotrinitrobenzene)	3058-38-6	3058-38-6			
Trimethylbenzene[1,2,4-]	95-63-6	95-63-6	2.40E+01	Deer mouse (Mammalian omnivore)	Benzene
Trimethylbenzene[1,3,5-]	108-67-8	108-67-8	2.40E+01	Deer mouse (Mammalian omnivore)	Benzene
Tris(o-cresyl) phosphate	1330-78-5	1330-78-5			



CASRN	Chemical	Synonyms	Soil ESL Status	Receptors (Toxicity Data Source)	Release 2.5 Status	Literature Search Status
110-91-6	Benzyl alcohol	Benzylmethanol	New ESL	Mammal	Incorporated	Completed
111-44-4	Bis(2-chloroethyl)ether		No ESLs		No ESLs	Need to search IRIS and ATSDR. 8 toxicity papers need to be reviewed and reviewed for relevance.
74-83-9	Bromomethane	Methyl Bromide	No ESLs New ESL	Mammal	No ESLs Incorporated	Need to search ATSDR; need to update color screen for relevance 1 IRIS, 2 DART and 4 TOXLINE papers; note: don't include articles with inhalation exposure routes
74-87-3	Chloromethane		No ESLs		No ESLs	Need to search ATSDR: 1 toxline and 2 DART papers need to be reviewed and reviewed for relevance; note: don't include articles with inhalation exposure routes
14797-55-8	Nitrate (expressed as NO3)		No ESLs New ESL	Mammal	No ESLs Incorporated	Need to complete review of DART abstracts; need to do TOXLINE literature search; note: don't include articles with drinking water exposure medium
86-71-4	Nitrosodiaz[2]		No ESLs New ESL	Mammal	No ESLs Incorporated	Completed
14797-73-0	Psithoxane		No ESLs New ESL	Mammal	No ESLs Incorporated	Need to do USACHPM, ATSDR, DART and TOXLINE literature search
75-69-4	Tribromofluoromethane	Freon 11	No ESLs New ESL	Mammal	No ESLs Incorporated	Completed
1330-78-5	Tri(n-octyl) phosphite		No ESLs New ESL	Bird	No ESLs Incorporated	Need to retrieve and review DART and TOXLINE papers for relevance
58-05-3	Benzo(a)anthracene		No ESLs New ESL	Bird	No ESLs Incorporated	Completed
129-00-0	Pyrene		No ESLs New ESL	Bird	No ESLs Incorporated	Completed
122-39-4	Diphenylamine		New ESL	Bird (PM)	Incorporated	Applicable data for birds. No data for mammals, plants or worms. IRIS, TOXNET, DART, and TOXNET searches in progress.
74-88-4	Iodomethane		New ESL	Bird (PM)	Incorporated	Applicable data for birds. No applicable data for plants. No data for mammals or worms. Literature searches in progress for IRIS, TOXNET, DART, and TOXNET searches.
591-78-6	Hexanon[2]	Methyl n-Butyl Ketone	New ESL	Mammal (LANL)	Incorporated	Applicable data for birds and mammals. No applicable data for plants. No data for worms. Literature identified in ATSDR and TOXNET; TOXLINE has not been reviewed.
95-48-7	Methylthano[2]	o-cresol	New ESL	Plant (ECOTOX)	Incorporated	Applicable data for plants. No data for birds, mammals or worms. Literature search in progress for IRIS. Literature searches not started for TOXNET, DART, and TOXNET searches.
108-39-4	Methylthano[3]		New ESL	Plant (ECOTOX)	Incorporated	Applicable data for plants. No data for birds, mammals or worms. Literature search in progress for IRIS. Literature searches not started for TOXNET, DART, and TOXNET searches.
319-86-8	BHC[alpha]		Surrogate BHC[gamma]		Surrogate BHC[gamma]	No data was found for birds, mammals, plants or worms. TOXNET, DART, and TOXNET searches have not been reviewed.
91-58-7	Chloronaphthalene[2]		Surrogate Naphthalene		Surrogate Naphthalene	No applicable data for birds, mammals, plants or worms. TOXNET, DART, and TOXNET searches have not been reviewed.
618-01-1	Dibromodiv[3,5]		No ESLs		No ESLs	No applicable data for mammals. No data for birds, plants and worms.
1031-07-8	Endosulfan sulfate		Surrogate Endosulfan		Surrogate Endosulfan	No applicable data for birds. No data for mammals, plants or worms. ATSDR literature search in progress. TOXNET, DART, and TOXNET searches have not been reviewed.
7421-93-4	Endrin aldehyde		Surrogate Endrin		Surrogate Endrin	No applicable data for birds. No data for mammals, plants or worms. ATSDR literature search in progress. TOXNET, DART, and TOXNET searches have not been reviewed.
5349-70-5	Endrin ketone		Surrogate Endrin		Surrogate Endrin	No applicable data for birds. No data for mammals, plants or worms. ATSDR literature search in progress. TOXNET, DART, and TOXNET searches have not been reviewed.
100-41-4	Ethylbenzene		Surrogate Benzene		Surrogate Benzene	No applicable data for plants or worms. No data for birds or mammals. ATSDR and IRIS literature searches in progress. TOXNET, DART, and TOXNET searches have not been reviewed.
1024-57-3	Hepachlor epoxide		Surrogate Hepachlor		Surrogate Hepachlor	No data was found for mammals, plants, or worms. Literature identified in CalEcoTox for birds has not been reviewed. ATSDR and IRIS literature searches in progress. TOXNET, DART, and TOXNET searches have not been reviewed.
FE	Iron		Use ECOSL report	Plant (EcoSL)	Incorporated	Searches supported by EPA EcoSL report for iron. Refer to <a href="http://www.epa.gov/ecoslp/eco-slp/efeco-slp_iron.pdf">http://www.epa.gov/ecoslp/eco-slp/efeco-slp_iron.pdf</a>
98-02-8	Isopropylbenzene		Surrogate Benzene		Surrogate Benzene	No applicable data for birds, mammals, plants or worms. TOXNET, DART, and TOXNET searches have not been reviewed.
98-07-6	Isopropylbenzene[4]		Surrogate Toluene		Surrogate Toluene	No data for birds, mammals, plants or worms. Literature search in progress for IRIS. Literature searches not started for TOXNET, DART, and TOXNET searches.
106-44-5	Methylthano[4]		Surrogate Phenol		Surrogate Phenol	No data for birds, mammals, plants or worms. Literature searches not started for TOXNET, DART, and TOXNET searches.
103-65-1	N-propylbenzene		Surrogate Benzene		Surrogate Benzene	No applicable data for mammals. No data for birds, plants and worms.
3058-38-6	TATB (triaminotrinitrobenzene)		No ESLs		No ESLs	No data for birds, mammals, plants and worms. Literature searches not started for TOXNET, DART, and TOXNET searches.
95-63-6	Trimethylbenzene[2,2,4]		Surrogate Benzene		Surrogate Benzene	No data for birds, mammals, plants and worms. Literature searches not started for TOXNET, DART, and TOXNET searches.
108-67-8	Trimethylbenzene[3,3,5]		Surrogate Benzene		Surrogate Benzene	No data for birds, mammals, plants and worms. Literature searches not started for TOXNET, DART, and TOXNET searches.

On-line Database	URL	Data Provided for ESL Development	Database Description
CEE-TV	<a href="http://www.pwrc.usgs.gov/contaminants-online/pages/CEETV/CEETVintro.htm">http://www.pwrc.usgs.gov/contaminants-online/pages/CEETV/CEETVintro.htm</a>	Not Evaluated. Residue and biomarker data reported, but no data for deriving ingested dose. No plant and worm data.	USGS Contaminant Exposure and Effects-Terrestrial Vertebrates database (CEE-TV) contains contaminant exposure and effects information for terrestrial vertebrates (birds, mammals, amphibians and reptiles) that reside in estuarine and coastal habitats along the Atlantic, Gulf and Pacific Coasts including Alaska and Hawaii and in the Great Lakes Region.
EPA OPPALB	<a href="http://www.epa.gov/oppefed1/ecorisk_de rs/aquatic_life_benchmark.htm">http://www.epa.gov/oppefed1/ecorisk_de rs/aquatic_life_benchmark.htm</a>	Not Evaluated. Aquatic Data ONLY.	USEPA Office of Pesticide Programs' Aquatic Life Benchmarks.
EXTOXNET	<a href="http://extoxnet.orst.edu/">http://extoxnet.orst.edu/</a>	Not Evaluated. Risk Assessment Definitions ONLY.	The EXtension TOXicology NETwork sponsored by the University of California-Davis, Oregon State University, Michigan State University, Cornell University, and the University of Idaho. No toxicity data, only risk assessment definitions. Resource not evaluated.
PAN	<a href="http://www.pesticideinfo.org/Search_Ecotoxicity.jsp">http://www.pesticideinfo.org/Search_Ecotoxicity.jsp</a>	Not Evaluated. Aquatic Data ONLY.	The Pesticide Action Network (PAN) Pesticide Database is your one-stop location for toxicity and regulatory information for pesticides. The PAN Pesticide Database brings together a diverse array of information on pesticides from many different sources, providing human toxicity (chronic and acute), ecotoxicity and regulatory information for about 6,400 pesticide active ingredients and their transformation products, as well as adjuvants and solvents used in pesticide products. Only aquatic ecotoxicity data reported.
TOXNET_ITER	<a href="http://toxnet.nlm.nih.gov/cgi-bin/sis/htmlgen?iter">http://toxnet.nlm.nih.gov/cgi-bin/sis/htmlgen?iter</a>	Not Evaluated. Reports cite IRIS and ATSDR for non-cancer toxicity data.	TOXNET (TOXicology Data NETwork) is a cluster of databases covering toxicology, hazardous chemicals, environmental health and related areas. It is managed by the Toxicology and Environmental Health Information Program (TEHIP) in the Division of Specialized Information Services (SIS) of the National Library of Medicine (NLM). International Toxicity Estimates for Risk (ITER) - Risk information for over 600 chemicals from authoritative groups worldwide.
USGS	<a href="http://www.pwrc.usgs.gov/wwtc/">http://www.pwrc.usgs.gov/wwtc/</a>	Not Evaluated. Links to other databases ONLY.	USGS Whole Wildlife Toxicology Catalog with website links to 40+ databases. No queries performed from this URL or from links not subsequently identified for investigation.
EPA ECOTOX	<a href="http://cfpub.epa.gov/ecotox/">http://cfpub.epa.gov/ecotox/</a>	Toxicity Values for Birds, Mammals, Plants and Worms.	USEPA ECOTOX Database. The ECOTOX (ECOTOXicology) database provides single chemical toxicity information for aquatic and terrestrial life. Values reported include LC50s, NOEC, LOEC, LOEL, NOEL, ED50, etc. Toxicity data for available substances is reported in worksheet "ECOTOX." Only terrestrial data for Growth, Mortality, Reproduction and Population queried from database. Searched by CASRN.
ORNL Invertebrates	<a href="http://www.esd.ornl.gov/programs/ecorisk/documents/tm85r3.pdf">http://www.esd.ornl.gov/programs/ecorisk/documents/tm85r3.pdf</a>	Toxicity Values for Worms.	The ORNL (Oak Ridge National Laboratory) ecotoxicological screening benchmarks for earthworms and microbes/microbial processes are concentrations of chemicals in soil that are believed to represent acceptable concentrations with respect to selected ecological receptors. The benchmarks are based on NOECs and LOECs, which are reported for soil exposures to earthworm and microbes/ microbial processes. See PDF resource downloadable from website.
ORNL Plants	<a href="http://www.esd.ornl.gov/programs/ecorisk/documents/tm85r3.pdf">http://www.esd.ornl.gov/programs/ecorisk/documents/tm85r3.pdf</a>	Toxicity Values for Plants.	The ORNL (Oak Ridge National Laboratory) ecotoxicological screening benchmarks for terrestrial plants are concentrations of chemicals in soil or solution that are believed to represent acceptable concentrations with respect to selected ecological receptors. The benchmarks are based on NOECs and LOECs, which are reported for soil or solution exposures to plants. See PDF resource downloadable from website.
ORNL Wildlife	<a href="http://www.esd.ornl.gov/programs/ecorisk/documents/tm86r3.pdf">http://www.esd.ornl.gov/programs/ecorisk/documents/tm86r3.pdf</a>	Toxicity Values for Birds and Mammals.	The ORNL (Oak Ridge National Laboratory) ecotoxicological screening benchmarks for wildlife are concentrations of chemicals in ambient media that are believed to represent acceptable concentrations with respect to selected ecological receptors. The benchmarks are based on NOELs and LOELs, which are reported for oral exposures in birds and mammals. See PDF resource downloadable from website.

On-line Database	URL	Data Provided for ESL Development	Database Description
LANL PTSE (New Values)	Not available	Toxicity Values for Birds, Mammals, Plants and Worms.	The LANL (Los Alamos National Laboratory) Ecorisk Database contains Ecological Screening Levels for aquatic and terrestrial receptors in soil, sediment, water and air. The LANL ESLs are based on TRVs, which are derived using the Primary Toxicity Study Evaluation Process. TRVs reported are chronic NOAELs/NOECs for terrestrial birds, mammals, plants and earthworms.
ABC Birds	<a href="http://www.abcbirds.org/abcprograms/policy/pesticides/aims/aims/toxicity.cfm">http://www.abcbirds.org/abcprograms/policy/pesticides/aims/aims/toxicity.cfm</a>	Toxicity Values for Birds.	The American Bird Conservancy (ABC) Pesticide Toxicity Database contains acute pesticide toxicity data in birds.
Cal/ECOTOX	<a href="http://www.oehha.org/cal_ecotox/default.htm">http://www.oehha.org/cal_ecotox/default.htm</a>	Toxicity Data for Birds and Mammals for Oral Non-cancer Toxicity Data. References also reported.	The California OHHEA (Office of Environmental Health Hazard Assessment) Wildlife Biology, Exposure Factor, and Toxicity Database (Cal/ECOTOX) is a compilation of physiological and ecological parameters and toxicity data for a number of California fish and wildlife. Species, chemical, endpoint type, endpoint description, endpoint value, endpoint range, study description and reference are reported. Data for chemicals of interest reported in worksheet "CalECOTOX".
ERED	<a href="http://el.erdc.usace.army.mil/ered/Index.cfm">http://el.erdc.usace.army.mil/ered/Index.cfm</a>	Toxicity Values for Birds.	The U.S. Army Corps of Engineers/U.S. Environmental Protection Agency Environmental Residue-Effects Database (ERED) is a compilation of data, taken from the literature, where biological effects (e.g., reduced survival, growth, etc.) and tissue contaminant concentrations were simultaneously measured in the same organism. Currently, the database is limited to those instances where biological effects observed in an organism are linked to a specific contaminant within its tissues.
IPM Centers	<a href="http://www.ipmcenters.org/ECOTOX/DataAccess.cfm">http://www.ipmcenters.org/ECOTOX/DataAccess.cfm</a>	Toxicity Values for Birds, Mammals, Plants and Worms.	USEPA NATIONAL INFORMATION SYSTEM FOR THE REGIONAL IPM (Integrated Pest Management) Centers OPP Pesticide Ecotoxicity Database. The Ecological Fate and Effects Division of the USEPA Office of Pesticide Programs is continuing efforts to update the database with all EPA reviewed ecotoxicity endpoints for pesticides registered or previously registered in the U.S. Toxicity data on over 800 active ingredients, metabolites, and multi-ingredient formulations are presently included in the database. The toxicity data inputted into the database is compiled from actual studies reviewed by EPA in conjunction with pesticide registration or reregistration and studies performed by USEPA, USDA and USFWS laboratories which have been reviewed by Agency biologists and judged acceptable for use in the ecological risk assessment process. The database presently contains over 21,000 records for acute and chronic toxicity endpoints on terrestrial and aquatic plants, aquatic invertebrates, terrestrial invertebrates, insects, amphibians, fish, birds, reptiles, and wild mammals. The database is presented in Microsoft ACCESS and contains 35 fields per record entry. Each record entry sum
USACHPPM	<a href="http://chppm-www.apqea.army.mil/erawg/tox/index.htm">http://chppm-www.apqea.army.mil/erawg/tox/index.htm</a>	Toxicity Values for Birds and Mammals.	USACHPPM (US Army Center for Health Promotion and Preventive Medicine) complete chemical toxicological assessments/profiles for wildlife with reference list.
ATSDR	<a href="http://www.atsdr.cdc.gov/">http://www.atsdr.cdc.gov/</a>	Reference Titles for Mammal Oral Non-cancer Toxicity Data.	ATSDR (Agency for Toxic Substances and Disease Registry) Toxicological Profiles for human health. These profiles succinctly characterize the toxicologic and adverse health effects information for a hazardous substance. Each peer-reviewed profile identifies and reviews the key literature that describes a hazardous substance's toxicologic properties. Other pertinent literature is also presented, but is described in less detail than the key studies. The references are generally for mammalian studies all routes.

On-line Database	URL	Data Provided for ESL Development	Database Description
IRIS	<a href="http://www.epa.gov/ncea/iris/search_key_word.htm">http://www.epa.gov/ncea/iris/search_key_word.htm</a>	Reference Abstracts for Mammals for Oral Non-cancer Toxicity Data.	<p>The USEPA Integrated Risk Information System (IRIS) is an electronic database containing information on human health effects that may result from exposure to various substances in the environment. IRIS is prepared and maintained by the EPA's National Center for Environmental Assessment (NCEA) within the Office of Research and Development (ORD). Noncancer effects: Oral reference doses and inhalation reference concentrations (RfDs and RfCs, respectively) for effects known or assumed to be produced through a nonlinear (possibly threshold) mode of action. In most instances, RfDs and RfCs are developed for the noncarcinogenic effects of substances. Cancer effects: Descriptors that characterize the weight of evidence for human carcinogenicity, oral slope factors, and oral and inhalation unit risks for carcinogenic effects. Where a nonlinear mode of action is established, RfD and RfC values may be used. Primary toxicity study references for mammalian test species are reported.</p>
TOXNET_DART/ETIC	<a href="http://toxnet.nlm.nih.gov/cgi-bin/sis/htmlgen?DARTETIC">http://toxnet.nlm.nih.gov/cgi-bin/sis/htmlgen?DARTETIC</a>	Reference Abstracts for Mammals for Oral Non-cancer Toxicity Data.	<p>TOXNET (TOXicology Data NETwork) is a cluster of databases covering toxicology, hazardous chemicals, environmental health and related areas. It is managed by the Toxicology and Environmental Health Information Program (TEHIP) in the Division of Specialized Information Services (SIS) of the National Library of Medicine (NLM). DART@ETIC (Development and Reproductive Toxicology/Environmental Teratology Information Center) is a bibliographic database covering literature on reproductive and developmental toxicology. DART is managed by NLM and funded by the EPA, the National Institute of Environmental Health Sciences (NIEHS) and NLM. DART/ETIC contains references to reproductive and developmental toxicology literature published since 1965.</p>
TOXNET_TOXLINE	<a href="http://toxnet.nlm.nih.gov/cgi-bin/sis/htmlgen?TOXLINE">http://toxnet.nlm.nih.gov/cgi-bin/sis/htmlgen?TOXLINE</a>	Reference Abstracts for Mammals for Oral Non-cancer Toxicity Data.	<p>TOXNET (TOXicology Data NETwork) is a cluster of databases covering toxicology, hazardous chemicals, environmental health and related areas. It is managed by the Toxicology and Environmental Health Information Program (TEHIP) in the Division of Specialized Information Services (SIS) of the National Library of Medicine (NLM). TOXLINE® is a bibliographic database providing comprehensive coverage of the biochemical, pharmacological, physiological, and toxicological effects of drugs and other chemicals from 1965 to the present. TOXLINE contains over 3 million citations, almost all with abstracts and/or index terms and CAS Registry Numbers.</p>

CASRN	Chemical	LANL PTSE	EPA ECOTOX	ORNL Wildlife	ORNL Plants	ORNL Invertebrates	ABC Birds	Cu/Cobalt	ERED	IPM Centers	USACHPM	ATSDR	IRIS	TONNET DARTTTC	TONNET TOXLINE	
100516	Benzyl alcohol	Chemical Not in Database	No Applicable Data for Plants - 2 results reported; age is the exposure medium - not an ecologically relevant exposure medium. No Bird, Mammal or Worm data for chemical in database.	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	2 applicable abstracts out of first 100 reviewed	5 applicable abstracts out of first 100 reviewed						
315-86-8	BHC[alpha]	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Literature Search Not Started	Literature Search Not Started	
114-44-4	BHQ-chlorothal[ether]	Chemical Not in Database	No Applicable Data for Worms - 2 results reported; FLT (filar paper) is the exposure medium - not an ecologically relevant exposure medium. No Bird, Mammal or Plant data for chemical in database. No Applicable Data for Plants - 56 results reported; 56 results not used; 50 results excluded because the exposure route is FJ (fumigation) - not an ecologically relevant exposure route. 1 result excluded because endpoint is NR (not reported). No Bird, Mammal or Worm data for chemical in database.	Chemical Not in Database	No Bird, Mammal, Plant or Worm data for chemical in database.	Chemical Not in Database	Chemical Not in Database	Need to search Database	Literature Search In Progress; Need to search Database	Chemical Not in Database	6 applicable abstracts out of first 100 reviewed					
74-83-9	Bromomethane	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Need to search Database	Human Health oral RFD for epithelial hyperplasia of the forestomach based on gavage study in rats - body weight changes were measured - 1 applicable abstract	2 applicable abstracts out of 25 reviewed	4 applicable abstracts out of first 100 reviewed	
86-74-8	Carbazole	Chemical Not in Database	No Applicable Data for Birds, Mammals, Plants, or Worms. 2 results reported for rodents - not a relevant test species.	Chemical Not in Database	No Applicable Data; Aquatic Only	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	2 applicable abstracts out of 60 abstracts reviewed					
75-10-0	Carbon disulfide	Chemical Not in Database	Intram EEL data for Mammals; 90 results reported; 32 results used; 20 results not used because exposure route is IP (intraperitoneal) - not an ecologically relevant exposure route. 5 results not used because exposure route TP (topical, general) - also not an ecologically relevant exposure route. 15 results not used because effect is MPH (morphology) - not an ecologically relevant effect. 5 results not used because endpoint not reported. 8 results not used because a single endpoint value not reported. 5 results not used because reported in units of mortality. No Applicable Plant Data - 20 results reported; 20 results not used; 23 results excluded because exposure route is SO (soaked or dipped) - not an ecologically relevant exposure route. No Bird or Worm data for this chemical in database. 67 results reported for insects/rodents - not relevant test species.	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Literature Search In Progress	No Applicable Data; Human Health oral RFD based on inhalation study in rabbits - Not an Applicable Exposure Route	Literature Search Not Started	Literature Search Not Started					
106-47-8	Chloroacetal[4]	Chemical Not in Database	Intram EEL data for Plants and Worms; 17 results reported for plants; 17 results used; 3 results reported for worms; 1 result used; No Bird or Mammal data for chemical in database.	Chemical Not in Database	No Applicable Data for Plants; exposure medium a solution, not considered an ecologically relevant exposure medium for soil TRV.	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Literature Search In Progress	0 applicable abstracts out of 6 abstracts reviewed	Literature Search In Progress					
74-83-9	Chloromethane	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Need to search Database	Chemical Not in Database	2 applicable abstracts out of first 100 sites reviewed	Literature Search Not Started	
91-06-7	Chlorophthalate[2-]	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Literature Search Not Started	Literature Search Not Started	
95-50-1	Dichlorobenzene[1,2]	Chemical Not in Database	Intram EEL data for Mammals; 13 results reported; 1 result used; 10 results not used because exposure route is IP (intraperitoneal) - not an ecologically relevant exposure route. 1 result not used because effect is MPH (morphology) - not an ecologically relevant effect. 1 result not used because endpoint not reported. No Applicable Plant Data - 2 results reported, but exposure medium hypothesis; Not an ecologically relevant exposure medium. No Applicable Worm Data - 2 results reported, but exposure medium filar paper - Not an ecologically relevant exposure medium. No Bird data for this chemical in database; 2 results reported for Fungus - not a relevant test species.	Chemical Not in Database	No Applicable Data; Aquatic Only	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Literature Search In Progress	Chemical Not in Database	Literature Search Not Started	Literature Search Not Started				
541-71-3	Dichlorobenzene[1,3]	Chemical Not in Database	Intram EEL data for Mammals; 6 results reported; 2 results used; 2 results not used because exposure route is IP (intraperitoneal) - not an ecologically relevant exposure route. 2 results not used because effect is MPH (morphology) - not an ecologically relevant effect. No Bird, Plant or Worm data for this chemical in database.	Chemical Not in Database	No Applicable Data; Aquatic Only	Chemical Not in Database	Chemical Not in Database	Literature Search In Progress	Literature Search In Progress	Literature Search Not Started	Literature Search Not Started					
618-87-1	Dihydroindole[3,5]	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	6 applicable abstracts out of 35 abstracts reviewed	
122-39-4	Dihydrofuran	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Need to search Database	Need to search Database	
101-07-4	Ethylphenyl sulfide	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	No Applicable Data	Chemical Not in Database	Literature Search In Progress	Chemical Not in Database	Literature Search Not Started	Literature Search Not Started	
7421-93-5	Endrin aldehyde	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	No Applicable Data	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Literature Search Not Started	Literature Search Not Started	
5304-70-5	Endrin ketone	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	No Applicable Data	Chemical Not in Database	Literature Search In Progress	Chemical Not in Database	Literature Search Not Started	Literature Search Not Started	
100-41-4	Ethylbenzene	Chemical Not in Database	No Applicable Data for Plants - 1 result reported; 1 result used and 1 result excluded because exposure route is SO (soaked or dipped) - not an ecologically relevant exposure route. No Applicable Data for Worms - 2 results reported, but exposure medium is filar paper - not an ecologically relevant exposure medium. No Bird or Mammal data for chemical in database.	Chemical Not in Database	See worksheet "Cu/Cobalt"; American Health, Double-crested Cormorant, Great Horned Owl, Peregrine Falcon	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Literature Search In Progress	Literature Search In Progress	Literature Search Not Started	Literature Search Not Started				
102-57-3	Heptachlor epoxide	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	No Applicable Data; Aquatic Only	No Applicable Data	Chemical Not in Database	Literature Search In Progress	Literature Search In Progress	Literature Search Not Started	Literature Search Not Started	
118-74-1	Heptachlorbenzene	Chemical Not in Database	Intram EEL data for Bird, Mammal, Plant and Worm; 174 results reported for birds; 1 result used; 84 results not used because the effect is MPH (morphology) - not an ecologically relevant effect. 84 results excluded because endpoint not reported. 4 results excluded because data only ppm. 3 results excluded because LC50s; 91 results reported for mammals; 1 result used; 6 results not used because effect is MPH (morphology) - not an ecologically relevant effect. 84 results reported for plants; 27 results used; 2 results excluded because exposure duration not reported; 2 results excluded because LC50s for CRD (growth) and an LC50 should be for mortality. 2 results reported for worms; 1 result used; 1 result excluded because exposure medium FLT (filar paper) - not an ecologically relevant exposure medium. No Mammal data.	Chemical Not in Database	No Applicable Data; Aquatic Only	No Applicable Data	Chemical Not in Database	Literature Search In Progress	IRIS oral RFD for liver effects in chronic feeding study in rats - pup mortality measured (POTENTIAL) APPLICABLE data in Animal, D.L., C.A. Moore, S.M. Chastrom, et al. 1985. Long-term toxicity of heptachlorobenzene in the rat and the effect of gender. Water, Air, & Soil Chem. Toxic 23(9): 779-793.	Literature Search Not Started	Literature Search Not Started					
591-76-6	Hexachlor[2]	Bird and Mammal Data; No Data for Plants and Worms	No Applicable Data for Plants - 2 results reported; age is the exposure medium - not an ecologically relevant exposure medium. No Bird, Mammal or Worm data for chemical in database.	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	0 applicable abstracts out of 6 abstracts reviewed.	3 applicable abstracts out of 466 abstracts reviewed; 3 of 3 mammal papers reviewed have applicable data.						
74-88-4	Iodobenzene	Chemical Not in Database	No Applicable Data for Plants - 2 results reported; FLT (filar paper) is the exposure medium - not an ecologically relevant exposure medium. No Bird, Mammal or Plant data for chemical in database.	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Need to search Database	Need to search Database						
74-89-4	Iron	Chemical Not in Database	No Applicable Data for Birds; 1 result reported; endpoint NR. No Applicable Data for Plants - 2 results reported; AQU (aqueous/hydrolysis) is the exposure medium - not an ecologically relevant exposure medium. No Mammal or Worm data for chemical in database.	Chemical Not in Database	No Applicable Data for Plants; exposure medium a solution, not considered an ecologically relevant exposure medium for soil TRV.	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Literature Search Not Started	Literature Search Not Started				
98-82-8	Isopropylbenzene	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Literature Search Not Started	Literature Search Not Started	
99-07-6	Isopropylbenzene[4]	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Literature Search Not Started	Literature Search Not Started	
74-83-5	Lithium	Mammal Data; No Applicable Data for Birds - data reported is for drinking water exposure medium - not considered an ecologically relevant exposure medium for a soil TRV. No Plant or Worm Data.	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Literature Search Not Started	Literature Search Not Started
95-49-7	Methylthio[2-]	Chemical Not in Database	No Applicable Data for Plants; 12 results reported for plants; 5 results used; 4 results excluded because the exposure medium HYP (hydroponic) - not an ecologically relevant exposure medium. 1 result excluded because the exposure medium AGR (agar) - not an ecologically relevant exposure medium. 2 results excluded because the exposure duration NR (not reported). No Bird, Mammal or Worm data for chemical in database.	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Literature Search In Progress	Literature Search Not Started	Literature Search Not Started					
108-30-4	Methylthio[3-]	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Literature Search In Progress	Literature Search Not Started	
106-44-5	Methylthio[4-]	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Literature Search In Progress	Literature Search Not Started	
74-83-5	Molybdenum	Chemical Not in Database	No Applicable Data for Mammals; 10 results reported; DR (drinking water) is the exposure route - not an ecologically relevant exposure medium. No Bird, Plant or Worm data for chemical in database.	Chemical Not in Database	See worksheet "Cu/Cobalt"; American Health, Double-crested Cormorant, Great Horned Owl, Peregrine Falcon	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Chemical Not in Database	Literature Search Not Started	Literature Search Not Started				

CASN	Chemical	LANL PTSE	EPA ECOTOX	ORNL Wildlife	ORNL Plants	ORNL Invertebrata	ABC Birds	CatEcotox	ERED	IPM Centers	USACHPPM	ATSDR	IRIS	TONNET_DARTIETC	TONNET_TOXLINE
14377-53-8	Nitrate (expressed as NO3)	Chemical Not In Database	No Applicable Data for Plants - 8 results reported; exposure medium - not an ecologically relevant exposure medium for birds. No Applicable Data for Birds - Data is reported without a literature source.	No Applicable Data for Mammals - drinking water is the exposure medium - not an ecologically relevant exposure medium for birds. No Applicable Data for Birds - Data is reported without a literature source.	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Sodium nitrate only, see worksheet "CatEcotox" - No Applicable Data	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	No Applicable Data. Human Health oral RFD based on study in humans measuring early clinical signs of methemoglobinemia in excess of 12% - Not an Applicable Test Species or Ecologically Relevant Effect.	in progress-reviewed through #5	Need to search Database
88-74-4	Nitroethane[2]	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Need to search Database
103-65-1	N-propylbenzene	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	No Applicable Data, Aquatic Only	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Literature Search Not Started	Literature Search Not Started
14397-73-0	Pentachlorate	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	No Applicable Data, Aquatic Only	Chemical Not In Database	Need to search Database	Need to search Database	No Applicable Data. Human Health oral RFD based on study in humans measuring radiocesium uptake inhibition (RADI) in the thyroid - Not an Applicable Test Species or Ecologically Relevant Effect.	Need to search Database	Need to search Database
100-42-5	Styrene	Chemical Not In Database	Interim ESL data for Mammals, 3 results used; 6 results excluded because exposure NR (not reported). No Applicable Data for Plants - 4 results reported; 2 results excluded because the exposure medium is PVP (theophylline) - not an ecologically relevant exposure medium; 2 results excluded because the exposure duration is NR (not reported); No Bird or Mammal data for chemical in database.	Chemical Not In Database	See worksheet "Interim TRV": interim ESL data for Plants	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Literature Search In Progress	Literature Search In Progress	Literature Search In Progress	Literature Search In Progress
3058-38-6	TATB (triaminotriazobenzene)	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	1 applicable abstract out of 11 abstracts reviewed
75-85-4	Tetrafluoroethane	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Literature Search Not Started	Literature Search Not Started
95-83-6	Trimethylbenzene[1,2,4]	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Literature Search Not Started	Literature Search Not Started
106-67-8	Trimethylbenzene[1,3,5]	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Literature Search Not Started	Literature Search Not Started
1330-79-5	Tri(n)-cetyl phosphate	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	No Bird, Mammal, Plant or Worm data for chemical in database.	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	4 applicable abstracts out of 24 reviewed	12 applicable abstracts out of 100 reviewed
75-01-4	Vinyl chloride	Chemical Not In Database	Chemical Not In Database	Mammal Data, No Bird Data.	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Chemical Not In Database	Literature Search In Progress	IRIS oral RFD for low oral polychlorinated biphenyls in a chronic feeding study in rats - body weight and mortality were measured. POTENTIALLY APPLICABLE data in TL, HP; Interim, HR; Farn, VJ (1985) Lifespan oral carcinogenicity study of vinyl chloride in rats. Final report. CIVC Institute, THD Report No. 8.8.285(2)1000. TSCATS Document P-15-AJ-0184-0203. Farn, No. 0203 and TL, HP; Farn, VJ, Interim, HR. (1991) Lifetime (400 week) oral carcinogenicity study of vinyl chloride in rats. Food Chem Toxicol 29:713-716.	Literature Search Not Started	Literature Search Not Started

Chemical of Concern	Receptor Group	TRV	Units	Description	Data	
					Source	Status
Benzyl alcohol						
(Benzenemethanol)	Bird			None. No data.		
Benzyl alcohol						
(Benzenemethanol)	Plant			None. No applicable data.		
Benzyl alcohol						
(Benzenemethanol)	Worm			None. No data.		
BHC[delta-]	Bird			None. No data.		
BHC[delta-]	Mammal			None. No data.		
BHC[delta-]	Plant			None. No data.		
BHC[delta-]	Worm			None. No data.		
Bis(2-chloroethyl)ether	Bird			None. No data.		
Bis(2-chloroethyl)ether	Mammal			None. No data.		
Bis(2-chloroethyl)ether	Plant			None. No data.		
Bis(2-chloroethyl)ether	Worm			None. No applicable data.		
Bromomethane	Bird			None. No data.		
Bromomethane	Mammal			None. No data.		
Bromomethane	Plant			None. No applicable data.		
Bromomethane	Worm			None. No data.		
Carbazole	Bird			None. No applicable data.		
Carbazole	Plant			None. No applicable data.		
Carbazole	Worm			None. No applicable data.		
Carbon Disulfide	Bird			None. No data.		
Carbon Disulfide	Plant			None. No applicable data.		
Carbon Disulfide	Worm			None. No data.		
Chloroaniline[4-]	Bird			None. No data.		
Chloroaniline[4-]	Mammal			None. No applicable data.		
Chloromethane	Bird			None. No data.		
Chloromethane	Mammal			None. No data.		
Chloromethane	Plant			None. No data.		
Chloromethane	Worm			None. No data.		
Chloronaphthalene[2-]	Bird			None. No data.		
Chloronaphthalene[2-]	Mammal			None. No data.		
Chloronaphthalene[2-]	Plant			None. No data.		
Chloronaphthalene[2-]	Worm			None. No data.		
Dichlorobenzene[1,2-]	Bird			None. No data.		
Dichlorobenzene[1,2-]	Plant			None. No applicable data.		
Dichlorobenzene[1,2-]	Worm			None. No applicable data.		
Dichlorobenzene[1,3-]	Bird			None. No data.		
Dichlorobenzene[1,3-]	Plant			None. No data.		
Dichlorobenzene[1,3-]	Worm			None. No data.		
Dinitroaniline[3,5-]	Bird			None. No data.		
Dinitroaniline[3,5-]	Mammal			None. No data.		
Dinitroaniline[3,5-]	Plant			None. No data.		
Dinitroaniline[3,5-]	Worm			None. No data.		
Diphenylamine	Mammal			None. No data.		
Diphenylamine	Plant			None. No data.		
Diphenylamine	Worm			None. No data.		
Endosulfan sulfate	Bird			None. No data.		
Endosulfan sulfate	Mammal			None. No data.		
Endosulfan sulfate	Plant			None. No data.		
Endosulfan sulfate	Worm			None. No data.		
Endrin aldehyde	Bird			None. No data.		
Endrin aldehyde	Mammal			None. No data.		
Endrin aldehyde	Plant			None. No data.		
Endrin aldehyde	Worm			None. No data.		
Endrin ketone	Bird			None. No data.		
Endrin ketone	Mammal			None. No data.		
Endrin ketone	Plant			None. No data.		
Endrin ketone	Worm			None. No data.		
Ethylbenzene	Bird			None. No data.		
Ethylbenzene	Mammal			None. No data.		
Ethylbenzene	Plant			None. No applicable data.		
Ethylbenzene	Worm			None. No applicable data.		
Heptachlor epoxide	Bird			None. No data.		
Heptachlor epoxide	Mammal			None. No data.		
Heptachlor epoxide	Plant			None. No data.		
Heptachlor epoxide	Worm			None. No data.		
Hexanone[2-]	Plant			None. No applicable data.		

Chemical of Concern	Receptor Group	TRV	Units	Description	Data	
					Source	Status
Hexanone[2-]	Worm			None. No data.		
Iodomethane	Mammal			None. No data.		
Iodomethane	Plant			None. No data.		
Iodomethane	Worm			None. No data.		
Iron	Bird			None. No applicable data.		
Iron	Mammal			None. No data.		
Iron	Plant			None. No applicable data.		
Iron	Worm			None. No applicable data.		
Isopropylbenzene	Bird			None. No data.		
Isopropylbenzene	Mammal			None. No data.		
Isopropylbenzene	Plant			None. No data.		
Isopropylbenzene	Worm			None. No data.		
Isopropyltoluene[4-]	Bird			None. No data.		
Isopropyltoluene[4-]	Mammal			None. No data.		
Isopropyltoluene[4-]	Plant			None. No data.		
Isopropyltoluene[4-]	Worm			None. No data.		
Lithium	Bird			None. No applicable data.		
Lithium	Worm			None. No data.		
Methylphenol[2-]	Bird			None. No data.		
Methylphenol[2-]	Mammal			None. No data.		
Methylphenol[2-]	Worm			None. No data.		
Methylphenol[3-]	Bird			None. No data.		
Methylphenol[3-]	Mammal			None. No data.		
Methylphenol[3-]	Worm			None. No data.		
Methylphenol[4-]	Bird			None. No data.		
Methylphenol[4-]	Mammal			None. No data.		
Methylphenol[4-]	Plant			None. No data.		
Methylphenol[4-]	Worm			None. No data.		
Molybdenum	Mammal			None. No applicable data.		
Molybdenum	Worm			None. No data.		
Nitrate	Bird			None. No applicable data.		
Nitrate	Mammal			None. No applicable data.		
Nitrate	Plant			None. No applicable data.		
Nitrate	Worm			None. No data.		
Nitroaniline[2-]	Bird			None. No data.		
Nitroaniline[2-]	Plant			None. No data.		
Nitroaniline[2-]	Worm			None. No data.		
N-propylbenzene	Bird			None. No data.		
N-propylbenzene	Mammal			None. No data.		
N-propylbenzene	Plant			None. No data.		
N-propylbenzene	Worm			None. No data.		
Perchlorate	Bird			None. No data.		
Perchlorate	Mammal			None. No data.		
Perchlorate	Plant			None. No data.		
Perchlorate	Worm			None. No data.		
Styrene (Ethenylbenzene)	Bird			None. No data.		
Styrene (Ethenylbenzene)	Mammal			None. No data.		
Trichlorofluoromethane	Bird			None. No data.		
Trichlorofluoromethane	Plant			None. No data.		
Trichlorofluoromethane	Worm			None. No data.		
Trimethylbenzene[1,2,4-]	Bird			None. No data.		
Trimethylbenzene[1,2,4-]	Mammal			None. No data.		
Trimethylbenzene[1,2,4-]	Plant			None. No data.		
Trimethylbenzene[1,2,4-]	Worm			None. No data.		
Trimethylbenzene[1,3,5-]	Bird			None. No data.		
Trimethylbenzene[1,3,5-]	Mammal			None. No data.		
Trimethylbenzene[1,3,5-]	Plant			None. No data.		
Trimethylbenzene[1,3,5-]	Worm			None. No data.		
Tris(o-cresyl) phosphate	Bird			None. No applicable data.		
Tris(o-cresyl) phosphate	Mammal			None. No data.		
Tris(o-cresyl) phosphate	Plant			None. No data.		
Tris(o-cresyl) phosphate	Worm			None. No data.		
Vinyl Chloride	Bird			None. No data.		
Vinyl Chloride	Plant			None. No data.		
Vinyl Chloride	Worm			None. No data.		

	<u>Bird or Mammal :</u>	<u>Invertebrate or</u>
		<u>Plant:</u>
Chronic	> 90 days	> 6 days
Subchronic	14 to 90 days	3 to 6 days
Acute	< 14 days	< 3 days

LANL, 2010. Toxicity Reference Value Development Methods for the Los Alamos National Laboratory. LA-UR-10-4922. EP2010-0279

UF applied to derive a TRV that is  
a Chronic NOAEL (NOEC)-based  
EL:

Type of effect level available	
C-CL or chronic NOAEL (NOEC)	1
C-CL or chronic LOAEL (LOEC)	10
C-CL or chronic LD50 (LC50)	100
C-CL or chronic ED50 (EC50)	100
Subchronic NOAEL (NOEC)	10
Subchronic LOAEL (LOEC)	100
Subchronic LD50 (LC50)	100
Subchronic ED50 (EC50)	100
Acute or Single Dose NOAEL (NOEC)	100
Acute or Single Dose LOAEL (LOEC)	100
Acute or Single Dose LD50 (LC50)	100
Acute or Single Dose ED50 (EC50)	100

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