BIOACCUMULATION RISK ASSESSMENT MODELING SYSTEM (BRAMS)

USERS GUIDE

Prepared By:
Kelsie Baker, John Thomas Vogel II, and Alex Tkachuk
U.S. Army Engineer Research and Development Center, Environmental Lab
Igor.Linkov@usace.army.mil

Prepared For:
U.S. Army Corps of Engineers
U.S. Environmental Protection Agency
Table of Contents

Table of Contents ................................................................................................................................... 2

List of Figures ......................................................................................................................................... 4

Introduction ........................................................................................................................................... 5

PART I: Getting Started .......................................................................................................................... 5
  1. What is BRAMS? ................................................................................................................................. 5
  2. Background ......................................................................................................................................... 5
  3. Installation and StartUp ...................................................................................................................... 8
    3.1 System Requirements .................................................................................................................... 8
    3.2 Installation .................................................................................................................................... 8
    3.3 StartUp .......................................................................................................................................... 8
  4. User Interface ..................................................................................................................................... 8
    4.1 Main Window ................................................................................................................................. 8
    4.2 Menu ........................................................................................................................................... 9
    4.3 Toolbar ......................................................................................................................................... 11
    4.4 Notes Pane .................................................................................................................................. 12
    4.5 Explorer Pane ............................................................................................................................... 13
    4.6 Table Pane ................................................................................................................................... 14
    4.7 Properties Pane ............................................................................................................................ 14
    4.8 ReportViewer Window .................................................................................................................. 16

PART II: Building the Model ................................................................................................................ 18
  1. User Identification ............................................................................................................................. 18
  2. Create or open a project or template ............................................................................................... 18
  3. Add a new item .................................................................................................................................. 18
    4. Add or edit project and item properties ......................................................................................... 18
      4.1 General ....................................................................................................................................... 19
      4.2 By model type and item category .............................................................................................. 20
      4.3 Import Data From Excel (BEST Model) ...................................................................................... 26
  5. Using Templates .................................................................................................................................. 27
  6. Locking and Unlocking BEST Model Properties ............................................................................ 27
  7. Saving Permanently .......................................................................................................................... 28

Part III: Results ..................................................................................................................................... 29
  1. Generate Report ............................................................................................................................... 29
  2. Outputs ............................................................................................................................................. 29

Part IV: Step by Step Model Creation ................................................................................................. 35
  1. BEST Model ..................................................................................................................................... 35
    STEP ONE: Getting Started ............................................................................................................. 35
    STEP TWO: Open the appropriate BEST template file ..................................................................... 35
    STEP THREE (Optional): Add or Edit Items and Properties ............................................................. 36
    STEP FOUR: Import 28 Day Bioaccumulation Reference Data ....................................................... 36
    STEP FIVE: Name and save the reference site project ..................................................................... 37
    STEP SIX: Import 28 Day Bioaccumulation Test Data ..................................................................... 37
    STEP FIVE: Name and save the test site project .............................................................................. 37
    STEP SEVEN: Specify Reference Project .......................................................................................... 37
List of Figures

Figure 1: Main BRAMS Window ................................................................................................... 9
Figure 2: BRAMS main toolbar .................................................................................................. 11
Figure 3: Notes Pane with example text .................................................................................... 13
Figure 4: TT (left) and BEST (right) Explorer Panes ................................................................. 14
Figure 5: Example BEST model Table pane showing defined values of characteristics of invertebrates including their Name, Lipid Content, and Normalization .................................................. 14
Figure 6: Example TT model Table pane showing defined characteristics of invertebrates including their Name, Environment, Lipid, and Diet Pathway ........................................................................ 14
Figure 7: Example TT model Properties pane showing input properties of an invertebrate including its Type, Name, Environment, Lipid content, Diet Pathway, and Bioaccumulation Test results ........................................................................................................... 15
Figure 8: Example BEST model Properties pane showing input properties of an invertebrate including its Type, Name, Lipid content, Normalization, and Bioaccumulation Test results ...... 16
Figure 9: Example TT model ReportViewer window page 1 .................................................... 17
Figure 10: Excel Chemicals Input Sheet, Columns A-E. .......................................................... 26
Figure 11: Excel Chemicals Input Sheet, Columns F-K ............................................................ 27
Figure 12: Excel Bioaccumulation Test Results Input Sheet .................................................... 27
Figure 13: TT ecological receptor properties report section .................................................... 29
Figure 14: TT ecological receptor Risk List report section ..................................................... 30
Figure 15: TT Exposure Concentration report section ............................................................ 30
Figure 16: TT Risk Figures report section for ecological receptor .......................................... 31
Figure 17: BEST Summary Report sections .............................................................................. 32
Figure 18: BEST Full Report sections ..................................................................................... 33
Figure 19: BEST Model Report Identification section ............................................................. 34
Introduction

This manual describes the Bioaccumulation Risk Assessment Modeling System (BRAMS) program components and features, guides the user in creating both Trophic Trace (TT) and Bioaccumulation Evaluation Screening Tool (BEST) models and details the assumptions and equations used in each model.

PART I: Getting Started

1. What is BRAMS?1*

The Bioaccumulation Risk Assessment Modeling System (BRAMS) is a stand-alone tool for calculating the potential human health and ecological risks associated with bioaccumulation of contaminants in dredged sediments. It contains two separate models, Trophic Trace (TT) and the Bioaccumulation Evaluation Screening Tool (BEST), which use separate equations and inputs to calculate risks. The stand-alone version of Trophic Trace was updated and recoded for the version implemented within BRAMS though it uses the same technical framework and retains all the functionality of the stand-alone Trophic Trace version 4.1. The BRAMS program is designed to provide health- and ecologically-protective estimates of potential risk using results from sediment chemistry tests and/or 28-day bioaccumulation tests. The program calculates the risks based on the characteristics of the site including the environment, species and chemicals involved, and food chain dynamics and contaminant concentrations. The user can edit inputs in an established template model or create entirely new models based on different trophic structures, human and ecological exposure scenarios and site-specific conditions. Model outputs include total carcinogenic and non-carcinogenic risks to humans and toxicity quotients for ecological receptors as well as risks from specific chemicals and dietary species. The program also compares risks to specified risk thresholds for easy screening. The algorithms incorporated into BRAMS follow USEPA and USACE risk assessment guidance (USEPA, 1989; 1997a; USEPA/USACE, 1998; Cura et al., 1999).

2. Background

Required by the Marine Protection Research and Sanctuaries Act (MPRSA), the current approach for evaluating dredged materials is outlined in the Ocean and Inland Testing Manuals (OTM; ITM) (EPA/USACE, 1991; 1998). It involves comparing the bioaccumulation test results of dredged materials with reference sediment test results and FDA action levels. These toxicological measures are indicative of a contaminant’s potential for adverse effects to human and ecological receptors. The OTM, commonly referred to as the “Green Book,” provides a general protocol for evaluating sediment toxicity and determining the suitability of dredged materials for open-water disposal. EPA and USACE share the responsibility for regulation of this dredged material. In

general, the Corps issues permits for dredged material disposal which are then subject to EPA review and concurrence before ocean disposal can occur. The tiered approach to evaluation of potential environmental impacts of ocean dumping is outlined in the Ocean and Inland Testing Manuals (OTM & ITM). BRAMS can be used in Tiers I through IV to provide information about potential risks associated with bioaccumulation. Specific guidance for sampling and testing in accordance with the OTM and ITM is provided in Regional Implementation Manuals.

Since 1999, EPA Region 1 has evaluated bioaccumulation test results using a screening tool that considers whether contaminants accumulated in test organisms might result in a risk to human or ecological receptors by direct or indirect consumption (Battelle, 2005). Also in 1999, Menzie-Cura & Associates, Inc. developed a mechanistic, process-based bioaccumulation risk model, *TrophicTrace*, to calculate the potential human and ecological impacts of bioaccumulation from sediment-associated contaminants (Bridges *et al.*, 2002; von Stackelberg *et al.*, 2004). The BRAMS program, released in 2012, includes two fully customizable models, *Trophic Trace* and BEST, based on the 2005 USACE *TrophicTrace* and the 1999 EPA Region 1 bioaccumulation risk assessment model frameworks, respectively.

**Trophic Trace (TT):**

In the *Trophic Trace* model, human and ecological receptors are exposed to potential contaminants in dredged materials via ingestion of prey. The model estimates expected concentrations using a sediment-based food web for organic compounds, via trophic transfer factors from invertebrates to fish for certain metals, and via bioconcentration factors from water to fish for the remaining metals and hydrophilic organic compounds. Water concentrations are estimated using a partitioning approach based on the user-specified sediment concentration or the user can input a water concentration directly (the model requires a freely dissolved concentration, but can estimate one from an input whole water concentration). Details of the model framework are discussed in Part V Section 1.

Uncertainty: *Trophic Trace* allows users to characterize uncertainty using trapezoidal fuzzy numbers (e.g., a minimum, a range of likeliest values or probable values, and a maximum) for each input parameter. These uncertainties are propagated throughout the analysis using principles of fuzzy arithmetic. Model results are also presented as trapezoidal fuzzy numbers representing a minimum value, a range of two most probable values, and a maximum value. Trapezoidal fuzzy numbers are explained in detail in Part V Section 3.

Included *Trophic Trace* Example: The BRAMS software includes a *Trophic Trace* example model that contains several human receptor population data libraries built into the demonstration form of the model, including recreational anglers (children and adult) in the New York and New Jersey (NY/NJ) area, and members of the general public (children and adult). The example exposure assumptions used for these demonstration receptor populations are obtained from the USEPA Exposure Factors Handbook (USEPA, 1997a; 1997b) as well as from the New Jersey Department of Health (NJDA, 1994). The values provided in the *Trophic Trace* model are for demonstration purposes only. All model runs should be based on site-specific information. The *Trophic Trace* example is also parameterized for several ecological receptors, including fish, osprey, bald eagle, mink, and otter. The example food web included in the model is sediment
based. Following the Gobas model, Trophic Trace assumes that organic compounds partition from organic carbon in sediment to the lipid fraction of benthic invertebrates. The example model is parameterized for a simple sediment based food web that is representative of a food web that might be found in the Northeast Region. The example invertebrate in Trophic Trace is the sandworm (*Nereis virens*). The model assumes that a forage fish represented by the mummichog (*Fundulus heteroclitus*), consumes sandworms and that a piscivorous fish represented by the summer flounder (*Paralichthys dentatus*) consumes the mummichog. The user can create additional food webs by modifying or adding invertebrates and fish species, such as pelagic invertebrates that derive the bulk of their exposure from the water column, and fish that consume both benthic and pelagic invertebrates.

**Bioaccumulation Evaluation Screening Tool (BEST):**

The BEST model is based on EPA Region 1’s Bioaccumulation Risk Assessment Model framework. This model was designed to evaluate dredged material for open water disposal in the New England region in accordance with Green Book testing protocol (EPA and USACE, 1991). BEST estimates expected concentrations in humans by (1) calculating the edible tissue concentration in human diet species including test organisms and their predators, (2) calculating an average daily dose to humans that consume these species, and (3) multiplying or dividing average daily dose by risk factors (e.g. oral cancer slope factor and oral reference dose) to determine potential risks. BEST model framework is discussed in detail in Part V Section 2. In the BEST model output, risk results for dredged sediments are compared to reference site risks, and acceptable risk thresholds. Calculated tissue concentrations are compared to FDA action levels and ecological effects levels. The model can be tailored to each specific project by modifying species, food webs and exposure scenarios in templates or creating entirely new templates and models.

Included BEST Template: The BEST template model included in BRAMS is based on the structure and input values of the 1999 EPA Region 1 Bioaccumulation Risk Assessment Model (Batelle, 2005). For the standard bioaccumulation risk assessment, the user enters 28-Day Bioaccumulation test results for the two invertebrate test species, *Macoma nasuta* and *Nereis virens*, which have been exposed to either the proposed dredge material or reference sediments. The model then calculates predator body burden in a fish fillet, lobster muscle, and lobster hepatopancreas. A lifetime daily average dose (LADD) is then determined using standard EPA default exposure factors (USEPA, 1989) and regional fish consumption rates (USEPA, 1988; Ruffle et al., 1994; RI Department of Health, 1997). Carcinogenic and non-carcinogenic risks to human health from fish, lobster, and molluscan shellfish consumption are calculated using standard risk factors (e.g. oral CSF, RfD, TEF) for contaminants present at the test site.
3. Installation and StartUp

3.1 System Requirements

<table>
<thead>
<tr>
<th>Central processor</th>
<th>Intel Pentium 1GHz or better</th>
</tr>
</thead>
<tbody>
<tr>
<td>Operational memory</td>
<td>512 MB or more</td>
</tr>
<tr>
<td>Free space on hard disk</td>
<td>20 MB</td>
</tr>
<tr>
<td>Operating system</td>
<td>Windows XP/Vista/7/10, macOS or Linux</td>
</tr>
<tr>
<td>Third-party software</td>
<td>Java Runtime Environment (JRE) Version 8 or later</td>
</tr>
</tbody>
</table>

3.2 Installation

- Insert the BRAMS disk into your computer.
- Locate the BRAMS CD on your Windows disk (D) drive or your Mac Desktop and double click it.
- Drag and drop or Copy and Paste the BRAMS-Dist Folder to the appropriate location on your hard drive or run the program directly from the CD.

3.3 StartUp

To open BRAMS, select BRAMS-Dist → bin → BRAMS.jar. Double clicking the file, BRAMS.jar, will open the program.

The StartUp window will then appear while BRAMS is loading.

Once BRAMS has finished loading, the User Identification window will appear prompting the user to enter their User Name. Enter your User Name and click Continue to continue to the main BRAMS window.

The user is now ready to create, edit, and run BRAMS models from the Main Window.

4. User Interface

4.1 Main Window

The main BRAMS window includes 9 general sections:
- Menu
- Toolbar
- Explorer Pane
- Table Pane
- Properties Pane
- Notes Pane
- Layer Pane
- Map Pane
- Graph Pane

These sections are explained in further detail below in sections 4.2 - 4.7.

Figure 1: Main BRAMS Window.

4.2 Menu

The menu consists of:

**File:** Here the user can create, open or save a project or template, change the user, import data from Excel files, lock and unlock model inputs and exit the program.

- **File:**
  - New Project
    - New TT Project
    - New BEST Project
    - New FishRand Project
  - Open Project or Template
  - Save Project
  - Save Project As
  - Save Permanently
  - Change User
  - Import
    - Bioaccumulation Test Results
    - Chemicals
  - Unlock Project
  - Lock Project
  - Exit
Edit: Here the user can access tools for editing script while inputting text into the properties pane.

- **Edit**
  - Undo
  - Redo
  - Copy
  - Paste
  - Cut
  - Delete

View: Here the user can change aspects of the main window by choosing to include or remove any of the panes (Notes, Explorer, Table, or Properties) or switch to and from full screen viewing.

- **View**
  - Full Screen
  - Panels
    - Table
    - Graph
    - Properties
    - Map
    - Layer
    - Explorer
**Notes**

**Run:** Here the user can choose to perform the TT or BEST model calculations or show the Report. The Show Report button will generate the model results and launch the ReportViewer window in front of the main window.

- **Run**
  - Calculate
  - Show Report

**Tools:** Here the user can edit several features of the software, including the output format and how the templates are generated.

- **Tools**
  - Options
    - Reports
    - Paths
  - Other

**Help:** Here you can access the BRAMS About window and this User Guide, which describes the program components and guides the user in creating, editing, and running bioaccumulation models.

- **Help**
  - User Guide
  - About
4.3 Toolbar

The toolbar consists of the following buttons:

![BRAMS main toolbar](image)

Figure 28: BRAMS main toolbar.

**Project Tools**

- **Create new TT project**: Displays the Create New Project window that asks whether the user would like to create a new TT project. Selecting *No* will return you to the main window with no changes. Selecting *Yes* will exit the current project and start a new blank TT project.

- **Create new BEST project**: Displays the Create New Project window that asks whether the user would like to create a new BEST project. Selecting *No* will return you to the main window with no changes. Selecting *Yes* will exit the current project and start a new blank BEST project.

- **Create new FishRand project**: Displays the Create New Project window that asks whether the user would like to create a new FishRand project. Selecting *No* will return you to the main window with no changes. Selecting *Yes* will exit the current project and start a new blank FishRand project.

- **Open project**: Displays the Open Project window where the user can search for, select and open an existing project. Once the project has been selected, click *Open* to open the project. To return to the Main window without opening a project, click *Cancel*. 
Save project: If the project has already been saved previously, clicking this button will overwrite the previous file and save the updated version under the same name and location. If the project has not been saved previously, clicking this button displays the Save Project as… window where the user can name and specify a location to save their current project. If the project has been saved permanently, this option will not be available because the project file cannot be overwritten.

Save project as: Displays the Save Project as… window where the user can name and specify a location to save the current project.

Editing Tools

- Back: Undo last action.
- Forward: Redo last action that was undone.
- Cut selected item: Cut selected text.
- Copy selected item: Copy selected text.
- Paste selected item: Paste last Cut or Copied text.
- Remove selected item: Delete selected text.

Reporting Tools

- Show Detailed Report: If all necessary model parameters have been entered, brings up Show Report window. If model is incomplete, information window will appear and explain what must be added or corrected before the report can be generated.
- Show Summary Report (BEST Model only): The summary report shows only a select set of results including total carcinogenic and non-carcinogenic risks and contaminant specific
risks for each invertebrate, as well as contaminant concentrations compared with FDA Action Levels, and Ecological Risk Levels.

**Show EPA Report (BEST Model only):** EPA Region 1 Report is designed to provide comparable results to the bioaccumulation risk analysis report that produced by EPA Region 1. Comparable table names from EPA’s report is shown in parentheses in table headings for easy reference.

### 4.4 Notes Pane

The Notes pane, located at the bottom of the main window, allows the user to enter text for each model item. Once an item is selected from either the Explorer or Table panes, notes about that item previously added by the user will appear in the Notes pane.

![Figure 39: Notes Pane with example text.](image)

### 4.5 Explorer Pane

The Explorer pane shows the model components and hierarchy. It allows the user to see the model items in each category and navigate between different items.

Each of these model components can or must include specific items such as invertebrates, chemicals, or environments. Selecting a category (e.g. Invertebrates) or an item (e.g. Worm) in the Explorer pane will display all items in that category and selected item properties in the Table pane. Selecting an item in the Explorer pane will also display its attributes in the Properties pane.

For the TT model, the Explorer pane includes:
For the BEST model, the Explorer pane includes:

- Project
- Invertebrates
- Fishes
- Birds
- Mammals
- Human
- Chemicals

4.6 Table Pane

The Table pane shows all the model items and their certain attributes in the selected category in the Explorer pane. Selecting one of the items in the Table pane will display its properties in the Properties pane.

<table>
<thead>
<tr>
<th>Name</th>
<th>Lipid</th>
<th>Normalization</th>
</tr>
</thead>
<tbody>
<tr>
<td>Worm</td>
<td>0.6146</td>
<td>1090</td>
</tr>
<tr>
<td>Clam</td>
<td>0.6124</td>
<td>1090</td>
</tr>
</tbody>
</table>

Figure 511: Example BEST model Table pane showing defined values of characteristics of invertebrates including their Name, Lipid Content, and Normalization.
Map pane brings GIS capabilities to BEST and FishRand projects. For FishRand project, Map pane is used to combine areas with the same parameters for calculations. For BEST project, you can specify locations using this pane and also specify concentration samples, used in 28-days bioaccumulation tests within locations for invertebrates.
Map pane consists of:

- Map toolbar contains buttons for the basic operations with spatial feature layers.
- Map area displays project spatial layers.
- Map status bar displays information about the current coordinates, the coordinate system that is used, etc.

Map toolbar consists of the following buttons:

![Map toolbar](image)

Figure 14: Map toolbar.

Map Tools

Create Layer: Creates a new spatial layer for the current project. Create Layer window appears asking user to enter the layer name, type, file name on the disk and coordinate reference system.

![Create New Layer window](image)

Figure 15: Create New Layer window.

Open Layer: Displays a standard Java dialog for the spatial layer file selection. It can be used to add not only for layers that were created by this tool, but also for arbitrary layers. Opened and created layers appears the Layer Management pane.

Remove Layer: Removes a current active spatial layer from Project.
Add feature: Adds a new spatial feature to a currently active layer. Required user actions depend on a layer type. In order to add a new feature to a point-type layer a user should click on a map where he/she wants to place a point feature; for a polygon-type layer he/she should click multiple times adding polygon points.

Select feature: Enables spatial feature selection mode that works only for an active layer. Click on a feature in order to select it. Selected feature is highlighted with a red color; also the selected feature properties will appear in Properties pane. In order to select multiple features, click somewhere on a map, and drag a red selection (rubber) selecting feature.

Move feature. Allows to move a feature to a new position. Press the left mouse button on the feature and drag and drop it to a new position.

Display full extent: Changes map scale to fit a map inside of Map pane.

Zoom in: Increases map scale.

Zoom out: Decreases map scale.

Exports bioaccumulation data from spatial layers and imports them BEST model. It can be used with BEST project only.

Use the following steps to export and import 28-days bioaccumulation results.

1. Create a new location layer.
2. Add location features to the new layer.
3. Create a concentration layer for each invertebrate. Connect the concentration layer with the corresponding invertebrate using the Properties pane.

4. Add concentration features and specify chemical concentrations using the Properties pane. The added concentration features should be inside of previously created location features.
5. Click the button.
6. Review the imported from the GIS layers locations and 28-day bioaccumulation results with mean and max values.
Figure 19: Generated locations.

Figure 20: Generated 28-days bioaccumulation tests.

4.8 Layer Pane

Layer Management pane allows to change the current active feature layer, change layer order, hide layers in Map pane. There are a number of parameters for every layer: visibility, name,
color, path to layer file. In order to make a layer active click its row; to hide a layer check off its check box.

![Layer](image)

Figure 21: Added layers to the project.

Selected layers have a number of setting in Properties pane. These settings depend on layer type. Using these settings user can change name, color and other layer properties depending on layer type.

![Properties](image)

Figure 22: Layer properties.

4.9 Graph Pane

**Graph pane shows food chain starting from invertebrates to humans for BEST and TT projects.** Food chain is not generated by default when project is opened. In order to generate food chain for a current project, the refresh button on the Graph pane toolbar should be clicked. The generated food chain is interactive, when user clicks on any item it’s getting selected in Explorer and Properties panels. Also, by clicking the button it is possible to export the food chain to an image file.
There are several appearance settings for food chain in the Option Dialog. Using these properties user can change food chain layout and species color.
4.10 Properties Pane

The Properties pane shows the properties of the selected item. Here, property values can be assigned and edited by the user. Editable attributes either have a button where the user must choose from a dropdown menu of possible values, or a button where the user must directly input the values. Once the user selects or inputs the value for an item, it will appear in the Table pane. The properties vary for different categories (e.g. Invertebrates vs. Mammals). If the project is locked (File → Lock Project), item properties, except for Bioaccumulation Test results, will not be editable. To edit these values, the user must unlock the project by selecting File → Unlock Project and entering the appropriate password.

For the TT model, the Properties pane includes properties under the categories:

- General – All items
- Specific – All items
- Toxicity- Fish, Birds, Mammals
- Chemical of Concern Table - Environment
- 28-Days Bioaccumulation Test - Invertebrates
- Diet – Fish, Birds, Mammals, Human
For the **BEST model**, the Properties pane includes properties under the categories:

- **General** – All items and Project
- **Specific** – All items
- **Calculation** – Humans and Chemicals
- **Diet** – Predators and Humans
- **28-Day Bioaccumulation Test** – Invertebrates
- **Risk Thresholds** – Project
- **Reference Project** – Project
- **Levels** – Chemicals
4.11 ReportViewer Window

The ReportViewer window shows the results of the risk assessment including risk endpoints such as the carcinogenic and non-carcinogenic risks for BEST and TT and NOAELTQ and LOAELTQ for TT. It also displays contaminant tissue concentrations and doses for model species and human and ecological receptors. Information included in the ReportViewer window depends on the model and report type. The ReportViewer appears when the user clicks the button or selects Run → Calculate. A summary report is also available for the BEST model, which can be accessed by clicking the button or by selecting Run → Show Summary Report. From the ReportViewer window, the user can navigate, adjust, save or print the risk assessment results.

4.12 Options Window

Options window allows to change the application general settings:

**Trophic Trace**
Show Graphs in Reports: Check on to show charts in the output report.
Logarithmic scale: Check on to use the logarithmic scale.

**BEST**
Show Zero Values: Check on to show zero rows in BEST reports.
Show Group Total: Check on to enable total group rows (see grouping) in BEST reports.
Show Group Detailed: Check on to enable total detailed rows (see grouping) in BEST reports.
**Graph**
Layout Option: Changes layout option (layout type) that is used automatically generating and placing food chain items.
Show Singles: Check on to show species that are not involved in food chain.
Human, Predator, Prey Color: Changes the color of the corresponding species nodes.

**Map**
Select Stroke Color: Changes the stroke color of selected features.
Select Fill Color: Changes the fill color of selected features.

![Options window.](image)
# BRAMS

## Table Of Contents

<table>
<thead>
<tr>
<th>Category</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fishes</td>
<td>1</td>
</tr>
<tr>
<td>Anchovy NSHR</td>
<td>1</td>
</tr>
<tr>
<td>Anchovy HCNYN</td>
<td>4</td>
</tr>
<tr>
<td>Anchovy CNYN</td>
<td>6</td>
</tr>
<tr>
<td>Anchovy SHLF</td>
<td>9</td>
</tr>
<tr>
<td>Sole NSHR</td>
<td>11</td>
</tr>
<tr>
<td>Sole HCNYN</td>
<td>14</td>
</tr>
<tr>
<td>Sole CNYN</td>
<td>18</td>
</tr>
<tr>
<td>Sole SHLF</td>
<td>21</td>
</tr>
<tr>
<td>Humans</td>
<td>24</td>
</tr>
<tr>
<td>Pier/Shore CTE</td>
<td>24</td>
</tr>
</tbody>
</table>

## Fishes

**Fish Name:** Anchovy NSHR  
**Body Weight:** 20  
**Lipid:** 3.8; 10.7; 24.6  
**Site Use Factor:** 1

### Risk List

<table>
<thead>
<tr>
<th>Chemical Name</th>
<th>Calculation Method</th>
<th>NOAELTQ</th>
<th>LOAELTQ</th>
<th>NOAELTQ Eggs</th>
<th>LOAELTQ Eggs</th>
</tr>
</thead>
<tbody>
<tr>
<td>DDE</td>
<td>Equilibrium Partitioning</td>
<td>0</td>
<td>0</td>
<td>0.0001</td>
<td>0.000006</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.000156</td>
<td>0.00011</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.00053</td>
<td>0.00033</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.00222</td>
<td>0.00139</td>
</tr>
<tr>
<td>DDE</td>
<td>Equilibrium Partitioning</td>
<td>0</td>
<td>0</td>
<td>0.00002</td>
<td>0.00001</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.00004</td>
<td>0.00002</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.00014</td>
<td>0.00003</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.00051</td>
<td>0.00039</td>
</tr>
<tr>
<td>DDD</td>
<td>Equilibrium Partitioning</td>
<td>0</td>
<td>0</td>
<td>0.00001</td>
<td>0.913E-6</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.00002</td>
<td>0.00001</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.00001</td>
<td>0.00005</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.00018</td>
<td>0.00024</td>
</tr>
<tr>
<td>Chlorothen</td>
<td>Equilibrium Partitioning</td>
<td>0.01209</td>
<td>0.02012</td>
<td>0.00054</td>
<td>0.00073</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>0.03174</td>
<td>0.01377</td>
<td>0.00462</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.06902</td>
<td>0.06902</td>
</tr>
<tr>
<td>Dichloro</td>
<td>Equilibrium Partitioning</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Endosulfan II</td>
<td>Equilibrium Partitioning</td>
<td>0.00002</td>
<td>0.00005</td>
<td>1.774E-6</td>
<td>5.368E-6</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.00005</td>
<td>0.00005</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.00012</td>
<td>0.00001</td>
</tr>
</tbody>
</table>

Friday, 27 April 2012
Figure 929: Example TT model ReportViewer window page 1.
PART II: Building the Model

For a quick, step by step guide to creating BRAMS models, see Part IV - Step By Step Model Creation.

1. User Identification

Users should identify themselves by entering their user name in the User Identification window. To access this window, select File → Change User. When opening BRAMS for the first time the User Identification window will appear automatically prompting the user to enter their User Name. This User Name will appear on the last page of the Reports.

2. Create or open a project or template

Create a new project:

**BEST**: From the main window select File → New Project → New BEST Project from the Menu bar or click in the toolbar. Then select Yes when the Create new project window appears.

**TT**: From the main window select File → New Project → New TT Project from the Menu bar or click in the toolbar. Then select Yes when the Create new project window appears.

**FishRand**: From the main window select File → New Project → New FishRand Project from the Menu bar or click in the toolbar. Then select Yes when the Create new project window appears.

To open an existing model or template:

From the main window select File → Open Project or Template or click , and then choose an existing model from the Open Project window.

5. Add a new item

In the Explorer pane, select the item category you wish to add to (e.g. Chemicals) then click in the Table pane to add a new item to that category. A new item will appear in the Explorer pane under the selected category and in the Table pane with default values for the item’s properties. Items can be added to the model in any order but item properties must be input in the order specified in section 4.1 below.
BEST model: Inputs other than Bioaccumulation Test results may be password protected if the project has been locked. If so, the project must be unlocked to edit inputs. To lock or unlock a project, see Part II Section 6.

6. **Add or edit project and item properties**

6.1 **General**

For each item, the user must specify values for each property in the Properties pane. Because some value choices depend on other inputs, add property values to items in each category in the Properties Pane in the order:

**BEST:** 1. Chemicals  
          2. Invertebrates  
          3. Predators  
          4. Humans  
          5. Locations

**TT:** 1. Chemicals  
         2. Environment  
         3. Invertebrates  
         4. Fish  
         5. Birds, Mammals, or Humans  
         6. Environment

BRAMS properties may be input using one of these four input methods:

**Excel Input (BEST Model Only):** Some Project, Chemical and Invertebrate items and properties are imported into the current BEST project directly from selected Microsoft (MS) Excel files with the appropriate format. See Part II section 4.3 for a complete description of the MS Excel Input method and input formats. MS Excel input templates are also included in the BRAMS software package.

**Text Box:** Click inside the text box directly right of the property name; delete any unwanted text and type new text.

**Drop Down Menu:**
Click [ ] to the right to of the property you wish to edit to view possible choices. Double click the appropriate choice.

**Add Button [ ]:**

Click [ ] to the right of the property you wish to edit. This will either display a drop-down window with four possible numerical inputs (for parameter uncertainty in TT models only) or a drop-down table with information relating the current item to another item type (e.g. Concentrations of specific Chemicals in specific Invertebrate’s tissue according to the 28-day Bioaccumulation test). After clicking [ ], add values depending on the property type:

**Independent Numerical Values** – Double click the specific input value in the dropdown window that you wish to edit, delete the current value and enter the new value.

**Other Item Dependent Values** - Click [ ] in the bottom right corner of the drop down table to add an existing item from within the related item category. Then click [ ] below the new item category to see the list of existing items. Choose one of the existing items by double clicking it. Next, specify values for each attribute by double clicking the cell corresponding to each and entering the desired value. Click [ ] to delete an item from the list.

6.2 By model type and item category

**4.2.1 BEST model:**

To edit an item’s properties:
- (1) In the Explorer pane, select the item category.
- (2) In the Table pane, select the specific item.
- (3) In the Properties pane, edit properties as described below.

**Project properties:**

**General:**
Specify the Project Name, Project Number by either, (1) clicking inside the text boxes to the right of each corresponding property name, deleting any contents, and entering the appropriate data or (2) importing the data from the 28-day Bioaccumulation Test Results Excel sheet (See Part II section 4.3 on Excel Inputs).

**Risk Thresholds:**
Enter carcinogenic and non-carcinogenic risk thresholds in the General Project Properties pane by clicking inside the corresponding text boxes and entering values. Calculated risk levels exceeding these thresholds will be highlighted in red in the Total Risks and Contaminant Specific Risks sections of the BEST Summary Report. Also, you can specify Number of Measurements that affects number of concentration samples in bioaccumulation test results for every location.
Reference: (For Optional Test and Reference site risk comparison):
Select reference location for project from the drop down list, which is populated with available locations from bioaccumulation results file.
List of which chemicals that should be included to risk estimation should be selected for each individual sediment location, except reference one. The list of chemicals that is used in calculations for reference location will reproduce the list used that is used for sampling location for which the reference is serving as a basis for comparison.

Daily Dose Settings: User can control whether Average Daily Dose, or Lifetime Average Daily Dose should be used for Summary and EPA reports.

![Figure 30: BEST project properties.](image-url)

**Item properties:**

For each item, the user must specify property values within the property categories including:

- General – All items
- Calculation – Humans and Chemicals
• Specific – All items
• Diet – Predators and Humans
• 28-Day Bioaccumulation Test – Invertebrates
• Levels – Chemicals

While the item you wish to edit is selected in the Table Pane, specify its properties by entering values into the boxes within the Properties pane.

**Chemicals:** All chemical data can be entered automatically by importing the data from the Chemical Input Excel sheet (See Part II section 4.3 on Excel Inputs). To enter chemical data manually, follow the directions below. If the value of any property is unknown, enter NA in the text box (default value).

**General:**
1. Name: Specify the name of the item by clicking inside the text box to the right of Name, deleting the current name, and typing a new one.

**Calculation:**
2. Calculating: Specify whether or not to include the chemical in the model calculations. Check the box to the right of to include the item, uncheck the box to exclude the item.

3. TEF Reference Chemical: Specify whether the chemical is a TEF Reference Chemical (ex. 2,3,7,8-TCDD and benzo(a)pyrene) by checking (Yes) or un-checking (No) the box to the right of Reference Chemical. Chemicals designated as TEF reference chemicals will then be added to the TEF calculation drop down menu.

4. TEF Relation: Specify if or how a Human Toxicity Equivalency Factor (TEF) applies. Select NA for all items that do not have a TEF (NA is the default value). For items such as individual dioxin congeners, coplanar PCB compounds and selected PAH compounds that have assigned TEF values, select the reference chemical their TEF applies to. This information can be entered manually or imported directly from a Chemical Excel file. To enter TEF information by importing from Excel, see Part II Section 3.3. To enter TEF information manually, click ☑ to the right of TEF to display a dropdown menu of possible TEF choices. See Part V Section 2.3 for additional information on TEF application to chemicals in the BEST model.

**Grouping:**
5. Total Chemical: Marks chemical as total group. Total group chemical is used only for grouping and does not appear in any calculations.

6. Grouping Chemical: Selects the chemical group from the drop-down menu, only total chemicals are shown there. Depending on the settings either every single non-total chemical is shown in the reports or chemicals results are grouped by total chemical.
Specific:

(7) Chemical Type: Click to specify whether the chemical is Organic or Metal. Organic and metal contaminant transfer are calculated differently.

(8) Cancer Slope Factor (mg/kg-day)^{-1}: Specify the Oral Cancer Slope factor associated with the chemical by clicking inside the textbox and entering the appropriate value. This value will be used to determine Carcinogenic Risk.

(9) Reference Dose (mg/kg-day): Specify the Reference Dose associated with the chemical by clicking inside the textbox and entering the appropriate value. This value will be used to determine Noncarcinogenic Risk.

(10) Biomagnification Factor: Specify the Reference Dose associated with the chemical by clicking inside the textbox and entering the appropriate value. This value will be used to determine contaminant transfer from invertebrates to predators.

(11) Human Toxicity Equivalence Factor: If a TEF does not apply, the selected chemical should have ‘N/A’ selected in the TEF Relation property field above and ‘NA’ will appear in the Human Toxicity Equivalence Factor text box. If a TEF does apply, click inside the textbox and enter the appropriate TEF value relating this chemical’s toxicity to a chemical of known toxicity.

(10) Steady State Correction Factor: Specify the steady state correction factor associated with the chemical by clicking inside the textbox and entering the appropriate value. This value will be used to correct for underestimates of contaminant transfer from sediment to invertebrates during 28-day bioaccumulation testing.

Levels:

(11) FDA Action Level: Specify the FDA Action Level (ppm) associated with the chemical by clicking inside the textbox and entering the appropriate value.

(12) Ecological Effect Level: Specify the Ecological Effect Level (ppm) associated with the chemical by clicking inside the textbox and entering the appropriate value.

(13) FDA Level of Concern: Specify the FDA Level of Concern Level (ppm) associated with the chemical by clicking inside the textbox and entering the appropriate value.

Invertebrates:

General:

(1) Name: Specify the name of the item by clicking inside the text box to the right of Name, deleting the current name, and typing a new one.
(2) Group Category: BEST can aggregate results; in order to indicate which items in trophic level should be summed in report, their group (shared) name should be assigned.

Specific:
(3) Lipid: Specify the invertebrate’s lipid content (g lipid/g tissue) by clicking inside the text box, deleting the current value and entering the new value.

28-Days Bioaccumulation Test:
(4) Bioaccumulation Test: There are two ways to enter 28-day bioaccumulation test data:
   a. Import data directly from a 28-day Bioaccumulation Test Results Excel sheet as described in Part II Section 4.3 or
   b. Enter the data manually by first clicking to display a drop down window where you can add chemicals from the Chemicals category and Mean and Max Tissue Concentrations. Then click in the bottom right corner of the drop down window to add a new row. Next, click in the Chemical column of the new row to choose from the existing chemicals. Finally, specify values for Mean Tissue Concentration and Max Tissue Concentration by double clicking the corresponding cells and typing their values. Organic tissue concentrations should be in units of ng/g, while metal tissue concentrations should be in units of µg/g. Click to delete a chemical from the table.

![Figure 31: BEST invertebrate properties.](image)

Predators:

General:
(1) Name: Specify the name of the item by clicking inside the text box to the right of Name, deleting the current name, and typing a new one.

Grouping:
(2) Group Category: BEST can aggregate results; in order to indicate which items in trophic level should be summed in report, their group (shared) name should be assigned.

Specific:
(3) Lipid: Specify the invertebrate’s lipid content (g lipid/g tissue) by clicking inside the text box, deleting the current value and entering the new value.

Diet:
(4) Predator Diet: First click to display a drop-down window where you can add each predator’s potential prey species. Click in the bottom right corner of the drop-down window to add a new prey species. Then click to the right of the new row to specify which of the existing Invertebrate species to add.

Humans:
General:
(1) Name: Specify the name of the item by clicking inside the text box to the right of Name, deleting the current name, and typing a new one.

Calculation:
(2) Calculating: Specify whether or not to include the item in the model calculations. Check the box to the right of Calculating to include the item, uncheck the box to exclude the item from the model calculations.

Specific:
(3) Body Weight: Specify the human’s body weight in kilograms by clicking inside the text box, deleting the current value and entering the new value.
(4) Averaging Time: Specify the averaging time in days to be used in the human’s average daily dose calculation by clicking inside the text box, deleting the current value and entering the new value.

Diet:
(5) Human Diet: Click to display a drop-down table where you can add species that make up the selected human’s diet. Click in the bottom right corner of the table to add a new row. Then click below the Prey column in the new row to choose from the existing lower trophic level species. Then specify values for fraction ingested, frequency.
(days/year), ingestion rate (kg/day), and exposure duration (years), by double clicking the space below each and entering the appropriate values. These values will be used to calculate the human’s average daily dose of contaminants in each diet species. Click to delete a row from the list.

![Diagram of BEST human properties](image)

**Figure 33: BEST human properties.**

### 4.2.2 TT Model:

For each item, the user must specify property values within three or more categories including:

- General – All items
- Specific – All items
- 28-Days Bioaccumulation Test - Invertebrates
- Diet – Fish, Birds, Mammals, Human
- Toxicity- Fish, Birds, Mammals
- Chemical of Concern Table - Environment

While the item you wish to edit is selected, specify its properties by entering values into the boxes within the Properties pane. For each Properties category, follow the directions below.

**General:**

Specify the name of the item by clicking the box to the right of Name, deleting the current name, and typing a new one. Specify the environment by clicking the then the appropriate environment type.

**Specific:**

Specify values in the Specific category by clicking to the right of the characteristic you wish to edit. This will display a drop-down window with four possible inputs.
Double click the input value you wish to edit, delete the current value and enter the new value. If the input value is known with certainty, simply enter the known value in all four spaces. If the input value is uncertain, enter four values representing the range of possibilities according to trapezoidal fuzzy numbering (See Part V Section 3). Or, for some input categories, (Invertebrates-Diet Pathway, Fish-Trophic Level and Reference Invertebrate, Chemicals-Chemical Type) click the 🔄 to the right of the characteristic you wish to edit to display a drop-down menu of pre-specified choices and click the desired choice.

28-Days Bioaccumulation Test:

For items in the Invertebrates category, specify Bioaccumulation Test values by clicking 🔄 to the right of Bioaccumulation Test. This will display a drop-down window where you can input chemicals, their concentrations and whether or not steady state applies. To add a chemical, click 🔄 in the bottom right corner of the drop-down window. Then click the 🔄 in the new row below Chemical, this will display the list of chemicals that have previously been added to the Chemicals category. Click the chemical you wish to add information about and it will appear on the list. Then double click the box corresponding to its concentration. Now you can either add one, certain value by typing it in the box or add four values for uncertain inputs by clicking 🔄 to the right of the box and entering the values according to trapezoidal fuzzy numbering (See Part V Section 3). Next, if steady state applies, check the box below Steady State Applies. If it does not apply, leave the box blank.

Diet:

For items in the Fishes, Birds, Mammals and Human categories, specify Diet properties by clicking 🔄 to the right of Diet. This will display a drop-down window where you can input the selected item’s prey species, and the percent of their diet each makes up. To add a prey species, click 🔄 button in the bottom right corner of the drop-down window. Then click the 🔄 in the new row below Prey, to will display the list of potential prey species that have previously been added to the model. Click the species that make up the selected item’s diet and they will appear on the list. Then double click the Percent box and type the percent of the selected item’s diet that each prey species makes up. Enter prey species until the total diet percent adds up to 100.

Toxicity:

For items in the Fishes, Birds, and Mammals categories, specify Toxicity properties by clicking 🔄 to the right of Toxicity and, for Fishes and Birds, Eggs Toxicity. For both, this will display a drop-down window where you can input chemicals, and their NOAEL and LOAEL Residue values. To add a chemical, click 🔄 in the bottom right corner of the drop-down window then 🔄 below the Chemical category in the dropdown window. This will display the list of chemicals that have previously been added to the Chemicals category. Click the chemical you wish to add information about and it will appear on the list. Then double click the box corresponding to NOAEL Residue. Now you can either add one, certain value by typing it in the box or add four values for uncertain inputs by clicking 🔄 to the right
of the box and entering the values according to trapezoidal fuzzy numbering (See Part V Section 3). Next add values to LOAEL Residue in the same way.

**Chemical of Concern Table:**

For items in the Environment category, specify Chemical of Concern properties by clicking ☐ to the right of Chemical of Concern. This will display a drop-down window where you can input chemicals, and their concentrations in water and sediment. To add a chemical, click the green Insert Row button in the bottom right corner of the drop-down window, and then click ☐ below the Chemical category in the dropdown window. This will display the list of chemicals that have previously been added to the Chemicals category. Click the chemical you wish to add information about and it will appear on the list. First specify whether you will enter the Total or the Dissolved concentration by clicking ☐ below Concentration Type in the dropdown window and then clicking either Total or Dissolved. Then, for both the Water and Sediment categories, double click the boxes corresponding to each and either add one, certain value by typing it in the box or add four values for uncertain inputs by clicking ☐ to the right of the box and entering the values according to trapezoidal fuzzy numbering (See Part V Section 3).

### 6.3 Import Data From Excel (BEST Model)

Invertebrates and their 28-day Bioaccumulation Test Results as well as Chemicals and all chemical properties can be entered directly into a BEST model by importing values from Microsoft Excel spreadsheets according to established templates. In order for BRAMS to read the values correctly, they must be uploaded in accordance with specific Excel templates included in the BRAMS software package. Templates for the 28-day Bioaccumulation Test Input and Chemicals Input are provided with the software package and shown below in figures 9-11. To add certain invertebrate or chemical items (e.g. *Macoma nasuta*) or properties (e.g. Mean Tissue Concentration from 28-day Bioaccumulation Test results for “Lead” in “*Macoma nasuta*”) the Chemical and Invertebrate names must match those in the model exactly, otherwise new items will be created.

Once all desired values have been correctly entered into the Excel Input templates, select File → Import → Import Bioaccumulation Results or Import Chemicals. The Open Excel window will appear where you can select the appropriate Excel file. Once you have selected the desired file, click Open. The window will then close and the data within the selected Excel file will be added to the model.

**Chemical Inputs** should be entered in the format shown in Figure 10 and Figure 11 below. Properties of each chemical (e.g. Oral Cancer Slope Factor, Oral Reference Dose, Biomagnification Factor, etc.) entered here will be automatically entered into the Chemicals section. NA will be assigned to properties, which either have the value, ‘NA’ or that have no contents in the corresponding Excel cells. If chemical properties that correspond to blank cells in Excel appear as ‘0’ instead of ‘NA’ once imported into BRAMS, go back to the cells and use the
Excel function ‘Clear Contents’ to empty the cells. An extensive chemical library is included in the BRAMS package and in the included BEST template file.

![Figure 3410: Excel Chemicals Input Sheet, Columns A-E.](image)

<table>
<thead>
<tr>
<th>A</th>
<th>B</th>
<th>C</th>
<th>D</th>
<th>E</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Chemicals</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Name</td>
<td>Chemical Type</td>
<td>Oral Cancer Slope Factor (mg/kg-day)</td>
<td>Oral Reference Dose (mg/kg-day)</td>
</tr>
<tr>
<td>3</td>
<td>c1</td>
<td>Organic</td>
<td>1.00</td>
<td>1.00</td>
</tr>
<tr>
<td>4</td>
<td>c2</td>
<td>Organic</td>
<td>1.00</td>
<td>1.00</td>
</tr>
<tr>
<td>5</td>
<td>c3</td>
<td>Metal</td>
<td>1.00</td>
<td>1.00</td>
</tr>
<tr>
<td>6</td>
<td>c4</td>
<td>Organic</td>
<td>1.00</td>
<td>1.00</td>
</tr>
</tbody>
</table>

![Figure 3511: Excel Chemicals Input Sheet, Columns F-K.](image)

<table>
<thead>
<tr>
<th>F</th>
<th>G</th>
<th>H</th>
<th>I</th>
<th>J</th>
<th>K</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Human Toxicity Equivalence Factor</td>
<td>TEF Reference Chemical</td>
<td>TEF Relation</td>
<td>Steady State Correction Factor</td>
<td>FDA Action Level (mg/kg)</td>
</tr>
<tr>
<td>2</td>
<td>1.00</td>
<td>NA</td>
<td>NA</td>
<td>100</td>
<td>10</td>
</tr>
<tr>
<td>3</td>
<td>1.00</td>
<td>NA</td>
<td>NA</td>
<td>100</td>
<td>10</td>
</tr>
</tbody>
</table>

The **28-day Bioaccumulation Test Results** must be input into the software in the format shown in Figure 12. The user must include the organism, contaminant, mean tissue concentration and maximum tissue concentration as well as the Project Name, Project Number and Location. For unit consistency, [Tissue]/Normalization Factor must equal units of mg/kg. For example if the Normalization Factor (Invertebrate property) = 1000, enter tissue concentration in ng/g. If the Normalization Factor (Invertebrate property) = 1, enter tissue concentration in µg/g.

![Figure 3612: Excel Bioaccumulation Test Results Input Sheet](image)

<table>
<thead>
<tr>
<th>A</th>
<th>B</th>
<th>C</th>
<th>D</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>28-days Bioaccumulation Test Results</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Project Name:</td>
<td>p_name1</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>Project Number:</td>
<td>p_number1</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>Location</td>
<td>f1</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>Organism</td>
<td>Contaminant</td>
<td>Mean Tissue Concentration [Tissue]/Normalization Factor = µg/g</td>
</tr>
<tr>
<td>7</td>
<td>o1</td>
<td>c1</td>
<td>1.00</td>
</tr>
<tr>
<td>8</td>
<td>o1</td>
<td>c2</td>
<td>1.00</td>
</tr>
<tr>
<td>9</td>
<td>o1</td>
<td>c3</td>
<td>1.00</td>
</tr>
<tr>
<td>10</td>
<td>o1</td>
<td>c4</td>
<td>1.00</td>
</tr>
<tr>
<td>11</td>
<td>o2</td>
<td>c1</td>
<td>1.00</td>
</tr>
<tr>
<td>12</td>
<td>o2</td>
<td>c2</td>
<td>1.00</td>
</tr>
<tr>
<td>13</td>
<td>o2</td>
<td>c3</td>
<td>1.00</td>
</tr>
<tr>
<td>14</td>
<td>o2</td>
<td>c4</td>
<td>1.00</td>
</tr>
</tbody>
</table>

### 7. Using Templates

Using templates as a base to build each project model from saves time and effort and ensures consistency between related projects. Instead of entering all items and properties for each new
project, templates allow the user to start with all or most of the project components, except for certain project specific data, already entered into the model. Included BRAMS templates can be found in the Examples folder. Once a template has been created, values can be protected by locking the model or saving it permanently.

8. Locking and Unlocking BEST Model Properties

The majority of BEST model users should base their risk assessment models on templates with established property values and should only need to edit the 28-days Bioaccumulation Test Results. To prevent users from changing the established values (e.g. Lipid content of N. virens, Oral Cancer Slope Factor of DDT), these values can be locked. Property values other than 28 Day Bioaccumulation Results will become grayed-out and password protected.

**Lock:** To lock a project, select File  Lock Project. This will display the Password Protection window where the user should enter a password and click continue.

**Unlock:** To unlock a project, select File  Unlock Project. This will display the Password Protection window where the user should enter the previously set password and click continue. Property values should once again be freely editable.

9. Saving Permanently

The Save Permanently feature allows users to save a project file so that it cannot be overwritten. A permanently saved file can be accessed and edited but then must be saved under a different name. This feature is designed to preserve templates for future use and finalized projects for future review.

To save a file permanently, select File  Save permanently. This will display the usual Save Project As… window where the user can name and specify a location to save the current project permanently.
Part III: Results

1. Generate Report

Once all items and their properties necessary for the risk assessment have been specified, click either Run → Calculate or Run → Show Report in the Menu or click the Show Report button in the Toolbar to generate the report. An additional summary and EPA reports are available for the BEST model. The summary report can be accessed by selecting Run → Calculate Summary Report or clicking the Show Summary Report button. EPA report can be accessed by selecting Run → Calculate EPA Report or clicking Show EPA Report button. The ReportViewer window will then appear showing results of the model analysis. The different outputs for TT and BEST are detailed in Section 2 below. Reports can be exported to multiple file formats by selecting File → Export Report or printed by selecting File → Print Report.

10. Outputs

Trophic Trace

Trophic Trace evaluates risks with different endpoints for humans and ecological receptors.

Every report contains a brief description of the model it is based on.

<table>
<thead>
<tr>
<th>Project name</th>
<th>TTMModel</th>
</tr>
</thead>
<tbody>
<tr>
<td>Description</td>
<td>Here comes the description!!</td>
</tr>
<tr>
<td>User name</td>
<td>Lex</td>
</tr>
</tbody>
</table>

Figure 37: TT Brief model description.

For a given ecological receptor, the Trophic Trace output report will first provide a description of the receptor including its name, body weight, lipid content and site use factor. For human receptors, the output first shows the human’s name, weight, lifespan and diet information.

<table>
<thead>
<tr>
<th>Fishes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fish Name</td>
</tr>
<tr>
<td>Body Weight</td>
</tr>
<tr>
<td>Lipid</td>
</tr>
<tr>
<td>Site Use Factor</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Fishes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fish Name</td>
</tr>
<tr>
<td>Body Weight (g)</td>
</tr>
<tr>
<td>Lipid (%)</td>
</tr>
<tr>
<td>Site Use Factor</td>
</tr>
</tbody>
</table>

Figure 3813: TT ecological receptor properties report section.
Next, the output displays the receptor’s “Risk List.” For ecological receptors, this section identifies the chemical names, calculation method, NOAEL TQ, LOAEL TQ, NOAEL TQ for eggs, and LOAEL TQ for eggs. For human receptors, the Risk List section shows the calculation method, incremental lifetime cancer risk (LCR), and noncarcinogenic hazard index (Hazard) from each chemical.

<table>
<thead>
<tr>
<th>Chemical Name</th>
<th>Calculation Method</th>
<th>NOAEL TQ</th>
<th>LOAEL TQ</th>
<th>NOAEL TQ Eggs</th>
<th>LOAEL TQ Eggs</th>
</tr>
</thead>
<tbody>
<tr>
<td>DDD</td>
<td>Equilibrium Partitioning</td>
<td>0</td>
<td>0</td>
<td>0.00006</td>
<td>0.00006</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DDE</td>
<td>Equilibrium Partitioning</td>
<td>0</td>
<td>0</td>
<td>0.00002</td>
<td>0.00002</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DDT</td>
<td>Equilibrium Partitioning</td>
<td>0</td>
<td>0</td>
<td>0.00001</td>
<td>8.91E-6</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chlordane</td>
<td>Equilibrium Partitioning</td>
<td>0.01299</td>
<td>0.00564</td>
<td>0.00001</td>
<td>0.00001</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.02012</td>
<td>0.00873</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.09174</td>
<td>0.01377</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.13117</td>
<td>0.05892</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dieldrin</td>
<td>Equilibrium Partitioning</td>
<td>0</td>
<td>0</td>
<td>0.00002</td>
<td>0.00002</td>
</tr>
<tr>
<td>Endosulfan II</td>
<td>Equilibrium Partitioning</td>
<td>0.00002</td>
<td>1.77E-6</td>
<td>0.00001</td>
<td>0.00001</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.00055</td>
<td>5.34E-6</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.00056</td>
<td>5.36E-6</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.00012</td>
<td>0.00001</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Figure 3914: TT ecological receptor Risk List report section.

The next set of information displayed for each ecological and human receptor is the “Exposure Concentration.” Included as part of this are diet items, the environment, diet percent, chemical, and concentration of that chemical.
### Exposure Concentration

<table>
<thead>
<tr>
<th>Diet Item</th>
<th>Environment</th>
<th>Diet Percent</th>
<th>Chemical</th>
<th>Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plankton NSHR</td>
<td>Nearshore</td>
<td>100.0</td>
<td>DDD</td>
<td>0.00575</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>DDE</td>
<td>0.00159</td>
</tr>
<tr>
<td>Plankton NSHR</td>
<td>Nearshore</td>
<td>100.0</td>
<td>DDT</td>
<td>0.00999</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Chlorodane</td>
<td>0.00127</td>
</tr>
<tr>
<td>Plankton NSHR</td>
<td>Nearshore</td>
<td>100.0</td>
<td>Dieldrin</td>
<td>0.00241</td>
</tr>
<tr>
<td>Plankton NSHR</td>
<td>Nearshore</td>
<td>100.0</td>
<td>Endosulfan II</td>
<td>0.00003</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Endosulfan sulfate</td>
<td>0.00004</td>
</tr>
<tr>
<td>Plankton NSHR</td>
<td>Nearshore</td>
<td>100.0</td>
<td>Endrin</td>
<td>0.00154</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Toxaphene</td>
<td>0.03867</td>
</tr>
</tbody>
</table>

---

### Exposure Concentration

<table>
<thead>
<tr>
<th>Diet Item</th>
<th>Environment</th>
<th>Diet Percent</th>
<th>Chemical</th>
<th>Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plankton NSHR</td>
<td>Nearshore</td>
<td>99.0</td>
<td>DDD</td>
<td>5.75E-3</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>DDE</td>
<td>1.59E-3</td>
</tr>
<tr>
<td>Plankton NSHR</td>
<td>Nearshore</td>
<td>99.0</td>
<td>DDT</td>
<td>6.86E-4</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Chlorodane</td>
<td>1.25E-2</td>
</tr>
<tr>
<td>Plankton NSHR</td>
<td>Nearshore</td>
<td>99.0</td>
<td>Dieldrin</td>
<td>2.41E-3</td>
</tr>
<tr>
<td>Plankton NSHR</td>
<td>Nearshore</td>
<td>99.0</td>
<td>Endosulfan II</td>
<td>3.1E-5</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Endosulfan sulfate</td>
<td>4.48E-6</td>
</tr>
<tr>
<td>Plankton NSHR</td>
<td>Nearshore</td>
<td>99.0</td>
<td>Endrin</td>
<td>1.54E-3</td>
</tr>
</tbody>
</table>

Figure 4015: TT Exposure Concentration report section.
The last section included in the Trophic Trace output report is a graphical representation of the risks including uncertainty bounds. For ecological receptors this includes NOAEL TQ, LOAEL TQ, NOAEL TQ eggs, and LOAEL TQ eggs, for each of the chemicals present at the site. For human receptors, this includes LCR and Hazard for each of the chemicals present at the site.

Figure 4116: TT Risk Figures report section for ecological receptor.
BEST

The BEST model has three options for output reports: the Summary Report, the Full Report, and the EPA Report. The Summary Report can be generated by clicking 🔧, while the Full Report is generated by clicking 🔧. And the EPA Report is generated by clicking 🔧.

**Summary Report**
Summary report compares risk estimates from sediment sample locations to those of the project reference site, both for total risk and contaminant specific risks.

**Important:** The Summary Report results for non-cancer risks are calculated using average daily dose whereas EPA Region 1 Report results are calculated using lifetime average daily dose. See Error! Reference source not found. for the formulas.

Summary report includes the following section for each location-invertebrate pair:

**Total Carcinogenic and Non-Carcinogenic Risks** – reports total cancer risk and non-cancer risks in each human diet species for both the test and reference sites

<table>
<thead>
<tr>
<th>Location: SW-1, 2, 4, 5</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Total</strong></td>
</tr>
<tr>
<td><strong>Carcinogenic</strong></td>
</tr>
<tr>
<td><strong>Non-Carcinogenic</strong></td>
</tr>
<tr>
<td><strong>Carcinogenic</strong></td>
</tr>
<tr>
<td><strong>Non-Carcinogenic</strong></td>
</tr>
<tr>
<td>Test</td>
</tr>
<tr>
<td>7.11E-4</td>
</tr>
<tr>
<td>2.06E-2</td>
</tr>
<tr>
<td>7.34E-2</td>
</tr>
<tr>
<td>Reference</td>
</tr>
<tr>
<td>7.98E-4</td>
</tr>
<tr>
<td>2.41E-2</td>
</tr>
<tr>
<td>8.59E-2</td>
</tr>
</tbody>
</table>

Figure 42: Total Carcinogenic and Non-Carcinogenic Risks report section.

**FDA Action and Ecological Risk Levels** – reports mean tissue concentration and steady state corrected mean tissue concentration for chemicals in the current invertebrate, along with three thresholds: FDA action level, ecological effect level, and FDA level of concern, for comparison purposes

<table>
<thead>
<tr>
<th>Location: SW-1, 2, 4, 5</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Chemical</strong></td>
</tr>
<tr>
<td><strong>Mean Tissue Concentration (mg/kg)</strong></td>
</tr>
<tr>
<td><strong>Steady State Corrected Mean Tissue Concentration</strong></td>
</tr>
<tr>
<td><strong>FDA Action Level (mg/kg)</strong></td>
</tr>
<tr>
<td><strong>Ecological Effect Level (mg/kg)</strong></td>
</tr>
<tr>
<td><strong>FDA Level Of Concern (mg/kg)</strong></td>
</tr>
<tr>
<td>Mercury</td>
</tr>
<tr>
<td>3.07E-3</td>
</tr>
<tr>
<td>3.07E-3</td>
</tr>
<tr>
<td>1E0</td>
</tr>
<tr>
<td>2E-1</td>
</tr>
<tr>
<td>NA</td>
</tr>
</tbody>
</table>

Figure 43: FDA Action and Ecological Risk Levels report section.
Contaminant Specific Carcinogenic and Non-Carcinogenic Risks – reports cancer and non-cancer risks from each individual chemical in each of human diet species for test and reference sites

**Important: risk estimates report on those chemicals that are selected by the user (see Select Chemicals of Concern)**
Important: threshold tables report on any chemical that has a threshold and appears in the bioaccumulation lab results, regardless if it is selected as a chemical of concern

**Total Estimated Risks From Organics (see EPA Table Xa)** - compares total risk estimates from sediment sample locations to those of project reference site

<table>
<thead>
<tr>
<th>Total Estimated Risks From Organics (see EPA Table Xa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Receptor: Adult Recreational Angler</td>
</tr>
<tr>
<td>Organism: Nereis virens</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>SW-1, 2, 4, 5</th>
<th>Cancer Risk</th>
<th>Non-Cancer Risk</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Lobster Muscle</td>
<td></td>
</tr>
<tr>
<td>Test</td>
<td>1.13E-3</td>
<td>6.29E-3</td>
</tr>
<tr>
<td>Reference</td>
<td>1.27E-3</td>
<td>8.39E-3</td>
</tr>
</tbody>
</table>

Figure 45: Total Estimated Risks from Organics report section.

**Seafood Non-Cancer Risks (see EPA Table 6a, Columns F & G)** – reports non-cancer risk from individual metal of concern along with reference location results

<table>
<thead>
<tr>
<th>Seafood Non-Cancer Risks (see EPA Table 6a, Columns F &amp; G)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Receptor: Adult Recreational Angler</td>
</tr>
<tr>
<td>Organism: Macoma nasuta</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>SW-1, 2, 4, 5</th>
<th>Arsenic</th>
<th>Non-Cancer Risk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Test</td>
<td></td>
<td>2.15E0</td>
</tr>
<tr>
<td>Reference</td>
<td></td>
<td>2.35E0</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>SW-1, 2, 4, 5</th>
<th>Cadmium</th>
<th>Non-Cancer Risk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Test</td>
<td></td>
<td>2.15E-2</td>
</tr>
<tr>
<td>Reference</td>
<td></td>
<td>1.55E-2</td>
</tr>
</tbody>
</table>

Figure 46: Seafood Non-Cancer Risks report section.

**FDA Action Limit/ Tolerance (see EPA Table 3, Columns D & E)** – reports the FDA threshold value of organics (and mercury) along with steady state corrected mean tissue concentration in the current invertebrate

<table>
<thead>
<tr>
<th>FDA Action Limit/Tolerance (see EPA Table 3, Columns D &amp; E)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Receptor: Adult Recreational Angler</td>
</tr>
<tr>
<td>Organism: Nereis virens</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>SW-1, 2, 4, 5</th>
<th>Contaminant</th>
<th>FDA Action Level (mg/kg)</th>
<th>Steady State Corrected Mean Tissue Concentration (mg/kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mercury</td>
<td>1E0</td>
<td>3.07E-3</td>
</tr>
<tr>
<td>SW-3, 6</td>
<td>Mercury</td>
<td>1E0</td>
<td>3.07E-3</td>
</tr>
<tr>
<td>SW-7, 8, 9 Upper</td>
<td>Mercury</td>
<td>1E0</td>
<td>3.07E-3</td>
</tr>
<tr>
<td>SW-7, 8, 9 Lower</td>
<td>Mercury</td>
<td>1E0</td>
<td>3.07E-3</td>
</tr>
</tbody>
</table>

Figure 47: FDA Action Limit/ Tolerance report section.

**Ecological Effect Level** - reports ecological effect threshold of individual chemicals along with steady state corrected mean tissue concentration in the current invertebrate
**Ecological Effect Level**  
**Receptor:** Adult Recreational Angler  
**Organism:** Nereis virens

<table>
<thead>
<tr>
<th>Contaminant</th>
<th>Ecological Effect Level (mg/kg)</th>
<th>Steady State Corrected Mean Tissue Concentration (mg/kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SW-1, 2, 4, 5</td>
<td>Benzo(a)pyrene 8E3</td>
<td>7.0E0</td>
</tr>
<tr>
<td>SW-1, 2, 4, 5</td>
<td>Anthracene 3.75E3</td>
<td>1.1E0</td>
</tr>
<tr>
<td>SW-1, 2, 4, 5</td>
<td>Atrin 2.96E2</td>
<td>4.1E-1</td>
</tr>
<tr>
<td>SW-1, 2, 4, 5</td>
<td>Arsenic 1.26E1</td>
<td>1.9E0</td>
</tr>
</tbody>
</table>

Figure 48: Ecological Effect Level report section.

**FDA Level of Concern** - reports FDA threshold value of metals along with steady state corrected mean tissue concentration in the current invertebrate

<table>
<thead>
<tr>
<th>Contaminant</th>
<th>FDA Level of Concern (mg/kg)</th>
<th>Steady State Corrected Mean Tissue Concentration (mg/kg)</th>
</tr>
</thead>
</table>

Figure 49: FDA Level of Concern report section.

**Full Report**  
Full report includes the following sections for each location-invertebrate pair:  
**Selected chemicals** – contains list of selected chemicals per location.

<table>
<thead>
<tr>
<th>MBDS</th>
<th>SW-1, 2, 4, 5</th>
<th>SW-3, 6</th>
<th>SW-7, 8, 9</th>
<th>SW-7, 8, 9</th>
</tr>
</thead>
<tbody>
<tr>
<td>128</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>138</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>153</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>170</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>18</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
</tbody>
</table>

Figure 50: Selected chemicals report section.

**Diet Report** – reports cancer risk and the non-cancer risk associated with each human diet species
Prey Report – reports cancer risk and non-cancer risk associated with each prey-predator-human exposure pathway

Chemical Report – reports cancer risk and non-cancer risk associated with each chemical of concern via each human diet chain, along with intermediate calculations; edible tissue concentration, average daily dose, and lifetime average daily dose
BEST reports are also stamped with the version of the software, the date, and the name of the user who ran that iteration of the software. This provides better record keeping for BRAMS users.

<table>
<thead>
<tr>
<th>Diet Report</th>
<th>Cancer Risk</th>
<th>Non-Cancer Risk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fish Fillet</td>
<td>2.339E-8</td>
<td>2.339E-8</td>
</tr>
<tr>
<td>Lobster Muscle</td>
<td>1.871E-7</td>
<td>1.871E-7</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Prey Report</th>
<th>Diet Name</th>
<th>Cancer Risk</th>
<th>Non-Cancer Risk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Worm</td>
<td>Fish Fillet</td>
<td>1.25E-8</td>
<td>1.25E-8</td>
</tr>
<tr>
<td>Clam</td>
<td>Fish Fillet</td>
<td>1.089E-8</td>
<td>1.089E-8</td>
</tr>
<tr>
<td>Worm</td>
<td>Lobster Muscle</td>
<td>1E-7</td>
<td>1E-7</td>
</tr>
<tr>
<td>Clam</td>
<td>Lobster Muscle</td>
<td>8.713E-8</td>
<td>8.713E-8</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Chemical Report</th>
<th>Diet Name</th>
<th>Edible Tissue Concentration</th>
<th>Average Daily Dose</th>
<th>Cancer Risk</th>
<th>Non-Cancer Risk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chem #1</td>
<td>Fish Fillet Worm</td>
<td>0.00103</td>
<td>1.25E-8</td>
<td>1.25E-8</td>
<td>1.25E-8</td>
</tr>
<tr>
<td>Chem #1</td>
<td>Fish Fillet Clam</td>
<td>0.0009</td>
<td>1.089E-8</td>
<td>1.089E-8</td>
<td>1.089E-8</td>
</tr>
<tr>
<td>Chem #2</td>
<td>Fish Fillet Clam</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Chem #1</td>
<td>Lobster Muscle Worm</td>
<td>0.00274</td>
<td>1E-7</td>
<td>1E-7</td>
<td>1E-7</td>
</tr>
<tr>
<td>Chem #1</td>
<td>Lobster Muscle Clam</td>
<td>0.00239</td>
<td>8.713E-8</td>
<td>8.713E-8</td>
<td>8.713E-8</td>
</tr>
<tr>
<td>Chem #2</td>
<td>Lobster Muscle Clam</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>
Software version: BRAMS 3.0
Last date: 04/27/2012
User name: JOHN DOE

Figure 5419: BEST Model Report Identification section.
Part IV: Step by Step Model Creation

This section presents simple, step by step instructions to create and run BRAMS models to conduct bioaccumulation risk assessments. A more detailed description of BRAMS components, and model building and editing is presented in Section II.

1. BEST Model

Below is an example of how to quickly, and efficiently run a Bioaccumulation Risk Assessment using the BEST model. In this example, the user will first create a BEST model for a reference site then create a BEST model for a project test site, and finally generate results that compare the two sites.

This example assumes the user has the following:

(1) The BRAMS software package saved to the computer. If not, see Part I Section 2.2.
(2) The appropriate BEST template file (.best) with all or most necessary project inputs except 28-day bioaccumulation testing results. See Part II Section 5 for an explanation of BRAMS templates. See Part II Section 4.2.1 for a detailed instruction of how to create a new model or template starting with a blank model.
(3) Completed Bioaccumulation Test Result Excel files for the project test and reference sites. If not, see Part II Section 4.3.

Once the user has these items, they may proceed to Step one.

STEP ONE: Getting Started

- Open the BRAMS software by running the executable file BRAMS.jar.
- The StartUp window will then appear while BRAMS is loading.
- If this is the first time the program has been opened, once BRAMS has finished loading, the User Identification window will appear prompting the user to enter their User Name. Enter your User Name and click Continue to continue to the main BRAMS window.
- If BRAMS has been previously opened on this computer, the Main Window will appear automatically and the previously entered User Name will be used.
- From the Main Window, the user is now ready to create, open, edit, and run BRAMS models.

STEP TWO: Open the appropriate BEST template file

- From the main menu select File → Open Project or Template or click in the toolbar.
The Open Project window will appear where the user should locate and select the appropriate BEST template file for the current project. Once the template file has been selected, click *Open* to load the BEST template file data.

**STEP THREE (Optional): Add or Edit Items and Properties**

If the project requires any changes to model items or properties from the selected template model, make changes in the following way:

- **Add or delete project items:**
  - In the Explorer Pane, click and highlight the item category you wish to edit. Items in this category will then appear in the Table Pane.
  - In the Table Pane, Click **[+]** to create a new item in the selected category. In the Properties Pane, enter the name of the item in the Name text box.
  - To delete an item, select it and click **[-]**.

- **Edit item or project properties:**
  - In the Explorer Pane, select the category of the item you wish to enter properties for or select the Project category.
    - Project category: Project properties will appear in the Properties Pane.
    - Item categories: Select the specific item you wish to edit properties for in the Table Pane. The selected item’s properties will then appear in the Properties Pane.
  - In the Properties Pane, edit any properties.
    - For those properties with a **[ ]**: Click the **[ ]** and select an item from the drop down menu.
    - For those properties with a **[ ]**: 
      - Numerical values: If the input value is known with certainty, simply enter the known value in the text box. If the input value is uncertain, click the **[ ]** and enter four values representing the range of possibilities according to trapezoidal fuzzy numbering (See section Part IV section 3).
      - Tables: Click the **[ ]** next to the property. A table will appear where you can input related items by clicking **[ ]** and selecting from a drop down menu of options. Then specify any remaining information in the table relating the items in the property table to the selected item.
    - For those properties with a **✓**: Click inside the box to check or uncheck the box.
  - Continue editing until all project specific item and property changes have been made.

**STEP FOUR: Import 28 Day Bioaccumulation Reference Data**

- In the main menu, select File → Import → Import Bioaccumulation Results.
• The Open Excel window will then appear. Locate and select the Bioaccumulation Test Result Excel file containing project reference data and click *Open*.

• The Import Mode window will then appear, click *Yes* to import the data.

• Reference site data within the imported Bioaccumulation Test Result Excel file should now appear in the 28-days Bioaccumulation Test properties for each Invertabrate included in the test and general project properties should appear in the Project Properties pane.

**STEP FIVE: Name and save the reference site project**

• From the main menu select File → Save Project As or click to name and save the project. Be careful not to overwrite the template file.

• The Save Project As… window will appear prompting the user to assign a name and location to save the new BEST reference project. Select the desired location and name the project, then click *Save*. This will save the project and return you to the Main Window.

**STEP SIX: Import 28 Day Bioaccumulation Test Data**

• In the main menu, select File → Import → Import Bioaccumulation Results.

• The Open Excel window will then appear. Locate and select the Bioaccumulation Test Result Excel file containing project test site data and click *Open*.

• The Import Mode window will then appear, click *Yes* to import the data.

• Test site data within the imported Bioaccumulation Test Result Excel file should now appear in the 28-days Bioaccumulation Test properties for each Invertabrate included in the test and general project properties should appear in the Project Properties pane.

**STEP FIVE: Name and save the test site project**

• From the main menu select File → Save Project As or click to name and save the project. Be careful not to overwrite the reference site file.

• The Save Project As… window will appear prompting the user to assign a name and location to save the new BEST reference project. Select the desired location and name the project, then click *Save*. This will save the project and return you to the Main Window.

**STEP SEVEN: Specify Reference Project**
• In the Explorer pane, click the main Project category (the first category in the Explorer Pane).
• In the Properties Pane, click  to the right of Reference Project in the Reference section.
• The Open window will appear where you can search for and select the file corresponding to the reference site, then click Open.
• The reference project file should now appear in the Reference project input box.

STEP EIGHT: Save the completed test site project

• From the main menu select File → Save Project or click  to overwrite the previous file and save the current project.

STEP NINE: Calculate and display results

• Click  to run the model calculations and view the detailed report or click  to run the model calculations and view the summary report. After the calculation has completed, the Report window will appear showing risk assessment results for the project.

STEP TEN: Save Results

• In the Report window, save the project risk assessment results by clicking  

11. Trophic Trace Model

This section presents simple, step by step instructions to create and run a TT model. In this example, the user will develop an entirely new TT model. Alternatively, if an appropriate template model is available, the user should begin with a template model similar to the method in the BEST model step by step example above.

STEP ONE: Getting Started

• Open the BRAMS software by running the executable file BRAMS.jar.
• The StartUp window will then appear while BRAMS is loading.
• If this is the first time the program has been opened, once BRAMS has finished loading, the User Identification window will appear prompting the user to enter their User Name. Enter your User Name and click Continue to continue to the main BRAMS window.
• If BRAMS has been previously opened on this computer, the Main Window will appear and the previously entered User Name will be used.
From the Main Window, the user is now ready to create, open, edit, and run BRAMS models.

STEP TWO: Open a new TT project

- Click to open a new Trophic Trace project. The blank Trophic Trace framework will then appear in the BRAMS panes.

- As mentioned previously, the user can also begin with an appropriate template model when available. To load a template model, simply click in the toolbar and select the template file in the Open window.

STEP THREE: Add Items and Properties

Add items and their properties to the project in by navigating the Explorer, Table and Properties Panes within the Main Window.

- Add the items you wish to include in the project to each of the item categories (Invertebrates, Fishes, Birds, Mammals, Human, Chemicals, Environment) in the Explorer Pane.
  - In the Explorer Pane, click and highlight the item category you wish to edit. Items in this category will then appear in the Table Pane.
  - In the Table Pane, Click to create a new item in the selected category. To delete an item, select it and click .
  - In the Properties Pane, enter the name of the item in the Name text box.
  - Continue adding and naming items until all model items have been entered into the project.

- Enter the properties of each item in the Properties Pane.
  - In the Explorer Pane, select the category of the item you wish to enter properties for.
  - In the Table Pane, select the item you wish to add properties to.
  - In the Properties Pane, add all properties.
    - For General and Specific properties:
      - For those properties with a Click the and select an item from the drop down menu.
      - For those properties with a If the input value is known with certainty, simply enter the known value in the text box. If the input value is uncertain, click the and enter four values representing the range of possibilities according to trapezoidal fuzzy numbering (See section Part V section 3).
    - For Toxicity and Diet properties:
      - Click the next to the property. A table will appear where you can input related items by clicking and selecting from a drop down menu of options. Then specify any remaining information in...
the table relating the items in the property table to the selected item.
- Continue selecting items and adding properties until all properties have been specified.

STEP FOUR: Save Completed Project

- Click ![File](file_icon) to name and save the project. The Save Project As… window will appear prompting the user to assign a name and location to save the new TT project. Select the desired location and name the project, then click Save. This will save the project and return you to the Main Window.

STEP FIVE: Calculate and display results

- Click ![Flash](flash_icon) to run the model calculations and view the report. After the calculation has completed, the ReportViewer window will appear showing risk assessment results for the project.

STEP SIX: Save Results

- In the ReportViewer window, save the project risk assessment results by clicking ![Save](save_icon)
Part V: BRAMS Modeling Framework

This part summarizes the theoretical foundation of the Trophic Trace and BEST models in terms of input parameters, assumptions, and risk calculations. The information also provides a basis for comparison between the two models in BRAMS.

1. Modeling Framework in Trophic Trace

11.1 Gobas Model for organics

The model used to estimate fish body burdens for hydrophobic organic compounds relies on a steady-state uptake model based on the approach of Gobas (1993 and 1995):

\[ Cf = \frac{k_1 C_{wd} + k d \cdot C_{diet}}{k_2 + k e + k m + k} \]  

(1)

Where:

- \( k_1 \) = gill uptake rate (L/Kg/d)
- \( C_{wd} \) = freely dissolved concentration in water (ng/L)
- \( k_d \) = dietary uptake rate (d\(^{-1}\))
- \( C_{diet} \) = concentration in the diet (\(\mu\)g/kg)
- \( k_2 \) = gill elimination rate (d\(^{-1}\))
- \( k_e \) = fecal egestion rate (d\(^{-1}\))
- \( k_m \) = metabolic rate (d\(^{-1}\))
- \( k_g \) = growth rate (d\(^{-1}\)) is not used in calculations (consider equal = 0)
- \( C_f \) = concentration in fish (\(\mu\)g/kg)

Several sources provide equations for the rate constants (\(k_2\), \(k_e\), \(k_m\) and \(k_g\)) and these are described in greater detail in von Stackelberg et al. (2002).

Biota-sediment accumulation factor (BSAF)

Biota-sediment accumulation factors (BSAFs) used in the model are ratios that describe the relationship between the concentration of a nonpolar organic chemical in the lipid phase in tissue of a sediment-dwelling organism to the concentration in the sediment organic carbon phase to which the organism is exposed. BSAFs are defined as:

\[ BSAF = \frac{(C_B / f_L)}{(C_S / f_{OC})} \]  

(2)

---

* This section is adapted from the TrophicTrace 4.0 Users Manual.
Where:
\[ C_B = \text{concentration of contaminant in biota, mg/kg wet weight} \]
\[ f_L = \text{the fraction lipid of the biota, kg lipid/kg wet weight} \]
\[ C_S = \text{the concentration of contaminant in sediment, mg/kg dry weight} \]
\[ f_{OC} = \text{the fraction organic carbon in sediment, kg organic carbon/kg dry weight} \]

\[ C_B = C_S \cdot \left( \frac{f_L}{f_{OC}} \right) \cdot BSAF \tag{3} \]

Where:
\[ C_B = \text{concentration of contaminant in biota, mg/kg wet weight} \]
\[ f_L = \text{the fraction lipid of the biota, kg lipid/kg wet weight} \]
\[ C_S = \text{the concentration of contaminant in sediment, mg/kg dry weight} \]
\[ f_{OC} = \text{the fraction organic carbon in sediment, kg organic carbon/kg dry weight} \]
\[ BSAF = \text{biota-sediment accumulation factor (typical assumption is 1.0) obtained from site-specific measurements or literature sources} \]

The model can also accept a measured invertebrate concentration resulting from the standard Tier 3 28-day bioaccumulation test results. To account for the fact that these measured concentrations may not have achieved steady-state, a \( K_{ow} \)-dependent adjustment is made (McFarland, 1984; Connell and Hawker, 1988) automatically within BRAMS based on the following formula:

\[
\log t_{ss} = 6.9 \times 10^{-3}(\log K_{ow})^4 - 1.85 \times 10^{-1}(\log K_{ow})^3 + 1.65(\log K_{ow})^2 - 5.34(\log K_{ow}) + 5.93 \tag{4}
\]

Where:
\[ t_{ss} = \text{time required to reach steady-state} \]

11.2 Trophic Transfer Factor (TTF) and Bioconcentration Factor (BCF) for inorganic and hydrophilic organic compounds

Estimates of fish burdens for inorganic and hydrophilic organic compounds rely on two different approaches, depending on data availability. The first approach is a trophic transfer factor (TTF) from prey to predator approach, and the second is a bioconcentration factor (BCF) approach. For some chemicals, there are data available on bioaccumulation from invertebrates to fish (Dillon, 1995). Currently, TTF are available for copper, cadmium, lead, zinc, and arsenic. In the BCF approach, water concentrations are multiplied by a bioconcentration factor to estimate fish body burdens. Water concentrations can either be provided by the user or estimated by the model assuming equilibrium partitioning from sediment.

Both the food web model for hydrophobic organic compounds and the BCF approach for inorganic and hydrophilic organic compounds require a freely dissolved water concentration as an input. Trophic Trace incorporates two approaches for estimating a freely dissolved water
concentration: 1) a user-specified freely dissolved water concentration from site-specific data; 2) from a subroutine (equation 5) using either a user-specified whole water concentration or an estimated whole water concentration (calculated by assuming equilibrium partitioning from a user-specified sediment concentration). The subroutine that estimates a freely dissolved water concentration is shown in equation 5:

\[ C_{wd} = \frac{1}{\text{DOC} \cdot \text{DE}_{oc} \cdot \text{K}_{oc}} \]  

(5)

Where:
- \( C_{wd} \) = freely dissolved concentration in water (ng/L)
- \( \text{DOC} \) = dissolved organic carbon (mg/L)
- \( \text{DE}_{oc} \) = density of organic carbon (0.041 mg OC/mg)
- \( \text{K}_{oc} \) = organic carbon/water partition coefficient (L/kg OC)
- \( \text{POC} \) = particulate organic carbon (mg/L)
- \( C_{ww} \) = whole water concentration (ng/L)

If a whole water concentration is not available, the program uses equilibrium partitioning with sediment to estimate a freely dissolved water concentration. The equation for organic contaminants is:

\[ C_w = C_{oc} \cdot \text{K}_{oc} \]  

(6)

Where:
- \( C_w \) = concentration of freely dissolved chemical in the water (\( \mu \)g/L)
- \( C_{oc} \) = the organic carbon-normalized sediment concentration (\( \mu \)g/kg dry wt sediment) and \( \text{K}_{oc} \) = organic carbon-water partition coefficient (L/kg organic carbon)

The \( \text{K}_{oc} \) for each chemical can be estimated from its octanol-water partition coefficient, \( \text{K}_{ow} \), according to the following regression relationship (Connell and Hawker, 1988):

\[ \log \text{K}_{oc} = 0.00028 + 0.983 \log_{10}\text{K}_{ow} \]  

(7)

11.3 Risk Assessment Formulas

**Human Health Risk**
The estimates of fish body burdens represent point estimates of concentrations to which humans are exposed via fish ingestion. These fish tissue concentrations are used along with exposure assumptions specific to each human receptor population to calculate carcinogenic risk and non-carcinogenic hazard indices. Carcinogenic risk is calculated as follows:

\[
\text{Risk} = \frac{\text{CSF} \times \text{IR}_f \times \text{C}_f \times \text{ED}}{\text{BW} \times 1000000 \times \text{AT}}
\]

is always equal 70 (constant)

Where:
- Risk = incremental lifetime cancer risk
- CSF = cancer slope factor (mg/kg-day)^{-1}
- \(\text{IR}_f\) = annualized fish ingestion rate (g/day)
- \(\text{C}_f\) = concentration in fish (\(\mu\)g/kg)
- ED = exposure duration (days)
- BW = body weight (kg)
- AT = averaging time (days)

Non-carcinogenic hazard indices are calculated as follows:

\[
\text{HI} = \frac{\text{IR}_f \times \text{C}_f \times \text{ED}}{\text{RfD} \times \text{BW} \times 1000000 \times \text{AT}}
\]

Where:
- HI = hazard index
- RfD = Reference dose (mg/kg-day)
- \(\text{IR}_f\) = annualized fish ingestion rate (g/day)
- \(\text{C}_f\) = concentration in fish (\(\mu\)g/kg)
- ED = exposure duration (days)
- BW = body weight (kg)
- AT = averaging time (days)

**Ecological risk**

Potential ecological risks are evaluated by comparing predicted contaminant concentrations in tissue and/or daily dose estimates to appropriate toxicity reference values (TRVs). These comparisons are based on predicted tissue concentrations in mg/kg for fish, and on predicted daily dose estimates for the higher order ecological receptors.

TRVs are levels of exposure associated with either Lowest Observed Adverse Effects Levels (LOAELs) or No Observed Adverse Effects Levels (NOAELs). They provide a basis for...
judging the potential effects of measured or predicted exposures that are above or below these levels. TRVs are contaminant- and species-specific and are developed based on laboratory or field studies.

Use of both LOAELs and NOAELs provides perspective on the potential for risk as a result of exposure to contaminants in dredged materials. LOAELs are values at which effects have been observed in either laboratory or field studies, while the NOAEL represents the lowest dose or body burden at which an ecologically relevant effect was not observed. Exceedance of a LOAEL indicates a greater potential for risk.

Some studies examine toxicity endpoints (such as lethality, growth, and reproduction) that are thought to have greater potential for adverse effects on populations of organisms than other studies. Other studies examine toxicity endpoints such as behavior, disease, cell structure, or biochemical changes that affect individual organisms, but may not result in adverse effects at the population level. For example, toxic effects such as enzyme induction may or may not result in adverse effects to individual animals or populations. The procedure in Trophic Trace is to develop TRVs from studies that examine the effects of contaminants on lethality, growth or reproduction. Studies that examined the effects of contaminants on other sublethal endpoints are not used to select TRVs unless no other studies are available. Lethality, growth, and reproductive-based endpoints typically present the greatest risk to the viability of the individual organism and therefore of the population’s survival. Thus, these are considered to be the endpoints of greatest concern.

When exposures are expected to be long-term, data from studies of chronic exposure are preferable to data from medium-term (subchronic), short-term (acute), or single-exposure studies (USEPA, 1997c). Bioaccumulative substances are by definition persistent, and exposure of ecological receptors to these contaminants from dredged materials is expected to be long-term, and therefore studies of chronic exposure are preferentially used to select TRVs. Long-term studies are also preferred since reproductive effects of contaminants are typically studied after long-term exposure.

Dose-response studies compare the response of organisms exposed to a range of doses to that of a control group. Ideally, doses that are below and above the threshold level that causes adverse effects are examined. Toxicity endpoints determined in dose-response and other studies include:

- **NOAEL (No-Observed-Adverse-Effect-Level)** is the highest exposure level shown to be without adverse effect in organisms exposed to a range of doses. NOAELs may be expressed as dietary doses (e.g., mg contaminant consumed/kg body weight/d), as concentrations in external media (e.g., mg contaminant/kg food), or as concentrations in tissue of the affected organisms (e.g., mg chemical/kg egg).

- **LOAEL (Lowest-Observed-Adverse-Effect-Level)** is the lowest exposure level shown to produce adverse effect in organisms exposed to a range of doses. LOAELs may also be expressed as dietary doses (e.g., mg contaminant consumed/kg body weight/d), as
concentrations in external media (e.g., mg contaminant/kg food), or as concentrations in tissue of the effected organisms (e.g., mg chemical/kg egg).

- **LD<sub>50</sub>** is the Lethal Dose that results in death of 50% of the exposed organisms. Expressed in units of dose (e.g., mg contaminant administered/kg body weight of test organism/d).

- **LC<sub>50</sub>** is the Lethal Concentration in some external media (e.g., food, water, or sediment) that results in death of 50% of the exposed organisms. Expressed in units of concentration (e.g., mg contaminant/kg wet weight food).

- **ED<sub>50</sub>** is the Effective Dose that results in a sublethal effect in 50% of the exposed organisms (mg/kg/d).

- **EC<sub>50</sub>** is the Effective Concentration in some external media that results in a sublethal effect in 50% of the exposed organisms (mg/kg).

- **CBR** or Critical Body Residue is the concentration in the organism (e.g., whole body, liver, or egg) that is associated with an adverse effect (mg contaminant/kg wet wt tissue).

- **EL-effect** is the effect level that results in an adverse effect in organisms exposed to a single dose, rather than a range of doses. Expressed in units of dose (mg/kg/d) or concentration (mg/kg).

- **EL-no effect** is the effect level that does not result in an adverse effect in organisms exposed to a single dose, rather than a range of doses. Expressed in units of dose (mg/kg/d) or concentration (mg/kg).

Most USEPA risk assessments typically estimate risk by comparing the exposure of receptors of concern to TRVs that are based on NOAELs. Example TRVs included in Trophic Trace are developed on the basis of both NOAELs and LOAELs to provide perspective on the range of potential effects relative to measured or modeled exposures.

Differences in the feeding behavior of aquatic and terrestrial organisms determine the type of toxicity endpoints that are most easily measured and most useful in assessing risk. For example, the dose consumed in food is more easily measured for terrestrial animals than for aquatic organisms since uneaten food can be difficult to collect and quantify in an aqueous environment. Therefore, for aquatic organisms, toxicity endpoints are more often expressed as concentrations in external media (e.g., water) or as accumulated concentrations in the tissue of the exposed organism (also called a “body burden”). In some studies, doses are administered via gavage, intraperitoneal injection into an adult, or injection into a fish or bird egg. If appropriate studies are available, TRVs in Trophic Trace are selected on the basis of the most likely route of exposure, as described below:

- **TRVs for fish** are expressed as critical body residues (CBR) (e.g., mg/kg whole body weight and mg/kg lipid in eggs).
• TRVs for terrestrial receptors (e.g., birds and mammals) are expressed as daily dietary doses (e.g., mg/kg whole body wt/d).

• TRVs for birds are also expressed as concentrations in eggs (e.g. mg/kg wet wt egg).

The toxicity quotient in ecological risk assessments is calculated as follows:

\[
TQ = \frac{\sum IR_f \times C_f \times frac}{TRV \times BW}
\]

Where:

- TQ = Toxicity Quotient
- IR_f = annualized ingestion rate (kg/day)
- C_f = concentration in prey (mg/kg)
- frac = fraction in diet
- TRV = toxicity reference value (mg/kg/day)
- BW = body weight (kg)
12. Framework for BEST Model

The BEST model framework is based on the 1999 EPA Region 1 Bioaccumulation Risk Assessment Model.

12.1 Invertebrate tissue concentration

The BEST model predicts invertebrate tissue concentration using 28-day bioaccumulation test results and chemical properties in the following equations for metals and organics respectively.

Equation 1: BEST Formula for Invertebrate Edible Tissue Concentration (Metals)

\[
[\text{Edible Tissue}_{\text{Invertebrate, Metal}}] (\text{mg/kg}) = [\text{C}_{\text{prey}}] \times \text{SSCF}
\]

Where:
\[
[\text{Edible Tissue}_{\text{Invertebrate, Metal}}] = \text{Concentration of metal contaminant in edible tissue of invertebrate species (µg/g)}
\]
\[
[\text{C}_{\text{prey}}] = \text{concentration of contaminant in the invertebrate in the maximum replicate of five replicates from the 28-day bioaccumulation test data depending on the user’s choice (µg/g)}
\]
\[
\text{SSCF} = \text{steady state correction factor (unitless)}
\]

Equation 2: BEST Formula for Invertebrate Edible Tissue Concentration (Organics)

\[
[\text{Edible Tissue}_{\text{Invertebrate, Organic}}] (\text{mg/kg}) = \frac{[\text{C}_{\text{prey}}] \times \text{SSCF}}{1000}
\]

Where:
\[
[\text{Edible Tissue}_{\text{Invertebrate, Organic}}] = \text{Concentration of organic contaminant in edible tissue of invertebrate species (mg/kg)}
\]
\[
[\text{C}_{\text{prey}}] = \text{concentration of contaminant in the invertebrate in the maximum replicate of five replicates from the 28-day bioaccumulation test data (ng/g)}
\]
\[
\text{SSCF} = \text{steady state correction factor (unitless)}
\]
\[
1000 = \text{Unit Normalization Factor [Edible Tissue}_{\text{Invertebrate, Organic}} (\text{ng/g} \rightarrow \mu\text{g/g})}
\]

12.2 Trophic Transfer

The BEST model predicts the chemical concentrations in predator species by applying a trophic transfer model to the measured contaminant concentrations in the benthic invertebrate test species using the following equation:

Equation 3: BEST Formula for Body Burdens in Predator Tissue

\[
[\text{Edible Tissue}_{\text{Predator}}] (\text{mg/kg}) = [\text{Edible Tissue}_{\text{Invertebrate}}] \times \text{BMF} \times \left( \frac{\text{Lipid}_{\text{pred}}}{\text{Lipid}_{\text{prey}}} \right)
\]
Where:

[Edible Tissue\textsubscript{Predator}] = Concentration of contaminant in edible tissue of predator (mg/kg)
[Edible Tissue\textsubscript{Invertebrate}] = Concentration of contaminant in edible tissue of invertebrate species (mg/kg)
Lipid\textsubscript{Pred} = predator mean lipid fraction (g lipid/g tissue)
Lipid\textsubscript{Prey} = invertebrate mean lipid fraction (g lipid/g tissue)
BMF = biomagnification factor (unitless)

12.3 Human Health Risk

Once the edible tissue concentration is calculated using the trophic transfer model, the result is used to determine the dose to humans that consume these species. The lifetime average daily dose (LADD) is calculated for cancer risks and non-cancer risks.

The LADD is calculated using the following equation:

\textbf{Equation 4: BEST Lifetime Average Daily Dose Equation}

\[
\text{LADD (mg/kg-day)} = \frac{\text{ETC x FI x F x IR x ED}}{\text{BW x LT}}
\]

Where:

LADD = Lifetime Average Daily Dose (mg/kg-day)
ETC = Edible Tissue Concentration of diet species (mg/kg)
FI = Fraction Ingested (unitless)
F = Frequency (days/year)
IR = Fish/shellfish Ingestion Rate (kg/day)
ED = Exposure Duration (years)
BW = Body Weight (kg)
LT = Lifetime (days)

To determine the carcinogenic risk level, the LADD is multiplied by an oral cancer slope factor (CSF) according to the following equation:

\textbf{Equation 5: Standard Cancer Risk Equation}

\[
\text{Cancer Risk} = \text{LADD x CSF}
\]

Where:

LADD = Lifetime Average Daily Dose (mg/kg-day)
CSF = Oral Cancer Slope Factor (mg/kg)

To determine the non-carcinogenic hazard (i.e., hazard quotient), the LADD is divided by
reference dose (RfD) according to the following equation:

Equation 6: Standard Non-cancer Risk Equation

\[ \text{Non-cancer Risk} = \frac{\text{LADD}}{\text{RfD}} \]

Where:
- \( \text{LADD} \) = Lifetime Average Daily Dose (mg/kg-day)
- \( \text{RfD} \) = Oral Reference Dose (mg/kg-day)

Non-cancer risks greater than 1 are considered indicative of potential health effects. For cancer risks, an acceptable risk range of \(1 \times 10^{-6}\) to \(1 \times 10^{-4}\) is typically applied. Risk thresholds can be set in the Risk Thresholds section of the General properties.

12.4 Model Assumptions

**Trophic Transfer**
As previously described, the BEST risk model predicts edible tissue concentrations in predator species by applying a trophic transfer model to the measured concentrations in invertebrate test species (Equation 1) based on the results of the 28-day bioaccumulation testing. This approach assumes that the test species represent the prey species typically consumed by predators, and that 100 percent of the organisms consumed are exposed to the dredged material.

**Steady State Correction Factor**
A steady state correction factor (SSCF) is used to correct for underestimates of prey tissue concentrations in standard 28-day bioaccumulation testing. Because the standard testing period (28 days) might not be long enough for the exposed organisms to reach a state of equilibrium with their environment, the SSCF is applied to further estimate prey tissue concentrations under natural exposure periods that are longer than the standard testing duration.

**Biomagnification Factor**
The biomagnification factor (BMF) accounts for accumulation of chemicals in predator tissues from consumption of invertebrate prey. Chemicals that biomagnify, or increase concentration up the food chain, will have BMFs >1 while those that do no biomagnify will have BMFs ≤ 1.

**Toxicity Equivalency Factor**
To address potential toxicity for dioxin compounds and for selected PCBs and PAHs, a toxicity equivalency factor (TEF) approach should be applied by the user. In this approach, specific compounds for which toxicity information are uncertain or unavailable are assigned a TEF value that estimates their toxicity relative to a compound of known toxicity (called a TEF Reference Chemical in BEST). For example, individual dioxin congeners, as well as coplanar PCB compounds, should each be assigned a value ranging from 0 to 1 indicating their toxicity relative to 2,3,7,8-TCDD. The measured concentrations of these compounds are multiplied by the risk
factors (CSF and RfD) of the TEF Reference Chemical they are related to in order to determine their cancer and non-cancer risks.

**Risks**

As previously discussed, the BEST model estimates the carcinogenic and non-carcinogenic risk to human health based on the consumption of species directly exposed to contaminants in the sediment and their predators using standard risk equations (EPA, 1989b) based on the lifetime average daily dose (LADD) and average daily dose (ADD), respectively.

13. Interval Analysis of “Fuzzy Math” to Characterize Parameter Uncertainty In Trophic Trace³*

13.1 Trapezoidal fuzzy numbers

A trapezoidal fuzzy number is simply four numerical values \([A, B, C, D]\) where \(A\) is less than or equal to \(B\), \(B\) is less than or equal to \(C\), and \(C\) is less than or equal to \(D\). For the fuzzy parameter \(F=[A, B, C, D]\) the interval \([A,D]\) represents the plausible range of the parameter. The number \(A\) is the minimum possible value of the parameter, and \(D\) is the maximum possible value of the parameter. The range \([B,C]\) is the most likely range of the parameter \(F\). So, fuzzy results yield both “worst case” and “best estimates” simultaneously.

Trapezoidal fuzzy numbers is an example of a fuzzy set and could be represented via its membership function showing the degree of membership for each value of the parameter (see Figure 20).

Degree of membership is a number between 0 and 1. The range of certain values of the parameter have a membership level equal to one (green line on Figure 19 corresponding to the interval \([B,C]\)), restricted values with a degree of membership equal to zero are shown in red. All other values are more or less possible in proportion to their membership degree. This approach allows us to consider the fuzzy set as a measure for possibility (Zimmermann, 1991). Note that the y-axis does not represent a probability or likelihood. The degree of membership in the fuzzy set is proportional, however, such that if the degree of membership = 1 (B to C, also called the likeliest or probable range), then the parameter value, given the inputs, will definitely be within that range. The parameter may take on values from the sides of the trapezoid (A to B and C to D, also called the full or possible range), but these values are only “possibilities” with the degree of possibility reflected in the degree of membership. For example, a value that has a degree of membership = 0.5 (A to B and C to D) is a “possibility” with a degree of possibility of 0.5. The y-axis does not represent a probability or likelihood.

³* This section is adapted from the TrophicTrace 4.0 Users Manual.

membership of 0.8 is much more possible than a value with a degree of membership that is only 0.1.

13.2 Example: Interval

In the case when all possible values of the parameter are equally plausible (e.g., equivalent to a uniform distribution), then the range of the parameter can be described by an interval and interval analysis is used to analyze a model with such parameters. The membership function for an interval is a stepped function (see Figure 21).

This approach also provides the possibility to consider the interval \([A,D]\) as a trapezoidal fuzzy number \([A,B,C,D]\) with \(B=A\) and \(C=D\), i.e. as the fuzzy number \([A,A,D,D]\), and to use such parameters in modeling. In this case, the short representation \([A,D]\) can be used instead of \([A,A,D,D]\).

13.3 Triangular Fuzzy Numbers as a Particular Case of Trapezoidal Fuzzy Numbers

Trapezoidal fuzzy numbers also include fuzzy numbers with a triangular shape for the membership function. A triangular fuzzy number is evaluated as a trapezoidal fuzzy number \([A,B,C,D]\) with \(B=C\), i.e. \([A,B,B,D]\). Such a fuzzy number could be used for a quantitative description of a parameter for which a possible range is known together with a single most likely value. This is shown graphically in Figure 22. The short notation for a triangular fuzzy number is \([A,B,D]= [A,B,D,D]\)

13.4 Exact Parameter Value

It might be possible to know or only have information for one value for some parameters in the model. The approach to treat them as a trapezoidal fuzzy number \([A,B,C,D]\) with \(A=B=C=D: [A,A,A,A]\) allows the model to include such parameters simultaneously with other parameters that are more uncertain. Zadeh (1965) provides an implementation for processing of fuzzy numbers by the extension principle. Note, that all following notations are equivalent: \(A=[A]=[A,A]=[A,A,A]= [A,A,A,A]\); all denote standard (not fuzzy) numbers as the particular case of the fuzzy number.
Trophic Trace performs the extension principle for the model equation process, but approximates results by trapezoidal shapes, too. The approximation approach uses the vertex method (Dong and Shah, 1987) for computing a function of fuzzy variables.

13.5 Arithmetic of Trapezoidal Fuzzy Numbers

3.5.1 Addition

According to the extension principle, the sum of two trapezoidal fuzzy numbers is also a trapezoidal number. The following formula provides the exact value used by Trophic Trace.

\[
[A_1, B_1, C_1, D_1] + [A_2, B_2, C_2, D_2] = [A_1 + A_2, B_1 + B_2, C_1 + C_2, D_1 + D_2]
\]  (10)

3.5.2 Subtraction

As for subtraction, the extension principle provides an exact solution for this operation, as shown in the following formula.

\[
[A_1, B_1, C_1, D_1] - [A_2, B_2, C_2, D_2] = [A_1 - A_2, B_1 - B_2, C_1 - C_2, D_1 - D_2]
\]  (11)

3.5.3 Multiplication

Trophic Trace uses the following approximate formula for multiplication of fuzzy numbers.

\[
[A_1, B_1, C_1, D_1] \times [A_2, B_2, C_2, D_2] \sim [A_1 \times A_2, B_1 \times B_2, C_1 \times C_2, D_1 \times D_2]
\]  (12)

The vertex method is based on \(\alpha\)-cut conception and interval analysis. It can be shown that the exact solution of multiplying trapezoidal fuzzy numbers has a curvilinear trapezium shape (so the result of multiplication of trapezoidal fuzzy numbers is not trapezoidal itself: this is the purpose of using the “approximate” sign “\(\sim\)”. We approximate the result by the trapezoidal fuzzy number anyway. The vertexes of this curvilinear relationship are calculated using formula (12) from above.

3.5.4 Division

\(\alpha\)-cut conception and interval analysis provides the following formula used in the Trophic Trace model for operation of division of positive trapezoidal fuzzy numbers.

\[
[A_1, B_1, C_1, D_1] / [A_2, B_2, C_2, D_2] \sim [A_1 / D_2, B_1 / C_2, C_1 / B_2, D_1 / A_2]
\]  (13)

As in interval analysis, the multiplication and division of fuzzy numbers are inverse to each other only for the case when all fuzzy parameters are exact values (all four components are equal). If parameter F has plausible range \([A_1, D_1]\) and the plausible range for parameter Y is \([A_2, D_2]\), then...
to obtain the minimum value for the parameter F/Y one needs to divide the minimum value of the parameter F by the maximum value of the parameter Y. The maximum value of F/Y is obtained by dividing maximum F by minimum Y.

### 3.5.5 Power operations

The extension principle provides an exact solution for extending the exponent function for fuzzy numbers.

\[
\text{EXP} [A, B, C, D] = [\text{EXP} (A), \text{EXP} (B), \text{EXP} (C), \text{EXP} (D)]
\]  

(14)

This function is the particular case of a power function for which the extension principle also provides the exact solution, as shown below.

\[
= [A_1^{A_2}, B_1^{B_2}, C_1^{C_2}, D_1^{D_2}]
\]  

(15)
PART VI: Appendices

APPENDIX A: References


New Jersey Marine Sciences Consortium (NJMSC) and New Jersey Department of Agriculture (NJDA). (1994). Fish consumption patterns by New Jersey Consumers and Anglers. Prepared for New Jersey Department of Environmental Protection and Energy, Division of Science and Research. August.


United States Environmental Protection Agency (USEPA). (2000). Proposed changes to the bioaccumulation testing evaluation framework and response to scientific peer reviewers comments on the existing framework for determining the suitability of dredged material to be placed at the Historic Area Remediation Site (HARS). USEPA, Region 2, New York, October.


## APPENDIX B: ACRONYMS

<table>
<thead>
<tr>
<th>Acronym</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ADDAMS</td>
<td>USACE fate and transport modeling system</td>
</tr>
<tr>
<td>BEST</td>
<td>Bioaccumulation Evaluation Screening Tool</td>
</tr>
<tr>
<td>BCF</td>
<td>Bioconcentration Factor</td>
</tr>
<tr>
<td>BSAF</td>
<td>Biota Sediment Accumulation Factor (here used only as a benthic sediment accumulation factor)</td>
</tr>
<tr>
<td>BRAMS</td>
<td>Bioaccumulation Risk Assessment Modeling Software</td>
</tr>
<tr>
<td>CSF</td>
<td>Cancer Slope Factor (mg/kg-day)$^{-1}$</td>
</tr>
<tr>
<td>DOC</td>
<td>Dissolved Organic Carbon</td>
</tr>
<tr>
<td>ITM</td>
<td>Inland Testing Manual</td>
</tr>
<tr>
<td>$K_{oc}$</td>
<td>Log</td>
</tr>
<tr>
<td>$K_{ow}$</td>
<td>Log-octanol water partitioning coefficient</td>
</tr>
<tr>
<td>LADD</td>
<td>Lifetime Average Daily Dose</td>
</tr>
<tr>
<td>LOAEL</td>
<td>Lowest Observed Adverse Effect Level for ecological receptors</td>
</tr>
<tr>
<td>NJDA</td>
<td>New Jersey Department of Agriculture</td>
</tr>
<tr>
<td>NOAEL</td>
<td>No Observed Adverse Effect Level for ecological receptors</td>
</tr>
<tr>
<td>NY/NJ</td>
<td>New York/New Jersey</td>
</tr>
<tr>
<td>OTM</td>
<td>Ocean Testing Manual</td>
</tr>
<tr>
<td>PCBs</td>
<td>Polychlorinated Biphenyls</td>
</tr>
<tr>
<td>POC</td>
<td>Particulate Organic Carbon</td>
</tr>
<tr>
<td>RfD</td>
<td>Reference Dose (mg/kg-day)</td>
</tr>
<tr>
<td>TBP</td>
<td>Theoretical Bioaccumulation Potential</td>
</tr>
<tr>
<td>TT</td>
<td>Trophic Trace</td>
</tr>
<tr>
<td>TTF</td>
<td>Trophic Transfer Factor (from invertebrates to fish)</td>
</tr>
<tr>
<td>TRV</td>
<td>Toxicity Reference Value for ecological receptors</td>
</tr>
<tr>
<td>USACE</td>
<td>United States Army Corps of Engineers</td>
</tr>
<tr>
<td>USEPA</td>
<td>United States Environmental Protection Agency</td>
</tr>
</tbody>
</table>