Starting Soon: Bioavailability of Contaminants in Soil: Considerations for Human Health Risk Assessment



- ► Access online document: http://bcs-1.itrcweb.org/
- ▶ Download PowerPoint file
 - CLU-IN training page at http://www.clu-in.org/conf/itrc/bcs/
 - Under "Download Training Materials"
- ▶ Download Decision Process Flowchart, BCS-1 Definition of Terms, and Review Checklist, for reference during the training class
 - https://clu-in.org/conf/itrc/bcs/ITRC-BCS-TrainingHandouts.pdf
- ▶ Using Adobe Connect
 - Related Links (on right)
 - Select name of link
 - Click "Browse To"
 - Full Screen button near top of page





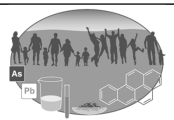
Poll Question: What is Your Experience Level with Soil Contaminant Bioavailability?

- little or no experience
- · some knowledge and experience
- expert

Welcome – Thanks for joining this ITRC Training Class



Bioavailability of Contaminants in Soil: Considerations for Human Health Risk Assessment



Bioavailability of Contaminants in Soil: Considerations for Human Health Risk Assessment (BCS-1) ITRC Technical and Regulatory Guidance document

Sponsored by: Interstate Technology and Regulatory Council (www.itrcweb.org)
Hosted by: US EPA Clean Up Information Network (www.cluin.org)

Training Course Overview:

ITRC Guidance: Bioavailability of Contaminants in Soil: Considerations for Human Health Risk Assessment (BCS-1) http://bcs-1.itrcweb.org/

Risk-based cleanup goals are often calculated assuming that chemicals present in soil are absorbed by humans as efficiently as the chemicals dosed during the toxicity tests used to determine regulatory toxicity values (such as the Reference Dose or Cancer Slope Factor). This assumption can result in inaccurate exposure estimates and associated risks for some contaminated sites because the amount of a chemical absorbed (the chemical's bioavailability) from contaminated soil can be a fraction of the total amount present. Properly accounting for soil-chemical interactions on the bioavailability of chemicals from soil can lead to more accurate estimates of exposures to soil contaminants and improve risk assessments by decreasing uncertainty.

The basis for this training course is the ITRC guidance: <u>Bioavailability of Contaminants in Soil: Considerations for Human Health Risk Assessment (BCS-1)</u>. This guidance describes the general concepts of the bioavailability of contaminants in soil, reviews the state of the science, and discusses how to incorporate bioavailability into the human health risk assessment process. This guidance addresses lead, arsenic, and polycyclic aromatic hydrocarbons (PAHs) because evaluating bioavailability is better understood for these chemicals than for others, particularly for the incidental ingestion of soil.

The target audience for this guidance and training course are:

- Project managers interested in decreasing uncertainty in the risk assessment which may lead to reduced remedial action costs.
- Risk assessors new to bioavailability or those who want additional confidence and training in the current methods and common practices for using bioavailability assessment to more accurately determine human health risk at a contaminated site.

As a participant in this training you should learn to:

- Value the ITRC document as a "go-to" resource for soil bioavailability
- Apply the decision process to determine when a site-specific bioavailability assessment may be appropriate
- Use the ITRC Review Checklist to develop or review a risk assessment that includes soil bioavailability
- Consider factors that affect arsenic, lead and PAH bioavailability
- Select appropriate methods to evaluate soil bioavailability
- Use tools to develop site-specific soil bioavailability estimates and incorporate them into human health risk assessment

Learners can envision themselves implementing the ITRC guidance through case study applications. Training participants are encouraged to view the associated ITRC guidance, <u>Bioavailability of Contaminants in Soil: Considerations for Human Health Risk Assessment (BCS-1)</u> prior to attending the class.

ITRC (Interstate Technology and Regulatory Council) www.itrcweb.org
Training Co-Sponsored by: US EPA Technology Innovation and Field Services Division (TIFSD) (www.clu-in.org)
ITRC Training Program: training@itrcweb.org; Phone: 402-201-2419

Housekeeping



- ► Course time is 2¼ hours
- ► This event is being recorded
- ▶ Trainers control slides
 - Want to control your own slides? You can download presentation file on Clu-in training page

- Questions and feedback
 - Throughout training: type in the "Q & A" box
 - At Q&A breaks: unmute your phone with #6 to ask out loud
 - At end of class: Feedback form available from last slide
 - Need confirmation of your participation today? Fill out the feedback form and check box for confirmation email and certificate

Copyright 2019 Interstate Technology & Regulatory Council, 1250 H Street, NW, Suite 850, Washington, DC 20005

Notes:

I'm sure that some of you are familiar with these rules from previous CLU-IN events, let's run through them quickly for our new participants.

We have started the seminar with all phone lines muted to prevent background noise. Please keep your phone lines muted during the seminar to minimize disruption and background noise. During the question and answer break, press #6 to unmute your lines to ask a question (note: *6 to mute again). Also, please do NOT put this call on hold as this may bring unwanted background music over the lines and interrupt the seminar.

Use the "Q&A" box to ask questions, make comments, or report technical problems any time. For questions and comments provided out loud, please hold until the designated Q&A breaks.

Everyone – please complete the feedback form before you leave the training website. Link to feedback form is available on last slide.

ITRC (<u>www.itrcweb.org</u>) – Shaping the Future of Regulatory Acceptance



- Host organization
 - Network
 - State regulators
 - All 50 states, PR, DC
 - Federal partners









- Academia
- · Community stakeholders
- ▶ Follow ITRC







Disclaimer

- Full version in "Notes" section
- Partially funded by the U.S. government
 - ITRC nor US government warranty material
 - ITRC nor US government endorse specific products
- ► ITRC materials available for your use – see <u>usage policy</u>
- Available from www.itrcweb.org
 - Technical and regulatory guidance documents
 - Online and classroom training schedule
 - More...

The Interstate Technology and Regulatory Council (ITRC) is a state-led coalition of regulators, industry experts, citizen stakeholders, academia and federal partners that work to achieve regulatory acceptance of environmental technologies and innovative approaches. ITRC consists of all 50 states (and Puerto Rico and the District of Columbia) that work to break down barriers and reduce compliance costs, making it easier to use new technologies and helping states maximize resources. ITRC brings together a diverse mix of environmental experts and stakeholders from both the public and private sectors to broaden and deepen technical knowledge and advance the regulatory acceptance of environmental technologies. Together, we're building the environmental community's ability to expedite quality decision making while protecting human health and the environment. With our network of organizations and individuals throughout the environmental community, ITRC is a unique catalyst for dialogue between regulators and the regulated community.

For a state to be a member of ITRC their environmental agency must designate a State Point of Contact. To find out who your State POC is check out the "contacts" section at www.itrcweb.org. Also, click on "membership" to learn how you can become a member of an ITRC Technical Team.

Disclaimer: This material was prepared as an account of work sponsored by an agency of the United States Government. Neither the United States Government nor any agency thereof, nor any of their employees, makes any warranty, express or implied, or assumes any legal liability or responsibility for the accuracy, completeness, or usefulness of any information, apparatus, product, or process disclosed, or represents that its use would not infringe privately owned rights. Reference herein to any specific commercial product, process, or service by trade name, trademark, manufacturer, or otherwise does not necessarily constitute or imply its endorsement, recommendation, or favoring by the United States Government or any agency thereof. The views and opinions of authors expressed herein do not necessarily state or reflect those of the United States Government or any agency thereof and no official endorsement should be inferred.

The information provided in documents, training curricula, and other print or electronic materials created by the Interstate Technology and Regulatory Council ("ITRC" and such materials are referred to as "ITRC Materials") is intended as a general reference to help regulators and others develop a consistent approach to their evaluation, regulatory approval, and deployment of environmental technologies. The information in ITRC Materials was formulated to be reliable and accurate. However, the information is provided "as is" and use of this information is at the users' own risk.

ITRC Materials do not necessarily address all applicable health and safety risks and precautions with respect to particular materials, conditions, or procedures in specific applications of any technology. Consequently, ITRC recommends consulting applicable standards, laws, regulations, suppliers of materials, and material safety data sheets for information concerning safety and health risks and precautions and compliance with then-applicable laws and regulations. ITRC, ERIS and ECOS shall not be liable in the event of any conflict between information in ITRC Materials and such laws, regulations, and/or other ordinances. The content in ITRC Materials may be revised or withdrawn at any time without prior notice.

ITRC, ERIS, and ECOS make no representations or warranties, express or implied, with respect to information in ITRC Materials and specifically disclaim all warranties to the fullest extent permitted by law (including, but not limited to, merchantability or fitness for a particular purpose). ITRC, ERIS, and ECOS will not accept liability for damages of any kind that result from acting upon or using this information.

ITRC, ERIS, and ECOS do not endorse or recommend the use of specific technology or technology provider through ITRC Materials. Reference to technologies, products, or services offered by other parties does not constitute a guarantee by ITRC, ERIS, and ECOS of the quality or value of those technologies, products, or services. Information in ITRC Materials is for general reference only; it should not be construed as definitive guidance for any specific site and is not a substitute for consultation with qualified professional advisors.

Meet the ITRC Trainers





Geoff Siemering
University of Wisconsin –
Madison
Madison, WI
608-262-9969
geoff.siemering@wisc.edu



Valerie Hanley California DTSC Sacramento, CA 916-255-6440 Valerie.Hanley@dtsc.ca.gov



Anita Meyer
US Army Corps of
Engineers
Omaha, NE
402-697-2585
Anita.K.Meyer@usace.army
.mil



Barrie Selcoe Jacobs Houston, TX 281-246-4322 barrie.selcoe@jacobs.com



Anita.K.Meyer@usace.army .mil Kevin Long Terraphase Engineering Inc. Princeton, NJ 609-462-2855 kevin.long@terraphase.com

Read trainer bios at https://clu-in.org/conf/itrc/bcs/

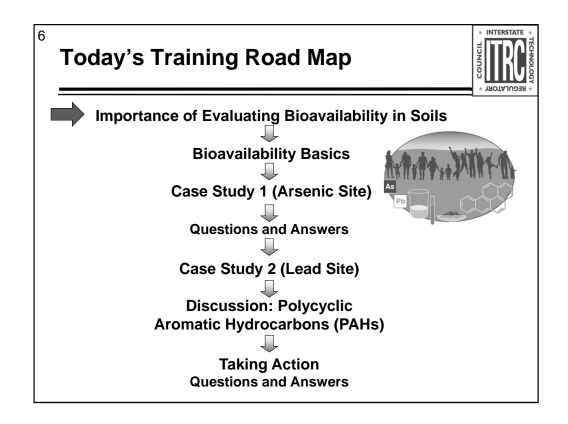
Geoffrey Siemering is a researcher with the Department of Soil Science at University of Wisconsin in Madison. Beginning his work with UW-Madison in 2014, Geoff conducts research and develops outreach programming on soil contaminant issues at the interface of public health and environmental regulation. Recent projects include bioavailability of lead in urban soils, reuse of lead and zinc mine-scarred agricultural land, quantification of cheesemaking and vegetable processing facility wastewater soil denitrification, and determination of anthropogenic polycyclic aromatic hydrocarbon baseline values for urban Wisconsin. He also has experience with triad-approach monitoring of aquatic herbicide impacts, and radionuclide waste disposal. Prior to UW-Madison, Geoff worked for the San Francisco Estuary Institute and Lawrence Livermore National Laboratory. Since 2015, Geoff has contributed to ITRC's Bioavailability of Contaminants in Soil. Geoff earned a bachelor's degree in geochemistry from Pomona College, Claremont, California in 1994 and a master's degree in soil science from the University of California, Berkeley in 1999.

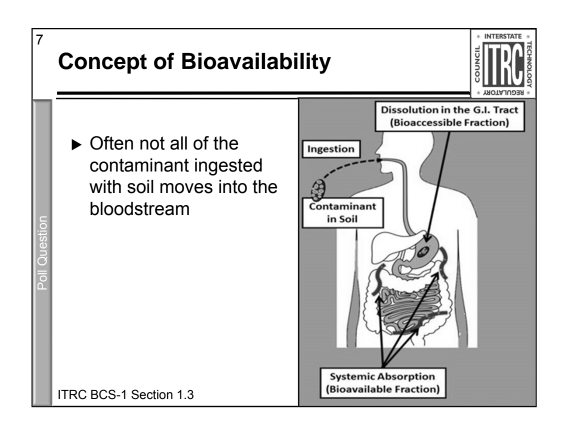
Anita Meyer is a risk assessor and toxicologist with the Army Corps of Engineers Environmental and Munitions Center of Expertise. She works for the Huntsville Center and is located in Omaha, Nebraska. Since 1997 Anita has gained experience with CERCLA and RCRA risk assessments on formerly used defense sites, military munitions response program sites, former Manhattan Project sites, Army and Air Force active sites and on EPA Superfund projects. Beginning in 2009, Anita has supported the Department of Defense (DoD) Chemical and Material Risk Management Program Directorate, leading DoD interagency reviews of EPA toxicological assessments, as well as regulatory risk assessments for TSCA. Anita represents the Army and the Corps of Engineers on interagency committees and workgroups related to environmental investigation and cleanup. She has been a member of four ITRC technical teams, Bioavailability in Contaminated Soils, Incremental Sampling Methodology, Risk Assessment, and Risk Assessment Resources. She provides risk assessment expertise on Corps of Engineers projects and has utilized bioavailability assessments on former skeet target ranges. Anita also consults on DoD and Army policy, writes Corps of Engineers guidance, and teaches Corps of Engineers courses on risk assessment and in systematic planning. Prior to joining the Corps of Engineers Anita performed cancer and drug development research. Anita earned a bachelor's degree in biological sciences in 1984 and a master's degree in cell biology and genetics in1987 from the University of Nebraska in Lincoln, Nebraska. She is certified by the American Board of Toxicology (DABT).

Kevin Long is a Principal Consultant in Terraphase's Princeton, NJ office. Since 2000, he has applied risk assessment and risk management strategies to support site characterization, risk management, and redevelopment at hazardous waste and brownfield sites under Superfund, RCRA, and various state and provincial cleanup programs. Working on such projects, he has helped to control unacceptable human exposures at dozens of sites, including those that may pose an imminent and substantial danger to human health. Such projects have involved addressing contamination in all sorts of environmental media and, in many cases, have required complex exposure assessment, fate and transport modeling, statistical analysis, risk management design, and risk communication. He has been a member of the ITRC Risk Assessment team since 2012. Kevin earned a bachelor's degree in 2000 and master's degree in 2006, both in Civil and Environmental Engineering, from Princeton University in Princeton, NJ.

Dr. Valerie Hanley is a Staff Toxicologist in the Human and Ecological Risk Office at the California Department of Toxic Substances Control (DTSC) in Sacramento, CA. Valerie has been with DTSC since 2008. She recently authored a Human Health Risk Assessment Note on how to evaluate Arsenic contaminated sites with a specific emphasis on how and when to use bioavailability in those site evaluations. Valerie has been involved in the study of arsenic bioavailability since 2009 when DTSC was awarded funding from US EPA to evaluate and develop new methods to determine arsenic bioavailability in mining soils. Through this work Valerie helped develop the California Arsenic Bioaccessibility (CAB) Method, which is now recommended for use in sites throughout California. Valerie joined the ITRC Bioavailability in Contaminated Soils Team in 2015 and is one of the lead authors on the arsenic chapter of the document. In addition to her work on arsenic, Valerie evaluates Human Health Risk Assessments for a variety of sites and is involved in DTSC's Safer Consumer Products program. Valerie earned a Bachelor's degree in Molecular, Cellular, and Developmental Biology from The University of California (UC) Santa Cruz in 2001 and her PhD in Comparative Pathology from UC Davis in 2007. She completed a postdoctoral fellowship at UC Davis in Respiratory Toxicology in 2008.

Barrie Selcoe is a Principal Technologist with Jacobs in Houston, Texas. Barrie has worked at Jacobs since 2018, specializing in human health risk assessment. She is responsible for planning and overseeing human health risk-based activities at hazardous waste sites across the U.S. and internationally. She utilizes numerous federal (USEPA and Department of Defense) and state guidance documents in risk assessment projects, and is involved in all stages of site planning, investigation and reporting, cleanup level identification, and remedial action planning. She has been involved in risk assessments in 40 states and about 20 countries. She has worked on risk assessments incorporating incremental sampling and site-specific bioaccessibility studies. She has provided risk assessment services for numerous Comprehensive Environmental Response, Compensation, and Liability Act (CERCLA)/Superfund sites, Resource Conservation and Recovery Act (RCRA) facilities, state-program sites, voluntary actions, and international projects. She has prepared risk assessments for various types of sites, including industrial and commercial facilities, industrial and municipal landfills, bulk fuel terminals, rivers, U.S. Department of Defense facilities, and residential areas. Prior to Jacobs (which purchased CH2M in 2018), she worked as a human health risk assessor for 19 years with CH2M, 7 years with Philip Environmental, and 3 years with O'Brien & Gere Engineers. Since 2012, Barrie has contributed as a team member on ITRC's Risk Assessment team, Bioavailability in Contaminated Soil team, TPH Risk Evaluation at Petroleum-Contaminated Sites team, and PFAS team. She earned a bachelor's degree in microbiology from San Diego State University in San Diego, California in 1986, and a Master's of Public Health from the University of Pittsburgh Graduate School of Public Health in Pittsburgh, Pennsylvania in 1999.





Poll: If a contaminant is ingested and passes through (is not absorbed FROM) the human gastrointestinal tract (G.I. Tract), DOES IT CONTRIBUTE TO SYSTEMIC RISK?

Yes

No

I don't know

Answer is NO because our risk assessment process for ingestion of contaminated soil focuses on risks from systemic exposure to contaminants in soil. The next sections of this training and the ITRC document address *exactly* this issue.

You Should Learn to...



- ► Value the ITRC document as a "go-to" resource for soil bioavailability
- ► Apply decision process to determine when a site-specific bioavailability assessment may be appropriate
- ▶ Use the ITRC Review Checklist to develop or review a risk assessment that includes soil bioavailability
- ► Consider factors that affect arsenic, lead and polycyclic aromatic hydrocarbons (PAH) bioavailability
- ▶ Select appropriate methods to evaluate soil bioavailability
- ▶ Be able to incorporate soil bioavailability into human health risk assessments

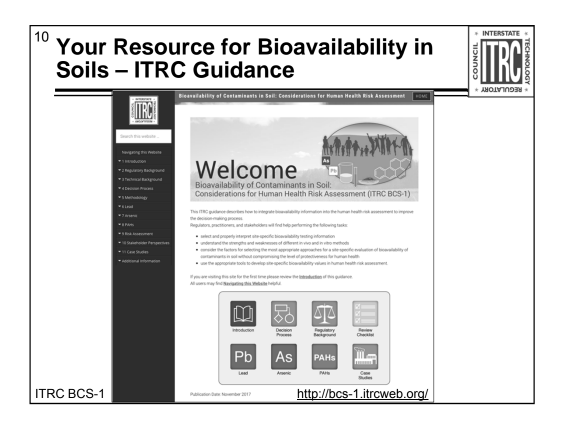
Why You Should Consider Evaluating Bioavailability in Soils



- Reduces uncertainty, provides a more accurate understanding of chemical exposures and associated risk
- ► Leads to a more effective use of resources without compromising health protection
- May reduce remedial action costs and increase flexibility of remedial options
- Risk assessment allows for modifying exposure factors to better represent site conditions



Photo courtesy of Geoff Siemering, University of Wisconsin, 2017



Focus of ITRC Training and Guidance



- ▶ Bioavailability of contaminants in soil to humans
 - Bioavailability in sediment or in reference to ecological receptors (see ITRC Guidance: http://www.itrcweb.org/contseds-bioavailability/)
- ► Specifically covers As (arsenic), Pb (lead), and polycyclic aromatic hydrocarbons (PAHs)
 - Although guidance can be used for assessing bioavailability of other contaminants
- ► Focuses on the soil ingestion pathway
- ▶ Limited dermal bioavailability information as it relates to PAHs

Bioavailability Tools



- ▶ Web-based Guidance Document ITRC BCS-1
 - The go-to guide for bioavailability assessments

(Provided in the Webinar Handouts)

- ▶ Decision Process Flow Chart Section 4.1
 - · Will be presented in both case studies
- ▶ Definition of Terms
- ▶ Review Checklist
 - Can be used as a tool to review a bioavailability assessment
 - Can be used to prepare a bioavailability study

ITRC BCS-1 http://bcs-1.itrcweb.org/

¹³ A Regulator's Experience with Bioavailability – Learning Opportunities



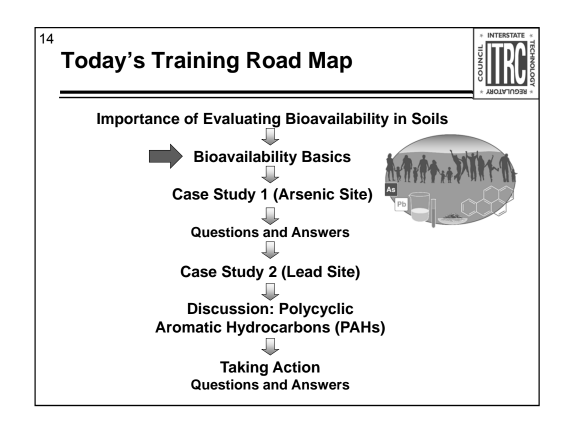
- Regulator with limited experience in bioavailability overseeing arsenic cleanup project
- ► Consultant recommends assessing bioavailability of arsenic at site
- ▶ Project manager and team toxicologist agree to using bioavailability in risk assessment
- ► Risk assessment presented much lower risk than previous estimates
- ➤ Significantly reduced remedial action costs
- Increased the accuracy of the risk estimate



Photo source: Red Rock Road ECSI #1855, OEQ, 2009

Poll Question: (was originally shown as participants log on slide 1) and brought back here to show results and discuss

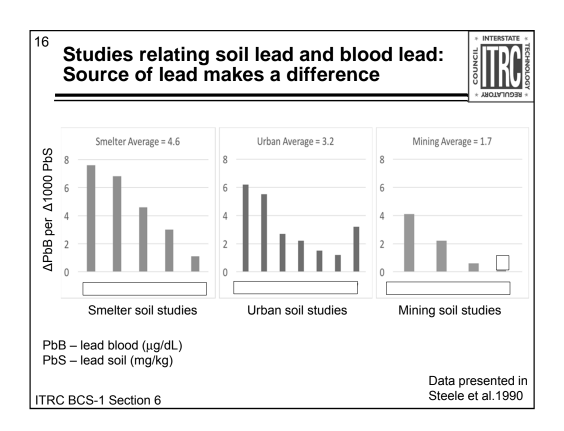
- little or no experience
- some knowledge and experience
- expert



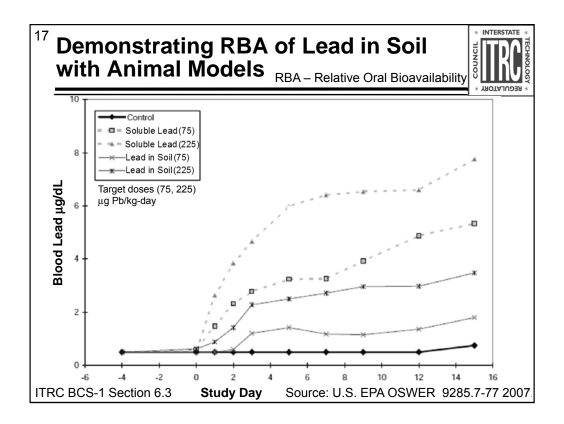
⁵ Bioavailability of Contaminants in Soil Basics



- ▶ History: how we recognized the issue
- ▶ Relevance to Human Health Risk Assessment
- ► Concepts with applicability to all chemicals
- ► Key definitions
- ▶ In vivo in vitro correlation (IVIVC)
- ▶ Soil properties that influence bioavailability

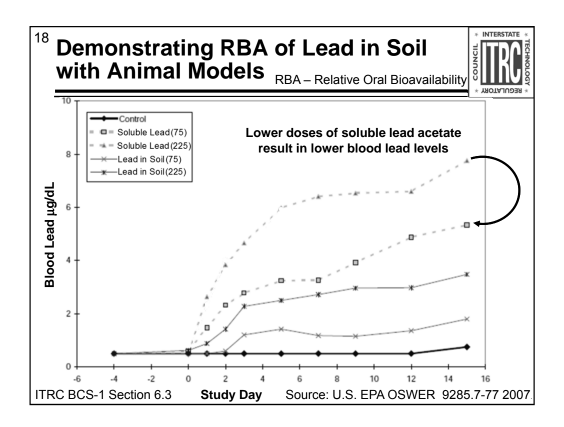


Steele, M. J., B. D. Beck, B. L. Murphy, and H. S. Strauss. 1990. "Assessing the Contribution from Lead in Mining Wastes to Blood Lead." Regulatory Toxicology and Pharmacology, 11: 158-190.

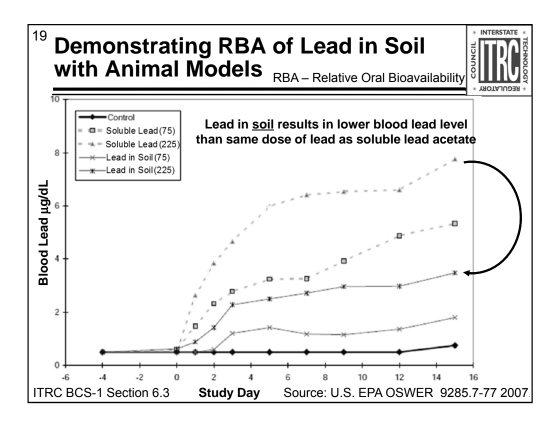


Range: 1% (Galena-enriched soil) to 105% Lowest site soil was 6%, for Tailings sample from California Gulch Highest value California Gulch Fe/Mn PbO

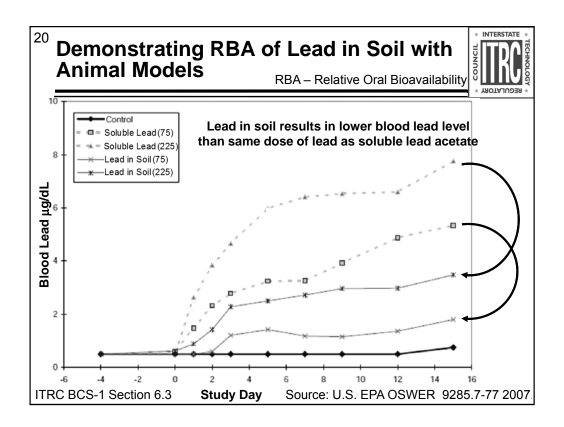
Target lead dose (75, 225) – expressed in units of <u>micrograms</u> of lead per kg of body weight per day.



Range: 1% (Galena-enriched soil) to 105% Lowest site soil was 6%, for Tailings sample from California Gulch Highest value California Gulch Fe/Mn PbO



Range: 1% (Galena-enriched soil) to 105% Lowest site soil was 6%, for Tailings sample from California Gulch Highest value California Gulch Fe/Mn PbO



Graph of concentration of lead in blood over time.

Regulatory Recognition of Using Bioavailability for Risk Assessment

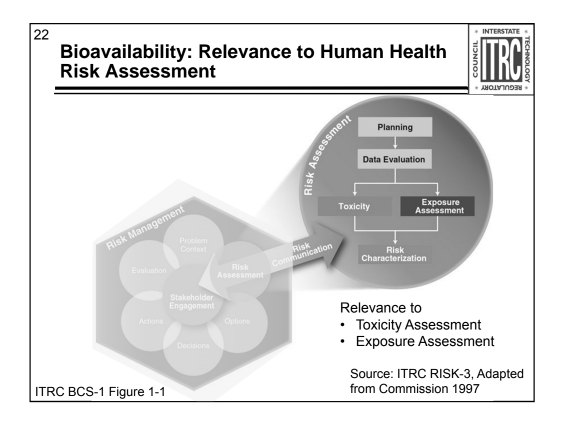


"If the medium of exposure [at] the site... differs from the medium of exposure assumed by the toxicity value... an absorption adjustment may... be appropriate."

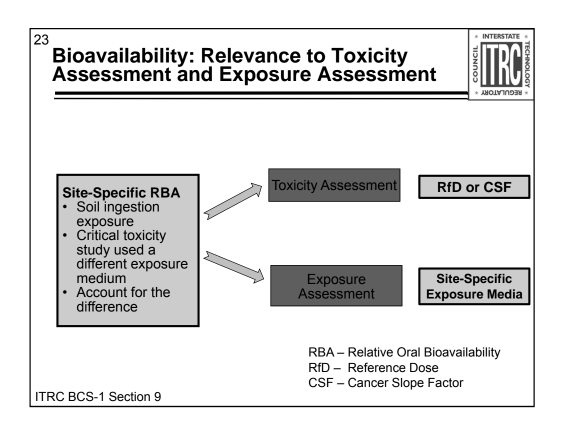
"[to] adjust a food or soil ingestion exposure estimate to match a RfD or slope factor based on... drinking water..."

USEPA 1989 "Risk Assessment Guidance for Superfund (RAGS)" EPA/540/1-89/002

USEPA. 1989. "Risk Assessment Guidance for Superfund. Volume I: Human Health Evaluation Manual (Part A). Interim Final." *EPA/540/1-89/002*. Washington, D.C.: Office of Emergency and Remedial Response. U.S. Environmental Protection Agency.



Commission, Presidential/Congressional. 1997. "Framework for Environmental Health Risk Management." Final Report, Volume 1. Washington, D. C.: The Presidential/Congressional Commission on Risk Assessment and Risk Management.



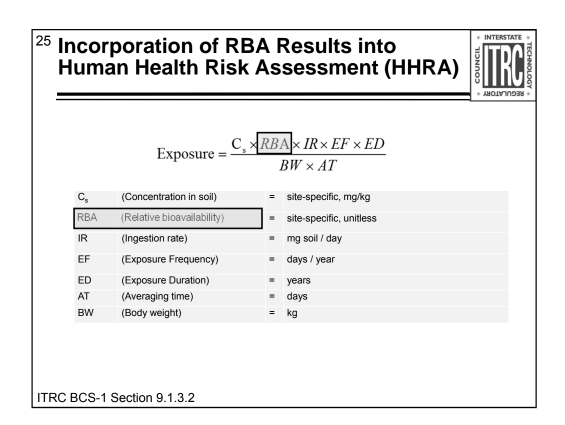
Definition:Relative Oral Bioavailability (RBA)



► Comparison of bioavailability of a chemical in different dosing media

► RBA = Absolute Bioavailability from Soil
Absolute Bioavailability from form dosed in critical toxicity study

ITRC BCS-1 Section 1.3



²⁶ Bioavailability Evaluation Can Apply to All Chemicals



► Including priority listed chemicals
The ATSDR 2017 Substance Priority List

2017 Rank	Substance Name
1	ARSENIC
2	LEAD
3	MERCURY
4	VINYL CHLORIDE
5	POLYCHLORINATED BIPHENYLS
6	BENZENE
7	CADMIUM
8	BENZO(A)PYRENE
9	POLYCYCLIC AROMATIC HYDROCARBONS
10	BENZO(B)FLUORANTHENE

https://www.atsdr.cdc. gov/SPL/index.html#c ontent-main

▶ Although current default assumes RBA of 100% for all chemicals in soil except arsenic and lead (default 60%)

ATSDR 2017. "Substance Priority List." Agency for Toxic Substances and Disease Registry. https://www.atsdr.cdc.gov/SPL/index.html#content-main

Definition:
Bioaccessibility

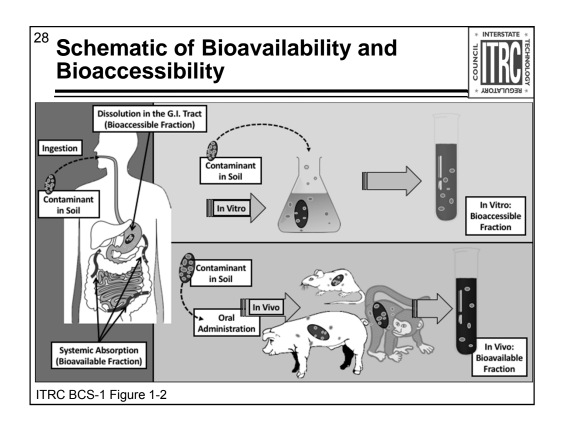


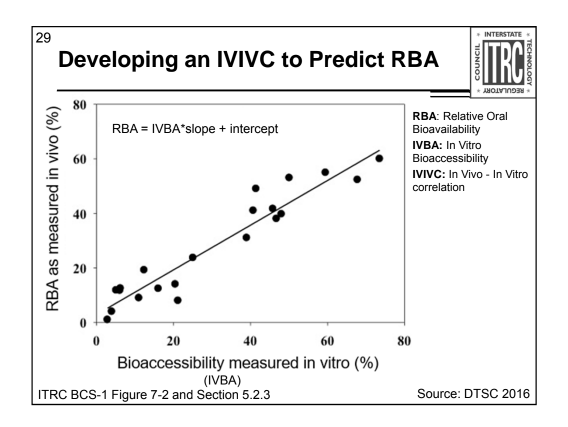
X 100

Bioaccessible Fraction (%) = $\frac{\text{Mass of chemical soluble from soil}}{\text{Total mass of chemical present in soil}}$

- ► Fraction of total amount of chemical present that is soluble / available for uptake
- ► In vitro methods attempt to characterize this parameter
 - In vitro bioaccessibility (IVBA)

ITRC BCS-1 Section 5.2

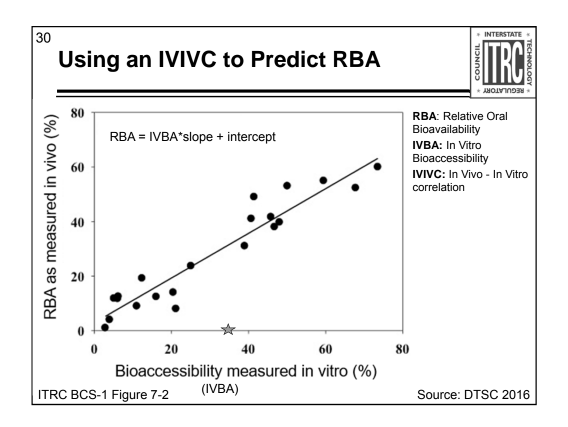




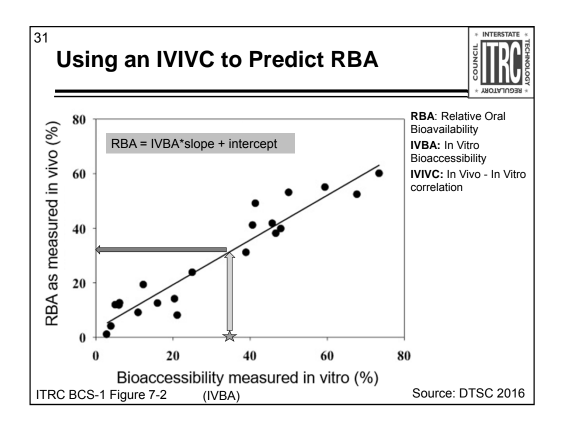
"The goal of IVIVC is to promote an in vitro IVBA test method to replace in vivo RBA feeding studies. Successful IVIVC has been established when the RBA of a test soil can be determined using a predictive model (for example, simple linear regression), and meet the USEPA requirement (2007b) that "the in vitro result (entered as input) will yield an estimate of the in vivo value (as output)." If a good IVIVC has been established, then the in vitro data for soils can be used as the sole basis for adjusting RBA in a human health risk assessment" (BCS-1 document, Section 5.2.3).

The IVIVCs for lead and arsenic that currently have approval from regulatory agencies were developed either by aggregating information about several (or many) soils that were investigated over several different studies, or were part of a large-scale study that included many soils.

Generally, IVIVC development requires significant research. So IVIVCs for use in risk assessment are generally either developed and published in the peer-reviewed literature, or developed with the involvement of regulatory agencies – and frequently both!



See notes on Slide 29

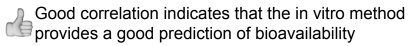


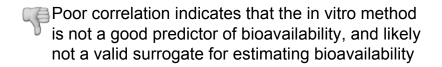
See notes on Slide 29.

Definition:In Vivo - In Vitro correlation (IVIVC)

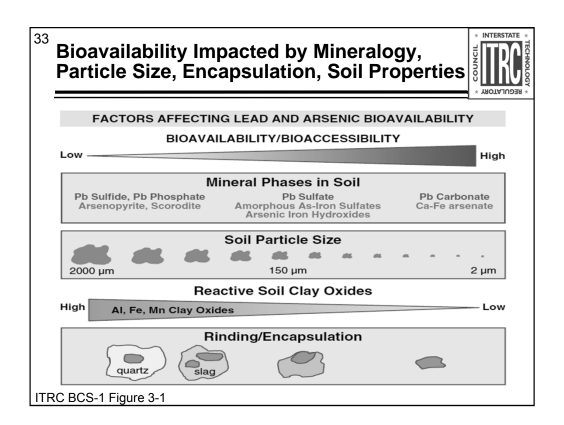


▶ Refers to a correlation between in vitro bioaccessibility results and in vivo bioavailability results





ITRC BCS-1 Section 5.2.3



Regulatory Recognition of Using Bioavailability for Risk Assessment

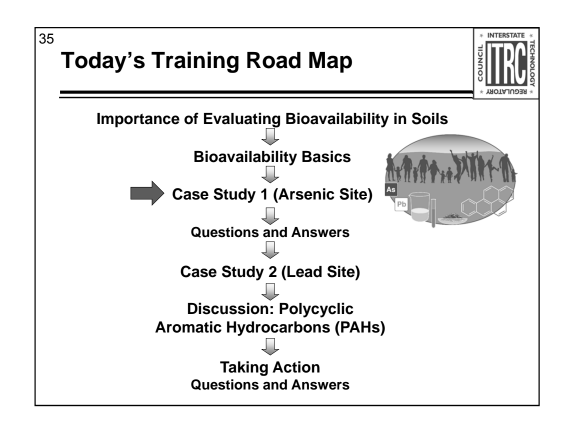


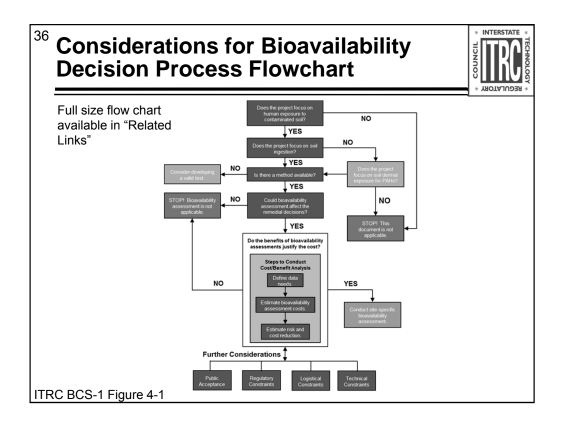
- ► Lead: specific guidance on using bioavailability in the risk assessment of lead-contaminated sites (USEPA 2007)
- ▶ Arsenic: Significant efforts to summarize and evaluate the bioavailability of arsenic from soil (USEPA 2012, USEPA 2017a,b,c)
- Completed a review of the available information on dioxins (USEPA 2015)
- ► Guidance to evaluate arsenic from California and Hawaii (DTSC 2016, Hawaii DOH, 2010, 2012)
- ► Several site-specific precedents
 - Pb, As, Cd, dioxins, PAHs.

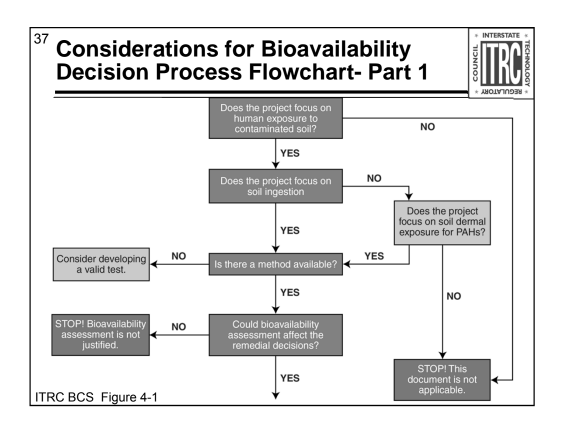
ITRC BCS-1 Section 2

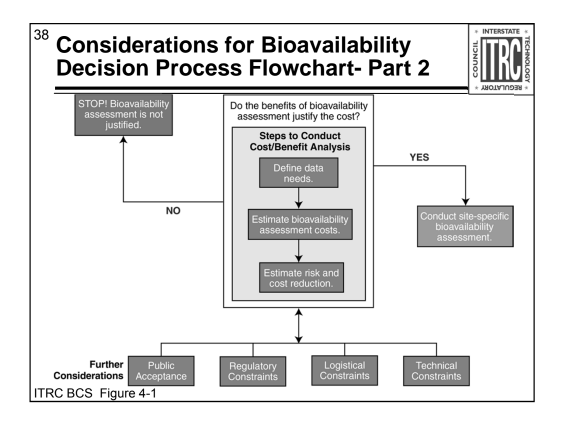
Poll Question: Does the state you work in use bioavailability when assessing risk?

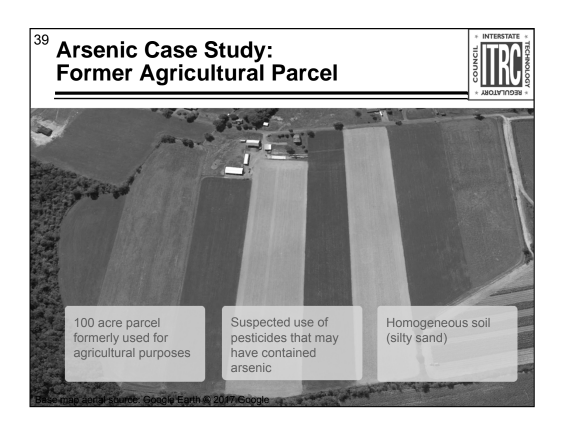
- Yes
- No
- -Don't Know

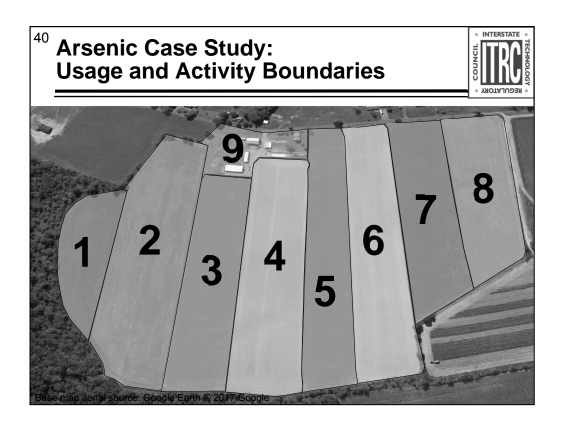


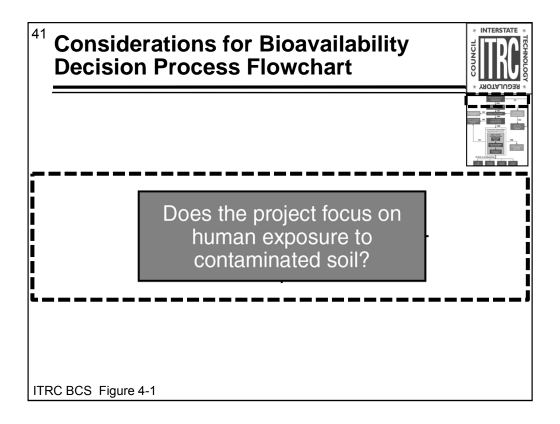


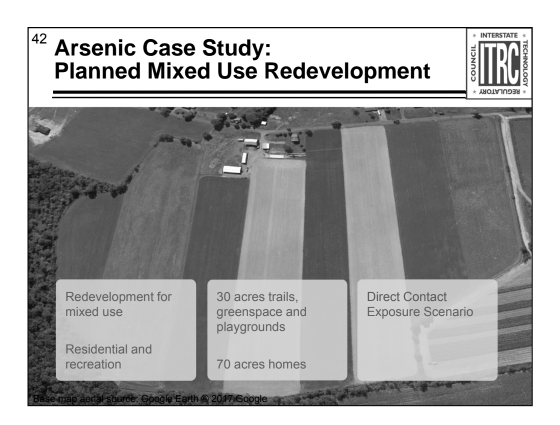


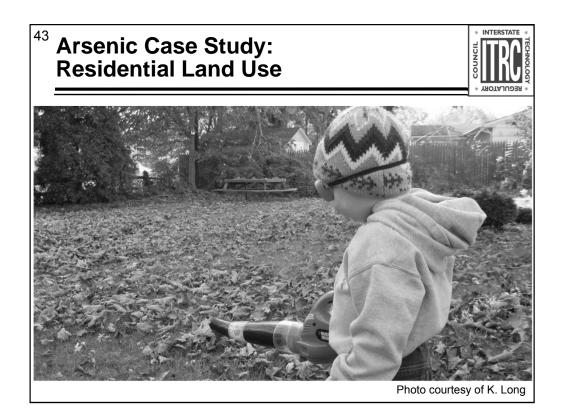


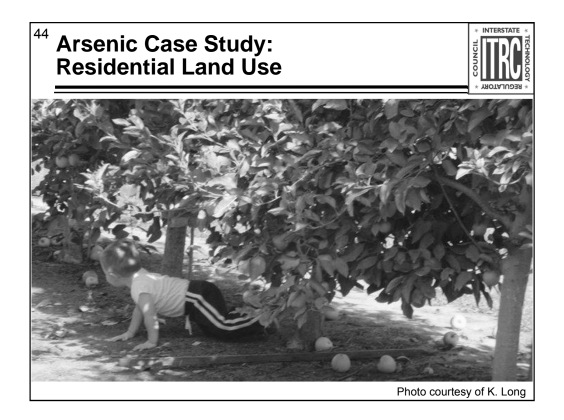










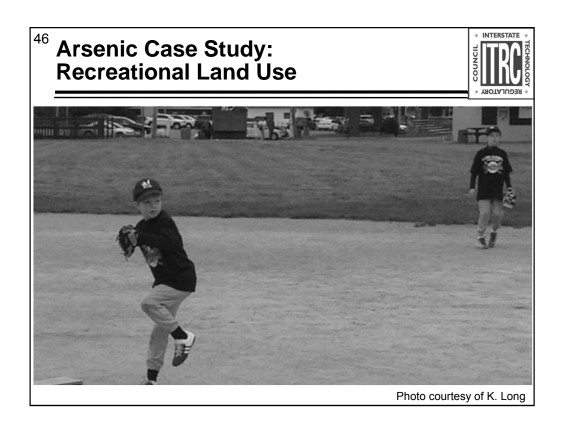


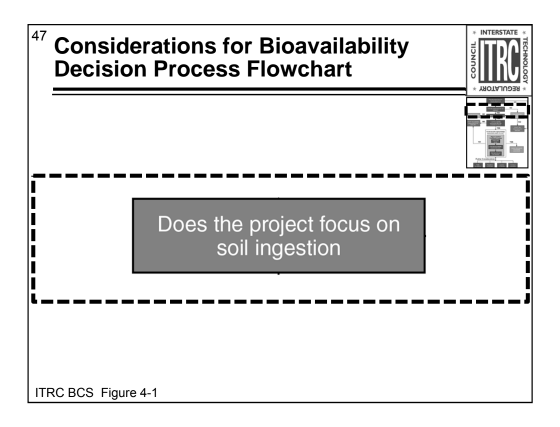
⁴⁵ Arsenic Case Study: Residential Land Use

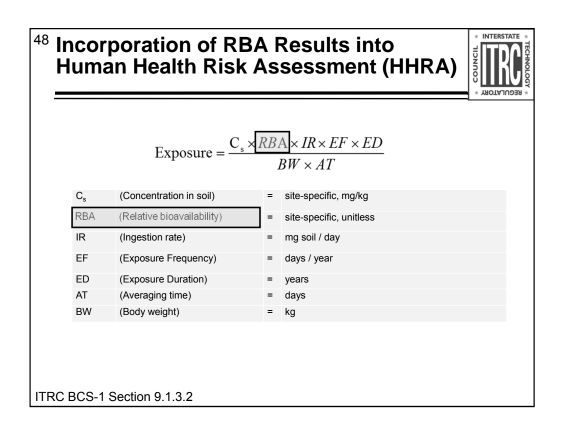


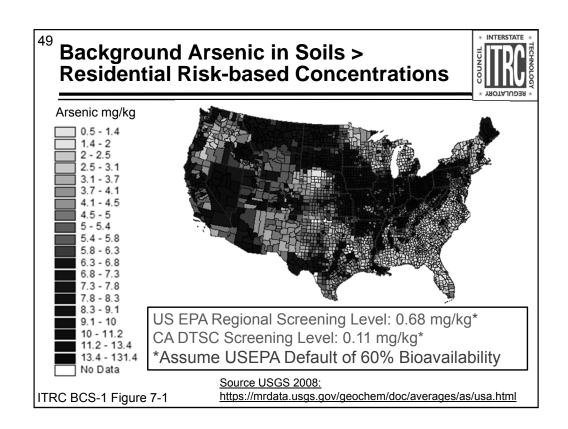


Photo courtesy of V. Hanley

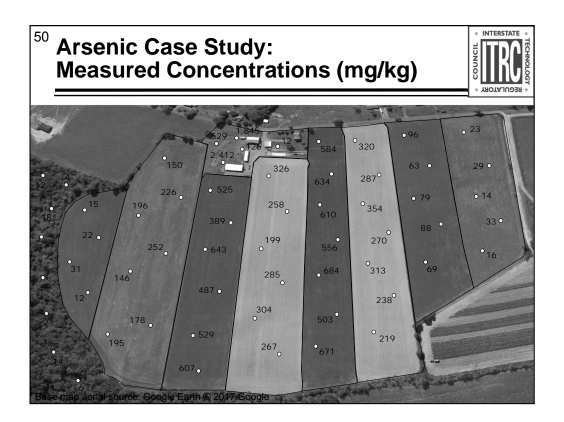


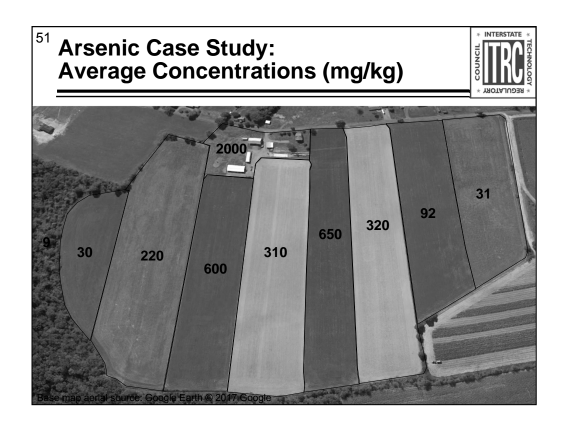


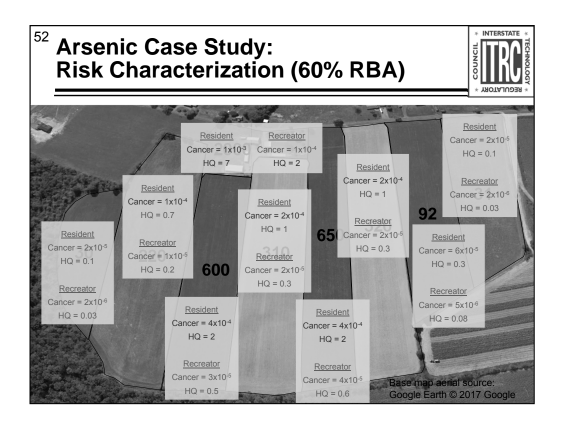


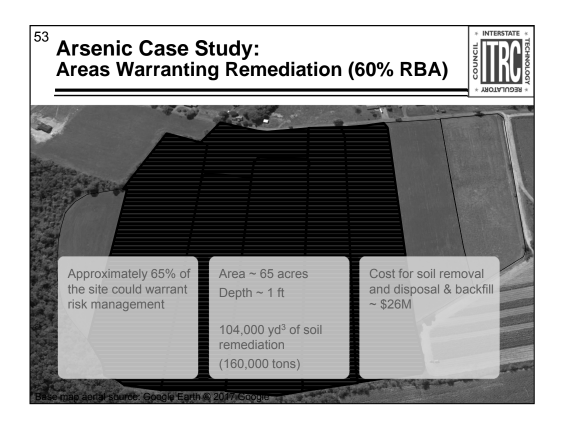


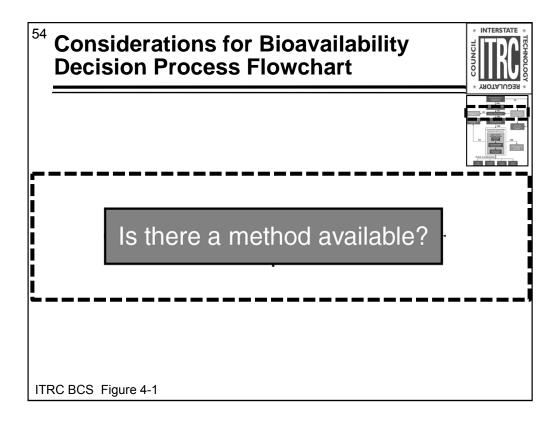
USGS (United States Geological Survey). 2008. "National Geochemical Survey, Geochemistry by County." https://mrdata.usgs.gov/geochem/doc/averages/countydata.htm.











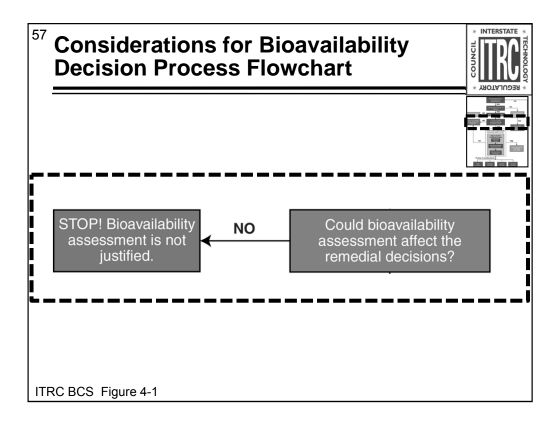
⁵⁵ Available Methods for Determining Arsenic Bioavailability In Vivo

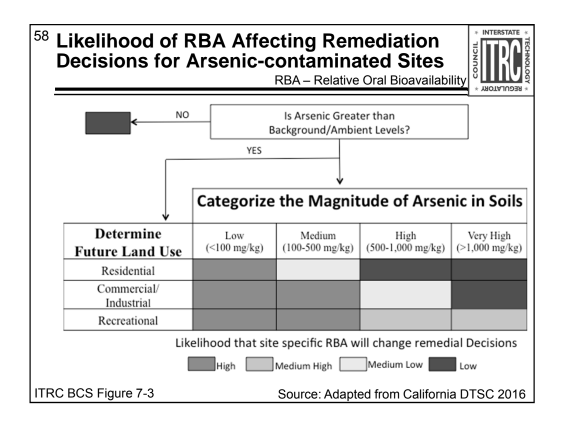


Animal model	Biomarkers of arsenic exposure	Reference
Juvenile Swine	Steady state urinary excretion	Rodriguez et al. 1999; Casteel et al. 2006; Weis and LaVelle, 1991; Basta et al. 2007; Denys et al. 2012; Brattin and Casteel 2013
	Single dose blood AUC	USEPA 1996; Juhasz et al. 2007, 2008
Mice (C57BL/6)	Steady state urinary excretion	Bradham et al. 2011
Monkeys (Cebus, Cynomolgus)	Single dose urinary excretion	Freeman et al. 1995; Roberts et al. 2002, 2007; USEPA 2009

ITRC BCS-1 Table 7-1

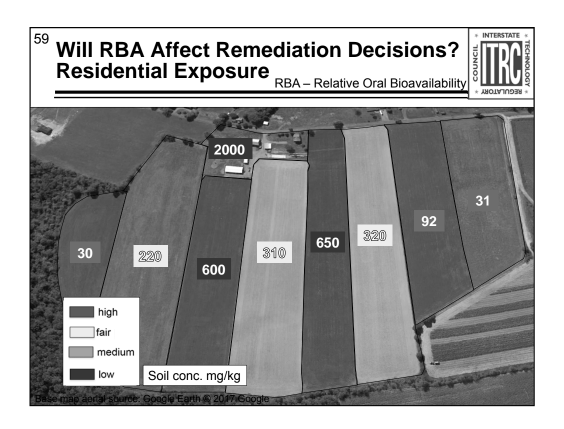
Available Methods for Determining Arsenic Bioavailability In Vitro				
Method USEPA Method 1340	Key Reference Diamond et al. 2016	Notes Method adopted by USEPA. Guidance issued May 2017		
Also known as RBALP, SBRC, and USEPA 9200	2010	https://semspub.epa.gov/work/HQ/196750.pdf		
California Arsenic Bioaccessibility Method (CAB)	Whitacre et al. 2017	Method adopted by California DTSC Guidance issued Aug. 2016 http://www.dtsc.ca.gov/AssessingRisk/upload/H HRA-Note-6-CAB-Method-082216.pdf		
Unified BARGE Method (UBM)	Wragg et al.2011 Denys et al. 2012	ISO certification (17924) – widely used throughout Europe. https://www.bgs.ac.uk/barge/home.html		
In Vitro	Basta et al. 2007	No regulatory guidance exists to support this		
Gastrointestinal Method (IVG)	Rodriguez et al., 1999	method. First published method to report strong IVIVC, but did not include interlaboratory round robin study necessary for regulatory guidance and approval by USEPA.		
Physiological Based Extraction Test (PBET)	Ruby et al. 1996	No regulatory guidance exists to support this method.		

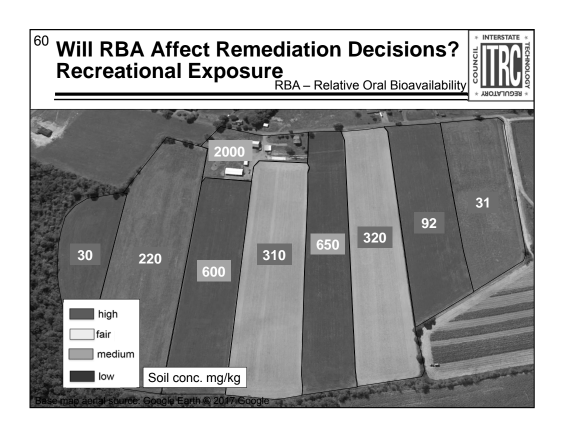


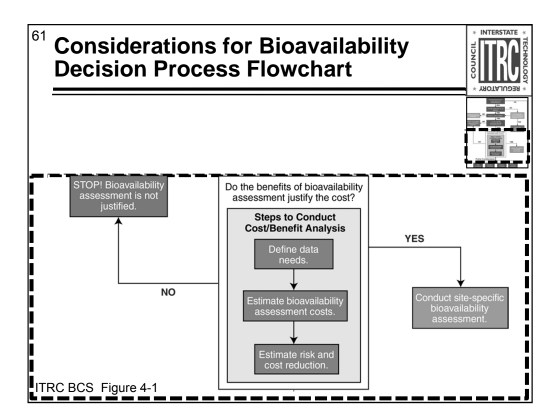


DTSC. 2016. "Human Health Risk Assessment Note 6: Recommended Methodology for Evaluating Site-Specific Arsenic Bioavailability in California Soils." Sacramento, CA: California Environmental Protection Agency.

https://www.dtsc.ca.gov/AssessingRisk/upload/HHRA-Note-6-CAB-Method.pdf







Poll Question



- ► How much do you think the in vitro bioavailability study would cost for this site?
 - \$1,000
 - \$20,000
 - \$100,000

Poll Question: How much do you think the in vitro bioavailability study would cost for this site?

\$1,000

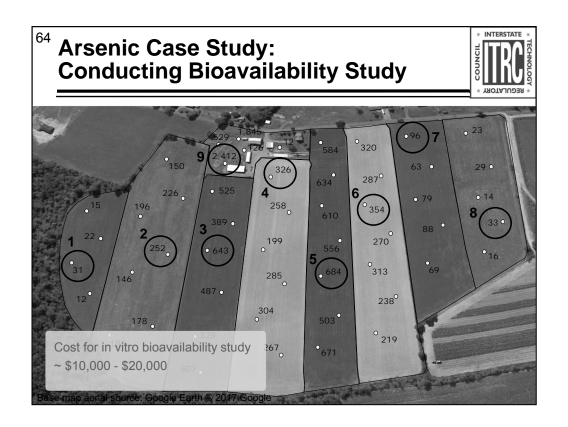
\$20,000

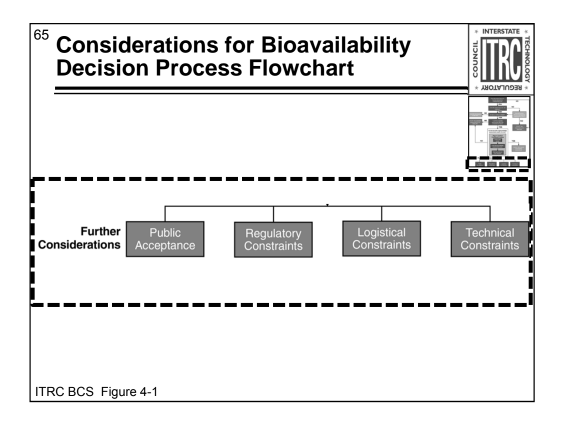
\$100,000

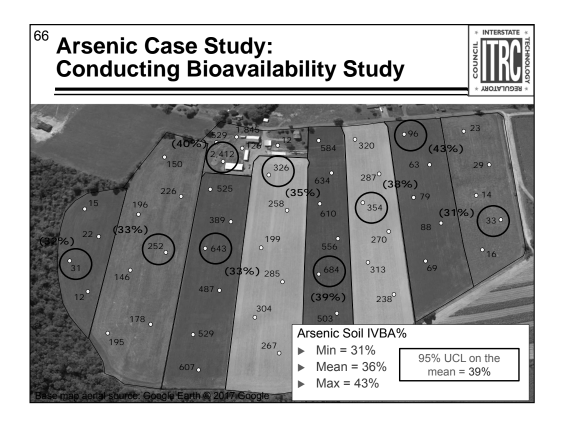
Approximate Costs for Bioavailability Analysis

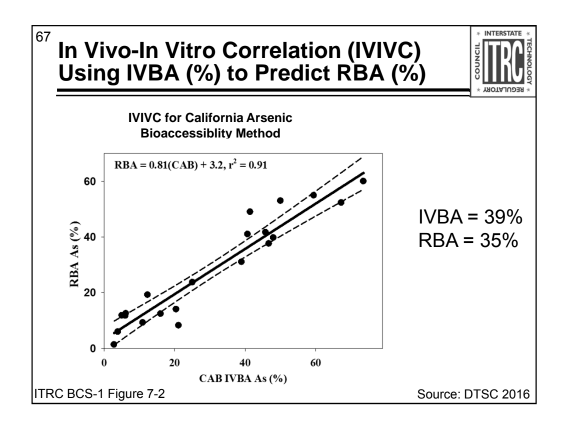
* INTERSTATE *				
* COUNCIL	MOTA LUDER *			

Analysis	Approximate Unit Cost Per Sample (USD)	Provider
Soil properties	\$500-\$1,000 (per sample)	Commercial labs
Soil mineralogy	\$200-\$1,000 (per sample)	Academic and commercia labs
IVBA for Pb or As	\$150-\$1,000 (per sample)	Academic and commercia labs
IVBA for PAHs	\$350 - \$1000 (per sample)	Academic and commercia labs
In vivo (mouse, rat)	\$25,000-\$30,000 (per study)	Academic or government labs
In vivo (swine)	\$75,000 (for 3 soils, metals only)	Academic labs
In vivo (primate)	\$90,000 (for three soils, metals only)	Academic labs
C BCS-1 Table 4-1	Co	st data collected in 2015-1





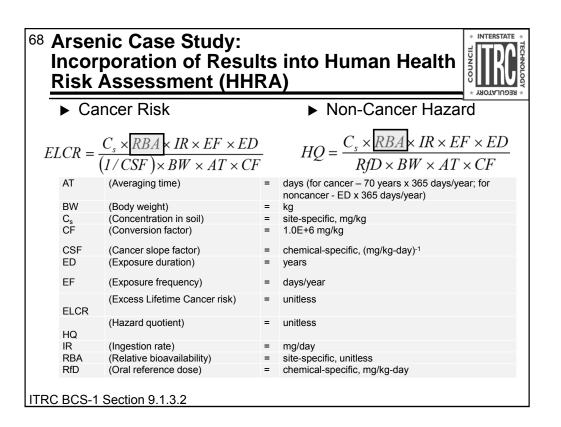


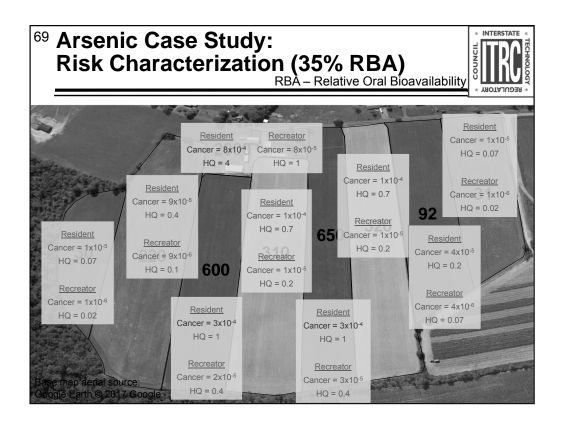


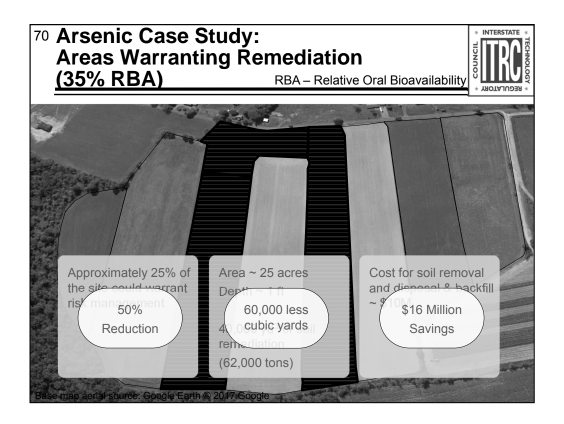
In vitro analysis gives IVBA %, which can be used to determine RBA using a validated IVIVC. This is an example from the CAB method

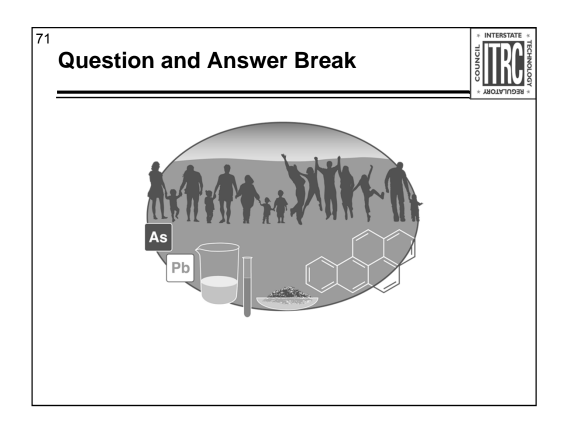
DTSC. 2016. "Human Health Risk Assessment Note 6: Recommended Methodology for Evaluating Site-Specific Arsenic Bioavailability in California Soils." Sacramento, CA: California Environmental Protection Agency.

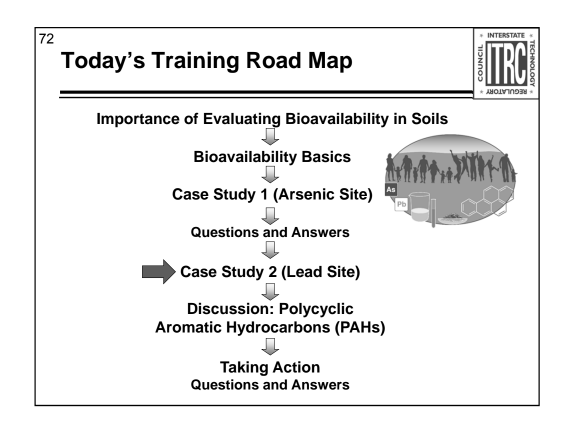
https://www.dtsc.ca.gov/AssessingRisk/upload/HHRA-Note-6-CAB-Method.pdf











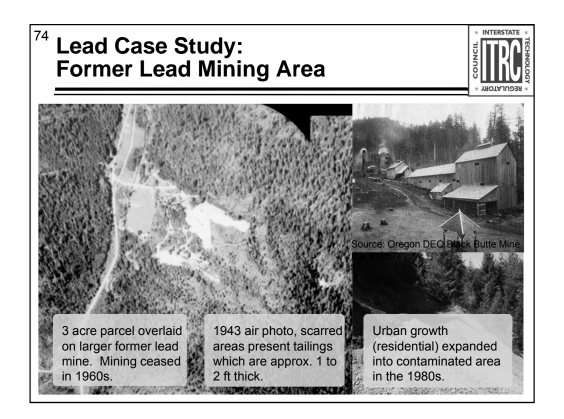
Lead Case Study

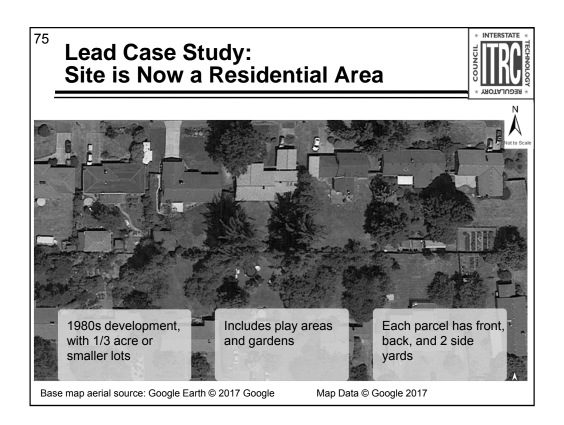


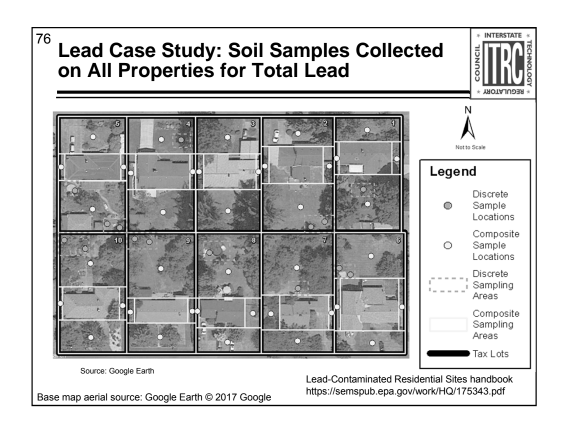
- ► Case study is presented as a series of meetings between regulator and consultant
- ► Historic lead mining area
- ► Contaminant source lead tailings
- ▶ Residential area
- Future land uses are residential and commercial



Source: Pixnio.com





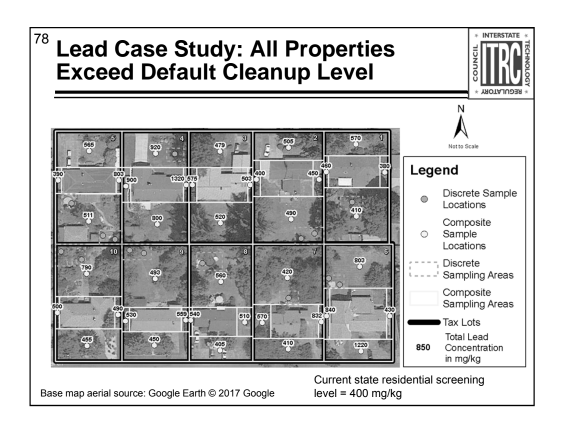


USEPA. 2003. "Superfund lead-contaminated residential sites handbook." *OSWER 9285.7-50*. Washington, D.C.: Office of Emergency and Remedial Response.

⁷⁷ Lead Case Study: Total Lead Sampling Complete

- ▶ Available samples for nature & extent
 - 10 properties; 4 yards each (1 composite sample/yard) = 40 samples
 - 5 properties with gardens (2 discrete samples/garden) = 10 samples
 - 5 properties with play areas (1 discrete sample/play area) = 5 samples
- ► Total lead concentrations
 - 380 to 1,321 mg/kg, arithmetic mean = 850 mg/kg, low standard deviation
- ► Background 30 mg/kg
- ► Soil type Well graded gravel with fines and thin organic silt at surface

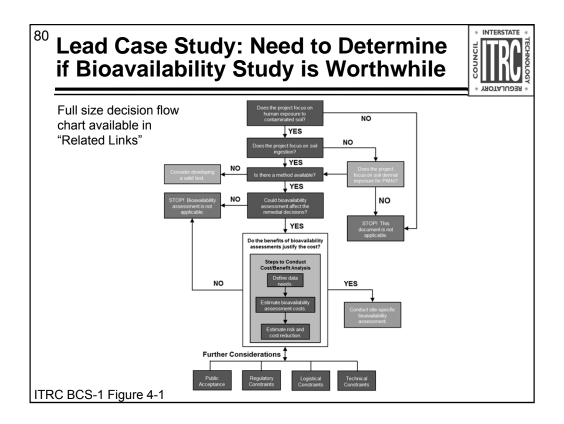


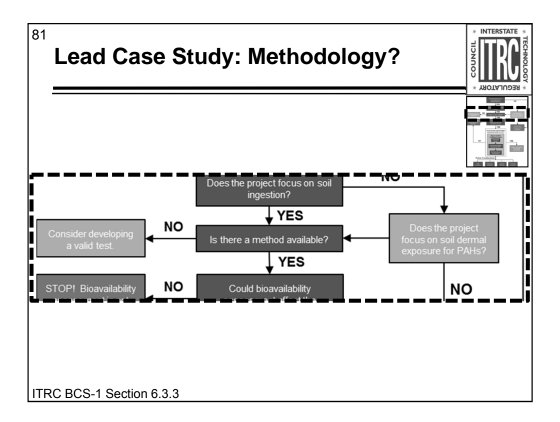


Lead Case Study: Estimated Costs Could Justify a Site-Specific Bioavailability Study



- ► Excavation volume based on nature & extent sampling
 - 3 acres
 - 1 to 2 ft depth
 - ~5,000 cy
- ► Estimated excavation cost = \$700,000
- ➤ ~250 truck trips @ 20 yards each during remediation
- ▶ Disposal is large portion of \$
- ▶ ~2 weeks for excavation and yard restoration





Lead Case Study: USEPA Recent Guidance on Lead IVBA Testing



- ► USEPA "Standard Operating Procedure for an In Vitro Bioaccessibility Assay for Lead and Arsenic in Soil" (2017) Method 1340
- ► <u>Soil Bioavailability at Superfund Sites Web Page</u> https://www.epa.gov/superfund/soil-bioavailability-superfund-sites-guidance

Apparatus used in USEPA Method



ITRC BCS-1 Section 6.3.3

Photo courtesy of Geoff Siemering, University of Wisconsin, 2017

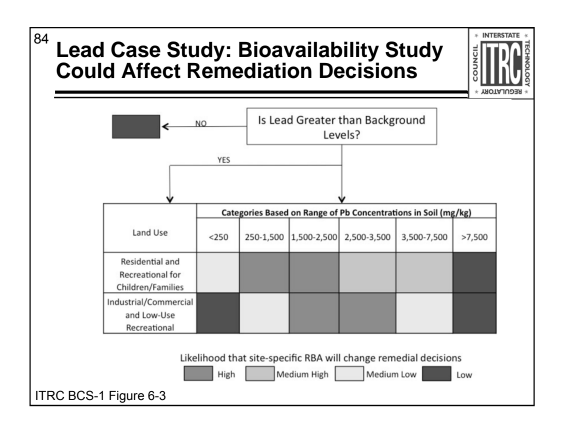
USEPA. 2017c. "Method 1340 In Vitro Bioaccessibility Assay for Lead in Soil." SW-846 Update VI. Washington, D. C.

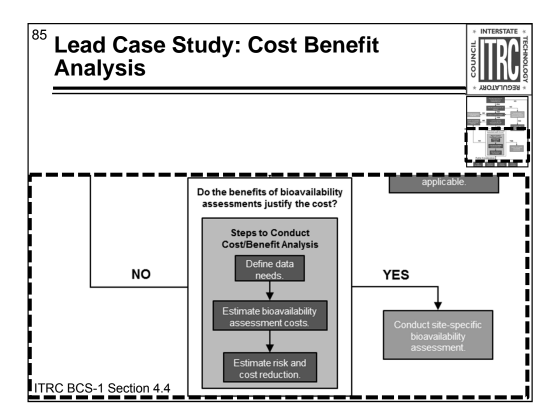
https://www.epa.gov/superfund/soil-bioavailability-superfund-sites-guidance

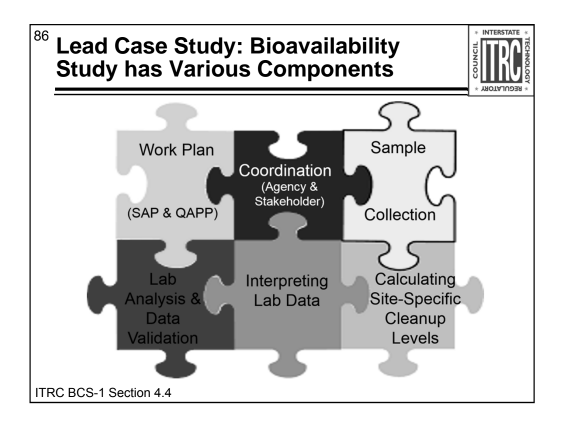
Lead Case Study: Should Studies be In Vitro or In Vivo?



- ▶ Reasons we don't need in vivo
 - Lead has been well studied with a variety of soils with good in vivo - in vitro correlation
 - · Site soil is well-characterized
 - Site soil type & waste type are similar to those tested by USEPA
 - Site soil type has an established in vivo in vitro correlation







Poll Question



► How many samples should be collected for bioavailability testing (not including duplicate samples) at this 3-acre site? (Note: nature & extent sampling is complete)

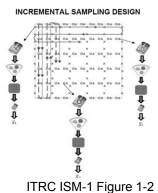
- 1 incremental sample across 3 acres
- 2 incremental samples across 3 acres
- 10 incremental samples across 3 acres
- 1 discrete sample per property
- · 2 discrete samples per property



Lead Case Study: Guidance on Lead Sampling for IVBA Testing



- ▶ USEPA "Guidance for Sample Collection for In Vitro Bioaccessibility Assay for Lead (Pb) in Soil" (2015)
 - "2 composites made up of 30 increments"
 - "In general, for most risk assessment applications, acceptable Type I error rate can be expected if ITRC (2012) recommendations are followed (30 increments per composite"
- ► Equal representation (volume, depth) from all increments
- ► Collected in triplicate
- ► ITRC ISM guidance at www.itrcweb.org/ism-1



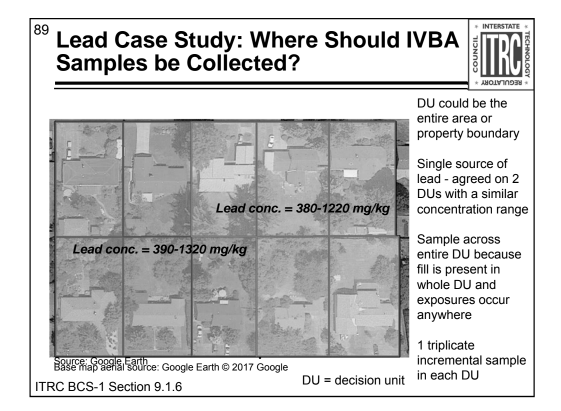
ITRC BCS-1 Section 9.1.6

USEPA. 2015a. "Guidance for Sample Collection for Bioaccessibility Assay for Lead (Pb) in Soil." OSWER 9200.3-100.

https://www.epa.gov/superfund/soil-bioavailability-superfund-sites-guidance#lead

ITRC. 2012. "Incremental Sampling Methodology." *ISM-1*. Washington, D.C.: Interstate Technology & Regulatory Council, Incremental Sampling Methodology

https://www.epa.gov/superfund/soil-bioavailability-superfund-sites-guidance Team. www.itrcweb.org/ism-1.



Lead Case Study: Use USEPA Guidance on Soil Sieving



- ► USEPA "Recommendations for Sieving Soil and Dust Samples at Lead Sites for Assessment of Incidental Ingestion" (2016)
- ► Sieve soil to <150 µm
- Reasonable upper-bound estimate of the soil/dust fraction that is most likely to stick to hands/ objects and be ingested
- ► Potential for lead enrichment in <150 µm particles at some sites
- Size fraction recommended for IVBA studies



Photo courtesy of Geoff Siemering, University of Wisconsin, 2017

USEPA. 2016e. "Recommendations for sieving soil and dust samples at lead sites for assessment of incidental ingestion. ." *OLEM Directive 9200.1-128*. Washington, D.C.: USEPA.

https://www.epa.gov/superfund/lead-superfund-sites-guidance#sampling

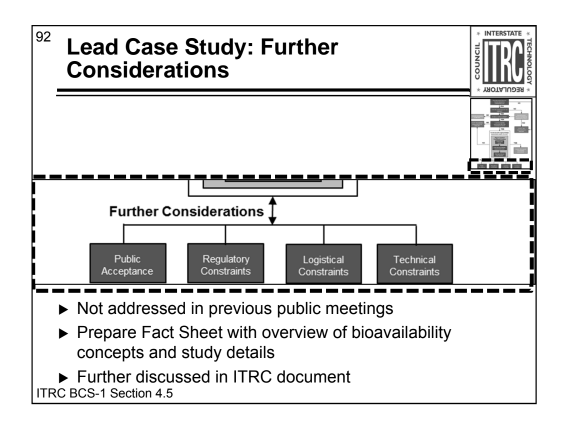
Lead Case Study: Potential Cost Impacts on the Project



- Without bioavailability study (based on existing nature & extent sampling only)
 - excavation volume = 5000 cy (1-2 ft. depth, 3 acres)
 - ~\$700,000
- ► After bioavailability study (potentially)
 - Possible RBA = 20 to 30%
 - Excavation volume = 0 cy
 - ~\$30,000 (cost of study)
 - Work planning
 - Sampling & analysis
 - Reporting
 - Remedy will be protective



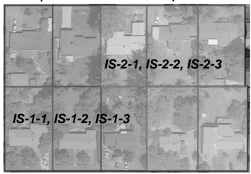
Photo Source: Oregon DEQ Black Butte Mine File #1657



Lead Case Study: Planning Meeting Resolved Path Forward



- ▶ Use USEPA Method 1340
- ▶ Divide site into 2 decision units
- ► Collect an incremental sample in triplicate from each decision unit
- ▶ Calculate site-specific soil cleanup levels using results



Base map aerial source: Google Earth © 2017 Google

Lead Case Study: Follow-up Meeting Held to Discuss Study Results



- Work Plan was submitted and approved
- ► Bioavailability study samples were collected
- ► Laboratory provided results for the samples
- Meeting between agency and consultant

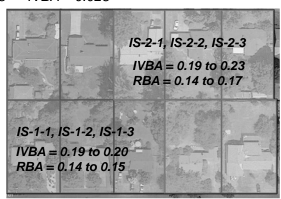


Source: Pixnio.com

Lead Case Study: RBA Predicted from IVBA

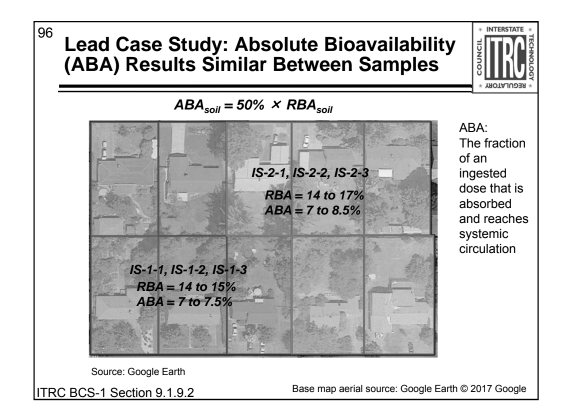


- ▶ Laboratory measured in vitro bioaccessibility (IVBA)
- ▶ Used data to predict relative bioavailability (RBA)
- ► Linear regression model established by USEPA (2007): RBA = 0.88 × IVBA - 0.028



ITRC BCS-1 Section 9.1.9.2

Base map aerial source: Google Earth © 2017 Google



Poll Question



► What RBA % would you use in a site-specific risk-based cleanup level calculation?

- Maximum of 6 values (17%)
- Average of 6 values (15%)
- Higher 95% UCL on the mean of the 2 triplicate samples (16.5%)

98 Lead Case Study: Site-specific Bioavailability Data Incorporated into Lead Models



- ▶ Pharmacokinetic models are used to evaluate lead exposures
- ▶ Residential land use Integrated Exposure Uptake Biokinetic (IEUBK) Model
- ► Commercial land use Adult Lead Methodology
- ▶ Default RBA in models is 60%
- ► Guidance document discusses methodology to incorporate site-specific RBA
- ► Site-specific RBA data reduces uncertainty

ITRC BCS-1 Section 9.1.9.2

USEPA Recently Published Guidance on Target Blood Lead Levels



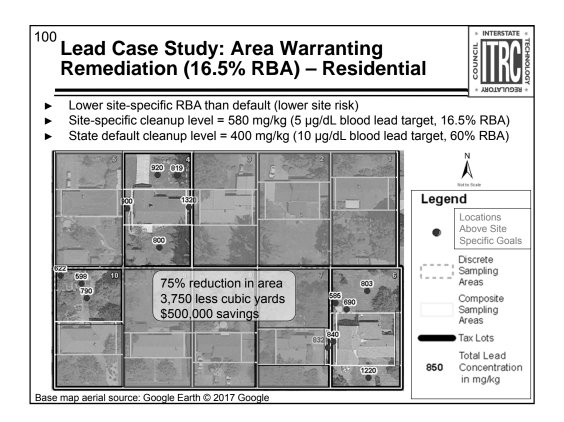
▶ USEPA "Update of the Adult Lead Methodology's Default Baseline Blood Lead Concentration and Geometric Standard Deviation Parameters and the Integrated Exposure Uptake Biokinetic Model's Default Maternal Blood Lead Concentration at Birth Variable" (2017)

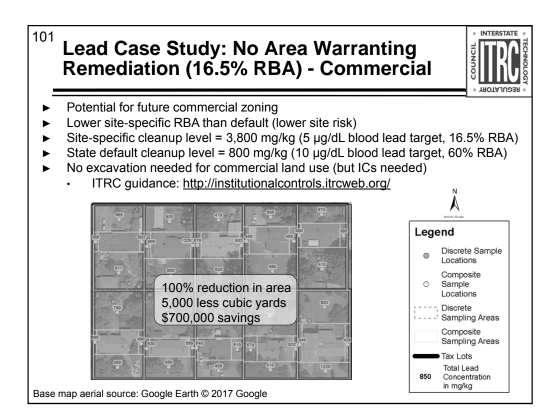
"OLEM recognizes adverse health effects as blood lead concentrations below 10 ug/dL. Accordingly, OLEM is updating the soil lead strategy to incorporate this new information."

(OLEM = USEPA Office of Land and Emergency Management)

► ITRC RISK-3 (2015) – Section 5.1.5 addresses lead toxicity and blood lead levels

https://www.epa.gov/superfund/lead-superfund-sites-guidance#adultlead





Lead Case Study: Site-Specific Bioavailability Results Useful for Decisions

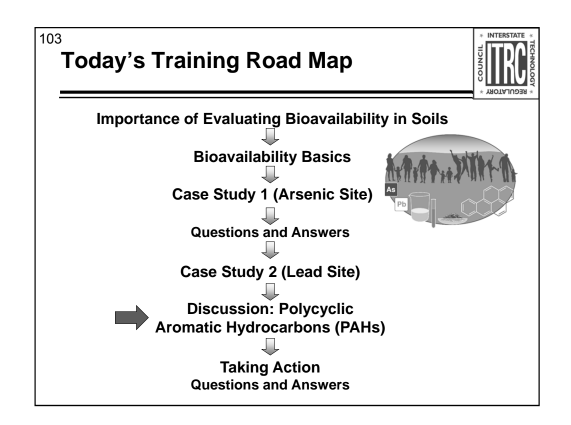


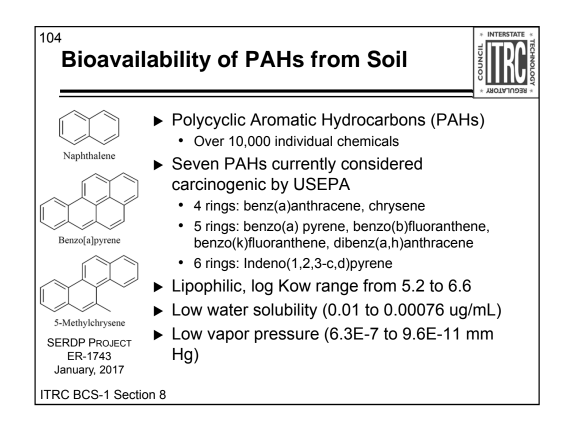
▶ Reduces:

- · Uncertainty in site risk and risk-based cleanup
- Disruption of residents
- Remediation-related risks (e.g., truck traffic, tree damage)
- · Remedial action costs

▶ Provides:

- Additional site-specific data to supplement nature and extent sampling
- · Decisions protective of human health
- Achievement of same target risk level
- Flexibility of remedial options
- Stakeholder outreach is important throughout





Images from Final Report SERDP Project ER-1743 "PAH Interactions with Soil and Effects on Bioaccessibility and Bioavailability to Humans."

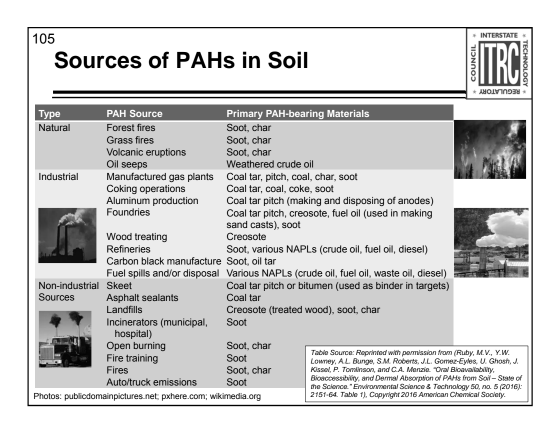


Table Source: Reprinted with permission from (Ruby, M.V., Y.W. Lowney, A.L. Bunge, S.M. Roberts, J.L. Gomez-Eyles, U. Ghosh, J. Kissel, P. Tomlinson, and C.A. Menzie. "Oral Bioavailability, Bioaccessibility, and Dermal Absorption of PAHs from Soil – State of the Science." Environmental Science & Technology 50, no. 5 (2016): 2151-64. Table 1), Copyright 2016 American Chemical Society.

State of the Science: Bioavailability of PAHs from Soil

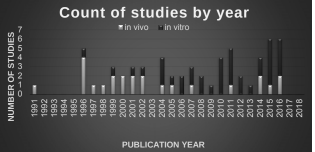


- ► Among the most common chemicals of concern at contaminated sites
- ► Current regulatory default is to assume that the RBA of PAHs in soil is 100%
 - Assumes absorption of PAHs from soil equivalent to absorption from PAH-spiked food

State of the Science: Bioavailability of PAHs from Soil



- ► Considerable interest in incorporating bioavailability estimates in HHRA
- ► Over 60 studies performed (including in vivo and in vitro studies)
- ► Studies have supported site-specific RBA values for use in HHRA
- Elucidating factors controlling binding of PAHs to soil
- Still no consensus on in vitro nor even in vivo methods



Source: Alloy 2017

Alloy 2017. http://www.cleanupconference.com/wp-content/uploads/2017/09/CleanUp_2017_Proceedings_Low-Res.pdf

Evaluating RBA of Organics from Soil



Studying RBA of organic chemicals is harder than metals!

- Methods for estimating bioavailability
 - Lagged behind metals such as lead and arsenic
 - Assessment is complex
- ► Chemical Mixture
- ▶ Analytical costs

- ▶ Metabolism
 - Hepatic (in the liver)
 - Target tissue
 - Microbial
 - Multiple metabolites
- ► Enterohepatic recirculation
 - Most absorbed PAHs are returned to the GI tract through bile and some are reabsorbed
- ► IVBA requires simulated intestinal environment

ITRC BCS-1 Section 8

109 Key Considerations in Study Design ITRC Document Provides Useful Information to Assess Studies



- ► Appropriate soil particle size
- ▶ Relevant comparison group
- Linearity of pharmacokinetics
- ► Repeated versus single dose
- Measurement of parent compound, metabolites, or both
- Adequate number of subjects

- Relevant concentrations/doses, number of different doses
- ► Ability to demonstrate full range of RBA
- Average versus individual subject RBA measurements
- ▶ Mass balance

ITRC BCS-1 Section 5

110

Key Considerations in Study DesignITRC Document Provides Overview Specific to PAHs



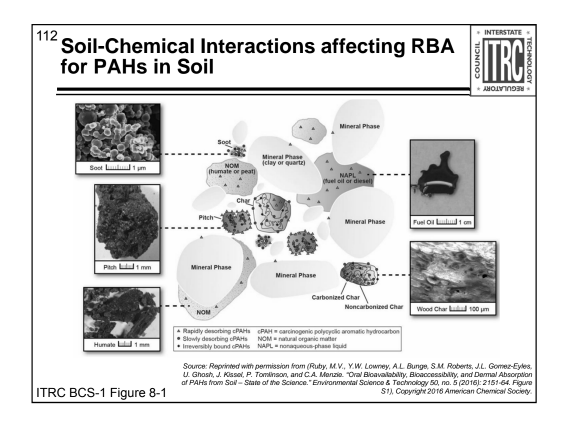
- ▶ Sources
- **▶** Toxicity
- ▶ Factors influencing RBA from soil
- ▶ In vivo and in vitro methods
- ► Summary of research conducted to date
- ► Considerations for dermal absorption
- ► Case study

ITRC BCS-1 Section 8

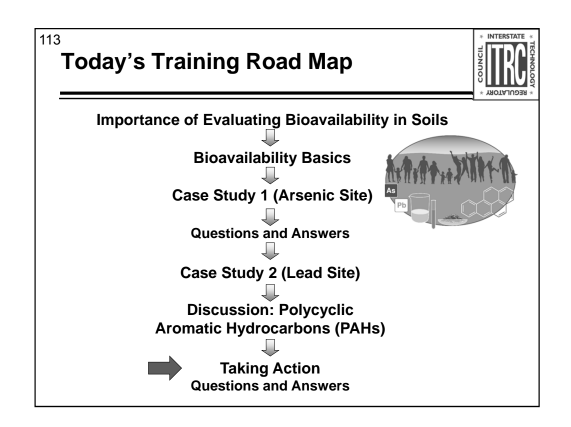
Bioavailability of PAHs from Soil What We Know



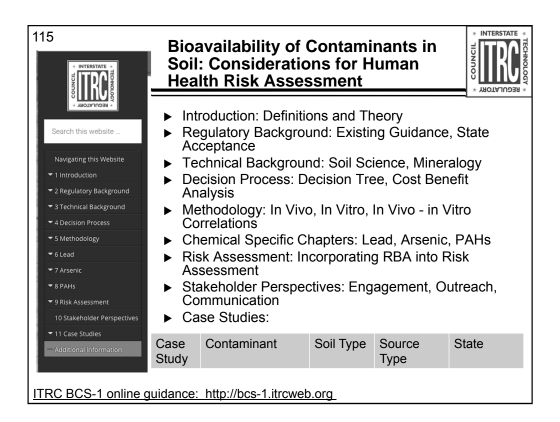
- ➤ Source of PAHs to soil dominates partitioning (in vitro) and RBA (in vivo)
- ► Some sources have higher RBA, others significantly reduced relative to soluble forms
 - · Lower RBA: Soot, Skeet, Pitch
 - Higher: Fuel oil, Non-aqueous phase liquid (NAPL)
- ► Soil characteristics are less important to controlling RBA (peat, clay content)
- ▶ Addition of charcoal to the soil reduces RBA
- Dermal exposure pathway important to calculated exposures
- ▶ More work to be done and is being done!

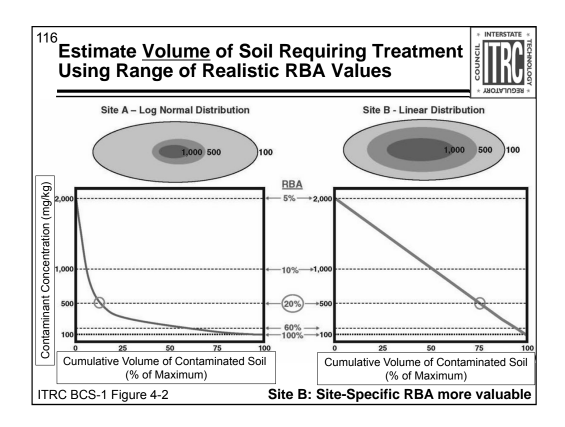


Source: Reprinted with permission from (Ruby, M.V., Y.W. Lowney, A.L. Bunge, S.M. Roberts, J.L. Gomez-Eyles, U. Ghosh, J. Kissel, P. Tomlinson, and C.A. Menzie. "Oral Bioavailability, Bioaccessibility, and Dermal Absorption of PAHs from Soil – State of the Science." Environmental Science & Technology 50, no. 5 (2016): 2151-64. Figure S1), Copyright 2016 American Chemical Society.









Two sites are shown in Figure 4-2, each with a maximum concentration of 2,000 mg/kg of a contaminant that has a cleanup level of 100 mg/kg (at an RBA of 100%). The RBA values are overlaid, to illustrate the cleanup levels corresponding to a given RBA.

As an example, the green circles indicate the volumes impacted if an RBA of 20% were accepted, effectively raising the cleanup level to 500 mg/kg.

At Site A, only 15% of the total contaminated soil volume is above 500 mg/kg, (contaminant distribution is log normal) and therefore would require cleanup. In contrast, with a different distribution (linear distribution) of the contaminant concentrations (Site B), 75% of the total volume would still require remediation at an RBA of 20%.

Site-specific conditions will vary, but some key features of the analysis of volume and RBA in Figure 4-2 are worth pointing out: Risk-based criteria, such as cleanup levels, increase significantly at RBA values of approximately 25% or less. For example: an RBA of 25% yields a cleanup level that is 4x higher an RBA of 10% yields a cleanup level that is 10x higher

The typical default value of a 60% RBA results in a relatively modest increase in cleanup levels: 1.67x higher.

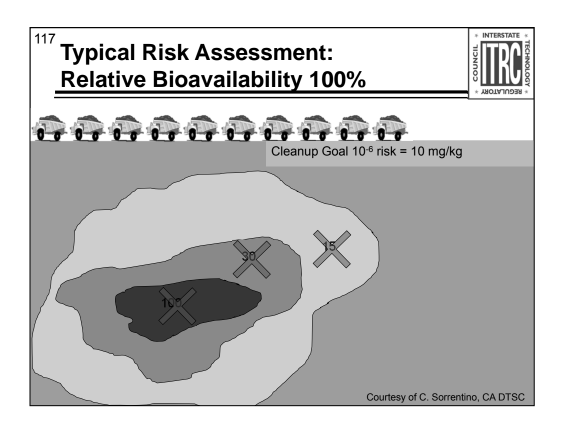
Estimating the volume requiring treatment at a range of realistic RBAs before beginning a site-specific bioavailability study may be valuable.

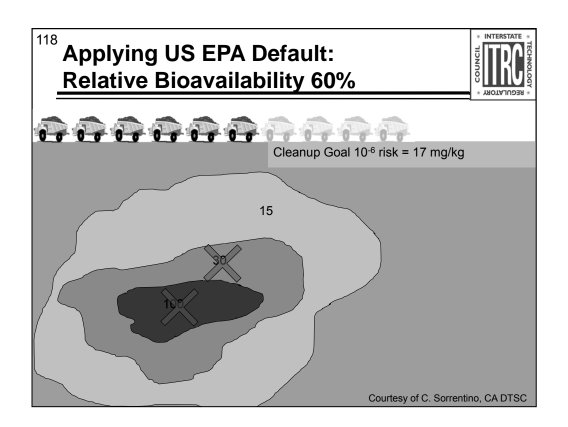
Some general observations regarding the value of incorporating site-specific RBA values include the following:

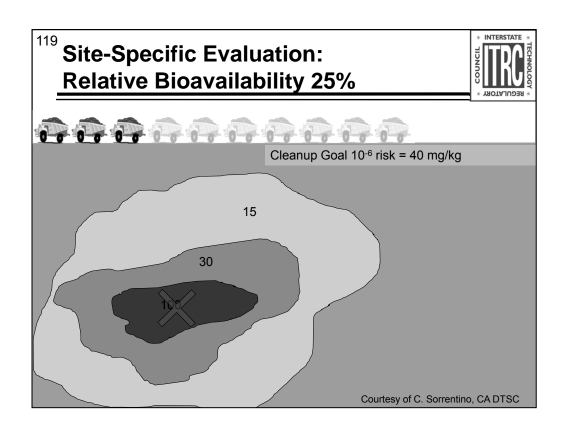
Small sites may not justify the expense of testing and increased regulatory costs.

At sites where discrete hot spots account for most of the risk (like Site A), or at sites with only a small volume of soil above cleanup goals, site-specific bioavailability assessment may be less valuable.

Bioavailability assessment is more valuable at sites with relatively high volumes of soil, and where most of the soil is contaminated at concentrations between the default cleanup levels and cleanup levels that incorporate an estimated RBA value (based on prior literature or experience with the specific soils or waste materials).





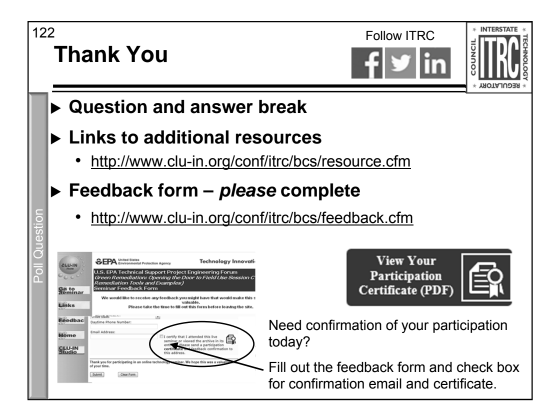


120 ITRC Document: Review Checklist * INTERSTATE http://bcs-1.itrcweb.org Bioavailability of Contaminants in Soil: Considerations for Human Health Risk Assessment номе **Review Checklist** This checklist summarizes elements that should be considered when developing or reviewing a risk assessment that uses a site-specific bioavailability or relative oral bioavailability (RBA) value. The checklist can be completed by a risk assessor or project manager or used by a reviewer to document that the information contained in the bioavailability assessment is complete and justified. Each site will vary depending on the chemical of interest, objectives, and purpose of the risk assessment. ☐ Are the methods used for soil sampling, chemical analysis and bioavailability testing including rationale for their selection and limitations, adequately described? [Lead, Arsenic, and PAH] • What soil sampling methods (for example, discrete, ISM) were used? What sieving was performed and what sieve size was used, if What analytical methods for the contaminants were used? • Identify the bioavailability and bioaccessibility methods (type of in vivo, in vitro, or combination models) used. • Identify the in vivo - in vitro correlation (IVIVC) used \square Is bioavailability assessment beneficial (feasibility, logistical and technical constraints)? [Decision Process] and Stakeholder • Is the site-specific bioavailability likely to affect the remedial decisions? • Is the cost of the bioavailability assessment justified with respect to the cost of remediation? Are validated bioavailability methods available? Has the use of site-specific bioavailability been accepted by the regulatory agency?

121 Site-Specific RBA Evaluation Take Home Messages



- ▶ Decrease the uncertainty of the risk assessment
- ► Maintains the Target Risk Level
- ▶ Improve Remedial Decision Making
- ► Often lead to significant savings of the resources available for remediation
- ▶ Multidisciplinary: Involve the Whole Team Early!
 - Regulatory: Project Managers, Geologists, Risk Assessors/Toxicologists
 - Consultants
 - · Stakeholders: Responsible Parties, Public



Links to additional resources:

http://www.clu-in.org/conf/itrc/bcs/resource.cfm

Your feedback is important – please fill out the form at:

http://www.clu-in.org/conf/itrc/bcs/feedback.cfm

The benefits that ITRC offers to state regulators and technology developers, vendors, and consultants include:

- ✓ Helping regulators build their knowledge base and raise their confidence about new environmental technologies
- √ Helping regulators save time and money when evaluating environmental technologies
- ✓ Guiding technology developers in the collection of performance data to satisfy the requirements of multiple states
- ✓ Helping technology vendors avoid the time and expense of conducting duplicative and costly demonstrations
- ✓ Providing a reliable network among members of the environmental community to focus on innovative environmental technologies

How you can get involved with ITRC:

- ✓ Join an ITRC Team with just 10% of your time you can have a positive impact on the regulatory process and acceptance of innovative technologies and approaches
- √Sponsor ITRC's technical team and other activities
- ✓ Use ITRC products and attend training courses
- ✓ Submit proposals for new technical teams and projects