NOTE TO READERS

The Canadian Council of Ministers of the Environment (CCME) is the primary minister-led intergovernmental forum for collective action on environmental issues of national and international concern.

This document was developed by the Contaminated Sites Working Group. It revises Federal Contaminated Sites Action Plan (FCSAP) Ecological Risk Assessment Guidance (Government of Canada n.d.), which was originally developed by Azimuth Consulting Group Inc. under contract with Environment and Climate Change Canada. The FCSAP Ecological Risk Assessment Guidance focusses on federal sites. CCME has modified the FCSAP Ecological Risk Assessment Guidance document to ensure it is applicable to all jurisdictions and aligns with CCME’s Framework for Ecological Risk Assessment: General Guidance (1996).

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<th>Definition</th>
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<tr>
<td>AEL</td>
<td>acceptable effect level</td>
</tr>
<tr>
<td>ANOVA</td>
<td>analysis of variance</td>
</tr>
<tr>
<td>APEC</td>
<td>area of potential environmental concern</td>
</tr>
<tr>
<td>ASTM</td>
<td>American Society for Testing and Materials</td>
</tr>
<tr>
<td>AVS</td>
<td>acid volatile sulfide</td>
</tr>
<tr>
<td>BAF</td>
<td>bioaccumulation factor</td>
</tr>
<tr>
<td>BCF</td>
<td>bioconcentration factor</td>
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<tr>
<td>BPJ</td>
<td>best professional judgment</td>
</tr>
<tr>
<td>BLM</td>
<td>biotic ligand model</td>
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<tr>
<td>CABIN</td>
<td>Canadian Aquatic Biomonitoring Network</td>
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<td>CCME</td>
<td>Canadian Council of Ministers of the Environment</td>
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<tr>
<td>CEC</td>
<td>cation exchange capacity</td>
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<tr>
<td>CEQG</td>
<td>Canadian Environmental Quality Guideline</td>
</tr>
<tr>
<td>COC</td>
<td>contaminant of concern</td>
</tr>
<tr>
<td>COPC</td>
<td>contaminant of potential concern</td>
</tr>
<tr>
<td>CSM</td>
<td>conceptual site model</td>
</tr>
<tr>
<td>DAF</td>
<td>dose adjustment factor</td>
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<tr>
<td>DQO</td>
<td>data quality objective</td>
</tr>
<tr>
<td>EC</td>
<td>Environment Canada</td>
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<tr>
<td>ERA</td>
<td>ecological risk assessment</td>
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<td>FCSAP</td>
<td>Federal Contaminated Sites Action Plan</td>
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<td>GIS</td>
<td>geographic information systems</td>
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<tr>
<td>HC_p</td>
<td>hazardous concentration (affecting percentage ( p ) of population)</td>
</tr>
<tr>
<td>HI</td>
<td>hazard index</td>
</tr>
<tr>
<td>HQ</td>
<td>hazard quotient</td>
</tr>
<tr>
<td>K_{ow}</td>
<td>octanol-water partition coefficient</td>
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<tr>
<td>LOAEL</td>
<td>lowest-observed-adverse-effect concentration / level</td>
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<tr>
<td>NOAEC / NOAEL</td>
<td>no-observed-adverse-effect concentration / level</td>
</tr>
<tr>
<td>NRC</td>
<td>National Research Council</td>
</tr>
<tr>
<td>MECP</td>
<td>Ontario Ministry of the Environment, Conservation and Parks</td>
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<td>PAH</td>
<td>polycyclic aromatic hydrocarbon</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Definition</td>
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<td>--------------</td>
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<tr>
<td>PCB</td>
<td>polychlorinated biphenyl</td>
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<tr>
<td>QA</td>
<td>quality assurance</td>
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<td>QC</td>
<td>quality control</td>
</tr>
<tr>
<td>RCA</td>
<td>reference condition approach</td>
</tr>
<tr>
<td>SAB-CS</td>
<td>Science Advisory Board for Contaminated Sites in British Columbia</td>
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<tr>
<td>SAP</td>
<td>sampling and analysis plan</td>
</tr>
<tr>
<td>SEM</td>
<td>simultaneously extracted metal</td>
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<tr>
<td>SSD</td>
<td>species sensitivity distribution</td>
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<td>SWAMP</td>
<td>Surface Water Ambient Monitoring Program</td>
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<td>TEQ</td>
<td>toxic equivalents</td>
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<td>TIE</td>
<td>toxicity identification evaluation</td>
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<tr>
<td>TRA</td>
<td>tissue residue approach</td>
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<td>TRG</td>
<td>tissue residue guideline</td>
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<td>TRV</td>
<td>toxicity reference value</td>
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<tr>
<td>UCLM</td>
<td>upper confidence limit of the mean</td>
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<tr>
<td>US EPA</td>
<td>United States Environmental Protection Agency</td>
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<tr>
<td>VEC</td>
<td>valued ecosystem component</td>
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<tr>
<td>WOE</td>
<td>weight of evidence</td>
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</table>
GLOSSARY

Abiotic medium is any environmental medium not associated with biological tissue (e.g., soil, sediment, water, air).

Acute relates to a small increment of time required to elicit an adverse environmental response. With respect to toxicity testing, the term describes tests applied over a short duration, typically less than 10 per cent of an organism’s lifespan. Note, however, that some short-term tests may be defined as chronic rather than acute if they are conducted using a sensitive life stage; definitions of acute versus chronic vary widely by jurisdiction.

A priori refers to prior knowledge about a condition, rather than that estimated by recent observation. In ecological risk assessment, the term a priori describes knowledge or models of biological systems the risk assessor considers before the analysis phases of the risk assessment.

Acceptable effect level (AEL) is the magnitude (or rate) of effects that would be acceptable for a specific measurement endpoint or assessment endpoint. The AEL operationalizes a protection goal.

Analysis of variance (ANOVA) is a statistical method used for a single dependent variable that performs comparisons and tracks the effects of one or more discrete factors (independent variables), each of which may have a number of levels and may interact to affect the dependent variable.

Application factor: see uncertainty factor.

Area of potential environmental concern (APEC) is a portion of a site where contamination is suspected or confirmed.

Assessment endpoint is an explicit expression of the environmental value to be protected. An assessment endpoint must include an entity (typically a receptor or receptor group—i.e., a “thing” to be protected) and a specific property of that receptor (an attribute). For example, if the entity is a fish community, attributes could include the number of species and the trophic structure. An assessment endpoint may also have an explicit spatial or temporal component.

Assessment factor: see uncertainty factor.

Aryl hydrocarbon receptor (Ah receptor) is a member of the family of basic helix-loop-helix transcription factors. The Ah receptor binds to certain chemicals, such as dioxins and polychlorinated biphenyl (PCB) congeners, causing the receptor to translocate into the nucleus of organism cells, eventually leading to genetic damage. The mechanism of toxicity via the Ah receptor underpins the use of the toxic equivalents system for evaluating responses of chlorinated organic substances to vertebrates.

Attribute is a quality of an endpoint that reflects one aspect of its value for informing the risk assessment.

Best professional judgment (BPJ) is the thorough application of critical judgment in professional practice, in which an experiential, reflective, self-corrective and purposeful thinking process is
applied to consider knowledge, context, evidence, methods, conceptualizations and criteria. BPJ is a means by which a practitioner can incorporate a diverse range of information without articulating a mechanical process for processing the information.

**Bias** is a systematic tendency that distorts the interpretation of results. In ecological risk assessment, a bias occurs in two main forms. In the study design or interpretation, bias is a pejorative term that reflects the partiality of a practitioner and prevents them from objectively considering an issue or situation. In statistical measurement, bias reflects a systematic under- or over-prediction of a true parameter value. Both forms of bias introduce systematic error into risk estimates.

**Bioaccumulation** is the process by which substances accumulate in the tissues of living organisms. Bioaccumulation occurs when the concentration of a contaminant of concern in an organism is higher than the concentration in the surrounding environment. Most substances bioaccumulate to some extent, whereas few biomagnify.

**Bioaccumulation factor (BAF)** is the quotient obtained by dividing the concentration of a substance in an organism (or specified tissue) by its concentration in a specified exposure medium, for example, air, food, sediment, soil, water (ASTM 2011).

**Bioconcentration factor (BCF)** is equivalent to an uptake factor, for the case where water (only) is the abiotic exposure medium.

**Biomagnification** refers to the process by which chemical concentrations in plants or animals increase relative to food from transfer through the food web (e.g., predators have greater concentrations of a particular chemical than their prey).

**Biota-sediment accumulation factor** is equivalent to an uptake factor, where the abiotic medium is sediment, and where both the tissue and sediment concentrations are normalized to carbon pools (lipid and total organic carbon, respectively).

**Biotic medium** is any biological medium (e.g., tissue) where contaminants of concern may be found.

**Category of evidence** is a group of related lines of evidence within a weight of evidence framework.

**Causal pathway assessment** determines proximate causes and identifies their sources and, where possible, characterizes the causal pathways that connect them.

**Causation** is the act or fact of causing, or the production of an effect by a cause. Causation differs from association (correlation) in that the latter does not imply a mechanistic linkage between observations. An assessment of causation in an ecological risk assessment attempts to distinguish between associations that are coincidental or caused by external factors and associations that are driven by underlying predictable mechanisms.

**Chronic** relates to an extended duration. In the context of toxicity testing, the term is used to describe tests that expose organisms over a substantial portion of their life cycle, for example more
than 10 per cent of the life cycle or throughout a sensitive life stage. Definitions of chronic vary widely.

Cluster analysis is a class of statistical techniques that can be applied to data that exhibit “natural” groupings based on an assessment of interdependence. Cluster analysis sorts through the raw data and groups them into clusters of relatively homogeneous cases or observations. Whereas factor analysis reduces the number of variables by grouping them into a smaller set of factors, cluster analysis reduces the number of observations or cases by grouping them into a smaller set of clusters.

Coherence is a concept that relates to the way in which multiple lines of evidence are congruent; approaches to the assessment of coherence include evaluations of causation, ecological relevance, logical interpretations and best professional judgment. In a weight of evidence approach, coherence analysis is applied following the “face-value” interpretation of results to determine whether the lines of evidence are consistent or provide a unified interpretation of findings.

Concentration response show the relationship between an effects measure and exposure (measured as concentration) across a range of exposure concentrations.

Conceptual site model (CSM) is a narrative and graphical representation of the relationships between contaminant sources, fate, exposure pathways, and receptors.

Condition assessment detects chemical, physical and biological impairment by analyzing environmental monitoring data.

Conservative expresses the tendency to deliberately overstate the potential for environmental harm. The overestimate is intended to provide a margin of error to buffer against uncertainty in the analysis, and to provide increased confidence that estimates or predictions of risk are not understated. In ecological risk assessment practice, it is common to apply conservatism in parameter estimation. However, when conservatism is too great, either through unrealistic assumptions or through compounding of multiple conservative assumptions, an analysis is deemed to be ultra-conservative, and therefore suspect.

Contaminants of Concern (COCs) are contaminants that have been selected for evaluation in the ecological risk assessment.

Control as a noun is an aspect of a controlled scientific experiment conducted to determine the effect of a single variable of interest on a particular system, used to minimize the unintended influence of other variables on the same system. Negative controls confirm that the procedure is not causing an unrelated effect, and are intended to reduce incidence of false positives. The term control as a verb can also be used in experimental design to refer to manipulation of treatments intended to mitigate the confounding effect of external variables.

Correspondence analysis is a multivariate statistical technique that is conceptually similar to principal components analysis, in which data are scaled such that rows and columns are treated equivalently.
Critical body residue (CBR) is an internal body or tissue concentration that is associated with a toxicological response in a receptor.

Deterministic methods are methods in which all biological, chemical, physical and environmental parameters are assumed to be constant and can be accurately specified. Deterministic methods commonly apply to either a “most likely” value for a parameter or a conservative value intended to guard against uncertainty.

Dichotomous characterizes a parameter with only two possible states.

Dilution series is an experimental design and technique in which an abiotic medium is divided into multiple exposure magnitudes by diluting the full-strength medium using clean material. A series of concentrations is specified using graded dilutions, with responses characterized for each treatment on a volume/volume, mass/volume or mass/mass basis.

Diversity is an attribute referring to variation within an ecological community. In general, high diversity is associated with high richness (number of taxa) and evenness of abundance among taxonomic groups. Diversity is often used as a measure of ecosystem health. A number of numerical diversity indices have been developed, each of which has different theoretical underpinnings.

Dose-response is the relationship between an effects measure and exposure (measured as dose) across a range of dose values.

Ecological relevance is the degree to which a type of information used in an ecological risk assessment (e.g., a measurement endpoint or line of evidence) can be meaningfully extrapolated to the biological scale of interest (e.g., the assessment endpoint).

Ecological risk assessment (ERA) is the process of evaluating the potential adverse effects on non-human organisms, populations or communities in response to human-induced stressors. ERA applies a formal framework, analytical process or model to estimate the effects of human actions on natural organisms, populations or communities and interprets the significance of those effects in light of the uncertainties identified in each study component.

Effect size is the absolute or relative magnitude of response to a stressor for a measurement endpoint.

Exposure assessment is, for any line of evidence, the component of a risk assessment that quantifies the degree to which an organism encounters a stressor.

Exposure pathways are the routes through which a receptor of concern encounters contaminants of concern in environmental media (e.g., soil, water, air, sediment). Examples of exposure pathways include ingestion and inhalation.

Exposure point concentration is the value that represents a conservative estimate of the chemical concentration or dose available to an organism from a route of exposure.

Extrapolation is an inference or estimation done by extending or projecting known information to a domain (spatial, temporal, biological or chemical) that has not yet been studied. In statistics, extrapolation entails estimation (of a value of a variable outside a known range) from values within
a known range, and requires an assumption that the estimated value follows logically from the known values.

**Extrapolation factor**: see uncertainty factor.

**Factor analysis** is a class of statistical methods that analyzes the variability among observed correlated variables in order to potentially reduce the number of variables to a set of fewer unobserved variables called factors. Factor analysis reduces the number of variables by grouping them into a smaller set of factors, whereas cluster analysis reduces the number of observations or cases by grouping them into a smaller set of clusters.

**False negative** is the error (often called a Type II error) in which a response occurs but is not detected.

**False positive** is the error (often called a Type I error) in which a response is deemed to occur when in fact there was no response. The term is often used to describe a situation in which an inappropriate conclusion was rendered based on available information.

**Feeding guild** is a group of organisms that use the same ecological resource in a similar way for feeding (e.g., insectivores, granivores, detritivores, carnivores) or a group of species that overlap significantly in their niche requirements.

**Gradient** is a concept of experimental design in which treatments are planned to include a range of exposures from low to high, or within a spatial range (e.g., near to far).

**Guidelines** are generic numerical limits or narrative statements that are recommended to protect and maintain the specified uses of water, sediment or soil.

**Hazard assessment** is, for any line of evidence, the component of a risk assessment that characterizes the nature of effects elicited by each contaminant under an exposure condition that is relevant to each receptor of concern.

**Hazard quotient (HQ)**: is a numerical ratio that divides an estimated environmental concentration or other exposure measure by a response benchmark. Typically, the response benchmark is a value assumed to be protective of the receptor of concern. HQ values below 1.0 indicate negligible potential for harm, whereas HQ values above 1.0 indicate that an adverse response is possible and that more precise or accurate evaluation of risks may be warranted to address uncertainty.

**Hazardous concentration (HCp)** is a threshold concentration from a species sensitivity distribution. The concentration is derived considering a proportion of the species affected (p) and an effect size of interest (e.g., acceptable level of response).

**Hazard index (HI)** is the arithmetic sum of individual hazard quotients, used to aggregate the individual responses of multiple stressors. The HI implicitly assumes linear additivity of response. An HI is applied where the mode of toxic action is considered to be similar among contaminants of concern.

**Home range** is the geographic area to which an organism normally confines its activity. For exposure assessment the activity of interest is usually the foraging area over a defined period of time, such as feeding range during the reproductive period.
**Hypothesis** is a proposed explanation for an observable phenomenon. In experimental design, a hypothesis is set forth and subsequently tested (either singly or along with multiple alternate hypotheses) to determine if the new data support or contradict the hypothesis.

**Interpolation** is the process of estimating a value of (a function or series) between two known values. The term can also be applied more generically to the assignment of qualities to members of a group based on observations of other members of the same group. Interpolation requires the underlying assumption that members of a group are similarly influenced by the processes under investigation.

**Likelihood** is, in common usage, synonymous with the probability or frequency of an event. In statistical usage, likelihood is distinguished from probability, and refers to estimating unknown parameters based on known outcomes.

**Line of evidence** is any pairing of exposure and effects measures that provides evidence for the evaluation of a specific assessment endpoint. Typically, a line of evidence requires use of one or more measurement endpoints. If the focus of the line of evidence is an effects measure (e.g., a toxicity test), the paired exposure measure may be quantitative (e.g., contaminant concentrations) or categorical (e.g., on-site versus reference conditions).

**Linear model** is a category of statistical methods that underlies many of the statistical analyses that are used in applied sciences. It is the foundation for the Student’s t-test, ANOVA, regression analysis and many multivariate methods. Linear models assume that the relationship between a response variable and explanatory variables (or factors) is linear or can be approximated as linear following appropriate data transformation.

**Line of evidence group** is a cluster of closely related lines of evidence that have a particular measurement endpoint (or multiple endpoints) in common and therefore incorporate some redundancy in a weight of evidence evaluation. Individual lines of evidence in a group should individually contribute sufficient incremental information (e.g., informing the evaluation of the assessment endpoint) to warrant inclusion as separate lines. A line of evidence grouping provides organization of related lines of evidence and flags potential for redundancy.

**Lowest-observed-adverse-effect level (LOAEL)** is the lowest amount, dose or concentration of an agent, found by experiment or observation, that causes an adverse alteration of morphology, functional capacity, growth, development or life span in an organism, system or (sub)population. Methods vary for identifying a LOAEL, but often apply statistical significance as a criterion.

**Measurement endpoint** is a parameter that measures or describes exposure of, or an effect on, a receptor of concern. Alternatively, the term describes a change in an attribute of an assessment endpoint (or its surrogate) in response to a stressor to which it is exposed.

**Model** is a simplified description of a system, theory or phenomenon that accounts for its known or inferred properties and that may be used for further study of its characteristics. In all cases, a model is a simplification of a more complex system, and the details not represented by the model structure are considered to be errors or variations not central to the problem at hand. Models include statistical models (numerical processes used to simulate or approximate complex
processes) and conceptual models (graphical or schematic representation of key processes and pathways).

**Monte Carlo analysis** is a probabilistic analysis technique where parameter values are drawn at random from defined input probability distributions and combined according to a model equation. The process is repeated iteratively until a relatively smooth distribution of solutions results.

**Multivariate** is a form of statistics encompassing the simultaneous observation and analysis of more than one statistical variable. In ecological risk assessment, the most common multivariate methods are clustering, correspondence analysis, factor analysis, principal components analysis and multi-dimensional scaling.

**Narcosis** is a condition of deep stupor or unconsciousness produced by a drug or other chemical substance.

**No-observed-adverse effect level (NOAEL)** is an exposure level at which there are no statistically or biologically significant increases in the frequency or severity of adverse effects between the exposed organisms or population and the appropriate control. Some effects may be produced at this level, but they are not considered to be adverse. Methods for identifying a NOAEL vary, but often apply statistical significance as a criterion.

**Ordination** is a method in multivariate analysis, complementary to data cluster analysis. It orders objects on multiple variables such that similar objects are near each other and dissimilar objects are farther from each other. These relationships between the objects are plotted on multiple axes and can be characterized numerically or graphically.

**Outcome assessment** evaluates the results of a past management action, through estimation or direct measurement.

**Point estimate** is a single numerical value used to represent the state of a random variable. A point estimate collapses (or ignores) all of the variability and incertitude regarding a parameter or variable.

**Potentially responsible party** refers to all industries, site owners, point sources and legally responsible entities associated with contamination at a site. The term is commonly used as part of contaminated sites legislation in the United States (*Comprehensive Environmental Response, Compensation, and Liability Act*, or Superfund).

**Practitioner** is an investigator in an ecological risk assessment responsible for the design, implementation and interpretation of results. The practitioner, who may be a consultant, interacts with the responsible party for the site (client), the regulators and other interested parties. In this guideline, the practitioner is also referred to as the *risk assessor*.

**Precision** is the quality of being repeatable in degree or value, or the ability of a measurement to be consistently reproduced. Note that precise results are not necessarily accurate, as a precise measurement can be consistently biased.
**Predictive assessment** estimates environmental, economic, and societal risks and benefits associated with different management alternatives. Acceptability of actions may be determined through evaluating the risks in light of social, economic and legal considerations.

**Prescriptive** pertains to giving directives or rules, without flexibility or subjective analysis. Prescriptive approaches have a high degree of repeatability and consistency among investigators, but low degree of adaptability to site-specific conditions.

**Probabilistic** describes a procedure in which the state of a random variable is described not as a point estimate (fixed value), but rather as a distribution of possible values. Using probabilistic methods, important biological, chemical, physical and environmental parameters are assumed to vary or are uncertain and therefore are specified using distributions.

**Probability** is a mathematical way of expressing knowledge or belief that an event or outcome will occur or has occurred. In statistical usage, probability is distinguished from likelihood, and refers to the prediction of unknown outcomes based on known parameters.

**Problem formulation** is the first step in an ecological risk assessment and clarifies the nature of issues associated with contamination at a site and how those issues will be addressed.

**Protection goal** is a narrative statement that defines the desirable level of protection for a receptor or receptor group (see also acceptable effect level).

**Qualitative** describes an approach that is narrative, referring to the characteristics of something being described, rather than numerical measurement.

**Quantitative** describes an approach that is numerical (applies mathematical scores, probabilities or parameters) in the derivation or analysis of risk estimates.

**Receptor of concern** is any non-human individual organism, species, population or community that is potentially exposed to contaminants of concern and that is considered in an ecological risk assessment to evaluate the potential risks to a valued ecosystem component. Identification of an organism, species, population or community as a receptor of concern does not mean that it is being harmed, only that a pathway exists such that there is potential for harm.

**Reference (condition)** is a location, group of locations or experimental treatment designed to reflect the ambient physical and chemical conditions of a contaminated medium or location in the absence of the stressors of concern in the risk assessment. For example, in a study of soil contamination, the reference condition should reflect the climate, substrate and habitat factors relevant to the site but with no incremental contamination relative to background conditions. In some cases, the term reference may be used in the context of an altered local background condition (e.g., where the local conditions surrounding a site are not pristine due to non-point sources of contaminants). In other cases, the term reference is used to refer to pristine conditions in the absence of both site-specific contamination and non-point sources of contaminants.

**Regression** is a form of statistical modelling that attempts to evaluate the numerical relationship between one variable (termed the *dependent variable*) and one or more other variables (termed the *independent variables*).
Response profile is the relationship between contaminant of concern concentrations and ecological effects.

Richness is used in analysis of biological communities to refer to the variety of organisms present in a sample (e.g., the variety of plants or invertebrates). The value of richness can be determined by summing the number of unique taxa present in the sample.

Risk characterization is the process of estimating the magnitude (and where relevant, the probability) of adverse ecological impacts based on the information obtained from the exposure and hazard assessments. Risk characterization also translates complex scientific information into a format that is useful for risk managers by conveying the ecological consequences of the risk estimates along with the associated uncertainties.

Sensitivity is the quality of being able to reliably detect perturbations in a parameter.

Spatial relates to space, particularly in terms of the lateral (horizontal) dimension. In ecological risk assessment, the term spatial is often used to refer to level of resolution (grain) and extent (area).

Species sensitivity distribution (SSD) is a cumulative probability distribution of toxicity values for multiple species.

Standard refers to an environmental benchmark that may be subject to regulatory enforcement. Standards can be associated with specific environmental legislation that conveys the responsibilities of site owners.

Statistical power is the probability that a test will properly reject a false null hypothesis (i.e., that it will not make a Type II error). The probability of a Type II error is referred to as the false negative rate ($\beta$). Therefore, power is equal to $1 - \beta$. Although there are no formal standards for power, many researchers assess the power of their tests using 0.80 as a standard for adequacy. Factors influencing the power of a given test (or study design) include the statistical significance criterion for probability of a Type I error ($\alpha$), the magnitude of the effect of interest in the population, the sample size ($n$), and the variation of the underlying data as determined by measurement error and stochasticity.

Stochasticity is random natural variations. Stochastic processes can be simulated, but the variations cannot be reduced through additional analysis, only better described.

Stressor is any substance or process that may cause an undesirable response to the health or biological status of an organism.

Surrogate valued ecosystem component is a receptor of concern that is representative of a receptor type that has been chosen as a valued ecosystem component (VEC) (e.g., a shrew may be used as a surrogate VEC for insectivorous mammals). More than one surrogate receptor of concern may be used to represent a particular VEC.

Taxon (plural: taxa) is a grouping of organisms given a formal taxonomic name (biological classification) such as species, genus and family, and identified as genetically distinct from other organisms.
Temporal relates to time, particularly in terms of changes or variations observed over a time period of interest.

Threshold is the dividing line (in units of exposure concentration or dose) between a zone of potential response and a zone of negligible response. Thresholds may be estimated using theory, data or a combination of both. In nature, thresholds generally do not occur as precise or static entities, due to the variations among individuals and environmental factors that influence responses. Therefore, a threshold is usually expressed as a best estimate considered protective of most of the population, and often includes a margin of safety in the derivation.

Tissue residue guidelines (TRG) are criteria or guidelines that refer to an internal body or tissue concentration in a receptor.

Toxicity is the observation of a chemically induced physiological or biological response that impairs the health of an organism.

Toxicity identification evaluation (TIE) is a tool in which physical or chemical manipulation of a sample is conducted to isolate and to identify toxic substances in a test medium. A biological test, in this case a toxicity test, is used as the indicator to determine whether the manipulation changed toxicity.

Toxicology is the field of science that explores the relationship between substances of environmental concern and the responses elicited to organisms.

Toxicity reference value (TRV) is an exposure concentration or dose that is not expected to cause an unacceptable level of effect in receptor(s) exposed to the contaminant of concern. A TRV is a specific type of threshold, as defined above.

Type I error is synonymous with false positive, the error of rejecting a null hypothesis when it is actually true. A Type I error occurs when we observe an apparent difference when in truth there is no difference, thus indicating a test of poor specificity.

Type II error is synonymous with false negative, the probability that a test will not reject an invalid null hypothesis. The probability of a Type II error is referred to as the false negative rate (β). This is the error of failing to observe a difference when in truth there is one, thus indicating a test of poor sensitivity.

Uncertainty is a term used in subtly different ways in a number of scientific fields. Generally, it refers to imperfect knowledge regarding a given parameter, process or condition. In risk assessment, uncertainty is the state of having limited knowledge where it is impossible to exactly describe an existing state or future outcome. Uncertainties come in many forms, including measurement uncertainty, random variations, conceptual uncertainty and ignorance.

Uncertainty factor is called an application factor, extrapolation factor or safety factor. It is a numerical factor sometimes used in hazard assessment and applied to observed endpoints in order to derive an exposure concentration below which adverse effects are unlikely to occur. The factor is applied in the face of uncertainty and applied in order to not underestimate risk. As the quantity
and quality of test data increase and their relevance to the organisms of interest improves, the size of the extrapolation factor diminishes.

**Univariate** tests are statistical tests that address one variable at a time. The term also applies to statistical tests for comparing two or more groups with respect to a single property, including the Student’s t-test, ANOVA, sign test, Wilcoxon rank sum test and the Mann-Whitney U test.

**Upper confidence limit of the mean (UCLM)** is a statistical measure of the upper bound of a confidence interval for the mean value of an environmental parameter, such as the expected environmental concentration of a substance.

**Uptake factor** is used to extrapolate contaminant concentrations from a single abiotic exposure medium to a tissue concentration in an organism. Several types of uptake factors exist, including the bioconcentration factor, bioaccumulation factor and the biota-sediment accumulation factor.

**Valued ecosystem component (VEC)** is, for the purposes of an ecological risk assessment (ERA), a component of the ecosystem that is potentially adversely affected, either directly or indirectly, by the contaminants at a site and that is identified by the risk assessor as one for which the ERA is to be designed to protect. A VEC can be any non-human individual organism, species, population, community, habitat or ecosystem. A receptor of concern may be the same as a VEC but it can also be a surrogate for the VEC or be a useful element in a line of evidence but not a VEC. For example, a VEC may be a wetland complex. Several receptors of concern may be selected to evaluate key attributes of this wetland (e.g., specific species at risk, diverse aquatic plant community, nutrient processing, water retention) and these would be evaluated to determine the potential direct and indirect risk of contaminants to the VEC.

A VEC is identified as such through having one or more of the following qualities:

- intrinsic ecological significance
- importance to human populations
- economic and or social value
- ability to serve as a baseline from which effects of changes can be evaluated.

**Weight** is the degree of emphasis placed on a finding or line of evidence relative to others. The weight is a function of the overall value (information, reduction of uncertainty) in terms of addressing an assessment endpoint, and is determined by assessing the attributes relevant to the study.

**Weight of evidence (WOE)** is a systematic procedure used to aggregate or synthesize a number of different types of evidence, with the objective of developing a single unified conclusion or explanation to an environmental characterization. WOE is one of the tools applied during the risk characterization stage of an ecological risk assessment.

**Wildlife**, in the context of ecological risk assessment, generally applies to birds and mammals and sometimes defined to include reptiles and amphibians. Generally, it excludes fish and invertebrates.
1 INTRODUCTION

The Ecological Risk Assessment Guidance Document is intended for site managers and risk assessors. It provides general guidance for conducting ecological risk assessments (ERAs) for soils, sediments, surface water and groundwater, largely in the context of managing contaminated sites. On its own, this document does not provide detailed technical guidance for conducting risk assessments.

For technical guidance, please see the four Federal Contaminated Sites Action Plan (FCSAP) technical modules:

- Module 1: Toxicity Test Selection and Interpretation (Environment Canada [EC] 2010-a)
- Module 2: Selection or Development of Site-specific Toxicity Reference Values (EC 2010-b)
- Module 3: Standardization of Wildlife Receptor Characteristics (EC 2012)
- Module 4: Causality Assessment (EC 2013)

1.1 Background

The basic ERA framework has already been described in previous documents (Canadian Council of Ministers of the Environment [CCME] 1996-a, 1997-a). This guidance document provides additional technical guidance to support risk assessment practitioners when conducting ERAs. It is not intended to replace the existing CCME framework; rather, it provides additional information and clarifications that have come to light since the previous documents were developed. For example, the current document discusses lines of evidence and the weight of evidence (WOE) approach, concepts that were not developed in the previous guidance.

1.2 Why Conduct an ERA?

Once a site is classified as contaminated, and has contaminant concentrations above existing ecologically based guidelines or levels of potential ecological concern, the site may be remediated to generic standards or an ERA may be used to determine whether and to what extent remediation or other risk management efforts are warranted to mitigate current or future ecological risks. An ERA provides a more detailed basis for determining whether remediation or other risk management measures are warranted (e.g., are there ecological risks?) and to what extent (e.g., which parts of a site should be remediated?).

1.3 Using ERA at Contaminated Sites

There are numerous potential drivers for the use of ERA at contaminated sites, such as regulatory triggers (e.g., contamination of an off-site property), due diligence or divestiture. The required ERA process may be driven in part or entirely by provincial or territorial regulations and policy.
This guidance document does not consider the jurisdictional context but rather attempts to focus on technical aspects of ERA that are likely to be applicable in many contexts, depending on the complexity of the ERA. The risk assessor is responsible for ensuring that the use of this guidance document meets appropriate jurisdictional requirements.

1.4 Communicating with Stakeholders
ERAs involve complex iterative processes. Stakeholder support is needed to ensure that the results are understood and that the objectives are consistent with stakeholder values. Consequently, everyone involved in the risk assessment process should encourage communication and early involvement of the various parties in the ERA process. Open lines of communication with the relevant jurisdiction(s) early in the process is important to ensure that the risk assessment will meet the expectations of the jurisdictional authority. Sharing information with appropriate stakeholders early in the process and obtaining stakeholder feedback at key milestones in the project is important. The site-specific consultation needs generally include up-front dialogue before work begins, as well as dialogue at milestones during the ERA process (e.g., review of a problem formulation).

1.5 Introduction to the ERA Framework
The standard conceptual framework for ERA (Figure 1-1) on a contaminated site described in this document remains consistent with existing CCME risk assessment guidance. However, the science of ERA is constantly evolving, and the last two decades have seen a significant increase in the complexity of risk assessments and the number of tools and methods used to characterize risks. The conceptual framework appears simple, but its application to multiple receptor groups via multiple exposure pathways using various lines of evidences can be quite complex. Consequently, in practice, the ERA framework is often applied using a WOE approach (Figure 1-2). In application, the WOE approach may be simple (e.g., a couple of lines of evidence for a single assessment endpoint) or complex (e.g., for complex sites with multiple receptors and assessment endpoints).

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Key Concept
Most ERAs warrant a WOE approach, whereby multiple lines of evidences are used to support the assessment. The WOE approach is entirely compatible with the standard conceptual framework for ERA.

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1 The precise terminology and delineation of the components of ERA vary across different jurisdictions and applications, but the vocabulary outlined here is used relatively consistently.
The WOE approach (Figure 1-2) integrates with the standard ERA framework as follows:

- Problem formulation defines the problem to be addressed and develops the scope for the ERA. For each receptor group or assessment endpoint, one or more lines of evidence\(^2\) are used in the risk assessment.
- Each line of evidence must combine information on exposure and effects.
- The exposure information typically characterizes the extent to which receptors are exposed to contaminants via various exposure pathways.
- The effects information characterizes the nature of effects observed or expected at the site.

\(^{2}\) The relationship between assessment endpoints, measurement endpoints and lines of evidences is considered in detail in Section 2.

**Definitions**

A *receptor of concern* in ERA is any non-human individual organism, species, population, community, habitat or ecosystem that is potentially exposed to contaminants of potential concern and that is considered in the ERA. Examples: a meadow vole population and a benthic invertebrate community.

An *assessment endpoint* is an explicit expression of the environmental value to be protected. An assessment endpoint must include a receptor or receptor group (i.e., an entity to be protected) and a specific attribute of that entity. Spatial and temporal elements may also be included. Example: abundance and viability of small mammal populations.
Figure 1-2: WOE approach to ERA

1. Problem Formulation

2. Assessment Endpoint A
   - Line of Evidence 1
     - Exposure Assessment
     - Hazard Assessment
     - Line of Evidence Evaluation
     - Risk Characterization (WOE)

3. Assessment Endpoint B
   - Line of Evidence 2
     - Exposure Assessment
     - Hazard Assessment
     - Line of Evidence Evaluation
     - Risk Characterization (WOE)

4. Overall Risk Characterization
Once individual lines of evidence are evaluated, the findings across all lines of evidence are evaluated in an integrated fashion to characterize risks for a particular assessment endpoint or receptor group.

1.6 Iterative Approach to Risk Assessment

In many cases, risk assessment follows a tiered approach, in which screening tools are applied at an early stage to determine if further work is needed or to prioritize future investigations. Often, jurisdictions will already have criteria or schedules that can be referred to for screening-level risk assessments that will cover a broad range of common contaminants of concern (COCs). These criteria will generally include screening for potential ecological effects as well as other pathways or receptors. If risks are acceptable using these criteria, there is probably no need for further work. On the other hand, if potential effects are identified, a more detailed and accurate risk assessment may be warranted.

This guidance document does not categorize types of risk assessments according to scope or level of detail (e.g., screening-level versus detailed risk assessment). Some regulatory or policy frameworks may have specific requirements in this regard. In practice, the process of tiering an ERA and the appropriate level of detail for each iteration is driven by many factors and is case-specific (Hill et al. 2000). The parties involved in an ERA should agree on the expectations for each iteration of the ERA, particularly regarding the type and degree of uncertainty they expect to resolve at each stage of investigation. Generally, it is important for each iteration of an ERA to address issues and uncertainties that are important from a risk management or decision-making perspective. In other words, each iteration of an ERA should significantly advance the usefulness of an ERA to support sound environmental management of contaminated sites.

1.7 Level of Detail

This guidance document and the four technical modules available on the FCSAP website (Government of Canada, n.d.) contain a high level of detail regarding many aspects of ERA. Consequently, some of the methods and approaches presented may apply only to complex sites where a detailed ERA is warranted. Importantly, and in accordance with the iterative approach to ERA articulated in the previous section, the level of complexity in an ERA should match the level of complexity of the site and its associated risks, taking into account the role of the ERA in supporting risk management decision making. Practitioners must judge the appropriate level of detail for each ERA on a site-specific basis.

As mentioned in the previous section, this guidance document does not define which ERA activities can be classified under the different levels of ERA as defined in CCME (1996-a) (i.e., screening, preliminary quantitative and detailed quantitative ERAs). Rather, the three documents CCME developed in 1996 can be viewed as potential iterations in the process. A risk assessment practitioner should be able to initiate an ERA at any of the three levels according to the needs of the situation, with additional iterations being possible within a level. With respect to screening-
level ERAs, soil, groundwater, water quality and sediment quality guidelines have often been developed based on protocols that are similar to screening-level risk assessments. Where these are available, discussion with the jurisdiction is appropriate to determine whether and how these can be employed to reduce the level of effort required in an ERA.

1.8 Organization of Document

This guidance document is organized around the conventional ERA framework, with major sections addressing problem formulation, exposure assessment, hazard assessment and risk characterization. This is intentional because the types of tools used for exposure assessment and hazard assessment fall into major categories that can conveniently be discussed simultaneously. Because exposure, effects and risk characterization apply to each line of evidence in an ERA, the concepts in all of the sections of this guidance must be understood before undertaking an ERA.

The introductions in Sections 2 to 5 provide an overview of problem formulation, exposure assessment, hazard assessment and risk characterization. Those introductions, together with text boxes on “key concepts” scattered throughout the text provide a sense of the scope and content of the guidance without complex technical details.

This guidance document cannot be comprehensive in all aspects. Selected aspects of ERA are addressed in detail in the four technical modules on the FCSAP website (Government of Canada n.d.).

1.9 Other Sources of Guidance

Numerous publications describe the basic elements of ERA. Many documents have been developed in the context of particular regulatory regimes, so the policy aspects of such documents may not apply to all, or even any, sites in Canada. In Canada, the general framework has been described by CCME (1996-a, 1997-a, 1997-b). This guidance document does not replace these documents. Rather it builds on them to provide a comprehensive framework that considers all aspects of current ERA practice in Canada.

Guidance on ERA from Canadian provinces or territories tends to focus on specific aspects of ERA and not on the overall ERA framework. Such guidance is referred to as appropriate in Sections 2 to 5 of this guidance document in the context of particular technical issues. There are a few cases where provincial guidance is more comprehensive; one is the guidance document on detailed ERA developed in British Columbia (Science Advisory Board for Contaminated Sites in British Columbia [SAB-CS] 2008). Although the policy elements of that guidance document are not always relevant outside of British Columbia, the technical content is relatively detailed and reflects best available practice in ERA. Another relevant provincial guidance document is guidance in Ontario for implementation of risk assessment under the Environmental Protection Act (Ontario Ministry of the Environment, Conservation and Parks [MECP] 2005); the policy elements of that document would also not be relevant outside of Ontario. The province of Québec has also developed guidance on an overall ERA framework (CEAEQ 1998). Finally, although specific to sediments, the Canada-Ontario decision-making framework for assessment of Great Lakes contaminated sediment also covers the key aspects of an overall framework for ERA (EC and MECP 2008).
Numerous guidance documents on ERA have been developed in the United States, in particular by the United States Environmental Protection Agency (US EPA) and the National Research Council (NRC). The basic ERA framework is described in US EPA (1992) and US EPA (1998). Many of the other US EPA documents are specific to certain cases (e.g., Superfund sites), and most of the NRC documents address particular issues in the practice of ERA (e.g., NRC 2009). Practitioners are encouraged to consult the US EPA and NRC websites to evaluate the potential usefulness of these documents and others that will be developed over time. The US EPA has recently compiled a table listing documents relevant to ERA (Appendix C in US EPA 2011).

In addition to guidance provided by government agencies, many books address the process and technical elements of ERA. Two commonly used reference books are Suter (2007), which addresses the generic framework for ERA, and Suter et al. (2000), which focuses on ERA for contaminated sites. Advanced ERA practitioners interested in detailed technical guidance on particular aspects of ERA should refer to the four technical modules on the FCSAP website (Government of Canada, n.d.), the technical appendices of SAB-CS (2008) and Suter et al. (2000).

2 PROBLEM FORMULATION

Problem formulation is the important first step in ERA. It clarifies the nature of issues associated with contamination at a site and how those issues will be addressed. The specific objectives of problem formulation are to:

- Frame the issues, including the goals, context and nature of potential effects.
- Design and plan an approach to assess risks, specifying the tools that will be used and how the results will be evaluated.

2.1 Overview of Problem Formulation

Problem formulation generally entails the following steps:

- Describe the site-management goal(s) and the specific assessment goal of the ERA. For example, if a site-management goal is to reclaim a site as parkland, the assessment goal of the (initial) ERA may be to assess whether current conditions at the site will support the protection goals for parkland.
- Review the regulatory context for the site and the ERA, including applicable legal instruments and policy.
- Review existing site information. This should include, at a minimum, a list of relevant documentation, a site description and a summary of key findings from previous investigations. For some complex ERAs, such a review may warrant a stand-alone chapter or document attached to the problem formulation.
- Select contaminants of concern (COCs) and describe any of their characteristics that are relevant to the ERA (e.g., transport and fate).
• Select receptors of concern that could be affected by contamination and that will be evaluated in the ERA. Receptors can be identified at the level of individual organisms, species, populations, communities or habitats. Importantly, it is usually not feasible (or necessary) to include every possible species in an ERA. Therefore, a subset of candidates are selected as surrogate receptors of concern for particular types of receptors.

• Identify the exposure pathways by which COCs may come into contact with the receptors of concern. Examples of exposure pathways include water and food consumption (for wildlife) and direct contact (for invertebrates).

• Develop a conceptual site model (CSM) that shows the potential links between source of contaminants, exposure pathways and receptors of concern.

• Clarify protection goals and associated acceptable effect levels (AELs). Typically, protection goals and AELs may vary by land use or by receptor (e.g., species at risk are normally afforded organism-level protection, whereas other species are normally afforded population-level protection). Many jurisdictions have adopted policies that will specify protection goals and AELs.

Definitions

A contaminant of concern (COC) is a contaminant that has been selected for evaluation in the ERA.

A surrogate valued ecosystem component (VEC) is a receptor of concern that has been chosen to represent a VEC (e.g., a shrew may be used as a surrogate VEC for insectivorous mammals).

Exposure pathways are the routes of exposure from environmental media (e.g., soil, water, air, sediment) to the receptors of concern. Examples of exposure pathways include ingestion and inhalation.

A conceptual site model (CSM) is a narrative and/or graphical representation of the relationships between contaminant sources, exposure pathways and receptors.

Key Concept

An assessment endpoint describes an attribute of a receptor or receptor group, but does not articulate a desired state for that attribute.
Identify assessment endpoints, which are attributes of receptors (the entities that are to be protected), often with specific spatial and temporal components. An ERA may have one assessment endpoint for a receptor group (e.g., ecological function of the soil invertebrate community), or there may be more than one assessment endpoint for a receptor or group of receptors.

Identify measurement endpoints, which are tools that measure exposure to a receptor, effects on a receptor or changes in attributes of assessment endpoints.

Develop lines of evidence for each assessment endpoint, which specify how measurement endpoints will be used to evaluate potential risks.

Articulate the general strategy for the ERA, including how risk characterization will be conducted, and a sampling and analysis plan (SAP). In some cases, for example for complex ERAs with many components, the SAP may be prepared as a stand-alone document separate from the problem formulation.

The rest of Section 2 explores each of these problem formulation elements in more detail. Although the elements are presented in a linear fashion, in fact most elements need to be developed together using an iterative process. Furthermore, because almost all of the planning for an ERA occurs during problem formulation, this step must fully consider the contents of Sections 3 to 5.

**Definitions**

A protection goal is usually a narrative statement that defines the desirable level of protection for a receptor or receptor group. An acceptable effect level (AEL) operationalizes the protection goal by specifying the magnitude (or rate) of effects that would be acceptable for a specific measurement endpoint or assessment endpoint.

A measurement endpoint is a parameter that measures or describes exposure for, or an effect on, a receptor of concern, or that measures or describes a change in an attribute of an assessment endpoint or its surrogate in response to a stressor to which it is exposed.

A line of evidence is any pairing of exposure and effects measures that provides evidence for the evaluation of a specific assessment endpoint. Typically a line of evidence requires use(s) of one or more measurement endpoints. If the focal point of the line of evidence is an effects measure (e.g., a toxicity test), the paired exposure measure may be quantitative (e.g., contaminant concentrations) or categorical (e.g., on site versus reference).

**Key Concept**

It is helpful to view measurement endpoints as tools, and lines of evidence as the use of those tools in one or more ways.

**Key Concept**

Begin with the end in mind. A proper problem formulation does not simply result in a list of tools to be used for the ERA, but also specifies how the results will be evaluated.
2.2 Site-Management Goals

At a broad level, an ERA is guided by the overall site-management goals. In the context of ERA, a management goal for a contaminated site is the overall planning objective for the site, usually worded as a statement about the desired condition of the ecosystem or its components in the context of future site use. Site-management goals may be relatively generic and stated at a high level (e.g., “maintain a sustainable aquatic community adjacent to a ferry terminal”). In other cases, more specific management goals may be identified, such as:

- determine whether contaminants (COCs) present in the surface soil layer require remediation within the existing provincial and federal regulatory frameworks for a particular land-use category
- determine whether intrusive remediation is warranted at a contaminated wetland adjacent to an airport
- develop a management plan for a Department of National Defence facility, ensuring that the plan will protect a specific federally listed species at risk
- assure that new homeowners in a brownfield redevelopment project will be able to grow the range of plants expected at normal residential properties in the area.

These more specific site-management goals are generally preferable because they provide direction to risk assessors and site managers. Specific site-management goals are often developed through discussion with regulators, site managers, owners and stakeholders. Such dialogue can clarify how the ERA will be used to support risk management and decision making. For example, if there are only two management options for a site, the ERA could be tailored to inform a decision about which option is preferable.

Site-management goals provide the overall framework under which the components of the problem formulation are developed. A site-management goal should not be confused with a protection goal (which is related to the desired level of protection for ecological receptors; see Section 2.3.1), although protection goals are derived in part based on understanding of the site-management goals.

2.2.1 Determining the Broad Assessment Goals

One of the potential pitfalls of ERA in practice is that individual practitioners are prone to applying identical approaches at different sites, even where site-specific considerations (including management issues) require different techniques. Therefore, it is imperative that risk assessors do the following:

- consider the overall purpose of the risk assessment before selecting or interpreting measurement endpoints, tools or techniques
- plan the study design to consider the fundamental underlying questions of interest
• provide output in a format useful to the risk manager for making decisions.

A risk assessment is not a purely scientific endeavour. Rather, it is as a management tool in which the analysis should proceed only to the point at which it meets the information needs of risk managers. Barnthouse (2008) notes that “ERA is best viewed as a bridge between science and management,” rather than a conventional scientific discipline such as chemistry, toxicology or ecology. Management decisions accommodate multiple goals and constraints, with a need to reconcile information collected across disciplines, scales and types of evidence.

To assist in framing management needs, a framework derived by Cormier and Suter (2008) for environmental risk assessment\(^3\) conceptualizes four themes of, as depicted in Figure 2-1.

- **Condition assessment**: Purpose is to detect chemical, physical and biological impairment, through analysis of environmental monitoring data.

- **Causal pathway assessment**: Purpose is to determine proximate causes, identify their sources and, where possible, characterize the causal pathways that connect them.

- **Predictive assessment**: Purpose is to estimate environmental, economic and societal risks and benefits associated with different management alternatives. Acceptability of actions may be determined through evaluating the risks in light of social, economic and legal considerations.

- **Outcome assessment**: Purpose is to evaluate the results of a past management action, through estimation or direct measurement.

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\(^3\) The term *environmental risk assessment* is used to distinguish human and ecological risk assessments from broader environmental assessments that incorporate social, cultural and economic analyses. However, environmental assessments, as defined by Cormier and Suter (2008), include any science-based assessments used to inform environmental management decisions, where such decisions accommodate multiple goals and constraints. The framework depicted in Figure 2-1 applies to all types of environmental assessment, although we have emphasized the application to contaminated sites.
Assessment from Effect to Cause

Problem Detection

Condition Assessment

Causal Pathway Assessment

Problems Resolution

Outcome Assessment

Predictive Assessment

Assessment from Cause to Effect

Environmental Epidemiology

Environmental Management

Figure 2-1 ERA framework as developed by Cormier and Suter (2008)

Source: Cormier and Suter (2008) reproduced in accordance with the Creative Commons Attribution Non-commercial License.
The framework recognizes that all assessment types can potentially resolve an environmental issue or prompt a refined assessment in a subsequent tier of investigation.

These four broad themes of assessments are organized based on two key questions:

- Are we interested in explaining what has already occurred in terms of environmental effects (environmental forensics) or in extrapolating our knowledge to the prediction or optimization of future conditions (environmental management)?

- Are we interested in simply detecting an environmental response (condition assessment) or in assigning or allocating responsibility among sources (causal pathway assessment)?

This framework is useful for focussing the risk assessment objectives, particularly by asking the two key questions above and then organizing the study design and evaluation to answer them. It is also possible to tier an investigation such that resolving issues of causation and prediction is deferred pending the results of a preliminary risk assessment.

This framework is applicable to all ERA scenarios, but is particularly well suited to the assessment of contaminated sites, as discussed below:

- **Condition assessment**: This may determine whether a site is sufficiently contaminated to warrant further assessment, or it may include a biological condition assessment to determine whether there is evidence of impairment from the site. In general, the initial stages of condition assessment rely on chemical characterization, but progress to toxicological and biological tools at more detailed stages of investigation.

- **Causal pathway assessment**: Some level of causal assessment is incorporated into all contaminated site investigations because the preliminary or detailed site investigations identify sources (e.g., areas of potential environmental concern) and contamination pathways, at least at a broad level. However, the importance of causal assessment increases for some contaminated site scenarios, such as where multiple responsible parties contribute to contamination, or where non-contaminant stressors may influence the pattern of observed responses. In these cases, the causal assessment explores in more detail (and with increased emphasis on quantitative methods or mechanistic understanding) the linkage between exposure and effect.

- **Predictive assessment**: The degree of prediction required in a contaminated sites assessment is a function of the range of potential site uses contemplated over time. Where a site is proposed for divestiture, but without a foreseeable change in site use, risk assessments may rely on empirical information from existing site characterization. Conversely, scenarios of significant redevelopment or remediation often trigger the need to model or predict future conditions of contamination and their influence on risk estimates. Changes in site use may result in revised assumptions regarding exposure (e.g., revised calculations of exposure concentrations or doses), effects (e.g., revised toxicity estimates based on changing contaminant fingerprint over time) and risk management alternatives (e.g., administrative controls on site use).
• **Outcome assessment**: In a contaminated sites application, an outcome assessment entails evaluation of multiple “what if” scenarios, with the objective of determining whether risk estimates can be meaningfully influenced by actions taken by the risk manager. For example, where baseline risks are considered to be unacceptable, a remedial options analysis can be undertaken to evaluate the impact of different management alternatives, including monitored natural recovery.

Although the Cormier and Suter (2008) framework is simplified and conceptual, it is possible to frame the core risk assessment needs through consideration of site-specific issues. The four themes are not mutually exclusive, so it is possible to draw elements from multiple themes (as shown in the quadrants) to develop an assessment framework that is appropriately customized to the site context. To refine the broad assessment goals beyond the simple four-theme framework, and to make the framework relevant to site-specific issues, it is helpful to pose questions that inform the selection of tools. For example:

• Are there multiple potentially responsible parties associated with the contamination (industries, site owners, point sources, legally responsible entities)? If yes, consider the role of causation.

• Is there a need to extrapolate results to other parcels or conditions? If yes, consider the importance of predictive tools.

• Is the study intended to detect environmental changes in response to source control or other management actions, such as remediation of contaminated soils up-gradient of a harbour facility, or monitoring of tailings treatment at an abandoned mine site? If yes, consider outcome assessment tools.

• Is the site within a context of significant regional background contamination? If yes, consider the role of causation.

• Is the site large or complex in terms of physical, chemical and biological conditions? If yes, consider tools for extrapolating across space, time, or habitat or substrate type.

• Are the processes affecting contaminant transport, accumulation and toxicity already well understood? If yes, the need for causation assessment or refining predictive tools may be lower.

• Is the study designed to screen or rank priorities for future tiers of study, as opposed to detailed remediation design? If yes, consider initial condition assessment.

• Does the site contamination affect off-site parties or sensitive habitats, as opposed to being a site-specific management issue? If yes, the need for causation assessment or refining predictive tools may be greater.

• Are there known confounding factors to direct assessment of risks, such as physical habitat modifications or mechanical disturbance? If yes, consider the need for causation assessment or refining predictive tools.
• Is there potential for cost savings through the use of an adaptive management approach, and is there adequate time available for such an approach? If yes, consider initial condition assessment to optimize resources, followed by other approaches.

• Does the ERA require an evaluation of stressors that are either not contaminants or that may confound the assessment of a primary contaminant (e.g., biological influence of cattle grazing, regional organic enrichment or invasive species)? If yes, consider the need for causation assessment or refining predictive tools.

• Is the ERA intended to evaluate the relative risks or benefits of alternative management approaches at the site? If yes, consider the need for predictive tools and outcome assessment, with the use of a formal decision analysis framework to guide management decisions.

Predictive assessments that focus on future risk scenarios are common in ERA. Future risks may differ from current risks for many reasons, including:

• Implementation of risk management measures such as remediation or fencing
• Natural attenuation, which may occur due to physical or chemical processes (e.g., dechlorination, burial by settlement of relatively clean material)
• Changing human use of the site, including addition or removal of infrastructure
• Natural ecological succession. For example, if a site is no longer subject to human use, natural ecological processes may result in changes to ecosystem type along a gradient of disturbance level (e.g., from plowed fields associated with agricultural land to a forest type that may become established).

In cases where both current and future risks are estimated, it is possible to estimate the expected change in risks that may occur. This comparative approach can be useful for evaluating the likely effectiveness of risk management measures.

Note that the assessment type is not necessarily static, but rather may progress from one type to another based on feedback from early stages of investigation, as implied by the arrows in Figure 2-1. Site investigations often begin with condition assessments, in which the primary objective is to determine whether the current site conditions are acceptable. Depending on the results, subsequent tiers of analysis may require increased emphasis on causality or prediction of changes over time.

2.3 Regulatory Context
The regulatory context for an ERA is important for determining the scope and identifying technical constraints.

Legal instruments and policies: The problem formulation should acknowledge the various federal and other (e.g., provincial) legal instruments and policies that are applicable for a particular site, and should promote consistency of the ERA with those legal requirements and policies. Examples of relevant federal legislation include the Fisheries Act, the Canadian Environmental Protection
Act, the Species at Risk Act, the Migratory Birds Convention Act, the Canadian Environmental Assessment Act and the Canada National Parks Act. There are numerous other potentially relevant federal and provincial/territorial legal instruments (see, for instance, SAB-CS 2008 for some further discussion). Where a site falls under both provincial/territorial and federal jurisdiction, it is usually necessary to meet the requirements of both agencies, and hence the most stringent of the two jurisdictions’ requirements normally need to be met.

The regulatory context can have direct implications for the technical details of the ERA. For example, the protection goal defined for a species at risk (i.e., a rare or endangered species) may be much different than for a common species. The Species at Risk Act requires that species at risk be protected at an individual level, but an ERA may aim to protect common species at the population level. In other cases, certain aspects of contamination may not be the primary focus of an ERA if they are addressed by other regulations, although cumulative impacts should not be ignored. For example, some discharges of contaminants are permitted under certain regulations. Depending on the goals of the ERA, it may or may not be relevant to explicitly consider the effects of such discharges (e.g., a risk assessment of a federal water lot conducted in the vicinity of a municipal effluent discharge may need to consider the effects of the discharge in order to discriminate between potential sources of impairment). In all cases, ERA practitioners should consider due diligence as well as the legal and policy requirements of the relevant jurisdictions.

Land use: Land use designations are usually important in determining whether or not a terrestrial site is contaminated, because the screening guidelines for a given contaminant often vary by land use. In addition, land use (either designated or actual use) affects an ERA in other important ways. First, policies developed for technical aspects of ERA may be specific to land use. For example, protection goals may be different in a park compared to an industrial property. Second, actual land use at a site may limit the scope of risks (particularly exposure pathways) that need to be considered. For example, if a site does not have (in either current or potential future uses scenarios) any exposed surface soil, many receptor groups will not be present. Third, land use in the areas surrounding a site may also limit the scope of an ERA. For example, if a site exists in the middle of an urban centre, considering large mammal receptors may be unnecessary. Conversely, in a relatively remote setting, a similarly sized parcel of developed land may require consideration of large mammals that inhabit adjacent areas but use the site for food. At the same time, caution must be used in this approach, since land use might impact assessment for a specific valued ecosystem component (VEC) (e.g., terrestrial receptors) but may not impact other relevant VECs (e.g., those related to surface water, sediment or groundwater). The problem
formulation should, as appropriate, highlight relevant implications for the ERA of the current and/or potential future land use of any given site.

2.4 Review of Existing Site Information

Every problem formulation uses existing information about a site as its starting point. Although problem formulation is the first formal stage of risk assessment, from a practical perspective, different sites have varying degrees of baseline site investigation information from which to begin the problem formulation stage. Therefore, the purpose of the review is to summarize pertinent information on contaminant sources and distribution, transport pathways and biological attributes of the site.

The basic information includes:

- documentation: a list of relevant available documents about the site
- site description: location, setting, etc.
- review of previous environmental site assessments and findings (e.g., site chemistry, historical and ongoing contaminant sources, screening guidelines used)
- if applicable, review of risk-related data for the site (in cases where previous risk-related investigations have been conducted, or if the problem-formulation process has been iterative).

For some complex ERAs, the review of existing information may warrant a stand-alone chapter separate from the problem formulation.

Environmental site investigation data often differs greatly in abundance, type and quality. Therefore, it is important to decide whether supplemental site investigation is needed before undertaking formal risk assessment activities such as problem formulation. The potential need will be site-specific, depending on how the ERA will be used to support site-management goals. If major data gaps are identified, risk assessors and site custodians should consider holding off on finalizing the problem formulation until they collect the supplemental data. In other cases, some aspects of a project or spatial components may progress along different timelines.

2.5 Contaminants of Potential Concern

As previously defined, contaminants of concern (COCs) are those contaminants that have been selected for evaluation in the ERA. In some jurisdictions, terms such as contaminants of potential concern (COPCs) or potential contaminants of concern (PCOCs) refer to the initial list of substances considered, whereas the term contaminants of concern (COCs) refers to the final list after the selection process conducted as part of the problem formulation. In other jurisdictions the term COC is not used at all, and the final
chosen based on the phase 1 and phase 2 environmental site assessments and the subsequent failure of parts of the site to meet the appropriate numeric guidelines or standard. For ERAs, this is normally narrowed to those that exceed an ecological component of the numerical guideline or standard. However, some situations exist where an environmental site assessment is not available and the COCs need to be determined from other information. This section reviews the broad categories of sources of COCs that should be considered in an ERA, and then focusses on the COC selection process. Finally, this section reviews the characteristics of COCs that must be understood in order to proceed to subsequent components of the problem formulation.

2.5.1 Sources of COCs
Understanding of the sources of COCs is important for determining likely exposure pathways. Categories of sources of COCs at a site include:

- on-site point sources (e.g., historical spills, ongoing point source effluent discharges)
- on-site non-point sources (e.g., contaminated groundwater, sediments or water; surface water runoff)
- underground artificial conduits such as sewers, pipelines and buried structures that may contribute to contamination
- preferential natural pathways such as fractures in limestone geology that facilitate transport of COCs
- significant off-site sources (including via long-range air transport) that need to be considered for their potential to confound site-related contamination or as contributors to cumulative risks.

Identifying sources of contamination requires a thorough understanding not only of the site itself, but also of the surrounding land use and the location of the site; this is critical for identifying off-site sources in particular. Typically, the relevant information can be summarized from site investigation documents. In fact, it is common for site assessment documents to distinguish areas of a site based on historical use and other factors, and then to identify the specific COCs associated with each area of potential environmental concern (APEC). This level of resolution for COC sources is often relevant to the ERA as well.

2.5.2 Selecting COCs
Selecting COCs is an important early step in the problem-formulation process. The starting point, as per Section 2.2.4, is usually the list generated from site investigation reports, although it is important to confirm whether additional COCs may be relevant before proceeding with the risk assessment. Often, the initial list of contaminants generated by site investigation reports is referred

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list is referred to as the list of COPCs. In this guidance document, the term COC refers to the final list of substances retained for the risk assessment at the end of problem formulation.
to as a preliminary list of COCs or a list of contaminants of potential concern (COPCs), which is reduced to a final list of COCs during problem formulation. Although there may be regulatory requirements to consider all of the COPCs identified during site investigation, the final list of COCs for an ERA may be different for several reasons. The process used to select COCs should be agreed upon with site custodians and regulators as early as possible in the ERA process. COC selection is important for ensuring that the ERA does not miss any important contaminants while also preventing needless analysis of contaminants that do not warrant evaluation.

This section focusses on identifying the key considerations that should guide COC selection. Further discussion of many of these issues can be found in CCME (2016) and SAB-CS (2008).

- **Applicable guidelines:** Most jurisdictions within Canada have environmental quality guidelines for typical COCs. When the federal government is the only stakeholder, Canadian Environmental Quality Guidelines (CEQGs) or Federal Environmental Quality Guidelines should be used for screening.

  Within a set of guidelines, there may be multiple options according to land use, water use, soil texture, transport/exposure pathways or other factors. In such cases, rationale is needed for determining exactly which guidelines are applicable and which are not. For cases where guidelines are based on consideration of both ecological and human health components, it may be reasonable to exclude the human health component if rationale is provided.

  Also, if there are site data from multiple media, it may be appropriate for one medium to take precedence. For example, if there is a soil guideline for the soil-to-groundwater pathway as well as a rigorous data set for groundwater directly, it may be appropriate to not screen data using the soil guideline. Consult the jurisdiction in question to determine the applicable guidelines to apply to a screening-level risk assessment.

- **Substances for which there are no guidelines:** In some cases, site-related substances may be present at elevated concentrations but may not be addressed by guidelines within the jurisdiction or CEQGs. In such cases, the risk assessor must decide whether to include the substance in the ERA or use alternative methods of screening. Alternative screening methods may include adopting guidelines from other jurisdictions. In such cases, the risk assessor should understand the policy within the jurisdiction in question regarding alternative guidelines and the level of protection inherent in such guidelines. It is important in such cases to consult the jurisdiction in question early within the risk assessment process to ensure that the jurisdictional authority will support the guidelines chosen.

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5 The term guidelines is used loosely in this section to include numeric environmental quality guidelines, criteria, standards, or any other regulatory or policy benchmark that may be used for COC screening.
Another approach that has been used is screening COCs in one media using guidelines for another media (e.g., application of an uncertainty factor to surface water data to facilitate comparison to groundwater guidelines). This approach is not supported by all jurisdictions and is not recommended unless defensible rationale is provided. If no screening guidelines exist for a substance, the risk assessor should question why that is the case. Often, environmental quality guidelines may not have been implemented due to high uncertainty in the available scientific data. Consideration of substances should be extended beyond conventional chemical stressors to macronutrients (e.g., phosphorus), dissolved oxygen, or other important indicators of habitat quality and quantity that could be potentially important contributors to total risk at a site. If a substance has no guidelines, has been identified in the phase 1 environmental site assessment as a potential COC, and has been measured at above reasonable detection limited at the site, then in most cases this substance should be carried through the ERA. The report should explain how the ERA handled these substances and include an uncertainty assessment. If such substances are excluded, then the risk assessors should provide solid rationale for doing so. Likewise, rationale should be given if a substance was considered in the ERA but not in the site investigation.

- **Background concentrations**: In some cases, background concentrations of a substance may exceed generic guidelines. In such cases, it may be appropriate to compare to background concentrations rather than to guidelines. This may include comparison to regional background data or to more localized data. For example, metal mines are typically located in areas of naturally elevated metal concentrations. Where background concentrations are potentially elevated, reference conditions should be defined carefully, or gradient-based sampling methods should extend far enough off site to establish suitable local background concentrations of COCs. Since jurisdictions handle background concentrations in ERAs differently, the risk assessor should consult the jurisdictional authority regarding the requirements for use of background conditions within a risk assessment.

- **Food chain uptake**: If environmental quality guidelines are based on protecting lower-level receptors (e.g., invertebrates), it is important to determine whether and how to screen COCs for evaluation of higher-level receptors via food chain uptake. For example, Canadian interim sediment quality guidelines (CCME 1999-a) are not designed to protect higher trophic levels against potentially bioaccumulative substances. Canadian tissue residue guidelines (TRGs) for protection of wildlife consumers of aquatic biota are more appropriate (CCME 1999-b), but do not cover all relevant substances. Some substances are known to be bioaccumulative or biomagnifying (or are named as such in policy or regulatory documents), but for other substances the need to consider food chain uptake may depend on site-specific characteristics. Screening for food chain pathways usually focusses on receptors with a definable home range size, so one consideration in screening the risk.

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**Key Concept**

Off-site data can provide important context in cases where substances are present at naturally elevated concentrations that exceed guidelines.
for those receptors is whether to screen based on individual samples or summary statistics for an area (this issue is discussed in more detail below).

- **Using statistics:** Most risk assessors conduct a preliminary screen of data using maximum concentrations for a COC in a particular medium. However, if the maximum concentration exceeds a guideline, and the receptor is mobile, risk assessors should consider using summary statistics (e.g., for the terrestrial environment, it might be useful to compare the 95 per cent upper confidence limit of the mean [UCLM] or the 90th percentile of the concentrations to the guideline). Risk assessors may consider using summary statistics on a case-by-case basis, in light of factors such as the number of samples, spacing of samples, seasonality or timing of samples (particularly for water), and the characteristics of receptors. Rationale should be provided for any decisions made. As a default, for immobile receptors (e.g., plants, small invertebrates) the maximum concentration for a COC should be used as a conservative starting point. For mobile receptors exposed to an area characterized by multiple samples, the maximum could be used as a conservative starting point if there are fewer than 10 samples. For sample sizes of 10 or more, the jurisdictional authority should be consulted regarding use of statistics within the jurisdiction. Generally, it is considered that the 95th UCLM and the 90th percentile are reasonable estimates of risk for large sample sizes involving mobile receptors when the area in question is smaller than the home range and provided that there is accompanying rationale. However, risk assessors must ensure that the statistic is supported by the jurisdiction for the particular ecological environment. In the case of soil, practitioners should consult the local jurisdiction for use of statistics in characterizing exposure to contaminated soil. CCME provides some general guidance for characterizing a volume of contaminated material using statistics (CCME 2016).

These concepts provide some guidance on characterizing an area of contamination to which a given receptor may be exposed. In the case of water, the temporal nature of the data should be considered. If data were collected during two separate events, it may be appropriate to use summary statistics from each event separately. Finally, summary statistics that are based on all data may require consideration of data points where contaminant concentrations are below detection limits. In such cases, risk assessors should provide clear rationale, including statistical rationale as appropriate, for methods used to deal with those data points (see, for instance, CCME [2012] for additional guidance).
• **Sampling depths:** For soil and sediment (and less commonly, water\(^6\)), determining the applicable depth is not always straightforward. Jurisdictions may have policies that standardize sampling depths relative to various pathways and receptors. The depth of surface soils or sediments may be standardized for many sites (e.g., default value for rooting depth of plants in soil based on policy determination), but exceptions can be expected. For example, the relevant surface soil horizon may be deeper where tap roots are present. Alternatively, if a site lacks deep-rooting plants (or has a planned future use that excludes them), surface soil depth ranges could be shallower. As another example, some COCs or receptors may be associated only with the humic soil layer and not with the underlying inorganic soil layer. In that case, the depth used for screening may not be a fixed depth, but may vary depending on the thickness of the humic layer.

As a default, if there is no site-specific information available to define the depth of the surface soil layer, all data in the top 1.5 metres for soil should be used to screen for surface soil exposure. This approach is consistent with the Canada-wide standard for petroleum hydrocarbons in soil (CCME 2008-a). For sediment, as a default, all available data in the top 1 metre should be used to screen for surface sediment exposure. However, caution must be taken in applying these default depths to ensure that depth increments are appropriately capturing exposure and not diluting exposure through too large a sampling increment (see, for instance, Section 3). For example, in aquatic environments, sedimentation rates will affect sampling depths, and the use of deeper sediments may dilute effects seen in the surficial sediments. Similarly, airborne soil contaminants may affect only the first few centimeters of the soil profile and have little influence on the top 1 metre as a whole. In all cases, risk assessors should provide rationale and consult with the jurisdictional authority to ensure consistency with governing policies.

The depth of soils and sediments considered for screening COCs may not be the same depth that is considered during exposure assessment for each receptor group. Section 3 elaborates on consideration of soil and sediment depth during exposure assessment.

For some ERAs, soil or sediment at depth will be explicitly considered in the ERA under two scenarios:

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\(^6\) Depth of surface water sampling can be important for lakes, for example if there is stratification.
(a) there is a plan or a possibility for that soil or sediment to become exposed (e.g., through removal or erosion)

(b) there is a possibility that contaminants at depth will be mobilized and reach receptors (e.g., via groundwater transport, burrowing animals, transfer through plant roots).

- **Sampling density and coverage:** COC screening for an ERA usually begins after site investigation has characterized the horizontal and vertical extent of contamination at a reasonable resolution. However, the density and coverage (horizontal, vertical) of sampling will vary by site, and in some cases the ERA is initiated before site assessment activities are complete. Ideally, risk assessors get involved in site investigation planning early so that sampling density is sufficient for screening and characterizing exposure later during the ERA. Specifically, sampling density must be sufficient, and the samples must be representative of site conditions relevant to the ERA. If data are limited, risk assessors should consider whether sufficient data are available to warrant excluding certain COCs based on existing samples not exceeding guidelines. Jurisdictions may already have recommendations on sampling densities for various scenarios. Practitioners should consult the jurisdictional authority and current CCME sampling guidance when considering sampling density and coverage (CCME 2016).

- **Data quality:** Another important consideration when evaluating existing data is the quality of those data, particularly with respect to analytical detection limits. If many or all existing data do not have detection limits that are lower than relevant guidelines, then the utility of the data for ruling out COCs is diminished. Practitioners should ensure that SAPs specify data quality objectives (DQOs) that meet the needs of the ERA and are consistent with current jurisdictional requirements. CCME provides some sampling guidance regarding data quality. See, for example, CCME 2016 and US EPA 2006 for further discussion. In some instances, higher-resolution methods or specific extraction procedures may be required to achieve the accuracy and precision requirements of the ERA.

- **Form of contaminants:** Risk assessors should be diligent in identifying the relevant form of contaminants and specifying how contaminants are identified. The exposure pathways identified in the ERA will in part determine the relevant form(s) of the contaminant (e.g., total versus dissolved, oxidation state or adjustment for environmental conditions such as pH). The type of contamination and the availability of toxicological data may determine the form of a contaminant considered in the ERA. For example, in the case of polychlorinated biphenyl (PCB), it may be possible to conduct the ERA based on total PCBs, on one or more individual congeners, on selected homologs, or on Aroclor mixtures. Alternatively, or in addition, the dioxin-like PCB congeners may be evaluated using the toxic equivalents (TEQ) model whereby the combined effects of all dioxin-like compounds are evaluated together. The appropriateness of each approach depends on the receptor type, the chemical signature present at the site and the availability of matched effects data for each quantitation method.
2.5.3 Characteristics of COCs

The characteristics of COCs are important for identifying receptors, exposure pathways and endpoints in the problem formulation. Characteristics can be separated into two types: transport and fate (including bioavailability), and effects.

Transport and fate: The transport and fate characteristics of a COC determine how the contaminant will move from source(s) and partition into various environmental media such as soil, water, sediment and biota. The transport and fate characteristics help determine which receptors and exposure pathways are relevant in the ERA. For example, sediment benthic organisms may be relevant for contaminants that are transported from an upland site to the aquatic environment via groundwater. The transport and fate description is usually qualitative, but when possible quantitative metrics should also be used. For example, for organic compounds the octanol-water partition coefficient \( (K_{OW}) \) provides insight into potential for bioaccumulation and biomagnification (e.g., substances with a high \( K_{OW} \) tend to partition into organic matter). The CCME national classification system for contaminated sites (CCME 2008-b) uses a threshold \( \log(K_{OW}) \) of 4, above which exposure via food chain transfer is considered more likely. The Persistence and Bioaccumulation Regulations under the Canadian Environmental Protection Act (Government of Canada 2000) use a \( \log(K_{OW}) \) of 5 or higher as one method of classifying a contaminant as bioaccumulative. Other methods rely on magnitude of bioaccumulation factors (BAFs) or bioconcentration factors (BCFs). Whenever pathways are excluded from consideration on the basis of the transport and fate properties of a COC, rationale is essential.

Bioavailability is an important factor influencing the degree to which COCs will partition from abiotic media into tissues. A COC that is bound with soil particles may pass through the gut of a receptor, whereas a COC in dissolved form in water may be much more bioavailable.

Consideration of transport and fate characteristics should include potential degradation processes that are relevant to the substance. Some contaminants degrade into breakdown products that may be as or more toxic than the parent compounds. In some cases (e.g., polycyclic aromatic hydrocarbon [PAH] contamination in aquatic life), the metabolism of the parent product is highly receptor specific.

The transport and fate of COCs will depend on their physical and chemical properties, and on the specific characteristics of the environmental media at the site. For this reason, conventional parameters collected during chemistry programs are important; examples include soil pH, water hardness, organic content of soil or sediment, and sediment grain size. These parameters affect the fate of COCs not only among abiotic media but also between abiotic and biotic media (e.g., by influencing bioavailability).

Effects: The review of effects characteristics of a COC emphasizes the types of organisms that may be affected by the COC and the relevant mechanisms of action. It is seldom necessary for the
information to be compiled for every specific receptor. Rather, broad characteristics relevant to key receptor groups are usually adequate for the purposes of the problem formulation. In some cases, the concentrations or doses associated with particular adverse effects may be specified, helping to identify the effects endpoints that are expected to be most sensitive and therefore the potential candidates for the formal hazard assessment. The effects characteristics provide important information for selection of receptors (e.g., which receptor groups are known to be sensitive to the COC) and for the selection of endpoints (e.g., those known to be caused by the COC and that are relevant to the ERA). While it is important to understand the basic environmental fate and toxicity of COCs at the problem formulation stage, more comprehensive reviews of effects literature are typically conducted as part of the hazard assessment during the ERA (e.g., if needed for derivation of a dose-response relationship).

Sites can vary greatly in their degree and nature of contamination. As the nature of contamination changes from single COCs to several COCs to complex COC mixtures, so do the challenges of understanding potential effects. When multiple contaminants are present in an exposure medium, they may interact to produce antagonistic, additive or synergistic effects. Ultimately, not accounting for these interactions, or applying invalid models to account for such interactions, could lead to erroneous risk conclusions. Some tools used in hazard assessment are better suited than others to deal with contaminant mixtures (see Section 4) or specific media that are contaminated. Some contaminant interactions have been well characterized. Examples include the biotic ligand model (BLM) for metals (Di Toro et al. 2001; Paquin et al. 2003), the ΣPAH model for PAHs (e.g., Ozretich et al. 2000; Swartz et al. 1995) and TEQ approaches for dioxin-like effects (e.g., CCME 2002).

Understanding contaminant interactions in detail at the problem formulation stage is not usually warranted, and may not be warranted at all (even during hazard assessment) depending on the scope of the ERA and the tools used. At a minimum, risk assessors should attempt to identify potentially important interactions when documenting the modes of action of COCs during problem formulation (e.g., Menzie et al. 2009). The risk assessor can then determine the best approach to integrating that information into the ERA.

### 2.6 Valued Ecosystem Components

This section contains the technical guidance for receptor selection for use in ERA. Specifically, the section provides guidance for determining which types of valued ecosystem components (VECs) should be considered at a site, and for identifying appropriate surrogate receptors of concern as representatives of those VECs.

#### Key Concept

For wildlife receptors, receptors of concern can be used in the ERA as surrogate VECs to represent risks to a VEC, such as a group of receptors with common characteristics (e.g., small omnivorous mammals, piscivorous birds).
For the purposes of an ERA, VECs are the components of the ecosystem in question that the risk assessor has identified as those the ERA should be designed to protect. They are identified as such through having one or more of the following qualities:

- intrinsic ecological significance
- importance to human populations
- economic or social value
- ability to serve as a baseline from which effects of changes can be evaluated.

A VEC can be any non-human individual organism, species, population, community, habitat or ecosystem. A VEC need not exist at the site in its current state, but should be capable of being there in the absence of contamination or other anthropogenic effects.

As previously defined, for ERA a receptor of concern is any non-human individual, species, population or community that is potentially exposed to COCs. A receptor of concern can be identified as a subset of the VECs at the site. A receptor of concern may be the same as a VEC, but it can also be a surrogate for the VEC or a useful element in a line of evidence for determining effects to a VEC. For example, a VEC may be a wetland complex. Several receptors of concern may be selected to evaluate key attributes of this wetland: for example, specific species at risk, diverse aquatic plant community, nutrient processing and water retention. These receptors would be evaluated to determine the potential direct and indirect risk of contaminants to the VEC.

The level of biological organization at which a VEC is defined varies. In the case of lower trophic levels, the community is often identified as the VEC (e.g., zooplankton community, benthic invertebrate community). In the case of higher trophic levels, the VEC may be an individual receptor of concern or species (e.g., mink, eagle). In the latter case, a particular species may be selected for direct assessment of that species or for use as a representative (or surrogate) for similar organisms. As described in this section, where a surrogate organism is used, the risk assessor should articulate the groups of organisms that the receptor of concern is intended to represent. In most cases, the groups are selected on the basis of functional feeding groups (e.g., small omnivorous mammals, piscivorous birds, forage fish) rather than on taxonomic linkages. In selecting a specific surrogate receptor of concern, the risk assessor considers the degree to which the surrogate may be assumed to be protective of related species or the VEC on the basis of contaminant sensitivity and life history considerations (diet, foraging range, etc.). This section provides guidance on these issues and consists of the following subsections:

- information compilation
- identification of receptor types
- criteria for selection of surrogate (representative) receptors of concern for the VECs
- linking receptors of concern to problem formulation.
2.6.1 Compiling Information
Consideration of potential receptors is site-specific and begins with understanding the ecological attributes of the site. The risk assessor should start by compiling information such as:

- general site characteristics (e.g., forest cover, roads, watershed, wetland areas)
- regional and local habitat surveys and land use classifications
- records of environmental conditions and parameters measured on site that may be relevant to any level of biological organization
- species inventories (flora and fauna) and species range maps
- species that are at risk (i.e., listed as rare or endangered) or have some similar status (consult the Species at Risk Act and provincial lists); identifying the possibility of species at risk at this early stage provides an opportunity for specific consideration of these species in the ERA
- other jurisdictional lists of suggested or required VECs (e.g., “Paramètres d'exposition chez les mammifères” and “Paramètres d'exposition chez les oiseaux” in Québec [CEAEQ 1999-a, 1999-b] and Rationale for the Development of Generic Soil and Groundwater Standards for Use at Contaminated Sites in Ontario [MECP 2011])
- information from local experts and residents of the area or surrounding properties
- potential presence of domestic animals (livestock, cats, dogs) that may warrant a particular level of protection (e.g., protection of individual organisms) or consideration of particular endpoints not usually considered for wildlife (e.g., cancer).

If site information is limited or simply not available at this point, practitioners should consider conducting a site visit with a qualified professional to obtain site information (e.g., basic site characteristics, habitat types represented and receptors common to the site). Even if information is available, a site visit can be effective in confirming existing information and providing a better basis for identifying receptors. Evaluation of a site should take into account seasonality, as some potential receptors may use the site for only a portion of their life cycle. Methods for site-specific surveys for purposes of receptor identification are usually qualitative, but may also be quantitative.7

Key Concept
Rationale should be provided to support inclusion and exclusion of receptor types in an ERA. Table 2-3 and Table 2-4 are recommended for this purpose.

Key Concept
Habitat surveys by wildlife biologists can help risk assessors to identify relevant receptors of concern.

7 For example, the U.S. Fish and Wildlife Service has developed Habitat Suitability Index models for fish and wildlife (see National Wetlands Research Center 2015).
2.6.2 Identifying Receptor Types

Numerous types of receptors are relevant to ERA. Depending on circumstances, receptors of concern can represent a VEC or a surrogate VEC (see Table 2-1 for aquatic ecosystems and Table 2-2 for terrestrial ecosystems). For higher trophic levels, surrogate VECs are often particular species (e.g., birds and mammals). For lower trophic levels, VECs are often communities.

During problem formulation, the practitioner must initially consider all receptor types that could be included in the ERA and should then provide rationale for why particular receptor types are included in or excluded from the ERA. Table 2-3 and Table 2-4 are recommended templates for this purpose, for aquatic and terrestrial ecosystems respectively. Determining the VECs that drive the risk assessment will be a key component to determining receptor types to be included in the ERA. Once receptor types are selected, surrogate VECs should be selected for each receptor type.

Selection of receptors of concern should be based on all the information compiled about the site, and should consider:

- representation from the various trophic levels, habitats, feeding guilds and environments that are appropriate for the site
- receptors that can be found off site in adjacent properties, but that use the subject site or could be affected by on-site contamination
- receptors that are expected to be present during particular times or seasons
- receptors that are expected to be present under future scenarios or land use, if relevant for the ERA.

**Definition**

A feeding guild is a group of organisms that use the same ecological resource in a similar way for feeding (e.g., insectivores, granivores, detritivores, carnivores).
Table 2-1: Types of receptors and example surrogates for aquatic ecosystems

<table>
<thead>
<tr>
<th>Aquatic receptor group</th>
<th>Aquatic receptor type</th>
<th>Example receptors for surrogate(^2,3) VECs for aquatic ecosystems</th>
<th>Marine</th>
<th>Freshwater</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Primary producer</strong></td>
<td>phytoplankton</td>
<td>phytoplankton community, seaweed species, plant or algae community</td>
<td>phytoplankton community</td>
<td>periphyton community, algal species, aquatic plant community</td>
</tr>
<tr>
<td></td>
<td>periphyton plants and algae</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Pelagic invertebrate</strong></td>
<td>zooplankton</td>
<td>zooplankton community, shrimp, jellyfish</td>
<td>zooplankton community</td>
<td>shrimp</td>
</tr>
<tr>
<td></td>
<td>others</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Benthic invertebrate</strong></td>
<td>epifauna</td>
<td>mussel, crab, benthos community</td>
<td></td>
<td>crayfish, benthos community, bivalve, benthos community</td>
</tr>
<tr>
<td></td>
<td>infauna</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Fish</strong></td>
<td>benthivorous</td>
<td>stickleback, sculpin, herring, flatfish</td>
<td>salmonid (e.g. kokanee)</td>
<td>salmonid</td>
</tr>
<tr>
<td></td>
<td>planktivorous</td>
<td>minnow</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>piscivorous</td>
<td>salmonid</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Mammal</strong></td>
<td>herbivorous</td>
<td>seal, otter, racoon,* bear*</td>
<td>muskrat,* beaver, moose, mink,* otter*</td>
<td>racoon,* bear*</td>
</tr>
<tr>
<td></td>
<td>piscivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>omnivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Bird</strong></td>
<td>hermivorous</td>
<td>goose,* brant, shorebird, diving duck, grebe, cormorant, heron,* eagle, kingfisher*</td>
<td>goose*</td>
<td>shorebird, swallow, grebe, loon, merganser, heron,* osprey, eagle, kingfisher*</td>
</tr>
<tr>
<td></td>
<td>insectivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>piscivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>omnivorous</td>
<td>dabbling duck</td>
<td></td>
<td>dabbling duck,* diving duck*</td>
</tr>
<tr>
<td><strong>Amphibian</strong></td>
<td>carnivorous</td>
<td>frog, toad, salamander</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Reptile</strong></td>
<td>omnivorous</td>
<td>turtle</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Notes:**

1. Receptor types in lower trophic levels are classified by habitat, whereas those in higher levels are classified by feeding guild.

2. Examples of surrogate VECs that are commonly used to represent the receptor types; note that more than one surrogate VEC can be selected for a given receptor type.

3. Surrogates are not always needed, particularly for lower trophic levels where the receptor of concern is often defined at the community level. In this table, lower trophic level communities are listed to clarify what is typically evaluated, but the communities are not surrogates; rather, they are the receptors of interest.

4. A shorebird describes a number of bird species found frequently on beaches, coastlines and inland shores, though they are not confined to these areas. These birds feed largely on benthic organisms or insects located in sediments near the shallow waters or waterlines.

* Receptors that are recommended in the province of Québec. Refer to CEAEQ (1999-a; 1999-b) for comprehensive species-specific lists.
### Table 2-2: Types of receptors and example surrogates for terrestrial ecosystems

<table>
<thead>
<tr>
<th>Terrestrial receptor group</th>
<th>Terrestrial receptor type¹</th>
<th>Example receptors for surrogate²,³ surrogates for terrestrial ecosystems</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary producer</td>
<td>moss, grass, shrub, tree,</td>
<td>plant species, plant community</td>
</tr>
<tr>
<td></td>
<td>forb</td>
<td></td>
</tr>
<tr>
<td>Invertebrate</td>
<td>microorganism community</td>
<td>activity, diversity, nutrient cycling, energy cycling</td>
</tr>
<tr>
<td></td>
<td>ground-dwelling aerial</td>
<td>invertebrate community, particular species (earthworm, springtail,</td>
</tr>
<tr>
<td></td>
<td></td>
<td>beetle)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>dragonfly</td>
</tr>
<tr>
<td>Mammal</td>
<td>herbivorous</td>
<td>vole,* mouse,* squirrel,* hare, cattle, sheep, deer,* caribou</td>
</tr>
<tr>
<td></td>
<td>insectivorous</td>
<td>shrew,* mole,* bat</td>
</tr>
<tr>
<td></td>
<td>carnivorous</td>
<td>marten, weasel,* domestic cat, domestic dog, coyote,* bobcat</td>
</tr>
<tr>
<td></td>
<td>omnivorous</td>
<td>fox,* skunk,* raccoon,* bear*</td>
</tr>
<tr>
<td>Bird</td>
<td>herbivorous</td>
<td>Canada goose</td>
</tr>
<tr>
<td></td>
<td>insectivorous</td>
<td>warbler, flycatcher, swallow</td>
</tr>
<tr>
<td></td>
<td>carnivorous</td>
<td>owl, hawk,* falcon</td>
</tr>
<tr>
<td></td>
<td>omnivorous</td>
<td>blackbird, sparrow,* crow,* grouse,* chickadee,* robin*</td>
</tr>
<tr>
<td>Amphibian</td>
<td>carnivorous</td>
<td>frog, toad, salamander</td>
</tr>
<tr>
<td>Reptile</td>
<td>carnivorous</td>
<td>snake, lizard</td>
</tr>
</tbody>
</table>

**Notes:**

1. Receptor types in lower trophic levels are classified by habitat, whereas those in higher levels are classified by feeding guild.
2. Examples of surrogate VECs that are commonly used to represent the receptor types; note that more than one surrogate VEC can be selected for a given receptor type.
3. Surrogates are not always needed, particularly for lower trophic levels where the receptor of concern is often defined at the community level. In this table, lower trophic level communities are listed to clarify what is typically evaluated, but the communities are not surrogates; rather, they are the receptors of interest.

* Receptors that are recommended in the province of Quèbec. Refer to CEAEQ (1999-a; 1999-b) for comprehensive species-specific lists.

See MECP (2011) for a list of birds and mammals that were used to develop generic site condition standards in Ontario.
Table 2-3: Template for receptor of concern selection and rationale in aquatic ecosystems

<table>
<thead>
<tr>
<th>Aquatic receptor group</th>
<th>Aquatic receptor type&lt;sup&gt;1&lt;/sup&gt;</th>
<th>Included in ERA? (yes/no)</th>
<th>Rationale&lt;sup&gt;2&lt;/sup&gt;</th>
<th>Surrogate VEC&lt;sup&gt;3&lt;/sup&gt; (if applicable)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary producer</td>
<td>phytoplankton</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>periphyton</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>macrophyte</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pelagic invertebrate</td>
<td>zooplankton</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>others</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Benthic invertebrate</td>
<td>epifauna</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>infauna</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fish</td>
<td>benthivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>planktivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>piscivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mammal</td>
<td>herbivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>piscivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>omnivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bird</td>
<td>herbivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>insectivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>piscivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>omnivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Amphibian</td>
<td>carnivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reptile</td>
<td>omnivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Notes:

<sup>1</sup> Each receptor type should be represented in an ERA if relevant to the site.

<sup>2</sup> A rationale must be provided whether the receptor type is being represented or not.

<sup>3</sup> A surrogate VEC is a receptor of concern that is used to represent a VEC. Surrogates are usually identified for fish and wildlife, but less often for lower trophic levels where the VEC is often defined at the community level. Note that more than one surrogate receptor of concern can be selected for a given receptor type.
Table 2-4: Template for receptor of concern selection and rationale in terrestrial ecosystems

<table>
<thead>
<tr>
<th>Terrestrial receptor group</th>
<th>Terrestrial receptor type¹</th>
<th>Included in ERA? (yes/no)</th>
<th>Rationale²</th>
<th>Surrogate VEC³ (if applicable)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary producer</td>
<td>moss, grass, shrub, tree, forb</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Invertebrate</td>
<td>ground-dwelling aerial</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mammal</td>
<td>herbivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>insectivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>carnivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>omnivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bird</td>
<td>herbivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>insectivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>carnivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>omnivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Amphibian</td>
<td>carnivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reptile</td>
<td>carnivorous</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Notes:
¹ Each receptor type should be represented in an ERA if relevant to the site.
² A rationale must be provided whether the receptor type is being represented or not.
³ A surrogate VEC is a receptor of concern that is used to represent a VEC. Surrogates are usually identified for wildlife, but less often for lower trophic levels where the VEC is often defined at the community level. Note that more than one surrogate receptor of concern can be selected for a given receptor type.
2.6.3 Criteria for Selecting Receptors of Concern for Surrogate VECs

Based on the information review, numerous possible surrogate VECs could be selected for each receptor type. It is often appropriate to include multiple surrogates for each receptor type due to variability among species. However, assessing the ecological risks to an exhaustive list of potential receptors of concern is generally neither practical nor necessary. The following criteria should be used to select the appropriate types of receptors and representative surrogates:

Ecological relevance: An ecologically “relevant” organism is one that is an appropriate indicator of actual or potential exposures given the environmental conditions germane to the assessment. An ecologically relevant organism should be expected to be found at a site under reasonably foreseeable conditions (e.g., an Arctic fox at a site in the Arctic), whereas an ecologically irrelevant organism is one that would not be expected to be found at a site under normal circumstances (e.g., a wolf at a small urban site). An important distinction to be made is that an organism need not be actually observed to be considered ecologically relevant. If contamination is sufficiently great that the organism is extirpated, or if an organism is sufficiently secretive, there may be little or no evidence of its presence. A common error made in problem formulation is to assume that local absence of an organism equates with lack of ecological relevance. It is usual practice to select receptors of concern that represent key functional groups that are expected to be exposed to the COCs on site, or that would be expected to be present at the site in the absence of contamination. In addition, keystone species that are important to ecosystem stability may be preferentially selected as receptors of concern.

Degree and mechanism of exposure to the COCs on site: A number of factors have the potential to affect the degree to which receptors of concern are exposed to the COCs on the site, including:

- status of the receptor of concern (e.g., life stage, migratory versus resident)
- how the receptor of concern uses the site (e.g., feeding guild, feeding behaviour, metabolism)
- how much and how often the receptor of concern uses the site (e.g., home range size, habitat suitability, off-site habitat characteristics)
- number and types of exposure pathways (e.g., environmental media, indirect or direct contact or consumption, bioaccumulation and biomagnification processes).

It is therefore important to consult the life history and background information of the receptor of concern, consider the intended use of a site in terms of its influence on habitat quality and availability, and understand which exposure pathways are relevant. Individual receptors may be exposed to COCs through a number of pathways, all of which should be identified during problem formulation. For receptor selection, information on the relative importance of these exposure pathways is critical. For example, if groundwater flow to aquatic life is an important fate pathway, this may indicate that intertidal receptors (e.g., benthos, mussels, kelp) would be more appropriate than finfish as receptors of concern. Exposure pathways are considered “open,” “operable” or “complete” if a COC is present and there is a route of exposure by which a receptor of concern
comes into contact with the COC. A common error in risk assessment is to confuse the distinction between a pathway that is operable but with low exposure concentrations and a pathway that is inoperable due to lack of plausible transport pathway.

Relative sensitivity to the COCs: It is customary to include species or other receptor types that are relatively sensitive to the COCs if such information is known. For example, some birds are known to be sensitive to certain pesticides due to effects on eggshell thinning, some fish are known to be sensitive to selenium based on reproductive toxicity endpoints, and mink are known to be sensitive to PCBs and mercury. The principle for selecting a sensitive species is that demonstrating lack of harm for a sensitive organism is considered an indication of protection for the less sensitive taxa in the same functional group. However, selecting receptors of concern based solely on sensitivity considerations is questionable. Sensitivity must be considered in terms of both the magnitude at which responses are observed and the type of response elicited. In addition, sensitivity may occur for particular life stages, and therefore the mechanisms by which site-related COCs could affect that life stage must be considered.

Relative importance from a conservation perspective: If rare, endangered or threatened species (i.e., species at risk) or habitats are confirmed to be present, these species must be considered as potential receptors of concern. They should also be included if they are likely to be present in the future (based on information regarding geographic distribution, habitat preferences and site-specific habitat availability).

Relative social, economic or cultural importance: Any particular species or group that is of special importance would typically be included as a VEC and would be included in the receptor of concern selection. These include domestic pets, livestock, species of significance to Indigenous communities, and species of commercial or recreational importance. Because of their importance as VECs, such receptors may be subject to a different level of protection than other receptors of concern.

Availability of ecotoxicological and life history data: Where effects data will be literature-based, receptors of concern for which ecotoxicological data are readily available are preferentially selected (see Section 4 regarding sources). Otherwise, the ability of an ERA to assess effects on the receptor of concern may be reduced. The benefit of selecting highly-specific receptors of concern is offset where data related to toxicity thresholds or exposure information is limited.

Availability of appropriate measurement endpoints: It is important to assess receptors of concern at an ecological scale that is relevant to management goals for the site and to select measurement endpoints that are aligned with those goals. The wetland ecosystem is an example of an ecosystem-level receptor where the measure of effect reflects an ecosystem-level process such as nutrient cycling or primary productivity. The benthic invertebrate community is an example of a community-level receptor where the measure of effect is a community-level attribute such as species diversity. An additional consideration related to measurement endpoints is the ability to distinguish effects from natural variation. For example, abundance of benthic organisms is often highly variable, particularly where habitat and substrate conditions vary; in these circumstances,
the practitioner must consider the statistical and practical (financial) constraints to detection of site-related impacts.

In addition to the criteria described above, rationale for selecting surrogate VECs may be based on logistical considerations or other tools such as a site visit and habitat assessment by a qualified biologist. Local expertise and traditional knowledge may also be useful in identifying appropriate receptors of concern.

2.6.4 Carrying Receptors of Concern Forward in the Problem Formulation
A list of receptors of concern must be carried forward and linked to the VECs, surrogate VECs and other elements of problem formulation. This occurs in at least two ways, as explained in the subsequent sections of this guidance. First, receptors of concern are a component of a conceptual site model (CSM) and are linked to sources of COCs via exposure pathways. Second, specific attributes of receptors of concern are identified to formulate the assessment endpoints for the ERA.

A table can be useful for summarizing the receptors of concern that are carried forward. As recommended earlier, templates can help guide the risk assessor during selection of surrogates that are used to represent some types of receptors of concern (see Table 2-3 for aquatic ecosystems and Table 2-4 for terrestrial ecosystems). Rationale should be provided even for receptor types that are not carried forward in the ERA.

In some cases, the process of selecting receptors of concern may not be completed during the problem formulation. Instead, the process may be presented in conceptual form pending the results of more detailed study. For example, for remote sites it may be more efficient for the problem formulation to focus on selecting broad receptor groups and types only, deferring selection of specific surrogates until a wildlife biologist visits the site during an ERA field program. This approach may be particularly appropriate for receptors at higher trophic levels.

2.7 Exposure Pathways
This section provides guidance on identifying exposure pathways linking contaminant sources to receptors of concern. Identifying exposure pathways is interrelated with other elements of the problem formulation. Identifying pathways integrates information on:

- sources of COCs
- contaminant fate and transport
- receptors of concern and their general characteristics.

These elements have been discussed in previous sections.

Practitioners should provide rationale for including or excluding any potential pathways for each receptor group. The rationale may be based on quantitative considerations (e.g., magnitude of...
concentrations in groundwater and expected dilution before contact with the receiving environment), qualitative considerations (e.g., putative limitation of inhalation exposures to surface-dwelling wildlife), or a combination of these approaches. Rationales should indicate whether pathways are considered to be:

- complete (or operative/open), with a documented link between source and receptor
- incomplete (or inoperative/closed), with no documented or anticipated link between source and receptor.

The following exposure pathways should be considered (adapted from SAB-CS [2008]), although specific requirements may vary by jurisdiction:

- Soil invertebrates and terrestrial plants are in direct contact with elevated COC concentrations in soil.
- Mammals, birds, amphibians, reptiles, fish and invertebrate macrofauna ingest elevated COC concentrations via plants and organisms that ingest soil or sediment (e.g., via consumption of soil-covered plant roots).
- Mammals, birds, amphibians and reptiles ingest elevated COC concentrations via water ingestion.
- Mammals, birds, amphibians and reptiles ingest elevated COC concentrations via consumption of prey items (particularly for those chemicals known to bioaccumulate).
- Aquatic species (macrophytes, plankton, invertebrates, amphibians and fish) are in direct contact with elevated COC concentrations in surface water, sediment or sediment porewater.
- Some aquatic species (e.g., planktivores, piscivores) ingest elevated COC concentrations via consumption of prey items.
- Dermal exposure (direct contact with soil and sediment) of wildlife should be considered, when relevant, for COCs that can be absorbed readily through this pathway. Dermal exposure can also be a relevant exposure pathway for amphibians and reptiles. Detailed guidance on how to assess dermal exposure is limited (SAB-CS 2008; Suter 1996). Approaches for this pathway should be taken on a site-specific basis with appropriate rationale and consultation.
- Inhalation exposure through wind-blown dust or inhalation of vapours can be a relevant pathway for some mammals, birds, reptiles and amphibians. In practice, this pathway has not been commonly assessed, but may be required in some jurisdictions in future, and

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8 For example, policy in BC allows the ERA practitioner to exclude dermal exposure and inhalation for birds and mammals except for rare cases (SAB-CS 2008).
should be considered where the conceptual model indicates potential widespread exposure. For example, a site with high concentrations of volatile compounds and good small mammal habitat may warrant consideration of vapour inhalation. Inhalation toxicity data are currently lacking for most contaminants, but some jurisdictions are developing guidance and screening values for soil and vapour. In addition, because small mammals generally construct their burrows to allow for airflow, characterizing exposure may be challenging.

- Indirect pathways such as food source depletion by toxicity of COCs to invertebrates should also be considered.

Exposure data and toxicity reference values (TRVs) for amphibians and reptiles are limited, and the difficulties in assessing exposure may result in high uncertainties. The ERA should identify these limitations and uncertainties.

Tables are recommended for summarizing the pathway selection process. Table 2-5 provides an example template for aquatic ecosystems, and Table 2-6 provides an example template for terrestrial ecosystems.

**Key Concept**

Rationale should be provided to support inclusion and exclusion of exposure pathways for each receptor group in an ERA. Tables 2-5 and 2-6 are recommended for this.
<table>
<thead>
<tr>
<th>Receptor group</th>
<th>Exposure pathway</th>
<th>Included (yes/no)</th>
<th>Rationale</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary producer</td>
<td>direct contact (water)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>direct contact (sediment)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pelagic invertebrate</td>
<td>direct contact (water)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Benthic invertebrate</td>
<td>direct contact (water)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>direct contact (sediment)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>food consumption (for macrofauna)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fish</td>
<td>direct contact (water)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>direct contact (sediment)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>food consumption</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>incidental sediment ingestion</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mammal</td>
<td>water consumption</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>food consumption</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>incidental sediment ingestion</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bird</td>
<td>water consumption</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>food consumption</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>incidental sediment ingestion</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Amphibians and reptiles</td>
<td>direct contact (water)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>water consumption</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>food consumption</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>incidental sediment ingestion</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

This table should be adapted on a site-specific basis and in many cases should have additional detail for receptor types (e.g., benthic infauna, benthic epifauna) or additional pathways that may be relevant for particular contaminants (e.g., maternal transfer via eggs or lactation).
### Table 2-6: Example of tabular format for justifying exposure pathway selection in terrestrial ecosystems

<table>
<thead>
<tr>
<th>Receptor group</th>
<th>Exposure pathway</th>
<th>Included (yes/no)</th>
<th>Rationale</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary producer</td>
<td>direct contact (soil, soil porewater or groundwater)$^1$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Invertebrate</td>
<td>direct contact (soil, soil porewater or groundwater)$^1$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mammal</td>
<td>water consumption</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>food consumption</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>incidental soil ingestion</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>dermal exposure</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>inhalation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bird</td>
<td>water consumption</td>
<td></td>
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<tr>
<td></td>
<td>food consumption</td>
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<tr>
<td></td>
<td>incidental soil ingestion</td>
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<tr>
<td></td>
<td>dermal exposure</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>inhalation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reptiles and amphibians</td>
<td>water consumption</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>food consumption</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>incidental soil ingestion</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>dermal exposure</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>inhalation</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

This table should be adapted on a site-specific basis and in many cases should have additional detail for receptor types (e.g., benthic infauna, benthic epifauna) or additional pathways that may be relevant for particular contaminants (e.g., maternal transfer via eggs or lactation).

$^1$ For ERA purposes, this guidance defines any water in soil interstitial spaces in the biologically active zone as soil porewater. In other words, groundwater may be a source of contaminants, but in the biologically active zone of soil, that water is considered to be porewater.
2.8 Conceptual Site Model
A conceptual site model (CSM)\(^9\) guides implementation of an ERA by clarifying the relationships between:

- contaminant sources
- relevant fate and transport pathways
- VECs and surrogate VECs (if used)
- receptors of concern
- relevant exposure pathways.

The CSM is a core component of most ERA frameworks (e.g., ASTM 2008; CCME 1996-a; US EPA 1998; SAB-CS 2008; Suter 1996). The CSM can be expressed in a table, matrix, diagram or pictorial format. Importantly, the CSM should be supported with text that cross-references the rationale used to select receptors of concern and exposure pathways (e.g., the rationale detailed in Table 2-3 and Table 2-6).

Because risk assessment is an iterative process, a CSM should be updated as more information becomes available to refine the problem formulation.

The overall complexity of a CSM should be proportional to the complexity of the site. A simplified food web diagram (showing significant interactions between the different trophic levels and feeding guilds) is often a useful component of a CSM for identifying links between COCs, VECs and receptors of concern at all trophic levels. For example, a CSM with a food web diagram may indicate that elevated COC concentrations in soil may impact both soil invertebrates and insectivorous small mammals.

Two main types of CSMs are pictorial and box diagram, each with certain advantages and disadvantages as follows:

**Pictorial:** This is a graphical CSM that incorporates visual representations of the pathways and receptors. Pictorial CSMs should typically contain arrows and descriptive text to summarize linkages between sources, pathways and receptors. This style of CSM is well suited to communicating contaminant sources, exposure pathways, major fate processes and food web dynamics to a non-technical audience. A disadvantage is that some fate processes and indirect effects, as well as information on complete or incomplete pathways, cannot be represented easily.

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\(^9\) Guidance developed for environmental site characterization (CCME 2011) distinguishes between a CSM and a conceptual exposure model (CEM). ERA practitioners typically do not use the term CEM. The definition of a CSM used here is consistent with ERA practice and existing ERA guidance (CCME 1996-a; US EPA 1998; SAB-CS 2008).
in a pictorial fashion. This disadvantage can be mitigated by augmenting the pictorial CSM with a tabular summary of exposure pathways, indicating where pathways are complete and significant for each receptor group considered in the ERA (an example of such a tabular summary is part of Figure 2-4, discussed below). Figure 2-2 and Figure 2-3 show examples of pictorial-style CSMs.

**Box diagram:** This type of CSM uses a flowchart style. An advantage of this approach is that it facilitates a more rigorous examination of the pathways and connections among and between contaminant sources, fate and exposure pathways, and receptors. This type of model may incorporate a tabular summary indicating where pathways are complete and significant. The main disadvantage of this CSM form is that information is more difficult to interpret, especially for a lay audience. Figure 2-4 shows an example of this type of CSM.

For particularly complex sites, the use of both types of CSMs should be considered (rather than just one), as each type of CSM has unique advantages.

Importantly, while Figure 2-2 to 2-4 are typical examples of basic CSMs, additional information can be added to the CSMs, particularly for complex sites. For example, CSMs can be annotated with information about the COCs associated with each pathway, their chemical form in various media, or the types of effects that are considered for each receptor of concern. Figure 2-5 provides a simple example of a CSM that is specifically about the COCs, the receptor and food chain linkages, and also shows how the exposure and hazard assessments are conducted. Finally, a CSM may also be used to show indirect or secondary effects. For example, effects on food supply for piscivorous birds associated with a contaminant-related decline in fish population density.

Software choices for creating conceptual models vary, but in general include spreadsheet packages (e.g., Microsoft Excel), presentation packages (e.g., Microsoft PowerPoint), or graphic packages such as Corel Draw and Microsoft Visio. Ultimately, the software used will be determined by the presentation format and ease of use. Typically, box diagrams are easily constructed using spreadsheets or presentation packages, whereas pictorial diagrams usually require graphics packages.

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10 This list is not comprehensive but includes some of the more commonly used software packages.
Figure 2-2: Example of pictorial-style conceptual site model

ROCs
1. Aquatic Plants
2. Benthic invertebrates - infauna, epifauna
3. Fish - Includes (a) demersal fish with diverse diet, including benthic invertebrates, and (b) piscivorous fish feeding mainly on other fish
4. Birds - Includes (a) piscivorous species feeding mainly on fish, and (b) wading birds or waterfowl that may feed on invertebrates, plants and/or fish.
5. Mammals - Includes (a) marine mammals, and (b) terrestrial mammals that use the marine environment for food.
1. Accumulation of COCs by soil invertebrates (ingestion, direct contact) and plants (root uptake).
2. Consumption of plants and soil invertebrates by small mammals and birds.
3. Consumption of small mammals and birds by carnivores.
4. Movement and accumulation of COCs from soil to hard-bottom benthic organisms via groundwater and surface water runoff.
5. Movement and accumulation of COCs from soil to soft-bottom benthic organisms via groundwater and surface water runoff.
6. Uptake from COCs by fish from water and food.
Figure 2-4: Example of box diagram conceptual site model
Figure 2-5: Example of a customized conceptual site model
2.9 Designing and Planning the ERA
This section discusses aspects of problem formulation that are aimed at preparing for implementation of the ERA, with a focus on the tools and analyses that will be used to evaluate potential risks for each receptor of concern and exposure pathway. The design and planning stage includes (for terminology and key concepts, refer back to Section 2.1.2):

- establishing protection goals and (usually) associated acceptable effect levels (AELs)
- identifying the VECs attached to protection goals and linking these to the receptors of concern that are being measured as surrogate VECs
- identifying assessment endpoints, which are the attributes of the receptors that are to be protected (e.g., abundance or viability of a mammal population)
- identifying measurement endpoints, which are the tools used to measure changes in assessment endpoints
- developing lines of evidence for each assessment endpoint, which specify how measurement endpoints will be used to evaluate potential risks
- articulating the strategy for the ERA, as well as the SAP.

Importantly, as with the earlier sections of the problem formulation, the elements in this section are interrelated and therefore developed in an iterative manner.

2.9.1 Protection Goals and Acceptable Effect Levels
Most ERAs have a description of the type and level of protection that is intended for each receptor or receptor group at a site. This information may be used to “judge” the results of the risk assessment. A protection goal may be a narrative statement that is then operationalized through an AEL that clarifies the magnitude or rate of effects that would be acceptable for a specific measurement endpoint or a group of measurement endpoints.\(^\text{12}\)

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11 Not all ERAs require protection goals. Consistent with the CCME (1996-a) formulation of “detailed” ERAs, risks may simply be characterized, with all judgments about acceptability being made after the ERA is complete.

12 Given the interlinkages between AELs and endpoints, they are typically developed at the same time. An AEL can be applied directly to the assessment endpoint if the assessment endpoint is quantitative.
Figure 2-6: Conceptual relationships between assessment endpoints, measurement endpoints and lines of evidence

- **Conceptual Issues**
  - Valued Ecosystem Components: What biota or habitat do we want to protect?
  - Assessment Endpoints: What specifically about the biota or habitat do we want to protect?
  - Measurement Endpoints: What tools should we use to measure exposure or effects?
  - Lines of Evidence: How exactly will we use those tools to assess risks?

- **Operational Issues**
**Table 2-7: Example table of assessment endpoints, measurement endpoints and lines of evidence**

<table>
<thead>
<tr>
<th>Receptor group(s)</th>
<th>Assessment endpoint</th>
<th>Lines of evidence</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Benthic invertebrates</strong></td>
<td>Aquatic invertebrate community structure, and ecological function as food for fish and wildlife</td>
<td><strong>Line of evidence group</strong></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Line of evidence 1: sediment chemistry</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Line of evidence 2: benthic community analysis</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Line of evidence 3: amphipod toxicity test</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Birds, mammals, amphibians</strong></td>
<td>Abundance and viability of local bird, mammal and amphibian populations</td>
<td>Line of evidence 1: food chain model</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Line of evidence 2: small mammal trapping</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Line of evidence 3: wildlife survey</td>
</tr>
</tbody>
</table>
A protection goal usually differs for common species (where the population level is often of interest) relative to listed species\(^{13}\) (where individual organisms may need to be protected), and may differ according to land use or the overall site-management goals (see Section 2.2.1) of the ERA.\(^{14}\)

The following are examples of narrative protection goals:

- Maintain populations and associated demographics of small mammals that are similar to those at background conditions.
- No adverse organism-level impacts on the western toad (a listed species).
- Low level of significant ecological effects, defined to allow small structural or functional changes that may exceed natural variability provided that such do not threaten the sustainability of receptors\(^{15}\) (this is specified as a goal for commercial and industrial lands in Québec [CEAEQ 1998]).

Protection goals are not always operationalized immediately as AELs, but rather may be left in narrative form until measurement endpoints and lines of evidence are specified. Where they are applied, protection goals are intended to provide a degree of consistency across assessments, and as such are often influenced by policy determinations rather than by technical criteria. AELs may vary by receptor of concern, by endpoint or by site, depending on several considerations, including:

- Is protection aimed at individual organisms, populations or communities?
- Is the receptor of concern a common species or a species at risk (i.e., listed as rare or endangered)?
- Are there relevant federal or provincial laws, or pertinent policy determinations, that dictate appropriate AELs?
- Can appropriate AELs be inferred from methods used to derive national or provincial environmental quality guidelines?
- What effect size can be reasonably detected given natural variability?
- What effect size (at individual, population or community level) would be ecologically relevant for the particular receptor of concern?
- What are the spatial and temporal scales at which the effect will occur?

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\(^{13}\) Refers to species that are formally designated provincially/territorially or federally as, for example, rare, endangered or threatened.

\(^{14}\) For example, Québec guidance (CEAEQ 1998) specifies more stringent protection goals when protection of biodiversity is an overall objective for a site.

\(^{15}\) Original text in CEAEQ (1998): « Un faible niveau de réponses écologiquement significatives, c'est-à-dire un faible changement structurel ou fonctionnel pouvant excéder la variabilité naturelle mais ne mettant pas en cause la pérennité des récepteurs. »
• Would the effect be reversible?

• What are the environmental or economic consequences of a Type I error (false-positive) or Type II error (false-negative) in the risk assessment conclusions?

As the range of these considerations shows, deriving ecologically meaningful AELs can be complex. However, even if AELs are not explicitly articulated, they are often implicit. For example, any wildlife ERA that uses a published TRV to estimate a hazard quotient is assuming an AEL that is equal to the response size specified in the derivation of the TRV (see Section 4 for further discussion of TRVs). The risk assessor should ensure to the extent possible that the AEL implicit in a TRV is compliant with the protection goal.

For risk assessments where AELs are specified, it is preferable to base AELs on effect sizes that are defined in advance, as opposed to effect sizes that happen to be statistically significant based on hypothesis tests. Thresholds that are specified based on statistical significance in hypothesis tests are subject to large variation in ecological significance, depending on the level of statistical significance chosen, the specific study chosen, and details of the experimental design such as the range of treatments and sample sizes. An AEL based on a predefined effect size facilitates application of concentration-response methods (e.g., Allard et al. 2010) that provide a more standardized level of protection across contaminants, receptors and assessments.

**Key Concept**

AELs should be based on ecologically relevant effect sizes.

### 2.9.2 Assessment Endpoints

An assessment endpoint is an explicit expression of the environmental value to be protected. An assessment endpoint must include a receptor or receptor group (i.e., an entity to be protected) and a specific property or attribute of that receptor. For example, if the receptor is a fish community, candidate endpoint properties could include the community’s population demographics, biomass, genetic variability, physical condition or trophic structure.

The distinction between assessment endpoints and protection goals is subtle: an assessment endpoint describes the environmental attribute of interest, whereas a protection goal articulates the desired state of that attribute. To distinguish an assessment endpoint from a protection goal, practitioners should avoid using assessment endpoints that express an objective or a desired state (e.g., “healthy” or “functional”), and instead apply value-neutral terminology. The following examples illustrate this point:

• “benthic community diversity” (assessment endpoint) versus “maintenance of a diverse benthic community” (protection goal)

• “osprey reproduction” (assessment endpoint) versus “successful osprey reproduction” (protection goal)

• “marmot abundance” (assessment endpoint) versus “self-sustaining marmot population” (protection goal).
Checkai et al. (2002) identify other pitfalls during endpoint identification, including assessment endpoints that:

- are too vague (e.g., “stream integrity” rather than “abundance of juvenile salmonids”)
- evaluate an overly specific ecological entity (e.g., *Hyalella* growth instead of abundance of benthic fish prey); when assessment endpoints are too specific, they may be poorly aligned with the stressors of concern in terms of sensitivity and relevance
- are difficult to operationalize (e.g., endpoints based on the response of locally rare species)
- are not sufficiently sensitive given the management goals (e.g., if the management goal is to assess potential effects on wildlife, an assessment endpoint based on “presence versus absence of wildlife” would be too coarse to be useful).

### 2.9.3 Measurement Endpoints

A measurement endpoint\(^\text{16}\) is generally any measure of exposure or effects for a receptor of concern or any measure of change in the attribute of an assessment endpoint. Measurement endpoints form the basis for lines of evidence used to estimate risks (see Figure 2-6 and Table 2-7 above). Examples of measurement endpoints include:

- survival and growth of giant kelp gametophytes exposed to field-collected seep samples
- plant biomass per unit area
- Simpson’s diversity index for soil invertebrate samples
- abundance of mayflies, caddisflies and stoneflies per standard grab
- molar ratio of acid volatile sulfides to simultaneously extractable metals (SEM:AVS), as an indicator of potential bioavailability.

These measurement endpoints are measures of either exposure or effects, but not both. In general, to maintain the distinction between measurement endpoints and lines of evidence, these simple types of measurement endpoints are preferred. More complex formulations of measurement endpoints that attempt to incorporate both exposure and effects information (e.g., comparison of deer mice density on site and off site, or comparison of the daily ingested COC dose for deer mice at the site to a dose-based TRV that represents an AEL) are no longer measurement endpoints but lines of evidence. Measurement endpoints and lines of evidence must be developed at the same time, otherwise a measurement endpoint could be proposed without any understanding of how the information will be used.

\(^\text{16}\) The term *measurement endpoint* is preferred to *measure of effect* because the broad definition of measurement endpoint can include not only measures of effect (measurable change in an attribute), but also measures of exposure (measures of stressor existence, bioavailability and movement) and measures of ecosystem and receptor characteristics (characteristics that influence or mediate the relationship between exposure and effect) (Checkai et al. 2002).
2.9.3.1 Criteria for Selecting Measurement Endpoints
Measurement endpoints are selected in the context of particular receptor groups and assessment endpoints. Consequently, selection of measurement endpoints does not occur in isolation. Some of the criteria relevant to selecting receptors (Section 2.2.5.3) are therefore directly relevant to selecting measurement endpoints. Major technical criteria relevant to selecting measurement endpoints are reviewed in the context of lines of evidence in Section 2.3.4. While the risk assessor may need to consider other practical constraints such as cost, feasibility and time constraints, technical criteria will need to satisfy requirements from the jurisdictional authority in order for the risk assessment to be successful. However, in many ERAs, an iterative approach may be used, whereby the measurement endpoints that offer the best value (effectiveness per unit cost) are used first, and additional measurement endpoints are used in subsequent iterations as needed.

2.9.3.2 Level of Organization: Organism, Population, Community
It is desirable to maximize the correspondence between assessment and measurement endpoints, such that attributes are measured that are functionally related to the environmental property of interest. It is desirable, but not necessary, to align measurement and assessment endpoints across a common level of ecological organization.

For example, for the assessment endpoint “passerine abundance,” the measurement endpoint might be “density of adult breeding pairs of American robins.” The line of evidence could then be defined as “percentage difference in density of adult breeding pairs of American robins on site X compared to reference conditions.” In this case, a population-level attribute (density of breeding pairs) is applied to the local population of robins. In this example, the measurement and assessment endpoints are both expressed at the population level. However, if it is difficult to measure the number of breeding pairs, or if it is a highly variable measure, or if the measure is likely to be confounded by off-site immigration, alternative measurement endpoints that might be considered include “mortality rate and reproductive success of robins.” In that case, two organism-level attributes (mortality and reproductive success) are assumed to be representative measures that may be extrapolated to the local population of robins. Such extrapolation could be conducted qualitatively using a narrative or quantitatively using a population model.

A key property of any measurement endpoint should be the ability to interpret the results in relation to protection goals. If the protection goal is “minimal effects to a terrestrial mammal community,” ideally the changes in measurement endpoints can be related to potential effects on populations and ultimately to that community. In practice this is quite challenging; this issue has plagued ecotoxicologists for several decades due to the complex linkages and uncertainties in ecological systems, including density dependence, intraspecies sensitivity variations and confounding habitat factors. Although ERA practices have evolved to address some of the uncertainties, such as adjustments using extrapolation factors and uncertainty factors, there remains a significant degree
of difficulty in extrapolating across levels of organization, and complexity in understanding
dynamics at higher levels of organization.

Most measurement endpoints in ERAs address organism-level attributes of a population or
community (Suter et al. 2005), such as mortality rate, reproductive success and growth. Assess ment endpoints commonly address populations or communities, whereas measurement endpoints address organism-level attributes that are believed to be linked to the population- or community-based assessment endpoint (CCME 2006). The quantitative linkage between organism-level attributes and responses to populations or communities is seldom known with confidence. However, it is usually assumed that there will be no effects at the population or community level if an ERA predicts no effects at the organism level. If the ERA predicts effects to individual organisms, it is not easy to predict effect levels for populations or communities. For example, a population that is already at carrying capacity may be unaffected by a higher mortality rate among individual organisms. Conversely, a population that is barely able to sustain itself may be extirpated under any additional stress. Extrapolating from organism-level attributes to populations requires an understanding of factors controlling population dynamics. Extrapolating from populations to communities requires an understanding of community interactions (e.g., one species may increase in abundance if its associated predator decreases in abundance). Although there is a desire in the ERA community to develop methods for extrapolation, normally ERA practitioners describe possible community effects only qualitatively, if at all.

Some measurement endpoints are easy to interpret ecologically because they address true community-level or at least population-level attributes, but they have the power to detect only very large changes. Other measurement endpoints that address organism-level attributes are more powerful at detecting change, but are less easily extrapolated to populations and communities. It is partly for this reason that ERAs depend on multiple lines of evidence. Nevertheless, methods for evaluating effects on population and communities exist (see Suter 2007) and should be employed whenever possible.

A specific difficulty in evaluating population-level effects lies in defining the population of interest (i.e., the assessment population). From a pure biology perspective, an ecological population is defined as a group of organisms of a single species that interbreed and share a common habitat. From a risk assessment perspective, however, this definition is too broad, particularly for organisms that migrate across large areas (up to the continental scale). If assessment populations are defined across large spatial scales, then effects on local groups of individual organisms near a particular contaminated site might not have an impact on the assessment population, yet might still exert local impacts that are considered unacceptable in relation to protection goals.

Key Concept

Whenever populations are of interest, particularly for wildlife, the population of management interest (the assessment population) should be defined as clearly as possible.

A further issue with respect to defining populations is understanding the ecological context of the group identified as a local population. A small patch of forest in the middle of farmland or an urban center may play an important role (e.g., migration corridor) when overall habitat is fragmented,
whereas a similar-sized area located in an unfragmented wilderness area may be less sensitive to ecological disruption.

The general issues of spatial scale and the overall magnitude of effects are addressed in more detail in Section 5.

2.9.3.3 Types of Effects Used in Measurement Endpoints

For measurement endpoints that are direct measures of effects, there is general agreement that certain types of effects are more suitable than others. Specifically, effects that are measured need to be ecologically relevant and linkable back to assessment endpoints that focus most often on population- or community-level attributes.

CCME (2006) notes that effects measured at the organism level should be those that are critical for a species to complete a normal life cycle and produce viable offspring. Mortality and reproduction are the two types of effects that can be most easily related to population-level effects, but population dynamics are typically complex and there may be several direct and indirect mechanisms by which lethal and sublethal effects could impact at the population level. Direct measures at the population and community levels are ideal but not often feasible or practical to obtain.

Other types of effects can be applied as surrogates for population responses but are generally more difficult to relate to population- and community-level effects. This is particularly true for effects that are not truly a measure of an adverse effect but rather are a measure of the potential for adverse response (e.g., enzyme induction).

Various Canadian jurisdictions provide some guidance on the types of effects that should be used for measurement endpoints (see Table 2-8). There are no clear rules for using these types of effects in endpoint selection, so ERA practitioners must use their judgment on a case-specific basis. When considering the advantages and disadvantages of various types of effects (Table 2-9), preference should be given, whenever possible, to types of effects that are as closely tied to assessment
Table 2-8: Types of effects and their acceptability in various jurisdictions for use in measurement endpoint selection\textsuperscript{17}

<table>
<thead>
<tr>
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<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Mortality</td>
<td>acceptable</td>
<td>acceptable</td>
<td>acceptable</td>
<td>acceptable</td>
<td>acceptable</td>
<td>acceptable</td>
<td>acceptable</td>
</tr>
<tr>
<td>Reproduction</td>
<td>acceptable</td>
<td>acceptable</td>
<td>acceptable</td>
<td>acceptable</td>
<td>acceptable</td>
<td>acceptable</td>
<td>acceptable</td>
</tr>
<tr>
<td>Growth</td>
<td>acceptable</td>
<td>acceptable</td>
<td>acceptable</td>
<td>acceptable</td>
<td>acceptable</td>
<td>acceptable</td>
<td>acceptable</td>
</tr>
<tr>
<td>Behaviour</td>
<td>acceptable</td>
<td>acceptable</td>
<td>variable</td>
<td>variable</td>
<td>variable</td>
<td>variable</td>
<td>variable</td>
</tr>
<tr>
<td>Morphology or deformity</td>
<td>acceptable</td>
<td>variable</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tumours</td>
<td>acceptable</td>
<td>variable</td>
<td>variable</td>
<td>variable</td>
<td>variable</td>
<td>variable</td>
<td>variable</td>
</tr>
<tr>
<td>Physiological measures such as absorption efficiency, nutrient uptake, blood volume</td>
<td>acceptable</td>
<td>variable</td>
<td>variable</td>
<td>variable</td>
<td>variable</td>
<td>variable</td>
<td>variable</td>
</tr>
<tr>
<td>Enzyme activity</td>
<td>acceptable</td>
<td>not acceptable</td>
<td>variable</td>
<td>variable</td>
<td>variable</td>
<td>variable</td>
<td>variable</td>
</tr>
<tr>
<td>Histopathology (cellular changes)</td>
<td>acceptable</td>
<td>acceptable</td>
<td>not acceptable</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Development (some measures only, e.g., sexual development)</td>
<td>variable</td>
<td></td>
<td>acceptable</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Immunological response</td>
<td>not acceptable</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Population-level attributes (e.g., biomass, abundance)</td>
<td>acceptable</td>
<td>acceptable</td>
<td>acceptable</td>
<td>acceptable</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Community-level attributes (e.g., diversity)</td>
<td>acceptable</td>
<td></td>
<td>acceptable</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

\textsuperscript{17} This table is simplified. Some guidance documents make exceptions on a case-by-case basis, with a key criterion being whether a particular effect is likely to affect survival, reproduction or growth. Blanks in the table do not indicate whether the effect is acceptable or not; rather, no specific mention was made of that effect type.

\textsuperscript{18} MECP (2011) is for development of generic standards, but is used as the basis for many risk assessments. Acceptability varies by receptor group. There are exceptions in many cases, and ERA proponents can deviate from the above if full and appropriate explanations are given.
### Table 2-9: Advantages and disadvantages of using particular types of effects as measurement endpoints

<table>
<thead>
<tr>
<th>Type of effect</th>
<th>Key advantages (A) and disadvantages (D)</th>
</tr>
</thead>
</table>
| Direct measures at community level, such as diversity, species richness or biomass | A: Usually directly relevant to the assessment endpoint.  
D: Can be difficult to measure directly; power to detect change may be limited. |
| Direct measures at population level, such as abundance or biomass              | A: Usually directly relevant to the assessment endpoint.  
D: Can be difficult to measure directly; power to detect change may be limited, and ability to establish causal relationships with stressors may be limited. |
| Mortality rates                                                                | A: Easy to measure in some cases; response times usually fast; relatively easy to relate to population level.  
D: Often less sensitive than other endpoints; difficult to measure *in situ* (e.g., for wildlife); responses may be delayed. |
| Reproductive endpoints (e.g., fecundity, reproductive success)                 | A: Can be easy to measure depending on the specific effect; relatively easy to relate to population level depending on the specific effect; can be an indicator of other unknown effects.  
D: Some reproductive effects are not easy to measure (e.g., for wildlife) and have long response times for some receptors. |
| Growth                                                                         | A: Often more sensitive than mortality or reproductive endpoints; can be an indicator of other unknown effects.  
D: More difficult to relate to population level. |
| Behaviour (where the behaviour could be linked to mortality, such as predator avoidance, or to reproduction, such as mating frequency) | A: Often more sensitive than mortality or reproductive endpoints; can be an indicator of other unknown effects; can often be linked to reproduction and mortality.  
D: Links to population and community level may be vague, effects may be subtle and response times may be long. |
| All other types of endpoints (see examples in Table 2-8)                       | A: May be more sensitive to contaminants than other endpoints.  
D: Difficult to relate to population- and community-level assessment endpoints, or may not have a net adverse effect. |

Endpoints as possible. Measurement endpoints that are relevant to assessment endpoints include direct measures at the population and community levels, or mortality and reproductive effects that can be directly related to population-level attributes. Among the other types of effects, growth is generally the most preferred. This does not mean that other endpoints should be excluded from consideration. If behavioural effects with likely implications at the population level (e.g., decreased predator avoidance) are observed at low concentrations, those should be considered relevant.\(^\text{19}\) There is some discussion in the FCSAP TRV module (Module 2, EC 2010-b) regarding links between measurement endpoints and TRV selection.

\(^{19}\) CCME (2007) notes that for derivation of water quality guidelines, nontraditional endpoints such as behaviour can be used if ecological relevance can be demonstrated.
2.9.4 Lines of Evidence

The ways in which measurement endpoints are organized and applied define the lines of evidence that will be carried through the ERA. Lines of evidence are derived from assessment and measurement endpoints (Figure 2-6; examples shown in Table 2-7). Although the figure shows a stepwise process, in reality the lines of evidence should be developed nearly concurrently with the measurement endpoints (i.e., there is no point identifying a tool without thinking ahead to the proposed application of the results). As highlighted previously, the scope of measurement endpoints varies widely, and endpoints can be defined in a way that makes them functionally equivalent to a line of evidence. Generally, it is easier to define measurement endpoints as measures of exposure or effect so that they are clearly distinguished from lines of evidence. The expression of lines of evidence provides a bridge between the unprocessed data collected to inform the risk assessment (measurement endpoints) and the subsequent analysis and interpretation of those data in the analysis stage of the ERA.

For example, if we measure species diversity in a soil invertebrate community (measurement endpoint), the data could be applied in several ways, including:

- comparisons of mean diversity index on site versus in reference conditions (e.g., using ANOVA)
- comparison of a diversity index to predetermined thresholds for soil quality based on ecological principles
- modelling of the diversity index versus the soil concentration of a contaminant (e.g., using simple linear regression).

These would be considered as separate lines of evidence derived from the same measure of effect (diversity index). Two of the analyses measure the magnitude of potential risks, while the other focusses on establishing potential causal relationships with contamination. Each line of evidence carries different (but valuable) information for informing the assessment endpoint.

From a practical perspective, it may be appropriate to group closely related lines of evidence. For example, as illustrated in Table 2-7, all lines of evidence that use results of an amphipod toxicity test in one way or another may be grouped together for purposes of analysis and reporting, even though that toxicity test may have more than one specific measurement endpoint (e.g., growth and survival) and there may be several specific lines of evidence developed that use the results of the toxicity test.
In specifying lines of evidence, it is important to provide a clear expression of the relationship between exposure and effects measures. For some lines of evidence, the relationship is obvious (e.g., comparing soil chemistry to soil quality guidelines). In other cases, the relationship is less intuitive and requires explanation (e.g., benthic invertebrate community diversity as a function of proximity to a point source). In the latter example, the “proximity” could be a function of distance, direction, or both, and the line of evidence may need to specify groupings of stations, distance-based transects or other measures of exposure.

2.9.4.1 Organizing Lines of Evidence
To facilitate consistency in practice, it is helpful to conceptualize the following four major categories of lines of evidence:

- **Site-specific toxicological evidence**: Considers measurement endpoints related to studies of test organism exposure to contaminated site media under controlled conditions.\(^{20}\)

- **Indirect toxicology evidence**: Considers toxicological information gleaned from other sites, under an assumption that the concentration-response relationship is either similar to, or can be estimated from, the data collected at other sites.

- **Site-specific biological evidence**: Considers direct assessment of the site’s biological condition.

- **Indirect biological evidence**: Considers indirect assessment of biology, through extrapolation of knowledge obtained at other sites.

Framing lines of evidence in this manner streamlines the risk characterization (Section 5) and is consistent with the framing of tools in the hazard assessment (Section 4).

One or more of these categories are often omitted from any given stage of a risk assessment, depending on the scope and complexity of the study, the project tiering strategy, and the objectives of the risk assessment. For example, site-specific toxicology studies are infrequently conducted for birds and mammals, and are practically non-existent for endangered species. Similarly, site-specific biological investigations are seldom conducted during a preliminary quantitative risk assessment. Lack of representation of any of the four categories is in itself not a cause for criticism. However, the risk assessor should explicitly acknowledge the implications and uncertainties associated with emphasizing or omitting any of the above lines of evidence.

Each of these broad categories carries different uncertainties and evaluation methods. Furthermore, it is common to have multiple individual lines of evidence within a single broad category, as follows:

- Site-specific toxicity tests are commonly conducted as part of a test battery approach with multiple species, durations and endpoints.

\(^{20}\) Controlled conditions may be in the laboratory or in situ.
• Comparisons to guidelines or benchmarks may entail multiple comparisons (different jurisdictions or case studies).

• Community studies have a multitude of potential endpoints (e.g., total density and diversity, major taxa density and diversity, sensitive taxa density and diversity, diversity indices).

• Biological endpoints from other sites can be numerous in type.

• Biological and toxicological endpoints can be compared against many candidate exposure metrics (e.g., chemistry of individual COCs, chemical surrogates such as TEQs, multivariate chemistry exposure metrics [principal components], distance or direction metrics).

One of the reasons for grouping endpoints into the four broad categories is to explicitly acknowledge the partial redundancy of having multiple related endpoints. Of course, formally organizing lines of evidence is not important for simple ERAs where there may be only a limited number of lines needed to address risks, or for cases where a limited number of lines of evidence lead to a clear conclusion that risks are negligible.

2.9.4.2 Selecting Lines of Evidence

The rationale for selecting lines of evidence should be explicit as part of problem formulation. The lines of evidence used for an ERA (see example in Table 2-7) are derived from a long list of potential lines of evidence. The criteria that are relevant for selecting lines of evidence may include:21

• **Ecological relevance**: To what degree is the assessment endpoint represented by the line of evidence?

• **Sensitivity**: To what degree can the line of evidence detect changes or differences from reference conditions? Are results reported quantitatively or using broad categories such as low, moderate and high? Does the line of evidence typically suffer from a high degree of random error?

• **Specificity**: Will the line of evidence be specific enough to identify effects from the COCs over and above other factors present at the site?

• **Spatial representativeness and site specificity**: Does the line of evidence provide information at the appropriate spatial scale, and does the line of evidence take into account site-specific factors that may influence the results compared with other sites?

• **Temporal representativeness**: Does the line of evidence capture temporal variation relevant to potential ecological risks?

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21 Adapted in part from Menzie et al. (1996) and SAB-CS (2008).
• *Expected data quality*: Based on the practitioner’s experience, what is the likelihood that the quality of data generated by this line of evidence will be poor and result in reduced utility of the line of evidence?

• *Expected acceptability*: Does the line of evidence have standard test methods or a long history of use that provides confidence that regulators will accept the results?

At the least, practitioners should provide a list of lines of evidence that they considered for a particular ERA, with rationale for inclusion or exclusion of each. This can be done in text or in a table. The rationale should be based on appropriate criteria such as those listed above. For complex ERAs where a phased approach to implementation is used, rationale should also be provided to justify which lines of evidence are proposed initially and which are deferred.

2.9.4.3 Applying Lines of Evidence

Because lines of evidence are carried forward in the ERA to evaluate risks, it is important to cross-check selection to ensure that the selected lines of evidence represent all receptor groups and all exposure pathways. Receptor groups are inherently cross-checked during preparation of the lines of evidence table (Table 2-7). For cross-checking exposure pathways, the simplest approach is to add a single yes/no column to the templates in Table 2-5 and Table 2-6 to show whether or not at least one proposed line of evidence is relevant to each exposure pathway.

Lines of evidence are evaluated and implemented as part of the ERA. The specific approach that will be used to integrate the lines of evidence in a WOE framework should be described as part of the general strategy for the ERA (see next section). The level of detail that needs to be provided depends on the complexity of the ERA and the type of approach used for risk characterization. Section 5 discusses the range of options for conducting WOE assessments. The most detailed WOE approaches may require formal quantitative assessment of the lines of evidence during problem formulation (e.g., weighting or ranking of respective lines of evidence based on a multi-attribute assessment of each). While such approaches may be overly cumbersome for most ERAs, critically assessing lines of evidence for relevance before data analysis guards against gratuitous assumptions (made after the fact) that provide superficial appearance of a systematic decision-making process but are in fact arbitrary and impossible to discern from subjective interpretations. Regardless of the level of formality used, rationale must be provided for the selection of lines of evidence at the problem formulation stage, as discussed in the previous section. Criteria that do not change based on data collected after the problem formulation (e.g., ecological relevance) can be carried forward directly from the problem formulation to risk characterization in a WOE framework (see Section 5).
2.9.5 General Strategy for the ERA

At the same time that lines of evidence are developed, it is important to design the overall implementation strategy for the ERA. The strategy should not get into details about field methods, lab methods or data analysis methods, as those are best left to the SAP (see next section). The strategy focuses on big-picture issues, typically covering:

- **Phasing/iteration**: Will the ERA be implemented in phases? If yes, what lines of evidence will be pursued in which phases? Under what conditions (results) could the first phase be sufficient for the ERA to be considered completed?

- **Timeline**: Implications of phasing and other constraints should be presented as they relate to the overall timeline expected for the ERA.

- **Experimental design** (see 2.3.5.1 for more discussion on experimental design): Will field studies incorporate a gradient design or comparison of the site to a reference condition? What amount of field replication will be needed in order to have adequate power to detect effect sizes of interest or to establish correlations between exposure and effects? What is the general spatial scale of sampling for each type of data? While details are listed in the SAP, the conceptual design should be articulated as part of the general strategy.

- **Coordination with ongoing site investigation**: If supplemental site investigation work is ongoing, how will that work mesh with and support the ERA? How will the site investigation data be used in the ERA?

- **Approach to risk characterization**: Assuming there is more than one line of evidence for at least some of the assessment endpoints, the practitioner must describe during problem formulation how they will implement the WOE approach to risk characterization. This should include details regarding:
  - how lines of evidence will be summarized and integrated
  - how judgments about the magnitude of risks, uncertainty about risks, causation or other attributes will be made (a default table is provided in Section 5 for this purpose).

In short, the details of how risk characterization will be implemented should be fully understood and articulated at the problem formulation stage. Detailed discussion of risk characterization including WOE approaches is deferred to Section 5 for organizational
simplicity, but most of the content is relevant (i.e., must be considered) at the problem formulation stage.

- **Transparency**: How will the ERA results as a whole be presented? What mechanisms or tools will enable reviewers to understand how conclusions were drawn? What mechanisms or tools will enable reviewers to make independent evaluations of risk based on the information presented?

The general implementation strategy should be discussed with key stakeholders and regulatory authorities before proceeding in order to confirm that the ERA will fulfill expectations. Provided there is agreement on the strategy, an SAP can be prepared before the work begins.

2.9.5.1 Control-Impact versus Gradient Designs

Experimental design warrants careful consideration in ERA, because the design dictates what types of inferences can be drawn from the data collected. An important element of experimental design for the practitioner is deciding in advance how the potential effects of contamination will be evaluated. The classic “control-impact” design that is often used in ERA to compare a site to a reference site has fundamental problems because of natural variability among sites unrelated to contamination. Comparison to a reference condition (based, for example, on multiple reference sites) is preferable but is also confounded to some extent by natural variability among sites. In most cases, a gradient design should be considered, as it allows the practitioner to evaluate potential relationships between contamination and effects, and to understand any differences observed between areas of varying concentrations of contaminants. The following discussion provides rationale in this regard.

In a classic control-impact design, the effect of contamination would be interpreted by comparing site-related performance to control performance. For example, a practitioner may compare plant growth at a contaminated site to plant growth at a control or “reference site,” assuming the two sites are identical except for the contamination. Unfortunately, no two sites are identical, so comparison of a site to a single reference site is of limited value. If multiple samples are taken for both sites, the hypothesis that the two sites have similar plant growth can be tested statistically, but any difference between the two sites cannot be taken as evidence as a contaminant-related effect, because we should expect the two sites to be innately different even in the absence of contamination. A practitioner who incorrectly assumes that a statistically significant difference in this case is related to contamination is committing pseudoreplication (Hurlbert 1984), because the data provide evidence only of variability between those two particular sites, not variability between contaminated and uncontaminated sites in general. The samples at each site are considered pseudoreplicates, not true replicates in a test for the effect of contamination.
One approach to addressing the problem of inherent variation among sites is to define a “reference condition” against which a contaminated site could be evaluated. In the context of a contaminated site, a reference condition would usually be one that is assumed to represent a range of conditions that would occur in the absence of site-specific contamination. A reference condition could be established in various ways (Stoddard et al. 2006), the most common of which is to use multiple reference sites to establish a range of conditions that represent reference. For example, the reference condition approach (RCA), based on multiple reference sites, is used under the Canadian Aquatic Biomonitoring Network (CABIN) to evaluate potential impacts of stressors on freshwater aquatic systems (Government of Canada 2017). This is a vast improvement over comparison to a single reference site. However, since the contaminated site itself is not replicated, it is impossible to know to what extent any observed differences between the site and the reference condition are natural or related to contamination. Provided that practitioners understand this limitation, comparison of a contaminated site to reference conditions derived from multiple reference sites can be useful. Other approaches for deriving a reference condition include interpreting historical condition (if information exists for conditions at a site before contamination), extrapolating from empirical relationships relating biological indicators to contamination (e.g., from other sites) or using ecological principles to specify expected conditions in the absence of contamination (Stoddard et al. 2006).

Comparison of biological variables at a contaminated site to a reference condition will always be confounded by the inherent variation in biological systems. Because assessment endpoints for ERAs are often at population or community levels of organization, practitioners should expect that the population or community of interest at a site will be inherently different from the population or community associated with any other particular site or set of sites. Landis et al. (2011) provide detailed arguments as to why reference sites are not relevant for populations and landscapes. The implication is that the sampling design for ERAs should usually focus less on testing for a difference between the site and a reference condition, and more on evaluating patterns based on gradients of contamination and other factors that are likely to drive biological variability (Landis et al. 2011). This is particularly true for measurement endpoints that measure populations or communities directly.

Gradient designs should aim to capture the range of COC concentrations from highest (on site) to lowest (on site or off site). Some gradient designs may have a directional spatial element such as distance from a point-source of contamination. Most importantly, gradient designs should control for patterns in environmental variables that may be correlated with contamination. Confounding variables often limit the ability of practitioners to make links between observed biological patterns and site-related contaminants. Inherent in a gradient design is the objective of determining whether populations and communities of interest are correlated with and caused by contamination. As discussed at length in Section 5, this latter aspect—establishing causality—should be a key component of any ERA.

In short, whenever possible, the study design for ERAs should aim to characterize gradients in contamination and other factors that are likely to drive responses in populations and communities. Comparing a site to reference conditions is also useful, but conclusions based solely on such
comparisons are limited when relationships to contamination and other predictor variables are not understood.

2.9.6 Sampling and Analysis Plan

A sampling and analysis plan (SAP) describes details of how the ERA will be implemented. A SAP usually focuses on technical details rather than higher-level strategic issues outlined in the previous section. Therefore, it is common for a draft problem formulation to end after discussing the general implementation strategy, with the SAP added later if there is general agreement that the ERA should proceed. In that case, the SAP may be added to the problem formulation before it is finalized or may be developed as a stand-alone document.

The scope of an SAP will vary depending on the complexity of the ERA and the level of detail that has already been specified earlier in the problem formulation. The SAP may address all of the planned sampling and analysis details for the entire ERA, or it may be limited to plans for the first phase (or tier) of the ERA. In cases where no further field sampling is needed, the SAP will relate to analysis only.

Importantly, the SAP must demonstrate that it is fulfilling the information needs for each line of evidence that will be used in the ERA. A checklist is the best way to cross-check the completeness of the SAP and make sure that the field and data requirements of each line of evidence are met. Because field programs are normally implemented at discrete times, accidental omissions related to field data collection can have significant implications. Table 2-10 provides a checklist template. While the primary benefit of such a table is for the risk assessor (i.e., to ensure the SAP is complete), including it in the SAP submission shows reviewers that a cross-checking process was undertaken.

The rest of this section expands on some of the SAP requirements listed in Table 2-10. Before practitioners develop an SAP, they should consult jurisdictional guidance and current CCME (2012) guidance on sampling in contaminated sites.

Field safety plan: A field safety plan is important for every project involving field work. Whether it is part of the SAP or handled separately is not important, but the SAP should at least confirm that the plan is or will be in place.

Logistics: An important but often overlooked component of an SAP is review of big-picture logistical considerations. Logistical considerations may include:

- time required to get sampling permits
- permission for site access
- transportation and accessibility (particularly for remote sites)
- availability of key sampling equipment

Key Concept
The appropriate level of detail in a SAP varies depending on the scale and complexity of the ERA, as well as expectations of stakeholders.
seasonal considerations for biological sampling (e.g., sampling for berries, mushrooms, kelp, eelgrass, leaves)

- tide cycles (e.g., intertidal work may require a very low tide that occurs over multiple days during daylight hours).

Chemistry sampling: Whenever chemistry samples are included as part of an SAP (water, soil, sediment, tissues), the SAP should specify, at the least, the following:

- relevant COCs for each media
- the form(s) of each COC to be measured in each medium
- sampling locations and replicates
- sample collection methods (equipment, depth of samples, processing, volumes, jars to be used, etc.)
- critical aspects of sample handling (filtration, storage, holding times, etc.)
- expected lab methods including preparation (e.g., dry weight or wet weight) and reporting units
- expected lab detection limits
- the list of supporting or “conventional” parameters to be measured. The list of conventional parameters will vary by media type. The risk assessor should not assume that the list of conventional parameters collected during site investigation will suffice for the ERA. Conventional parameters should include relevant indicators of potential bioavailability of COCs, which vary depending on the COC and media.

Section 3 identifies typical tools for chemistry sampling (by media), typical ancillary data, cautions to be exercised in field sampling and selected guidance documents related to chemistry sampling (see Table 3-1).
### Table 2-10: Example checklist for an SAP

<table>
<thead>
<tr>
<th>Field component</th>
<th>Soil chemistry</th>
<th>Invertebrate bioassay</th>
<th>Soil invertebrate community</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>A. Planning checklist</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Field safety plan established?</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Logistics</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Permits and site access permissions</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Transportation and access</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Availability of major sampling gear</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Seasonality appropriate for data?</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sampling</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Core parameters (e.g., COCs) included?</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ancillary/supporting parameters included?</td>
<td></td>
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<td></td>
</tr>
<tr>
<td>On-site sampling locations selected?</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reference sampling locations selected?</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Detailed field sampling methods established?</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sample handling methods specified?</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Field QA/QC methods and objectives established?</td>
<td></td>
<td></td>
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<tr>
<td>Laboratory analyses</td>
<td></td>
<td></td>
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<tr>
<td>Lab methods specified?</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Lab detection methods specified and adequate?</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lab QA/QC methods and objectives established?</td>
<td></td>
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<tr>
<td>Data analyses and modelling</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Data expected to be adequate to support all analyses?</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>B. Line of evidence requirements checklist</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Line of evidence 1a: compare bioassay results on site vs reference</td>
<td>x</td>
<td></td>
<td>y/n</td>
</tr>
<tr>
<td>Line of evidence 1b: regression of bioassay results vs soil chemistry</td>
<td>x</td>
<td>x</td>
<td>y/n</td>
</tr>
<tr>
<td>Line of evidence 2a: compare abundance/diversity on site vs reference</td>
<td></td>
<td>x</td>
<td>y/n</td>
</tr>
<tr>
<td>Line of evidence 2b: regression of abundance/diversity vs soil chemistry</td>
<td>x</td>
<td>x</td>
<td>y/n</td>
</tr>
</tbody>
</table>
**Biological and other sampling:** As with chemistry sampling, for each other type of field sampling there should be a description of what will be sampled, how it will be sampled and how the laboratory (if applicable) will conduct analysis of the samples. Examples of field sampling details include mesh size for sieving benthic invertebrates, quadrat size for evaluations of vegetative cover, and design specifications and bait for small mammal traps. Examples of lab methods to be specified include taxonomic resolution for measures of invertebrate density, and plans for salinity adjustments for bioassays conducted using groundwater adjacent to the marine environment.

Section 3 identifies typical tools for biological sampling, typical ancillary data, cautions to be exercised in field sampling and selected guidance documents related to chemistry sampling (see Table 3-1).

**Quality assurance and quality control:** QA/QC methods and expectations should always be specified before sampling, so that the quality of data is ensured to the extent possible. If data quality objectives (DQOs) cannot be met, it may be necessary to select new measurement endpoints and lines of evidence.

Specific QA/QC mechanisms typically associated with collection of environmental chemistry programs include:

- prevention of contamination during field sampling (e.g., use of clean jars)
- decontamination procedures between sampling stations to prevent cross-contamination, and potential use of cross-contamination swipes of sampling gear to test for cross-contamination
- field homogenization procedures (e.g., for bulk soil samples), and potential use of field duplicate or triplicate samples to evaluate the effectiveness of homogenization
- sample storage, transport and chain-of-custody, and potential use of lab travel blanks
- lab replicates to test for measurement error (as relative percent difference)
- analytical methods blanks, certified reference materials and matrix spikes
- method-detection limits relative to screening guidelines and relevant to use in ERA.

In the case of toxicity tests, labs, negative and positive controls, replication, instrument calibration, and other QA/QC mechanisms are used (see Section 4 for more information on toxicity testing). Labs implementing invertebrate enumeration may use resorts and sample splitting as QA/QC mechanisms. CCME (2012) provides more detailed consideration of QA/QC procedures.

**Data analysis and modelling:** Data analysis plans may or may not be included in an SAP, depending on the scale and complexity of the ERA and expectations of regulators, site custodians or stakeholders for a particular site. The formulation of lines of evidence describes how data will be analyzed, but there may be cases where additional details are warranted. For example, for a complex site where a food chain model will be used to estimate wildlife exposures, it may be appropriate to outline the key aspects of the model design and assumptions.
2.9.7 Communication and Review

Practitioners should look for outside review of their SAPs before they embark on field data collection and analysis. Reviewers could be appropriate regulatory authorities, experts, other affected stakeholders or peers. For complex sites, it may be important to develop communication tools (e.g., figures) that reduce the complexity in the problem formulation to something that is understandable by readers without technical expertise in ERA.

2.10 Uncertainties and Data Gaps in Problem Formulation

Uncertainties are pervasive in ERA. A focussed discussion on the key uncertainties in a problem formulation is worthwhile. One major benefit of explicitly discussing uncertainties is that it may lead to identifying specific data gaps that could be addressed as part of, or parallel to, the ERA. Some of the sources of uncertainties and data gaps common at the problem formulation stage include:

- **COCs**: There may be uncertainty about the list of COCs relevant to the site, which may be associated with incomplete history for the site, uncertainty about potential off-site sources of COCs, or simply chance that site investigation failed to detect a COC that is actually present at elevated concentrations. There may also be uncertainty about characteristics of the COCs related to fate, transport and effects.

- **Transport pathways**: The CSM assumes that all relevant fate and transport pathways have been characterized. However, even well-designed site investigation work may fail to detect key pathways. For example, movement of contaminants through groundwater to the marine receiving environment may occur only under certain narrow tidal and seasonal windows; in such a case, the risk assessor will remain “ignorant” and the CSM will not fully capture all relevant pathways.

- **VECs and receptors of concern**: There is always uncertainty about selection of VECs during the problem formulation. Usually the major receptor types are captured, and uncertainties are associated with the selection of receptors of concern as surrogate VECs. For example, wildlife biologists may fail to recognize habitats for certain VECs, so these habitats may be prematurely excluded from the ERA. Alternatively, effects literature may be limited (this is usually the case), so the surrogate species selected to represent a given receptor type may not be the most sensitive species.

- **Measurement endpoints**: Measurement endpoints are imperfect, either because of uncertainty in the measurements themselves (e.g., variability reduces the power to detect differences), or because of uncertainty about how the measurement endpoints translate into effects on assessment endpoints (e.g., what does a reduction in invertebrate growth mean at the population or community level?). While there is no particular data gap to be addressed, the risk assessor should acknowledge these uncertainties up front.
Box 2.1: Types of uncertainty in ERA

ERA contains several types of uncertainties, such as:

- **Natural variability** that cannot be “reduced” (e.g., variability in COC concentrations across a site, spatial variability in the distribution of biota). Natural variability can be acknowledged, characterized and incorporated into an ERA (i.e., using probabilistic methods).

- **Random measurement error** associated with estimating a parameter, such as may result from limitations in the number of observations or imprecision in the measurement techniques. Estimates of most parameters in an ERA are imprecise – examples include average soil concentration on a site (i.e., statistical estimation error due to limited sample sizes and lab analytical error), or average dose rates used in a food chain model (i.e., due to imprecision about all of the input parameters related to ingestion rates, COC concentrations in dietary items, etc.). The precision of estimates of these parameters can be improved by increasing sample sizes.

- **Systematic measurement error** (i.e., bias) resulting from inaccurate estimation or analytical techniques. For example, a mark-recapture program to estimate the abundance of a fish population may systematically underestimate the true abundance if a subset of the fish is not susceptible to the fishing gear. In some cases biases may be known and can be adjusted for, but in other cases they may be unknown.

- **Structural or model uncertainty** that reflects our limited understanding of the mechanisms driving risks. For example, we may fail to understand how an exposure pathway works, and therefore our empirical or mechanistic models may not reflect reality very well. Structural uncertainty can be addressed in part through the use of alternative or flexible model forms. Even where underlying processes are well known, models are deliberately developed to be simplifications of reality.

- **Ignorance** reflecting our failure to recognize mechanisms driving risks. For example, we may fail to recognize a relevant exposure pathway completely. True ignorance is, by its definition, unknown, and will not be captured in CSMs or in quantitative models used to estimate risks.

For more detail on types of uncertainties see Finkel (1990) and Morgan and Henrion (1990).

- **Site investigation**: Data gaps in site investigation that could affect any aspect of the problem formulation should be identified and brought to the attention of site custodians and site investigators. In some cases, substantial data gaps (e.g., lack of surface soil data for a large portion of a site) may warrant delaying finalization of the problem formulation until the data are collected.

- **TRVs**: Although TRV determination often occurs during the hazard assessment phase, identifying available TRVs early on can result in seeing the TRV gaps in advance and provide for better planning for those gaps.

The problem formulation will always be based on uncertain information. The risk assessor should identify the key uncertainties, specify which data gaps are most critical and specify the assumptions made in moving forward with implementing the ERA.
3 EXPOSURE ASSESSMENT

The general purpose of exposure assessment is to characterize the mechanisms by which receptors are exposed to COCs, and to quantify or categorize the magnitude of those exposures. Exposure and effects are matched together in one or more ways for every line of evidence that is evaluated in an ERA. Consequently, exposure assessment is not a single step in ERA, but is carried out for every line of evidence. In many cases, the same exposure information is used in multiple lines of evidence (e.g., COC concentrations often make up the exposure information that is matched to several different types of effects information). Importantly, while the details of exposure assessment are discussed in this section, they must be fully understood and articulated at the problem formulation stage in order to support design and planning of the ERA.

3.1 Overview of Exposure Assessment

Exposure assessment used to support any particular line of evidence generally entails the following elements (all of which need to be fully contemplated during problem formulation):

- Determine which type(s) of exposure measures will be used from among the following four broad types:

  1. **External exposure media** are media such as surface water, porewater, sediment, soil or food items to which a receptor is exposed. For example, soil invertebrates are expected to be exposed to COCs in soil. In some cases where external exposure media are the measure of exposure, an ERA can rely on site investigation data without additional data collection. However, in other cases it may be preferable to have concurrent exposure data that can be more precisely matched to effects data.

  2. **Internal exposure media** are tissues where contaminant concentrations are measured to represent exposure within the receptor itself. For example, mercury concentrations in fish tissue can be used as an indicator of mercury exposure. In general, internal exposure media are more relevant than external exposure media for COCs that bioaccumulate or biomagnify up the food chain, and can be used whenever matching effects data to which the data can be compared are available.

Key Concept

Exposure information is an input for every line of evidence in an ERA.
3. **Estimation of total dose.** For example, a small mammal may be exposed to COCs by ingesting surface water, ground insects or other food sources, or through incidental ingestion of soil. The cumulative intake of COCs by all pathways forms the total dose. Typically, estimation of total dose is implemented with a food chain model.

4. **Categorical measures of exposure** do not explicitly rely on any information about contaminant concentrations, but instead categorize exposure in a simple manner. Common examples of categorical exposure measures are:

- on-site versus reference condition
- site versus lab control
- spatial gradient categories such as near-field, mid-field and far-field.

Categorical measures of exposure are often used implicitly, but risk assessors should be explicit about their use for any line of evidence that depends on the categorical measure. For example, if bird densities are used as an effects measure comparing on-site and reference conditions, the implicit assumption is that on-site exposure is different from the reference condition. There may be information on COC concentrations in some media (e.g., soil) but perhaps not for other media (e.g., food item tissues). In that case, exposure for the line of evidence may not be characterized as COC concentrations. Rather, exposure may be characterized using the implicit categories for on site and reference.

- Determine whether the exposure data will be directly measured or estimated. Usually, concentrations of COCs in abiotic media (e.g., soil, sediment, water) are measured directly, but in some cases they are estimated (e.g., using fate and transport models). Concentrations of COCs in biotic media (e.g., tissues) are more often estimated (e.g., predicted using uptake factors), but estimation methods are uncertain, so preference is for direct measurement whenever possible.
• Determine how the data will be packaged to represent exposures for various receptors of
cconcern. For example, will maximum values be used, or will some kind of statistical metric
of the data be used to represent exposures (e.g., 95 per cent UCLM)?

**Definitions**

*Biotic media* are biological tissues where COCs may be found, whereas *abiotic media* are any other
environmental media (i.e., soil, sediment, water, air).

*Uptake factors* are the ratio of COC concentrations in tissue to the COC concentrations in an abiotic
medium such as soil or water.

• Determine what ancillary data will be collected in addition to COC concentrations,
including data related to evaluation of bioavailability.

• Characterize uncertainties in exposure, evaluate the implications of uncertainty using
sensitivity analysis, and, if warranted, integrate uncertainties into the exposure assessment
using probabilistic methods.

The outcome of exposure assessment is information that can be matched with effects measures to
provide evidence in the form of a line of evidence. It is critical that the risk assessor conceptualize
the exposure and effects information at the same time (during problem formulation) to ensure that
they can be integrated effectively and to ensure that all information and ancillary data needs are
identified before data collection.

Section 3.2 compares direct measurement and estimation, which is an issue that applies to all of
the types of exposure measures with the exception of categorical measures. Section 3.3 explores
the four types of exposure measures in detail, focussing on how data will be used to represent
exposure for receptors of concern and what ancillary data will be collected in addition to COC
concentrations. Section 3.4 discusses options for moving beyond typical point estimates of
exposure.

### 3.2 Direct Measurement versus Estimation

Risk assessors must not only decide what types of exposure measure are
appropriate for a given line of
evidence, but whether to measure or estimate exposure in each case. This
section provides guidance in this
regard, for abiotic media and for
tissues.

**Key Concept**

Direct measurement of COC concentrations in any
medium is preferred over estimation, particularly for
detailed ERAs, because of the much lower uncertainty
associated with direct measurement. However, there are
cases where estimation may be suitable or may be the
only feasible option.
3.2.1 Direct Measurement versus Estimation for Abiotic Media

Whenever abiotic media such as soil, water and sediment are used as measures of exposure, data on COC concentrations must be either measured or estimated. Direct measurement is most common for abiotic media, and is generally preferred for detailed ERAs because:

- there is much less uncertainty regarding measured COC concentrations compared to estimated concentrations
- many informative ancillary variables cannot be practically predicted and must be measured (e.g., pH, SEM:AVS)
- there are usually significant data available for soil and other media as a result of site investigations
- the cost of collecting additional chemistry data in abiotic media is generally not prohibitive.

However, there are cases where measurement is not possible, not practical or not necessary, and estimation of COC concentrations is preferred. This may occur, for example, when:

- an ERA is evaluating a future scenario under which current measured values are not directly relevant
- chemistry data cannot be collected safely (e.g., sediment in a river with difficult access)
- it is anticipated that the added accuracy provided by direct measurement would not affect characterization of risks or decisions regarding risk management.

COC concentrations in abiotic media are estimated using simple or complex models that predict the fate and transport of contaminants in the environment. A simple model would be one that does not predict transport of COCs but simply predicts concentrations in one medium from concentrations in another medium based on chemical properties. For example, the partitioning of organic compounds from water into the organic matter of sediments can be predicted based on the octanol-water partitioning coefficient ($K_{OW}$).

More complex models take into account the complex interactions of contaminant loadings, movement and partitioning into various media (Cowan et al. 1995). An example of a complex fate and transport model would be one that predicts contaminant concentrations in a section of river based on information about loadings and water flows in upstream tributaries. Developing fate and transport models can be expensive, and their relative advantages and disadvantages should be carefully considered.
3.2.2 Direct Measurement versus Estimation for Tissues

Whenever tissues are used as either a measure of internal exposure or as a food item for a higher-trophic-level receptor, the tissue concentrations of COCs can be either measured directly or estimated. Normally, screening-level ERAs are dependent upon data obtained from the environmental site assessments, and since these rarely contain tissue concentrations, where tissue concentrations are needed, screening-level ERAs more often depend on methods of estimating tissue concentrations from those of abiotic media. Otherwise, direct measurement is relatively common for some tissue types, such as plants, invertebrates and fish, but less common for other tissue types, such as mammals and birds. Whenever feasible, direct measurement is usually preferred over estimation because there is much less uncertainty regarding measured COC concentrations in tissues. However, estimation may be appropriate in some cases, including:

- when time constraints for the ERA preclude waiting for seasonal tissues (e.g., berries, tree leaves, bird eggs)
- for organisms, sites or media for which it is considered inappropriate to sacrifice individual organisms for purposes of obtaining data
- when an ERA is evaluating a future scenario under which current measured values are not directly relevant
- for purposes of generating initial risk estimates on a limited budget.

Importantly, it may be efficient to use a combination of measurement and estimation for tissues at large sites. Specifically, if a relationship can be established between, for example, soil and tissue concentrations to characterize the soil media, that relationship could then be extrapolated to other samples where only soil is available.

For cases where COC concentrations in tissues are estimated, at least three methods are available, each with advantages and disadvantages as follows:

1. **Uptake factors**: the ratio of the contaminant concentration in tissue to the concentration in an associated abiotic medium (e.g., water, soil or sediment). Uptake factors based on water are commonly referred to as bioconcentration factors (BCFs) or bioaccumulation...
factors (BAFs).\textsuperscript{22} Uptake factors are generally very uncertain, and they should be avoided if bioaccumulation regression models (below) are available. Published uptake factors are available for a range of contaminants and tissue types (Sample \textit{et al.} 1998; Suter \textit{et al.} 2000 and references therein), but these should be viewed as examples only. Risk assessors should seek out the most recent scientific literature as part of any detailed ERA and determine which uptake factors are applicable to a given site. Importantly, the units used in uptake factors (e.g., wet weight, dry weight, lipid normalized) must be the same as the units for the site-specific data, or must be converted to be equivalent.

2. \textit{Bioaccumulation regression models}: these models are superior to simple uptake factors for two reasons. First, they allow for inclusion of variables other than contaminant concentrations (e.g., using multiple regression approaches), which ultimately are capable of explaining more of the variation in the tissue data. Second, regression models are capable of accounting for nonlinearity in the relationships between soil and tissue concentrations. Nevertheless, uncertainties in regression models are typically high. As with simple uptake factors, summaries of bioaccumulation regression models are available for a range of contaminants and tissue types (Sample \textit{et al.} 1998; Suter \textit{et al.} 2000 and references therein), but again these should be viewed as examples only. Risk assessors should seek out the most recent scientific literature as part of any detailed ERA and determine which models are applicable to a given site.

3. \textit{Mechanistic bioaccumulation models}: these models are based on details of the physiology of the organism (e.g., metabolic transformation) and the behaviour of the contaminant (e.g., solubility and partition coefficients). Mechanistic models are data-intensive and complex, and therefore can rarely be developed on a site-specific basis.\textsuperscript{23} Moreover, such models may suffer from larger uncertainties than simple empirical models, due to cumulative uncertainties in modelling several mechanistic processes that may be poorly understood.

In practice, many exposure assessments may use a combination of measured and estimated tissue concentrations simultaneously. For example, when exposure is based on evaluation of total dose, some food item tissue concentrations may be measured, while others may be estimated.

3.3 \textbf{Types of Exposure Measures}

The key decision in exposure assessment is determining what type of exposure measure to use for a particular line of evidence in an ERA.

This section distinguishes four broad types of exposure measures:

\textsuperscript{22} In strict terms, BAF is intended to apply to the ratio between tissue and exposure medium (i.e., water) where all exposure pathways are considered simultaneously, whereas BCF is intended to refer to an exposure condition that includes water only. In common usage, however, the term BCF is often used to refer to the quantity that is more correctly described as BAF.

\textsuperscript{23} A few examples of mechanistic bioaccumulation models are referenced by Suter \textit{et al.} (2000).
• external exposure media to which a receptor is exposed (e.g., surface water, porewater, sediment, soil, food item tissue)
• internal exposure media that describe contaminants within the receptor itself
• estimation of total dose (e.g., estimate of dietary intake through food chain modelling)
• categorical measures of exposure (e.g., on site versus reference condition).

The decision about what type of exposure measure to use should be based on the following:

• The level of effort needed to collect the data, balanced against the need for precise information. For example, collecting tissue data for input into a food chain model may not be warranted until potential risks are first evaluated using conservative measures that require less effort to collect.

• Availability of matched effects data against which the exposure tool outputs can be compared. For example, measures of contaminants in external exposure media such as soil can be compared to benchmark concentrations associated with effects on plants or invertebrates for that medium. Measures of contaminants in internal exposure media such as fish tissues can be compared to critical body residues as a measure for the aquatic medium.

Importantly, a single exposure measure may be used in several lines of evidence. The manner in which exposure measures are used should be defined up front in the problem formulation, as there is little value in identifying a measure without also clarifying how it will be used.

### 3.3.1 External Exposure Media

External exposure media are any media to which a receptor is exposed. For example, soil is an external exposure medium for terrestrial invertebrates. External exposure media include not only abiotic media such as soil, water, sediment and air, but also food item tissues. In the case of strongly bioaccumulative and biomagnifying substances, tissues are usually the most relevant external exposure medium for higher trophic level receptors, due to the high proportion contributed to total dose.

This section focusses on how external exposure data are used to represent exposure for receptors of concern, and what ancillary data will be collected in addition to COC concentrations. Table 3-1 provides an overview of typical methods of evaluating each external exposure medium, common ancillary parameters and key challenges.

This section does not review field methods of collecting soil, surface water, groundwater, sediment, porewater and tissue data, as these methods are addressed in detail elsewhere by CCME (2016), EC (2011), Mudroch and MacKnight (1994), US EPA (2007-b) and, State of Washington *et al.* (2015).
3.3.1.1 Soil

Soil contaminant concentrations are very commonly used as a measure of external exposure, in particular for characterizing exposure for plants and soil invertebrates, but also for characterizing some exposure pathways for wildlife. Soil data that are typically collected as part of site investigation are rarely completely adequate for risk assessment unless risk assessors have been involved up front during site investigation. Where soil data do not meet the needs of exposure assessment as defined in this section, supplemental data collection may be warranted, particularly for detailed assessments.

Defining surface soil: As a default, all soil data in the top 1.5 metres can be considered as surface soils for purposes of measuring exposure for plants and soil invertebrates, as well as for higher-level receptors. This approach is consistent with both the default approach used during COC selection and the Canada-wide standard for petroleum hydrocarbons in soil (CCME 2008-a). Where jurisdictions have specific requirements or definitions for sampling depths related to surface soils, these should be followed. The depth requirements for sampling and analysis may also need to be defined more precisely to be relevant for the specific receptor of concern. The risk assessor should take into account:

- The depth of bioturbation due, for example, to burrowing insects, burrowing mammals and plant root systems. For deep-rooting plants and trees, it may be necessary to consider exposure to COCs at soil depths greater than 1.5 metres, whereas for insects the depth may be much shallower.

- Applicable regulatory policy (for sites intended for divestiture) that require that a different depth be considered for surface soil exposure.

- Differentiation of soil layers for certain receptor groups. For example, some receptors may be limited to the humic layer rather than underlying mineral soil (e.g., organisms that play a role in decomposition of organic matter). If there are large differences in COC concentrations between these two layers (e.g., airborne mercury typically accumulates in the humic layer), it may be appropriate to use soil data from only the humic layer.

- The depth that is likely to have been affected given the sources and nature of contamination at the site. For example, for an air deposition source there may only be a shallow surface layer (e.g., top 2–5 centimetres) that is contaminated and that should be used exclusively for understanding particular exposure pathways such as incidental ingestion.

- Natural processes or planned activities at the site that will result in accumulation of soils or removal of surface soils that will expose soil at depths. In such cases, the soil layer relevant for current exposure for a particular receptor may not be the same as the soil layer that is relevant for future exposure.

**Key Concept**
As a default, all soil data in the top 1.5 metres can be considered as surface soils for purposes of measuring exposure. However, site-specific (and receptor-specific) depths should be defined when precision in soil exposure estimates is warranted.
### Table 3-1: Sampling considerations for external exposure media

<table>
<thead>
<tr>
<th>Exposure medium</th>
<th>Typical tools for measurement/estimation</th>
<th>Typical ancillary parameters</th>
<th>Cautions and key issues</th>
<th>Guidance</th>
</tr>
</thead>
</table>
| Soil            | • Measurement of bulk soil chemistry, based on collection by trowel or auger | • Site-specific, but may include organic matter content, pH, moisture content, soil texture and cation exchange capacity (CEC) | • Sample depth | • Suter et al. (2000)  
|                 |                                          | • Differentiation of soil layers | • Spatial design and resolution | • CCME (2016) |
| Surface water   | • Measurement of total or dissolved concentrations using typical water-sampling gear | • Site-specific, but may include hardness, pH, alkalinity, acidity, temperature, dissolved oxygen, anions, cations, nutrients, conductivity, salinity, total suspended solids and dissolved organic carbon | • Temporal variability, including seasonality | • EC (2011)  
|                 |                                          | • Understanding relevance of bulk sediment versus porewater for each receptor type | | • CCME (2012)  
|                 |                                          | • Sample depth | | • Suter et al. (2000)  
| Sediment and sediment porewater | • Measurement of bulk sediment chemistry using grabs, divers or cores  
|                 | • Measurement of sediment porewater chemistry (dissolved) by extraction from sediments, or directly (e.g., using push-point samplers) | • For sediment: organic carbon, particle size, pH, sulphides, SEM:AVS, possibly iron and manganese hydroxides  
|                 |                                          | • For porewater: redox, plus similar parameters to surface water | • Understanding and addressing oxygenation of porewater samples during collection and transport | • EC (2011)  
|                 |                                          | | | • CCME (2012)  
|                 |                                          | | | • Mudroch and MacKnight (1994)  
|                 |                                          | | | • Suter et al. (2000)  
|                 |                                          | | | • State of Washington et al. (2015)  
| Groundwater     | • Measurement of dissolved concentrations using typical groundwater sampling gear | • Redox, plus similar parameters to surface water | • Understanding groundwater flow characteristics | Can be used as surrogate for porewater (e.g., for direct effects on plants and soil invertebrates), or for predicting future surface water impacts |
| Air/vapour      | • Rare for ERA, but direct measurement and modelling are both used (see text) | | | |
| Tissues         | • Direct measurement preferred  
|                 | • Estimation using uptake factors or models | • Lipid content  
|                 |                                          | • Moisture content | | • Suter et al. (2000)  
|                 |                                          | • Consideration of whether to test whole organism or selected tissue types | | • CCME (2012)  
|                 |                                          | • Consideration of whether to depurate, depending on how tissue data will be used | | • Beyer and Meador (2011)  
|                 |                                          | | | • State of Washington et al. (2015)  
|
Using soil data as an exposure measure:
The key question for the risk assessor in using soil data is whether to measure exposure to soil based on single soil samples or using statistical measures, both horizontally and vertically. For plants and soil invertebrates, the default for spatial characterization should be to measure exposure on a sample-by-sample basis. Where there is a sufficient sample size for each area of environmental concern (e.g., >10), it may be appropriate to consider using summary statistics for each area (e.g., 95 per cent UCLM and 90th percentile). In this type of approach, the assessor will have to consider UCLM, an upper distribution for the population as a whole and the maximum observed concentration for all samples. This will prevent significant injury in any location within the area of environmental concern for the given land use. Vertically, the soil data that are used (either sample by sample, or with summary statistics) must be only the soil data that are relevant for a particular receptor group. There is no point considering deep soil data for shallow-rooting plants.

While plants and immobile soil invertebrates will be affected locally by elevated COC concentrations at a single soil sample location, the spatial scale at which potential major risk management measures would be implemented is also relevant. In other words, exposure (and risks based on a given line of evidence) for plants and soil invertebrates should be understood at more than one spatial scale, because the spatial scale is an important component of the magnitude of risk estimates (see Section 5 for further discussion).

For mammals, birds and other wildlife receptors exposed to soil, incidental soil ingestion can sometimes be the most important exposure pathway. Where acute effects are expected from exposure to high COC concentrations in a particular sampling location or where sample size is small (e.g., <10) relative to the home range for the organism, exposure to the receptor of concern should consider the maximum concentration observed within the sampling location. Otherwise, it is appropriate to consider exposure for the receptors of concern based on summary statistics such as the arithmetic mean, the 95 per cent UCLM or the 90th percentile. Interpretation of summary statistics should take into account current CCME (2012) guidance on sampling for contaminated sites. The risk assessor should also determine whether the spatial layout and density of soil samples collected during site investigation or other evaluations is adequate to support assessment of risks for each surrogate VEC.

Ancillary parameters: Ancillary parameters that are often relevant for soil include:

- **Percentage of organic matter:** Organic matter is important for organic compounds that partition predominantly into lipids (i.e., have a high $K_{OW}$). In such cases, soil concentrations of COCs may be more appropriately characterized using organic carbon-normalized concentrations.
• **pH**: pH data are important for understanding general soil conditions, including the likely solubility, speciation and complexation of metals. At extreme pH, the data can be useful in predicting plant stress as well as presence or absence of biota.

• **Moisture content**: Moisture data are important if soil data will be used in food chain models, because data related to incidental soil ingestion rates may be based on dry- or wet-weight concentration units.

• **Cation exchange capacity (CEC)**: CEC is the maximum quantity of total cations that soil can hold. Clay and humus typically have higher CEC than sandy soils. This property can be useful in determining the relative bioavailability of metals, because lower-CEC soils are more likely to release metals to biota.

• **Redox potential (Eh)**: Eh is an electrical measurement characterizing the transfer of electrons in soils to or from a reference electrode. Eh can be used to determine if soil is anaerobic (low Eh) or aerobic (high Eh), which can affect the dissolution or precipitation of various metals.

• **Soil texture and composition**: Texture is the relative proportions of sand, silt and clay in soil. Structure refers to the aggregation of soil particles into larger secondary clusters, typically developed through the action of microbes or invertebrates. Both texture and composition can affect contaminant dynamics in soils.


**Evaluating bioavailability**: ERAs may use soil data collected during site investigation to initially characterize contamination in soil (e.g., whole soil samples and analysis of bulk soil chemistry). However, where precise understanding of risks is warranted, risk assessors should consider using speciation analyses or other methods that will more precisely characterize the contaminants in the soil. In addition, extraction techniques may be considered for characterizing the fractions that are more likely to be bioavailable (Allen 2002; Suter et al. 2000). Alternatively, if soil porewater is considered to be the relevant exposure medium (e.g., for plant roots), then soil porewater can be either measured or estimated from bulk soil chemistry using equilibrium partitioning models. Further discussion on these approaches is found in Allen (2002) and Suter et al. (2000). Finally, studies that simulate bioavailability in the human gastrointestinal tract (referred to as bioaccessibility tests or physiologically based extraction tests) are now used in human health risk assessment (see early work by Ruby et al. [1996]), and are becoming more prevalent in ERA as well. Results from methods intended to simulate the human gut may be directly relevant to mammals that have similar anatomy and gut conditions (e.g., pH) to those of humans. Alternatively, specific test protocols may be modified for other species. While the jurisdiction will need to be consulted on acceptability of methods to determine bioavailability, standards are emerging for use of bioavailability in soil (see, for example, ISO [2008-a,-b] and Jensen and Mesman [2006]).
3.3.1.2 Surface Water

Surface water exposures occur through direct contact (e.g., for aquatic plants, fish or benthic epifauna) or by ingestion (e.g., for wildlife).

Defining surface water: For purposes of ERA, surface water is water that is above the sediment-water interface in any aquatic system. Surface water can also be temporary pools or watercourses that provide aquatic habitat, drinking water, or other potential exposure routes for terrestrial species that may come into contact with the contaminated pool. Surface water is distinguished from sediment porewater, which is water in the interstitial spaces within the sediment. Importantly, surface water rather than (or in addition to) sediment porewater may be a relevant external exposure medium for some organisms that live in the sediment. For example, clams are buried in sediment but are exposed to surface water via their siphons, which filter water directly from the sediment–surface water interface.

Using surface water data as an exposure measure: As with soil, a key challenge for the risk assessor is deciding whether to use maximum measured COC concentrations or some statistical metric over space or time, for each receptor of concern. For sessile organisms (e.g., aquatic plants), maximum concentrations may be appropriate to represent concentrations in small areas, but statistical measures (e.g., 95 per cent UCLM and 90th percentile) can also be used to characterize average exposures in particular areas. For mobile receptors, maximum concentrations are recommended as a default if there are few samples (e.g., < 10) in the area covered by their home range size, or if seasonal variability in COC concentrations is expected but has not been measured. In cases where sample sizes are large and seasonal variability is captured (if warranted), summary statistics can be used (e.g., 95 per cent UCLM and 90th percentile). Importantly, where surface water data are used to represent drinking water exposure for wildlife, the risk assessor should consider the number of nearby options for drinking water and the proportion of total exposure that is likely to come from any one source. In such cases, statistical measures of surface water COC concentrations may be based on averaging across the sources rather than averaging across pooled samples (e.g., if one drinking water source has three samples and another has 20 samples, a 95 per cent UCLM based on the pooled samples will bias towards the second source).

A second issue that the risk assessor must consider is whether to use dissolved concentrations, total concentrations, or both as the measure of exposure. This decision may be affected in part by regulatory requirements, but should also take into account relevance for ERA. If the surface water data will be used for more than one purpose (e.g., as external exposure media for fish, as well as drinking water for wildlife), the data should be appropriate for all purposes. Total concentrations are most relevant for ingestion pathways, whereas dissolved concentrations (see the following bioavailability discussion for more information) are more relevant for direct contact pathways.

The proportion of “total” versus “dissolved” can be a moving target depending on the parameter and site conditions. As a default, risk assessors should use total concentrations in water, as this measure is more conservative, but in some cases may use dissolved concentrations provided that
rationale is given. In either case, it is important to make sure that the exposure data will be comparable to available effects data (e.g., exposure data based on dissolved concentrations should not be compared to effects data based on total concentrations). The majority of toxicity guidance developed for aquatic life is presented as total concentrations.

Finally, in most cases the specific form of dissolved contaminants does not need to be quantified for ERA. However, for some sites where the relative toxicity of a COC is highly dependent on its form, speciation analysis and risk assessment based on speciation may be appropriate. For example, different forms of iron in sediment porewater may differ in toxicity by several-fold, and efforts to link toxic responses in porewater bioassays to a potential causal effect of iron may require understanding of the relative concentration of each iron species in the porewater samples.

Bioavailability: Dissolved contaminants in water are not necessarily bioavailable. For example, research over the last two decades regarding metals bioavailability and mechanisms of toxicity in the aquatic environment has led to development of the biotic ligand model (BLM) (Di Toro et al. 2001; Paquin et al. 2003; see Section 4.2.2 for more details). This model accounts for the roles of total suspended solids, pH, dissolved organic carbon, cations (Ca, Mg, Na, K), anions (SO₄, Cl), alkalinity, hardness and sulfide in determining free metal ion concentrations in affecting metals bioavailability (and ultimately toxicity) in freshwater. Many of the typical ancillary parameters listed below are used to support understanding of potential bioavailability, including through use of the BLM.

Ancillary parameters: Typical ancillary parameters that are measured in surface water depending on the site and the ERA include the following:

- hardness
- pH (pH may also be a COC)
- alkalinity
- acidity
- temperature
- dissolved oxygen
- anions and nutrients (e.g., chloride, bromide, fluoride, nitrite, nitrate, sulphide, sulphate)
- cations (e.g., Ca, Mg, Na, K)
- conductivity
- salinity (for sites at the interface of freshwater and marine)
- total suspended solids
- dissolved organic carbon.

This is not an exhaustive list. In general, any parameter that is expected to provide useful information should be considered. Some parameters such as pH can be measured in both the lab
and the field. Generally lab equipment will be more accurate, but there is a subset of field measurements that should always be taken in the field, as they can be expected to change during storage and transport (typically temperature, pH, dissolved oxygen and conductivity).

3.3.1.3 Groundwater

Defining groundwater: For purposes of ERA, groundwater is any water that is not surface water and is not considered to be within the biologically active layer of surficial soil or sediment.

Using groundwater data as an exposure measure: Groundwater should generally not be considered as an exposure medium in ERA. Although stygo fauna (small, aquatic organisms that live within groundwater systems, such as caves and aquifers) may be directly exposed to groundwater, their adoption as assessment endpoints is rare.

Groundwater may be applied as a surrogate for exposures to organisms that live in soil or sediment porewater or even surface water in certain cases, such as:

- a preliminary or screening-level assessment based on existing data, where groundwater was collected during upland site investigations but soil or sediment porewater was not collected
- a conservative assessment where groundwater chemistry is used to represent worst-case exposure
- for sites where soil or sediment porewater is very difficult to access (e.g., if a foreshore is covered by rip-rap)
- for sites where a groundwater plume is migrating towards a surface water body but has not yet reached the surface water body. In this case, the groundwater may be considered somewhat representative of potential future discharges to surface water.

Ancillary parameters: Most of the same ancillary parameters for surface water should be measured in groundwater, with the addition of redox potential. A key consideration for groundwater assessment is that ancillary parameters such as redox potential and pH are likely to differ from those measured as the groundwater enters the transition zone where it interfaces with surface water. These ancillary parameters can have significant effects on the bioavailability of contaminants (e.g., metals that are dissolved in groundwater may precipitate out quickly as the water becomes oxygenated in the transition zone).

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24 See Environmental Canada (2010-c) for exceptions and further discussion.
3.3.1.4 Sediment and Sediment Porewater

**Defining sediment and porewater:** For purposes of ERA, sediment is the substrate in an aquatic feature, and sediment porewater is the water found in the interstitial spaces of the sediment. Sediment and sediment porewater are the primary exposure media for benthic invertebrates, particularly benthic infauna, and also for many aquatic plants or algae. Direct exposure may also be relevant for the early life history stages for some higher-level organisms (e.g., fish eggs). Incidental ingestion of sediment is also an important exposure pathway for some higher-level receptors such as bottom-dwelling fish and aquatic birds.

**Defining surface sediment:** As with soil, the depth of sediment that is relevant to ecological receptors should be carefully considered. As a default, and consistent with the default for COC selection, all sediment data in the top 1 metre can be considered for purposes of measuring exposure. However, for cases where more precision is warranted, the depth of surface sediment should be defined on a site-specific basis, taking into account:

- the depth of bioturbation due to flora and fauna (e.g., worms, bivalves)
- applicable policy at provincial or other order of government (for sites intended for divestiture) that require that a different depth be considered for surface sediment exposure
- natural processes or planned activities at the site that will result in deposition, erosion or removal of surface sediments that will expose sediments at depths.

**Using sediment and porewater data as exposure measures:** Typically, bulk sediment is used as the initial indicator of external exposure. Almost all environmental quality guidelines are based on bulk sediment and not on porewater, and so initial characterization of sediments focuses on that medium. However, porewater is often the medium in which contaminants are most likely to be biologically available, as opposed to the portion bound to particulate matter. Sediment and sediment porewater may be appropriate for use as external exposure media in ERA. Bulk sediment is recommended as the default external exposure medium for most cases because:

- effects data are more commonly associated with sediment, so bulk sediment chemistry is more likely to contribute to lines of evidence for the ERA
- sediment concentrations are less likely to change on short time scales (e.g., tidal fluctuation) or even longer time scales (e.g., seasonality), with the exception of patterns of deposition and scouring
- sediment sampling and analysis is relatively straightforward compared to porewater sampling and analysis, as the latter is influenced by a specific extraction technique and sample handling and preservation methods.

**Key Concept**

An ERA may characterize exposure using bulk sediment, sediment porewater, or both.
However, porewater should be evaluated in many cases (usually in addition to bulk sediment), such as:

- when there is ongoing transport of COCs in dissolved phase to the aquatic environment via groundwater
- cases where COCs are likely to partition predominantly into water and not adsorb to sediments
- when increased precision is desired in relating effects measures to bioavailable (dissolved) contaminant concentrations
- when effects measures that will be matched to the exposure data are based on porewater (e.g., porewater bioassays).

Risk assessors should not assume that bulk sediment alone is sufficient for any particular ERA.

**Ancillary parameters and bioavailability:** Ancillary parameters of importance for porewater are the same as those listed earlier for surface water and groundwater. Ancillary parameters of importance for bulk sediment typically include:

- **Organic carbon content:** Organic carbon is the most important factor determining partitioning of organic compounds into sediments.

- **Particle size (e.g., percentage clay, silt, sand and gravel):** Because the ratio of surface area to sediment volume is higher for finer sediments, evaluation of patterns of COC concentrations in sediments can be confounded by the influence of particle size (e.g., COCs are more likely to be bound up in finer sediments).

- **pH:** In bulk sediment, pH is an indicator of the general environment and types of receptors that could be expected to be present.

- **Sulphides:** In anaerobic sediments, sulphides are normally the predominant binding phase, and measurement of SEM:AVS\(^{25}\) can provide insights into the potential bioavailability of cadmium, copper, lead, nickel and zinc. The SEM:AVS model can also incorporate organic carbon (US EPA 1999).

- **Iron and manganese hydroxides (for metals):** In aerobic sediments, iron and manganese hydroxides can be an important binding phase. Sequential extraction techniques employ a series of chemical fractionation steps to elucidate the relative importance of various binding phases (e.g., Tessier et al. 1979).

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\(^{25}\) AVS is *acid volatile sulfide*, and SEM is *simultaneously extracted metal*. If SEM − AVS < 0, then it is assumed that sufficient sulfides are available to bind the SEM metals. The SEM:AVS model does not apply to oxygenated sediments. More discussion and caveats to use of SEM:AVS are found in Paquin et al. (2003) and Suter et al. (2000).
3.3.1.5 Air and Vapour

Air is often not justified as an exposure medium in ERA. In many cases, the contribution of airborne COCs to total exposure for wildlife would be negligible, in part because the volatile compounds that are most likely to be inhaled volatilize rapidly to air and are dispersed rapidly. While inhalation exposure has been shown to be unimportant for several contaminants (US EPA 2003), relatively few studies have evaluated volatile organic compounds in detail. Nevertheless, toxicity data are available for several compounds, and screening values for evaluating potential ecological risks via inhalation have been developed and applied (Archbold et al. 2007; Gallegos et al. 2007; Markwiese et al. 2008).

While air may be ruled out as an exposure medium in many ERAs, it should be considered in certain cases, such as:

- where a site with wildlife receptors is characterized by very high concentrations of volatile organic compounds
- where a site with volatile organic compounds has wildlife receptors that burrow on the site
- where plant foliage is expected to accumulate certain contaminants (e.g., mercury, dichlorodiphenyltrichloroethane [DDT]) through uptake of vapours (Suter et al. 2000).

In such cases, as a starting point air can be sampled directly, including from existing burrows or artificial burrows (Markwiese et al. 2008), and compared to screening values such as those summarized or developed in existing literature (Archbold et al. 2007; Gallegos et al. 2007).

3.3.1.6 Tissues of Food Items

Analysis of contaminant concentrations in organism tissue is a relevant external exposure tool in cases where the tissues represent an important food item for a receptor of concern. As mentioned in Section 3.2.2, direct measurement of tissues is preferred over estimation.

**Defining food item tissues:** Food item tissues include any diet items of a receptor, but do not include incidental ingestion of soil or sediment.

**Using food item tissue as an exposure measure:** Using food item tissue as an external exposure measure is appropriate whenever there are matching effects data. For example, concentrations of biomagnifying substances (e.g., mercury, PCBs) in fish may be compared to CEQGs for tissue residue for protection of wildlife consumers of aquatic life (CCME 2001-a). In any case where food item tissues are used as an external exposure measure, the specific tissue type that is collected (e.g., muscle only or whole body) must match the effects data to which the comparison will be made.\(^{26}\) In cases where a whole organism body is used, the risk assessor should as a

---

\(^{26}\) Unless there are models available that establish relationships between concentrations in various tissues and the whole organism. If both a particular tissue type and whole body are relevant (i.e., for different purposes), the particular tissue and the remaining tissues may be submitted for analysis. In this way, a whole-body concentration can later be calculated if necessary (as a mass-weighted average).
default not have the organism depurated (e.g., intestines voided) or washed, unless the whole organism effects data were known to be based on depurated organisms.

**Key Concept**

Tissues of food items can be used in two ways.

1. If there are effects data based on the COC concentration in a food item (i.e., CEQGs for tissue residue for protection of wildlife consumers of aquatic life), then the measured or estimated tissue concentration itself is the measure of exposure that is compared to the effects data. This use of the tissue data is as an external exposure medium.

2. In contrast, and more commonly, effects data are based on total dose. This is the case for most wildlife TRVs. In this case, the COC concentration in a tissue is one input to total dose, along with tissue data for other diet items and all other relevant exposure pathways, such as drinking water and incidental soil or sediment ingestion.

**Ancillary parameters:** Ancillary parameters that are generally important with tissue sampling are lipid content and moisture content. Lipid content is particularly important for contaminants that partition strongly into the lipid fraction (e.g., PCBs), because concentrations among tissues can be meaningfully compared only when lipid-normalized. Moisture content is important so that comparisons can be made to effects measures specified in either wet-weight or dry-weight terms.

### 3.3.2 Internal Exposure Media

**Defining internal exposure media:** Internal exposure refers to measures of contaminant concentrations within the receptor itself. These include chemical concentrations in particular tissues where toxic effects occur (e.g., liver), in other tissues used as indicators of body burden (e.g., bone, hair or muscle tissue) or in whole animals. Measures of internal exposure are commonly referred to as “body burdens” or “residues.”

**Using internal (body burden) contaminant concentration as an exposure measure:** Body burdens of COCs can be used as measures of internal exposure whenever there are readily available effects benchmarks to which the exposure data can be compared. To determine whether it is possible to use internal exposure measures in an ERA, the risk assessor should:

- Review information on the behaviour of the COCs in receptors to determine if internal exposure measures would be useful. Typically, this information is summarized in the review of COC characteristics during problem formulation. Some COCs are not suitable for internal exposure analysis due to their behaviour or fate in receptors. For example, PAHs are metabolized by wildlife, and therefore body burden of PAHs may not be a useful indicator of exposure for these receptors.
• Review published studies that have derived effects thresholds based on body burdens. This requires a review of primary literature; some thresholds have been compiled by Beyer and Meador (2011) and Suter et al. (2000), but are not comprehensive.

• Review the Environmental Residue-Effects Database,\textsuperscript{27} jointly compiled by the U.S. Army Corps of Engineers and the US EPA, to determine if there are adequate data available from which effects thresholds could be derived (using methods described in Section 4 and Module 2, EC 2010-b).

• Ensure that there are practical methods of collecting the particular tissue type that would need to be matched to the effects data. If the effects data are based on whole-body concentrations or common tissue types, there may be uptake factors or bioaccumulation models that would allow estimation rather than measurement of internal exposure. For measured data, the risk assessor should as a default have the organism depurated (e.g., intestines voided) or washed to ensure comparability with available effects data.\textsuperscript{28}

If body burden data are used, results should be interpreted with caution. Organisms in field settings may be capable of acclimatizing or adapting to tolerate higher concentrations of COCs than would otherwise be expected. In such cases, actual risks may be lower than predicted risks. Conversely, because site-specific tissue data are generally collected from living organisms, risks may be underestimated if there are highly exposed organisms that have been eliminated from the population (e.g., through direct toxicity or reduced fitness).

\textit{Ancillary parameters:} Ancillary parameters of importance for measuring internal exposures are the same as those associated with measurement of tissues for purposes of characterizing food items (Section 3.3.1.6). However, any other parameters needed to support matching of the exposure data to effects data should also be considered.

### 3.3.3 Estimation of Total Dose

\textit{Defining total dose:} Exposure is often assessed for higher-level receptors (e.g., wildlife) as total dose or intake, which is the total intake of a contaminant from all exposure pathways. Total dose can be used as a measure of exposure whenever there are effects data to compare to, which may be a literature-derived dose-response relationship or a TRV.

\begin{table}[h]
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\begin{tabular}{|c|}
\hline
\textbf{Key Concept} \\
\textbf{Estimating exposure as total dose requires various types of data to characterize exposure for receptors.} \\
FCSAP Module 3 (EC 2012) provides default values for several common wildlife receptors in Canada. These can be used as a starting point, particularly for simple ERAs or initial risk estimates. For cases where more precise estimation of risks is warranted, site-specific information should be used. \\
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\end{tabular}
\end{table}

\textsuperscript{27} The Environmental Residue-Effects Database contains data from over 2,000 studies, and is available at https://ered.el.erdc.dren.mil/. This database is the most up-to-date, comprehensive source of tissue residue effects levels. It should be supplemented with current primary literature surveys to support a particular ERA.

\textsuperscript{28} In contrast, as explained in Sections 3.3.1.6 and 3.3.3, depuration is not usually appropriate as a default when tissue data are used as a food item for higher-level receptors.
Using total dose as an exposure measure: Total dose is the most commonly used exposure measure for higher-trophic-level organisms (e.g., wildlife). Total dose should always be considered for detailed risk assessments involving wildlife, unless other lines of evidence are judged sufficient to draw conclusions about risks. As explained in Section 2, ingestion pathways—water, diet items, and incidental ingestion of soil and sediment—are usually by far the most important pathways, and inclusion of dermal exposure and inhalation pathways is rarely necessary. For each ingestion pathway, the minimum data needed to estimate total dose, and the recommended sources of data, are as follows:\textsuperscript{29}

- **Ingestion rate for drinking water:** This is typically characterized as L/day or L/kg body weight/day. For receptors not covered by FCSAP Module 3 (EC 2012), water ingestion rates may be available in the primary literature or other sources (e.g., the US EPA’s *Wildlife Exposure Factors Handbook* [US EPA 1993], “Paramètres d’exposition chez les mammifères” [CEAEQ 1999-a] and “Paramètres d’exposition chez les oiseaux” [CEAEQ 1999-b]). Allometric scaling can be used for organisms for which data are not available, using equations specified, for example, by Nagy (1987).

- **Ingestion rate for food:** This is typically characterized as kg food/kg body weight/day. For receptors not covered by FCSAP Module 3 (EC 2012), food ingestion rates may be available in the primary literature or other sources (e.g., the US EPA’s *Wildlife Exposure Factors Handbook* [US EPA 1993], “Paramètres d’exposition chez les mammifères” [CEAEQ 1999-a] and “Paramètres d’exposition chez les oiseaux” [CEAEQ 1999-b]). Allometric scaling can be used for organisms for which data are not available, using equations specified, for example, by Nagy (1987). Alternatively, equations relating food ingestion to metabolic rate can be used (US EPA 1993).

- **Incidental ingestion rates for soil and sediment:** This is typically characterized as a percentage of total food intake. For receptors not covered by FCSAP Module 3 (EC 2012), incidental ingestion rates may be available in the primary literature or other sources (e.g., Beyer et al. 1996; see also CCME [2006] for discussion). These rates may vary depending on whether they account for soil and sediment contained in the digestive tract or trapped in fur (see bullet on contaminant concentrations below).

- **Body weight of each receptor:** For receptors not covered by FCSAP Module 3 (EC 2012), water ingestion rates may be available in the primary literature or other sources (e.g., the US EPA’s *Wildlife Exposure Factors Handbook* [US EPA 1993]).

- **Diet proportions for any receptor that consumes more than one type of food:** FCSAP Module 3 (EC 2012) provides default values for some common receptors. However, diet proportions are highly site-specific and vary seasonally. For sites where precision in risk

\textsuperscript{29} FCSAP Module 3 provides specific default values for many of the receptor characteristics in this list for a range of common wildlife receptors in Canada (EC 2012).
estimates is warranted, site-specific information should be collected (see FCSAP Module 3 [EC 2012] for discussion).

- **Contaminant concentrations in soil, sediment, water and each food item:** As discussed in Section 3.2, ideally contaminant concentrations in each media are measured, but they can also be estimated. The specific tissues that are sampled should match consumption patterns. There are at least three considerations in this regard:
  - Whether to submit a whole animal (e.g., small mammal) for analysis, or only parts of an animal. If receptors are unlikely to consume (and digest) certain tissues such as bones or feathers, it may be appropriate to exclude those tissues from lab analyses.
  - Whether to depurate (i.e., void the digestive tract) tissues before analysis. This applies to earthworms or filter feeders that take in volumes of soil or sediment but digest only the “food” components. It is most conservative not to depurate (except for bioaccumulative substances), and this should be the default approach. However, risk assessors should recognize that contaminants bound in the soil or sediment may not be bioavailable to higher-trophic-level consumers.
  - Whether to wash tissues before analysis. This applies to any organisms such as invertebrates, but particularly mammals with fur. Washing will remove soil trapped in fur, which is acceptable as long as estimated incidental soil ingestion rates account for this route of soil ingestion.

- **Moisture content of soil, sediment, and food items:** This will allow conversions (as necessary) between ingestion rates and COC concentrations in food items. Whether data are reported on a dry-weight or wet-weight basis, harmonizing units is essential when calculating total dose. Moisture content should be measured by labs or derived from primary literature.

- **Home-range size or forage-range size of each receptor, relative to the size of the site or relevant portion of the site:** The home-range size should be estimated based on an up-to-date literature review, but can be adjusted based on professional judgment of a wildlife biologist. For example, if habitat quality is low in the undisturbed or uncontaminated background, range size may be larger, but good-quality habitat may result in a smaller range size. A conservative screening assessment may assume that a receptor spends all of its time on a site. Where the home-range size is larger than the site, more realistic assessments could apportion exposure between on site and off site (which requires data for off-site exposures). This is particularly important for large mammals or other receptors that may spend only a very small portion of time on a site.

30 Depuration is also not recommended as a default when diet items are used directly as an external exposure measure, for example when comparing to CEQGs for tissue residue for protection of wildlife consumers of aquatic life. In contrast, when tissues are collected and evaluated as an internal measure of exposure for the organism itself, depuration is usually appropriate; see discussion in Section 3.3.2.
• *Other dose adjustment factors (DAFs):* These can account for partial bioavailability (or any other factor that is believed to affect actual dose) and may also be used in more realistic models. Most TRVs are developed from studies conducted using highly bioavailable forms of contaminant (e.g., soluble metal salts) that may overestimate actual availability from site media. In the absence of specific information about bioavailability, risk assessors should assume 100 per cent bioavailability, although this will typically contribute to overestimation of exposure.

*Food chain models:* Simple models to calculate total dose can be formulated in a spreadsheet. The Key Concept box at the end of this subsection provides example equations. Complex models covering several COCs, several receptors and several distinct areas of a site may warrant a more elaborate set-up than a series of equations in a spreadsheet. For example, to avoid repetition of formulae in a spreadsheet, it may be appropriate to host input data in a spreadsheet or data file, then use a programming language to read the data and perform all calculations (e.g., Visual Basic for Applications, if the input data are contained in Microsoft Excel), and then output the results back to the spreadsheet or a data file. Visual Basic for Applications can also call other software that is useful for particular functions (e.g., Crystal Ball for probabilistic models). Software packages designed specifically for risk assessment are also available (e.g., GoldSim). Using more than one method of estimating total dose can be valuable for detecting errors (e.g., QA/QC check of the model).

<table>
<thead>
<tr>
<th><strong>Key Concept</strong></th>
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<tr>
<td>Food chain models are a series of equations that can be set up in a spreadsheet, though more elaborate models warrant programming.</td>
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Key Concept

Food chain models can be formulated in a spreadsheet using a series of equations. The following is an example set of equations:

1. **Food ingestion rates**: Food ingestion rates \((FI, \text{ kg dw/kg ww/day})\), if not known for a given receptor, can be estimated using allometric equations such as those described in Nagy (1987) for various feeding guilds, i.e.,

   \[
   FI = a \times BW^b
   \]  
   \((\text{Eq. 1})\)

   Where:

   \(BW\) represents the organism’s mean body weight \((\text{g, ww})\)

   \(a\) and \(b\) are constants specific to various groups of terrestrial vertebrates

   These dry weight food ingestion rates can be converted into wet weights \((IF, \text{ kg ww/kg ww/day})\) following equation 2:

   \[
   IF = \frac{FI}{(1 - \text{Moisture}_{\text{diet}})}
   \]  
   \((\text{Eq. 2})\)

   Where:

   \(\text{Moisture}_{\text{diet}}\) (unitless fraction) represents the weighted average moisture content in the diet of the animal, based on measured contents in tissues from the site or values from the literature.

2. **Soil and sediment ingestion rates**: Soil and sediment ingestion rates \((IS, \text{ kg dw/kg ww/day})\) are based on an estimated fraction of incidental ingestion during foraging activities. If not known for a given receptor, they can be derived from the food ingestion rate according to:

   \[
   IS = FI \times \phi
   \]  
   \((\text{Eq. 3})\)

   Where:

   \(FI\) (\text{kg dw/kg ww/day}) is the dry food ingestion rate

   \(\phi\) is the fraction of incidental soil or sediment ingested during feeding.

3. **Drinking water ingestion rates**: Drinking water ingestion rates \((IW, \text{ L/kg ww/day})\), if not known for a given receptor, can be estimated using allometric equations such as those described in Nagy (1987):

   \[
   IW = a \times BW^b
   \]  
   \((\text{Eq. 4})\)

   Where:

   \(BW\) (\text{kg, ww}) represents the organism’s mean body weight

   \(a\) (\text{L/kg*kg/day}) and \(b\) (unitless) are constants specific to various groups of terrestrial vertebrates.

4. **Dose from food**: Intake dose of contaminants from food \((DF, \text{ mg/kg bw/day})\) determined from the dietary concentration following:

   \[
   DF = IF \times \sum_j(C_{F_j} \times p_{F_j})
   \]  
   \((\text{Eq. 5})\)

   Where:

   \(IF\) (\text{kg ww/kg bw/day}) represents the feeding ingestion rate

   \(C_{F_j}\) (\text{mg/kg ww}) represents the COC concentration in prey item \(j\) in the diet of the receptor of concern

   \(p_{F_j}\) (unitless) represents the proportion of prey item \(j\) in the diet of the predator

continued on next page
5. **Dose from soil intake (primarily terrestrial foragers):** The total dose from incidental ingestion of COC contaminated soil \((D_S, \text{mg/kg-bw/day})\) calculated using the following equation:

\[
D_S = I_S \times C_S
\]  
(Eq. 6)

Where:
- \(I_S\) (kg dw/kg bw/day) represents the ingestion rate of soil
- \(C_S\) (mg/kg dw) represents the COC concentration in ingested soil.

6. **Dose from sediment intake (primarily aquatic foragers):** The total dose from incidental ingestion of COC-contaminated sediment \((D_{SED}, \text{mg/kg-bw/day})\) calculated using the following equation:

\[
D_{SED} = I_S \times C_{SED}
\]  
(Eq. 7)

Where:
- \(I_S\) (kg dw/kg bw/day) represents the ingestion rate of sediment
- \(C_{SED}\) (mg/kg dw) represents the COC concentration in ingested sediment.

7. **Dose from drinking water:** The total dose from drinking water ingestion of COCs \((D_W, \text{mg/kg-bw/day})\) calculated using the following equation:

\[
D_W = I_W \times C_W
\]  
(Eq. 8)

Where:
- \(I_W\) (L/kg bw/day) represents the drinking water ingestion rate
- \(C_W\) (mg/L) represents the COC concentration in the water.

8. **Total unadjusted dose:** The unadjusted dose \((D_{UT}, \text{mg/kg ww/day})\) can be calculated by taking the sum of the doses for the separate media: food, soil, sediment, water:

\[
D_{UT} = D_F + D_S + D_{SED} + D_W
\]  
(Eq. 9)

Where:
- \(D_F\) (mg/kg wet/day) is the dose from food
- \(D_S\) (mg/kg wet/day) is the dose from soil
- \(D_{SED}\) (mg/kg wet/day) is the dose from sediment
- \(D_W\) (mg/kg wet/day) is the dose from water.

9. **Dose adjustment factor (DAF):** The DAF can be calculated as a function of territory/foraging range, habitat quality and bioavailability of the COCs.

\[
DAF = FRF \times \alpha
\]  
(Eq. 10)

Where:
- \(FRF\) (unitless) is the foraging range factor, which represents the surface area of the site that overlaps with the territory or foraging range of the species
- \(\alpha\) (unitless) is the dietary uptake efficiency of a given chemical and can be thought of as the proportion of chemical that is absorbed through the intestinal tract compared to the total amount ingested. Since many literature studies are based on dietary efficiencies that are less than 1, the value chosen will need to account for the dietary uptake efficiency relative to the study(s) on which the TRV is based. The value does not account for difference in availability between soil and different food types.

10. **Total adjusted dose:** The total adjusted dose \((D_{AT}, \text{mg/kg wet/day})\) calculated by multiplying the unadjusted dose and the DAF:

\[
D_{AT} = D_{UT} \times DAF
\]  
(Eq. 11)

Where:
- \(D_{UT}\) is the unadjusted total dietary dose of a given chemical (mg/kg wet/day)
- \(DAF\) is the dose adjustment factor (unitless)
3.4 Beyond Point Estimates of Exposure

For receptors that are relatively immobile (e.g., invertebrates and plants), exposure assessment is typically conducted on a spatially explicit basis. This may be conducted by directly applying station-specific measurements of exposure to represent a management unit (grid cell), or by using multiple measurements to generate a modeled surface of exposures.

For wildlife (birds, mammals) and mobile aquatic and semi-aquatic organisms (fish, amphibians), exposure estimation is more challenging. Screening-level risk assessments often apply the principle of the exposure point concentration (or estimated exposure concentration), which is a conservative point estimate of the chemical concentration (or dose) available from a particular medium or route of exposure. Simple models may use the maximum concentrations from each medium to represent the exposure point concentration, or some other statistical metric (e.g., 95 per cent UCLM or 90th percentile) depending on sample sizes.

Disadvantages of the simplified point estimate approach include:

- lack of consideration of the relative spatial positions of receptors and contaminated media (due to habitat preferences, migration patterns, etc.), which can strongly influence estimates of exposure
- overreliance on extreme values (maxima) in the calculations of exposure point concentrations.

The point estimation approach assumes that receptors have equal and random access to all areas of an exposure unit, and that they occur evenly throughout the exposure unit. These conditions rarely apply in natural environments.

Point estimates of exposure can be improved by using probabilistic methods (see Section 5.3.6 and Section 5.6.3) and by incorporating spatial information as discussed below.

3.4.1 Partially Spatially Explicit Approaches

Several methods are available for cases where using summary statistics yields unacceptable uncertainty. If more spatial realism is desirable, risk assessors can use more advanced methods, such as:

- Dividing the exposure into several sources depending on likelihood of use. For example, for drinking water, exposure could be divided among several sources that a receptor uses, based on evaluation of how likely the receptor is to use that drinking water.
- For soils or sediments where use may be spatially related, weighting based on the location of the soil sample may be useful. For instance, when estimating incidental soil ingestion, soil samples can be weighted by their spatial “area of influence” or by the relative probability that a receptor will use that area, based on evaluation of habitat preferences (e.g., less soil will be incidentally ingested in areas subject to low use). The result of this weighting may be a spatially weighted average concentration that is used in the ERA for
evaluating incidental ingestion. This approach would typically require overlay of soil data and habitat polygons using geographic information systems (GIS) software.

- When estimating ingestion of contaminants through food items, the food item concentrations measured in various areas of a site can be weighted according to their relative probability of consumption based on habitat preferences (e.g., a sample of insect tissue in one area of the site would receive twice the weight of another that occurs in habitat that is half as preferable for an insectivorous receptor).

- The curve model (Freshman and Menzie 1996) may be used to describe the exposure to wildlife that forage over the contaminated site. This approach is based on rank-ordering the contamination measurements and the home range (foraging area) of the species of interest. The approach considers the distribution of concentration measurements (both frequency and magnitude), but does not account for natural foraging patterns or habitat preferences.

All of these types of improvements are attempts to account for spatial information, and risk assessors should implement these refinements where the level of effort is justified by the increased precision in risk estimates.

3.4.2 Spatially Explicit Methods

None of the refinements in Section 3.4.1 result in a truly spatially explicit exposure model. Exposure models that are truly spatially explicit aim to simulate the spatial behaviour of individual animals on a site, in the context of the habitats and other factors that influence site use. This is the only way to realistically capture variability in exposure within a population of animals.

With advances in GIS, explicit consideration of the heterogeneous distribution of receptors, their habitats and contamination is increasingly feasible. Tools for incorporating such spatial considerations in ERAs are more available, although they tend to be applied on large, complex risk assessments.

Some models may be adaptable to particular sites (e.g., the spatially explicit exposure model and others reviewed by Loos et al. [2010] and Wickwire et al. [2011]). However, their flexibility is often limited, and they have not yet been widely applied. See Hope et al. (2011) and Wickwire et al. (2011) for further discussion.

4 HAZARD ASSESSMENT

The general purpose of hazard assessment is to characterize the nature of effects elicited by each COC under an exposure condition that is relevant to each receptor of concern. This characterization is often called a “response profile” and is required for each combination of COC and line of evidence. Note that for some lines of evidence (e.g., toxicity tests of contaminant mixtures) it may be possible to characterize only the response for that mixture.
Effects information can be used in a variety of ways, which are not mutually exclusive:

- Develop a TRV. TRVs are commonly used in the HQ method of risk characterization (see Section 5.3.1 for more details), where they are compared to exposure estimates.
- Develop concentration-response (or dose-response) relationships. These can be used directly to estimate effect levels for a particular exposure concentration, or they can be used to derive TRVs for specific effect levels.
- Develop a site-specific remediation objective for a site where an initial ERA indicates that risk management is warranted, using either a TRV (first bullet above) based on literature or site-specific data, or a concentration-response relationship (second bullet above) based on literature or site-specific data.

Exposure and effects are matched in one or more ways for every line of evidence evaluated in an ERA. Consequently, hazard assessment is not a single step in ERA, but is carried out for every line of evidence. Importantly, while the details of hazard assessment are discussed in this section, they must be fully understood and articulated at the problem formulation stage in order to support design and planning of the ERA.

4.1 Overview of Hazard Assessment

Hazard assessment used to develop any particular line of evidence generally involves the following elements (note that the first four elements will have been decided as part of the problem formulation):

- Determine which type(s) of hazard assessment measure will be used, among the four broad types:
  1. *Site-specific controlled studies:* Considers measurement endpoints related to studies of test organism exposures to contaminated site media under controlled conditions. This category includes toxicity tests conducted in the laboratory using media collected on site, in the field (*in situ*) or a combination of both. The category includes both standardized test protocols and exploratory techniques such as toxicity identification evaluations (TIEs).
  2. *Indirect controlled information:* Considers toxicological information derived from other sites (or laboratory studies), under an assumption that the concentration-response relationship...
relationship is either similar to or can be estimated from the data collected at other sites. Results are extrapolated to the site of interest through consideration of contamination profiles, habitat similarities and factors that may influence relative bioavailability (e.g., chemical speciation, organic carbon or lipid content, particle size, salinity). Indirect toxicological evidence can take many forms, ranging from generic environmental quality guidelines based on toxicity database information, to concentration-response relationships gleaned from the literature or drawn from focussed studies conducted at other sites.

3. **Site-specific field studies**: Considers direct assessment of the site’s biological condition relative to the exposure metric. This category may include endpoints at the suborganism level (e.g., histopathological indicators), organism level (e.g., mortality, growth, deformities, erosions, lesions and tumours), population level (e.g., numbers and proportions of indicator organisms, vital rates) and community level (e.g., diversity, distribution of taxonomic groups).

4. **Indirect field information**: Considers indirect assessment of biology, through extrapolation of knowledge obtained at other sites. As with toxicology studies, the biological evidence must be scaled to the site condition based on consideration of exposure levels and ecological relevance. Given natural ecological variability, indirect biological information alone would almost never be sufficient for characterizing risks as part of a detailed ERA.

- Determine whether the effects data will be interpreted relative to an AEL (i.e., to derive a TRV) or used without predetermined AELs (i.e., to estimate actual effect sizes and leave the determination of “acceptable” or “unacceptable” to risk managers).
- Determine how contaminant mixtures will be considered. While response profiles need to address each combination of COC and receptor of concern, a single response profile could address multiple COCs simultaneously when appropriate measures are used. Site-specific hazard assessment measures (e.g., toxicity testing or biological surveys) allow for explicit consideration of chemical mixtures present at a site, thus integrating all interactions. Consequently, site-specific approaches are usually recommended where feasible.
- Decide which type of response profile should be developed given the nature of the available effects data:
  - *Continuous response profile*: This type of profile documents how effects (e.g., magnitude of response) vary over the range of realistic exposure levels. The profile can...
be used directly in risk characterization (e.g., when estimating actual effect levels associated with a particular exposure level), or can be used to derive a TRV for a response magnitude of interest (e.g., what exposure level corresponds to a 20 per cent adverse response). Understanding the exposure-response relationship also helps to facilitate the interpretation of the potential effects should predicted exposure exceed a TRV in the risk characterization.

- **Discrete response profile**: This type of profile is used in situations where effects data are scarce (e.g., limited literature effects data for some COCs for wildlife) or when effects apply to particular exposure scenarios only (e.g., those occurring at specific locations on the site). This could arise in a control-impact study design (e.g., when determining if a contaminated area differs from a reference condition) or a gradient design with discrete levels of impact, or when testing complex contaminant mixtures using site-specific effects measures (e.g., toxicity testing or biological surveys). In the case of limited data, TRVs can still be derived from a discrete response profile, but they may not coincide with the desired effect magnitude or exposure condition.

- Develop response profiles for each combination of COC and receptor of concern, or as appropriate (e.g., for contaminant mixtures) if specific profiles are not feasible or appropriate.

- Characterize uncertainties in effects, evaluate the implications of uncertainty using sensitivity analysis and, if warranted, integrate uncertainties into the hazard assessment (e.g., using probabilistic methods).

The outcomes of hazard assessment are measures that can be matched with exposure estimates to provide evidence in the form of a line of evidence. It is critical that the risk assessor conceptualize the exposure and effects information at the same time (during problem formulation) to ensure that they can be integrated effectively.

### 4.2 Categories of Measures for Hazard Assessment

This section discusses measures for hazard assessment, categorized according to the four broad categories for lines of evidence that were introduced in Section 4.1 (examples provided in Table 4-1; more details provided in Section 5):

- site-specific controlled studies
- indirect controlled information
- site-specific field studies
- indirect field information.

These categories of measures are distinguished by two factors:

- Site-specific versus indirect: This distinction is based on whether the measure addresses effects in exposure media (or receptors) from the site, either *in situ* or *ex situ*, or if it relies
on published effects data (e.g., from published research or grey literature or from other contaminated sites).

- Controlled versus field: This distinction essentially relates to whether a measure involves experimental manipulations to control environmental variables so that treatments differ only in their exposure to COCs (e.g., testing fish growth in site water in a laboratory or field testing in enclosures) or whether it focusses on quantifying the effects associated with naturally occurring exposure situations (e.g., a benthic invertebrate community study or a small mammal population survey). The distinction becomes less clear in field tests with high realism, but such field tests are rarely used in practice.

The types of measures used for hazard assessment are not mutually exclusive. As described in Section 4.2.4, risk assessors are encouraged to use more than one type of effects measure even for the same assessment endpoint. The decision about what types of effects measures to use will be based in part on:

- level of rigour needed to inform decision making
- resources required to properly use a measure
- availability and quality of information (e.g., for published studies)
- confidence in likely measure performance (e.g., ease of extrapolation to assessment endpoint and associated uncertainties)
- availability of matching exposure data with which effects measure outputs would be combined or integrated.

Regardless of what types of measures are used for hazard assessment, it is critical that they be specified in the problem formulation when the lines of evidence to be used in the ERA are developed.
### Table 4-1: Examples and categorization of hazard assessment measures

<table>
<thead>
<tr>
<th>Study type</th>
<th>Source of dose-response information</th>
<th>Line of evidence type</th>
<th>Examples of relevant measures</th>
</tr>
</thead>
<tbody>
<tr>
<td>Controlled</td>
<td>Site of interest</td>
<td>(1) Site-specific controlled</td>
<td>Laboratory seed germination test conducted using site soils; caged mussel study; amphibian metamorphosis assay using larvae harvested in site vernal pool; <em>in-situ</em> test of survival and growth for <em>Hyalella</em>; laboratory test of early life stage fish growth and survival.</td>
</tr>
<tr>
<td>Controlled</td>
<td>Guideline</td>
<td>(2) Indirect controlled</td>
<td>Water quality guideline developed from most sensitive tested species; sediment quality guideline developed from co-occurrence database (biological effects database for sediment); soil quality guideline for protection of microbial processes.</td>
</tr>
<tr>
<td>Controlled</td>
<td>Literature or compendium</td>
<td>(2) Indirect controlled</td>
<td>EC$_x$ $^{31}$ threshold from US EPA ECOTOX database review; avian dose-based TRV from ecological soil screening levels; critical tissue burden from literature search; SSD.</td>
</tr>
<tr>
<td>Controlled</td>
<td>Other site</td>
<td>(2) Indirect controlled</td>
<td>Use of threshold for reproductive success from a captive mink feeding study conducted for another site using fish harvested from that site.</td>
</tr>
<tr>
<td>Field</td>
<td>Site of interest</td>
<td>(3) Site-specific field</td>
<td>Benthic community enumeration; evaluation of salmon reproductive success and output; small mammal survey (density, biomass, net migration); vegetation transect or quadrat enumeration.</td>
</tr>
<tr>
<td>Field</td>
<td>Literature or compendium</td>
<td>(4) Indirect field</td>
<td>Literature summary of water concentrations associated with reduced richness of epibenthic invertebrates; literature summary of relationship between average sediment COC concentrations and incidence of tumours in fish.</td>
</tr>
<tr>
<td>Field</td>
<td>Other site</td>
<td>(4) Indirect field</td>
<td>Reproductive study of tree swallows (using nest box assessment) at Site A that could be used to assess potential avian effects at Site B, assuming some consistency of response for a standardized measure of exposure.</td>
</tr>
</tbody>
</table>

$^{31}$ EC$_x$ = effect concentration, with percent X of organisms affected
4.2.1 Site-specific Controlled Studies
Site-specific controlled assessments are used to directly test whether exposure to contaminated media (e.g., water, sediment, soil) from a site elicits adverse effects in test organisms under controlled conditions. This is an important distinction from field studies in that by controlling environmental variables, the test medium becomes the primary independent variable (i.e., predictor), with test endpoints being the dependent variables (i.e., outcomes).

Although all options for this type of measure involve some form of experimental manipulation to help reduce the influence of non-chemical factors on the outcome of the test, they vary in how well they mimic reality. At one end of the spectrum are standardized laboratory toxicity tests (e.g., \textit{ex situ} exposure to site media), where site media are taken to a laboratory facility and tested under controlled conditions following a detailed protocol. These tests are by far the most commonly used and are the primary focus of this guidance. At the opposite end of the spectrum are highly customized \textit{in situ} toxicity studies.

Laboratory tests under controlled conditions are valuable in that they can help isolate a toxic mechanism that could be obscured in a natural environment. As a result of these controls, laboratory tests tend to be more precise, though not necessarily more accurate (relevant) in terms of describing the assessment endpoint. Because the type of errors in toxicity tests differ qualitatively from those in field studies, it is not appropriate to compare the concentration-response results using only a coefficient of determination ($r^2$) or other purely statistical measure. Rather, assessment of uncertainty of laboratory testing must consider both numerical measures of uncertainty (e.g., inter-replicate variability) and uncertainty associated with lab-to-field extrapolation.

Several options are available for cases where standard toxicity tests are not environmentally realistic enough to properly derive a response profile (e.g., when the physical test set-up is not appropriate or when sample handling of the target exposure medium might increase or decrease COC bioavailability). Examples of ways in which toxicity testing can be modified to increase environmental realism include:

- setting up temporary testing facilities at the site (e.g., a continuous flow-through set-up taking water directly from an area of interest)
- conducting an \textit{in situ} toxicity study (e.g., in enclosures such as pens or mesocosms)
- altering standard protocols such as physical test set-up to increase test realism in a laboratory setting (e.g., increasing the number of refreshes of overlying water to better represent a flowing environment).
Another type of specialized site-specific toxicity testing is toxicity identification evaluation (TIE). TIEs involve physical or chemical manipulation of a sample to try to isolate and identify toxic substances in a test medium. TIEs are applied in an iterative fashion to progressively pinpoint a specific toxicant or class of toxicants. Clearly identifying the specific cause of toxicity can reduce uncertainty and increase confidence in conclusions. Information on TIEs is provided in FCSAP Module 1 (EC 2010-a).

Guidance on toxicity test selection and interpretation is presented in FCSAP Module 1 (EC 2010-a). This comprehensive technical module covers the following:

- an overview of toxicity testing in risk assessment, with specific emphasis on how tests are used in a WOE approach and how they can also be used to develop a site-specific TRV (additional information on site-specific TRVs can be found in FCSAP Module 2 [EC 2010-b])
- procedures for test selection
- additional considerations specific to porewater
- a summary of key information for about 75 of the most commonly used toxicity tests in North America
- interpretation of toxicity test results.

Site-specific toxicity tests are considered more useful than indirect toxicity information for the following reasons (Suter et al. 2000):

- site-specific bioavailability of the contaminants is considered
- form of the contaminant is realistic
- interactions among contaminants are simultaneously addressed
- spatial distribution of toxicity can be determined
- remedial goals may be determined with higher confidence.

Key limitations of site-specific toxicity tests include (SAB-CS 2008; Suter et al. 2000):

- The medium may be modified by sample collection and test preparation (particularly for sediments, but also for water and soil), which could affect contaminant form and bioavailability.
- Differences in sensitivity between the test organism and the receptor of concern may not be known. This could be due to taxonomic or genetic differences (e.g., some strains of test organisms are known to be particularly sensitive), or to other factors like acclimation (e.g., where pre-test holding conditions affect organism sensitivity in the toxicity test for essential elements) or adaptation (e.g., where an organism’s natural detoxification systems may not be working optimally due to holding in low-metals water).
- The testing scenario (e.g., duration and set-up) may not fully reflect site-specific realities.
• The cause of toxicity is not known (unless a TIE or other method for establishing causal linkages is conducted).
• Apparent toxicity may be due to differences between reference and site media in factors other than contaminant concentrations (e.g., higher nutrients or substrate-based responses in reference).
• Variability of test endpoints, particularly for sublethal endpoints during chronic exposures, may reduce the statistical power to detect target effect sizes.
• High costs, particularly for chronic testing, may force trade-offs in spatial or temporal sampling coverage.
• Effects are measured on individual organisms, which may then need to be extrapolated to or used to predict population- or community-level assessment endpoints.

Many of these limitations directly become sources of uncertainty for this type of measure. Section 5.6 discusses approaches for addressing uncertainties.

4.2.2 Indirect Toxicity Information from Controlled Studies
Risk assessments can benefit from the substantial body of literature available from ecotoxicological research. The Internet allows access to this information, as one can search online data compilations or search and retrieve primary literature. Thus, for a relatively low cost compared to other types of measures, a wealth of information can be accessed to augment the hazard assessment in a number of ways, including:

• compiling preliminary effects information during problem formulation (see Section 2.2.4 for more details)
• identifying and sourcing published effects models (e.g., BLM; see below for more details)
• compiling response profiles and deriving TRVs (see FCSAP Module 2 [EC 2010-b] for details).

FCSAP Module 2 (EC 2010-b) provides guidance on using indirect toxicity information in the development of TRVs. TRVs are an important part of the response profile in that they represent a concentration or dose that is not expected to cause an unacceptable adverse effect (see Section 2.3.1 for more discussion on AELs). FCSAP Module 2 covers the following:

• types and use of TRVs in ERA
  o dose-based TRVs
  o concentration-based TRVs for exposure media
  o concentration-based TRVs for tissues

Key Concept
Indirect toxicity information taps into the wealth of knowledge available in published studies. Used judiciously, this can be a cost-effective source of relevant data to develop response profiles.
• options for TRV selection
• review of published TRVs
• general considerations for TRV derivation
• derivation of site-specific, literature-based TRVs
  o literature review
  o data quality and selection criteria
  o derivation methods
  o uncertainty and extrapolations
• modification of existing guidelines to develop site-specific TRVs.

In addition to the limitations inherent in extrapolating from the laboratory to the field (discussed in Section 4.2.1) for site-specific toxicity measures, using indirect toxicological information also requires considering the site-specific relevance of the data. Potential sources of bias in literature toxicity data that are uncertainties for this type of measure include (Suter et al. 2000):

• chemical form used in toxicity tests may be more toxic than the dominant forms found at a contaminated site
• contaminant interactions are rarely considered
• test species may not be representative of the sensitivity of receptors of concern at the site
• exposure test media may not be representative of those found at the site
• laboratory test conditions may not be representative of field conditions.

The relevance of indirect toxicological information can be improved by filtering the available data to include studies that most closely match the needs of the ERA. Depending on the contaminant, one or more of the ancillary parameters listed in Sections 3.3.1 and 3.3.2 may play a key role in determining its toxicity (e.g., by affecting bioavailability). Uncertainty can be substantially reduced by appropriately matching reported test conditions to actual exposure conditions. Although in many situations the risk assessor must perform the task of filtering (if possible and appropriate), ideally the key factors affecting bioavailability and toxicity would be understood sufficiently to support site-specific predictive modelling of toxicity.

Recent advances in supporting science may address some of the common limitations to indirect toxicity information highlighted above. These include the BLM and the tissue residue approach (TRA) for toxicity assessment. These are discussed below.

*Biotic ligand model (BLM)*: Research in recent decades (e.g., Meyer 1999; Pagenkopf 1983) has led to major progress in our understanding of metals bioavailability and mechanisms of toxicity in both aquatic ecosystems (see review by Paquin et al. [2003]) and terrestrial ecosystems (see review by Allen [2002]). The culmination of this research to date is the development of the BLM, which integrates key discoveries from several disciplines to consider a range of factors influencing metals
bioavailability and, ultimately, toxicity. The premise of the BLM is that toxicity is related to the metal binding to an active biochemical site on the organism (i.e., the biotic ligand) and that this binding is related to concentrations of free metal cations and complexing ligands in the water (or solution phase for soils). The complexing ligands compete with the biotic ligand (e.g., in fish gills or at root elongation sites for plants) for free metals and other cations in the water (or solution phase for soils), thus directly affecting toxicity by dictating metals concentrations at the target site. A major advantage of the BLM is that it explicitly considers a range of modifying factors (e.g., as competing cations), influencing the response profile of a particular endpoint.

Aquatic BLM: The aquatic BLM has successfully been used for predicting acute aquatic toxicity related to copper (Santore et al. 2001), silver (Paquin et al. 1999) and zinc (Santore et al. 2002). The success of the BLM in accurately predicting toxicity has already led to its use in developing water quality criteria; the BLM features prominently in the US EPA’s criteria for copper (US EPA 2007-a). More recently, research has focussed on the BLM’s application to chronic toxicity (Clifford and McGeer 2010; De Schamphelaere et al. 2005; Peters et al. 2011; Schroeder et al. 2010; Schwartz and Vigneault 2007) and metals mixtures (Kamo and Nagai 2008). This research should lead to increased use of the aquatic BLM in risk assessments.

Terrestrial BLM: More recent efforts have developed and validated BLMs specifically for terrestrial ecosystems. Thakali et al. (2006-a) initially applied a terrestrial BLM to predict copper and nickel toxicity to barley root elongation in a number of soils, then to an expanded suite of toxicity endpoints (plants, invertebrates and microbes) across a range of non-calcareous soils from the European Union. Terrestrial BLMs have also been used to predict cobalt toxicity to worms (Lock et al. 2006) and barley (Lock et al. 2007). These methods are likely to be refined and expanded to other metals and toxicity endpoints.

Tissue residue approach (TRA): Another rapidly advancing area is using the TRA for toxicity assessment. A Society of Environmental Toxicology and Chemistry (SETAC) Pellston Workshop in 2007 led to a series of “state-of-the-science” papers on this subject (Adams et al. 2011; Escher et al. 2011; McCarty et al. 2011; McElroy et al. 2011; Meador et al. 2011; Sappington et al. 2011). This approach works on the premise that for describing toxicity to organisms, whole-body or organ-specific concentrations (residues) are a better dose metric than external exposure media (Escher et al. 2011). While this is somewhat intuitive (because contaminant bioavailability is explicitly considered in the TRA), the approach is not without its challenges, largely due to difficulty correlating internal concentrations to ecotoxicological outcomes. Variability in ecotoxicological outcomes and species sensitivity is due in part to differences in toxicokinetics, which is composed of several key processes (absorption, distribution, metabolism and excretion) that influence internal concentrations (Escher et al. 2011). Where variability is high (i.e., where internal concentrations are not proportionate to the concentration or dose at the target site), toxicokinetic modelling may be useful to derive the target dose. One of the main challenges of using the TRA will be the availability of appropriate tissue residue–response data (Sappington et al. 2011). Given the developing nature of the science, it would be prudent to treat this as a complementary line of evidence for the time being (Sappington et al. 2011).
4.2.3 Site-specific Field Studies

Site-specific field studies directly assess receptor of concern attributes in the field, thus eliminating many of the uncertainties associated with toxicological information. These studies can target a range of attributes for individuals (e.g., growth, reproductive success, survival), populations (e.g., biomass, abundance, density, age structure) or communities (e.g., diversity, species composition, abundance, density, biomass), making it possible to directly estimate the assessment endpoint (Appendix D in CCME 1997-a; Carlsen et al. 2008; Menzie et al. 2008). Comparisons should be made to reference conditions or along gradients in exposure. Unlike toxicity studies where several environmental variables are controlled to help isolate an exposure-related “signal,” field studies can be clouded by natural variability due to the inherent complexity of natural systems. Some of this natural variability can be controlled through proper experimental design (including identifying covariates and categorical factors) and through increased sample size (either in single studies or multiple monitoring events).

Risk assessors should consider the following factors when deciding whether field studies are appropriate (Suter et al. 2000):

- **Scale**: These studies are usually most appropriate for receptors of concern who have small home-range sizes and are likely to remain mostly inside the boundaries of the assessment area. However, field studies may also be appropriate for highly mobile, wide-ranging receptors of concern, particularly when those receptors are of particular importance to stakeholders.

- **Interpretation**: Variation in the attribute of interest must be interpretable in the context of confounding factors such as habitat heterogeneity.

- **Difficulty**: Studies can vary greatly in the scale and time needed for implementation. This needs to be balanced against the chances of obtaining useful information.

- **Appropriateness**: The study design and methods need to match the task at hand.

- **Technical expertise**: Study complexity may require specialized expertise beyond the risk assessment team.

- **Survey consequences**: In some cases (e.g., destructive sampling of small populations or of rare species), biological studies may cause unacceptable harm.

- **Data availability**: Suitable surrogate data sets may be available (e.g., from broader environmental management initiatives; see Section 4.2.4).

Once a risk assessor has committed to site-specific field effects measures in the ERA, the following additional considerations may help to design and implement the study. The design has to be worked out as part of problem formulation (see Section 2.3). At a minimum, risk assessors undertaking
site-specific biological studies should consider the following during the study design phase (and seek out more specific information relevant to their unique situation):

- **Defining the question**: Where possible, the focus of the study should be direct estimation of the assessment endpoint. In other cases, study objectives and how the results will be extrapolated to the assessment endpoint should be determined in advance (i.e., during the problem formulation).

- **Defining the assessment population**: This question has important implications for how effects might be interpreted (e.g., the larger the assessment population relative to the site, the more effects may be “diluted”). As a starting point, consider defining the assessment population as those organisms inhabiting the site of interest. Scale can then be adjusted based on receptor of concern–specific considerations (see Menzie et al. [2008] for more discussion of assessment populations).

- **Selecting relevant attributes**: As discussed above, this should either match, or be easily extrapolated to, the assessment endpoint. Multiple attributes are recommended where practical to provide a more robust means of assessing the question (Appendix D in CCME 1997-a; Carlsen et al. 2008; Menzie et al. 2008).

- **Designing the study**: Appropriate scientific methods (e.g., EC 2011; Krebs 1989) should be used to optimize the design to answer the question. This will include having an understanding of the statistical methods that will eventually be used.

- **Field methods**: Methods for most study types have at least been published and may even have recommended survey methods or standard protocols (e.g., EC 2002; State of Washington et al. 2015; see SAB-CS [2008] for more references; also, the US EPA has a variety of methods posted at their Environmental Monitoring and Assessment Program website [see US EPA n.d.]).

- **Data quality objectives (DQOs) and QA/QC**: Data quality objectives (DQOs) define the specifications for the data set. Quality assurance (QA) steps are the actions taken to meet those objectives, and quality control (QC) measures are the benchmarks used to verify data quality (e.g., EC 2011).

- **Data analysis and interpretation**: The statistical methods used should be those selected during the design phase. Interpretation should consider key uncertainties. This is often done by reporting observed effect sizes with confidence limits for each attribute (e.g., EC 2011).

Advantages of well-planned, site-specific field studies include the following:

- assessment endpoints can be directly estimated
- they integrate exposure by accounting for complexities such as bioavailability and contaminant mixtures
- they have a high degree of ecological relevance
- they are complementary to toxicity data
• they can reduce uncertainty and reliance on some assumptions.

Limitations of site-specific field studies include the following:

• the scale and time needed to obtain robust data sets can be high, in which case such a study is warranted only when the likely value in informing management decisions is also high

• natural variation can make it difficult to detect contaminant-related changes, even in well-planned studies

• measured effects may not be due to COCs, but rather to confounding natural environmental variables or non-chemical stressors

• conducting the studies may cause direct adverse effects to the target receptors of concern

• studies usually have to rely on spatial comparisons (e.g., across exposure gradients) due to the scarcity of baseline data for the site of interest. Selecting appropriate reference areas can be challenging.

Some of these limitations can translate directly into uncertainties. High natural variability can mask the detection of target effects, thus potentially resulting in a false negative conclusion (i.e., type II error). In contrast, differences between exposure and reference conditions may result in measured effects that are not actually due to contaminant exposure, resulting in a false positive conclusion (i.e., type I error). Statistical methods (e.g., confidence limits on effects sizes and power analysis) can be used to help understand the scope for type II errors. Complementary use of site-specific toxicity testing can help establish causality (or lack thereof) for field studies (see Section 5.5.2.2 and FCSAP Module 4 [EC 2013] for more discussion of causality).

4.2.4 Indirect Field Information

This category of measures is analogous to indirect toxicity information based on controlled studies, but emphasizes transferring appropriate field studies from other sites (e.g., those published in the literature) that could be used to help inform a response profile for the site of interest.

Given the resource and technical challenges of designing and implementing useful field studies discussed in Section 4.2.3, the advantages of finding an appropriate study are clear. The main challenge, however, is overcoming the hurdle of establishing relevance at the site of interest. Risk assessors should consider the following when drawing inferences from studies conducted at other sites:
• **Type of contamination:** Both COCs and the factors driving their bioavailability would ideally be similar at both sites. Ensuring this would require comparing data from the exposure assessment to that reported in the literature study. This is much easier for sites with only one or two COCs.

• **Pattern of contamination:** This includes magnitude and spatial and temporal patterns. Ideally, the magnitude and scales of both studies would be similar.

• **Habitats and receptors:** Site-use patterns by receptors will vary according to available habitat types (i.e., due to their differing habitat characteristics, related to the animals’ ecological needs). The configuration of high-use habitat types relative to the pattern of contamination will affect receptor of concern exposure.

Once a study is deemed appropriate, its data should be extracted in a similar manner to that discussed for indirect toxicity information. For example, in the mercury study shown in the text box (Brasso and Cristol 2008), swallow reproductive success can be plotted against each of the mercury exposure measures (or simply the one most relevant to your risk assessment) to develop response profiles.

### 4.3 Receptor-Specific Considerations

This section focuses on linking measures to receptor groups. Table 4-2 shows the relative frequency of use of each of the major types of hazard assessment measures in risk assessments. It should be noted that any of these types of measures may be appropriate for a given risk assessment, so the table should be used only to provide initial insight into what is typically done. Selection ultimately depends on the specific needs of the risk assessment.

**Key Concept**

Field studies reported in the literature can provide valuable information with which to derive a response profile for specific combinations of COCs and receptors of concern. For example, Brasso and Cristol (2008) studied effects of mercury exposure on the reproductive success of tree swallows. The authors collected several measures of mercury exposure (blood and feather total mercury in the birds, and total mercury in the insects fed to nestlings) against their primary effect measure, the number of nestlings that left the nest (i.e., fledglings). With consideration of the points listed in the text, this study could provide highly relevant data for other sites where mercury is the primary COC.

**Key Concept**

The four types of hazard assessment measures are not used equally among receptor groups in ERA. The relative frequency of use reflects current reality, which may not be ideal but often reflects limitations and challenges in application.
Table 4-2: Frequency of use of types of hazard assessment measures for each receptor group.

<table>
<thead>
<tr>
<th>Receptor group</th>
<th>Site-specific toxicological information from controlled studies</th>
<th>Indirect toxicological information from controlled studies</th>
<th>Site-specific information from field studies</th>
<th>Indirect information from field studies</th>
</tr>
</thead>
<tbody>
<tr>
<td>Terrestrial primary producers</td>
<td>moderate</td>
<td>moderate</td>
<td>moderate</td>
<td>low</td>
</tr>
<tr>
<td>Aquatic primary producers</td>
<td>high</td>
<td>high</td>
<td>moderate</td>
<td>low</td>
</tr>
<tr>
<td>Terrestrial invertebrates</td>
<td>moderate</td>
<td>moderate</td>
<td>moderate</td>
<td>low</td>
</tr>
<tr>
<td>Aquatic invertebrates</td>
<td>high</td>
<td>high</td>
<td>moderate</td>
<td>low</td>
</tr>
<tr>
<td>Fish</td>
<td>high</td>
<td>high</td>
<td>moderate</td>
<td>moderate</td>
</tr>
<tr>
<td>Birds and mammals</td>
<td>rare</td>
<td>high</td>
<td>low</td>
<td>moderate</td>
</tr>
<tr>
<td>Amphibians and reptiles</td>
<td>low</td>
<td>moderate</td>
<td>moderate</td>
<td>moderate</td>
</tr>
</tbody>
</table>

Categories are defined as follows:

- **High**: This rating was applied to both controlled toxicological measures for aquatic receptor of concern groups to reflect the long-term establishment, protocol development and value for risk assessments. It was also applied to the indirect controlled information measure for birds and mammals, due to jurisdictions’ reliance on this measure in the face of cost and uncertainty of alternatives.

- **Moderate**: This was applied to both controlled toxicological measures for terrestrial receptor of concern groups to reflect the growing use of these measures. It was also ascribed to the indirect controlled toxicological measure for amphibians and reptiles, mainly due to data limitations and the exclusion of these receptors of concern from many risk assessments (although use is increasing over time). Finally, it was applied to site-specific field studies for most receptors to reflect the technical challenges associated with this type of measure.

- **Low**: This was applied to measures of indirect field information for all receptor of concern groups, largely reflecting the difficulty of identifying studies that extrapolate well to the conditions of the site of interest (contamination pattern and relevant biology). It was also applied to site-specific field studies for birds and mammals to reflect the cost and complexity of robust studies for discerning contaminant factors from physical or habitat factors. Note that use of reconnaissance surveys, habitat surveys and semi-quantitative field measurements are more common in ERA but convey greater uncertainty.

- **Rare**: This was applied only to site-specific toxicological studies under controlled conditions for birds and mammals. Although they are possible to conduct, they are rarely (if ever in Canada) used, due to a host of challenges, including animal welfare issues (Suter et al. 2000).
4.4 Beyond Point Estimates of Toxicity

In many situations, the outcome of a hazard assessment is the derivation of one or more thresholds for ecological effects. These thresholds are intended to represent the transition from an environmental exposure that does not elicit a meaningful ecological response to an exposure that conveys potential for ecological effects. Such thresholds can be developed for numerous media (soil concentration, sediment concentration, water concentration, tissue concentration, ingested dose), and are carried in the risk characterization where they are used to calculate HQs.

A common problem encountered in ERA is that a single threshold value is used to summarize the concentration-response relationship. In addition to the problem of oversimplifying a complex relationship, use of a point estimate is sensitive to the choice of statistical method or decision rule used to calculate the threshold. For example, use of a statistical significance criterion to discern between effect and no-effect levels of exposure can lead to substantial differences in the magnitude and significance of the threshold exposure level, in addition to other statistical and interpretative issues (Landis and Chapman 2011).

4.4.1 Considerations

It is desirable to move beyond the use of single point estimates for effects that commonly serve as the denominator in quotient methods. Although full quantitative integration of concentration-response relationships is not always possible, at minimum it is important for risk assessors to understand the true effect size (or range) that is represented by a TRV or other measure of effects, in part to facilitate selection of TRVs that are aligned with protection goals and AELs. Specifically, the risk assessment can be informed by consideration of:

- effect size associated with the study that “drives” the toxicity threshold (e.g., water quality guideline, wildlife TRV dose)
- the difference between the no-observed-adverse-effect level (NOAEL) and lowest-observed-adverse-effect level (LOAEL), or the steepness of the concentration-response where multiple exposure levels are tested
- the degree to which the “most sensitive study” represents a larger number of experimental results, or alternatively represents an outlying response
- concordance of sensitivity for different receptor groups (such as domestic species versus wild organisms, passerines versus raptors, cold-water fish versus warm-water fish)
- concordance of short-term versus chronic test endpoints, or differences in sensitivity among various sublethal endpoints.

Key Concept

Use of a point estimate, particularly if drawn from a single study, conveys high uncertainty. FCSAP Module 2 (EC 2010-b) provides guidance for reducing uncertainty in TRV development using relatively simple approaches applied to existing data.
The above considerations cannot always be addressed in a quantitative manner. However, integrations of the relevant ranges of potential response are preferable to point estimates.

According to Allard et al. (2010), a meta-analysis approach to TRV derivation is preferred to results from single studies. This entails simultaneous consideration of numerous study results on a graph of effect size versus chemical concentration (see FCSAP Module 2 [2010-b]). A graphical approach, while complicated by variations in endpoint type, exposure gradients and study designs, helps to convey the variations in response at each exposure level.

4.4.2 Species Sensitivity Distributions

The species sensitivity distribution (SSD) concept is an example of a statistical approach to hazard assessment that moves beyond the “traditional” approach to threshold development (e.g., use of the point estimate from the most sensitive study using a statistical significance criterion). For example, CCME (1996-b, 2007) recommends the SSD approach for deriving soil quality guidelines and water quality thresholds where a sufficient number, quality and variety of toxicity test data are available.

In its usual usage, an SSD is the cumulative probability distribution of some measure of toxicity of a certain chemical to a set of animal species (for more background see FCSAP Module 1 [EC 2010-a], CCME [2007], Posthuma et al. [2002] and SAB-CS [2008]). At increasing concentrations of a toxicant, the proportion of species affected (at a given level of effect, such as 20 per cent growth impairment or 50 per cent reduction of abundance) increases.

A site-specific SSD is one example of a specialized application of site-specific correlation between concentration and response methods, whereby a site-specific SSD metric is related to contaminant concentration. Figure 4-1 provides a hypothetical example of how the SSD concept may be applied.

The SSD approach recognizes that individual species may have highly variable sensitivities to a given COC (Kooijman 1987), and that protection of 100 per cent of species is not necessarily required to protect the functional attributes of a community (e.g., benthic community). By combining results from multiple tests and covering a range of test organisms, it is possible to construct a distribution of sensitivities.

<table>
<thead>
<tr>
<th>Key Concept</th>
</tr>
</thead>
<tbody>
<tr>
<td>An SSD is a numerical expression of the ranges of organism sensitivity to a COC. An SSD can characterize variations among species, within species and across taxonomic groups. Most importantly, the SSD concept conveys that individual taxa do not respond similarly to a single concentration.</td>
</tr>
</tbody>
</table>
Figure 4-1: Species sensitivity distribution

Figure 4-1 gives an example of an SSD for aquatic receptors. In the graphical example, the circles represent individual species\(^{32}\) (for the purpose of this example, they may be assumed to be various freshwater epifauna). The x-axis depicts the chemical concentration (in logarithmic scale) at which a threshold response size occurs. The response type could be a 20 per cent reduction in growth in a laboratory toxicity test, or it could be a 50 per cent reduction in species abundance in a benthic community study. The SSD entails derivation of a smoothed curve (solid line) and associated confidence limits (dashed lines) through the concentration-response curve. This allows an assessment of hazardous concentration at which a given proportion of species is affected (e.g., 20 per cent of epifauna affected at 0.6 mg/L in the example).

To derive an SSD, single-species toxicity data (e.g., LC\(_{50}\) values, IC\(_X\) values, or LOAEL or NOAEL data) for many species are fit to a distribution such as the log-normal or log-logistic. From this distribution of species sensitivities, a HC\(_p\) is identified at which a certain percentage (\(p\)) of all species is assumed to be affected (Posthuma et al. 2002). Risk assessors’ selections of both the

\(^{32}\) Depending on the derivation details, an individual data point may represent a single study for the species, or it may be a summary metric integrating multiple studies, such as a geometric mean of multiple measurements.
percentage of species and the effect level are in part matters of policy and may be considered as AEL determinations prior to risk characterization.

In addition to the more conventional application of SSD (application to literature data), the SSD procedure can also be applied to resident biological communities. In this case, it is necessary to identify a subset of the enumerated taxa for which there are sufficient numbers of organisms to assess potential concentration response. Next, each retained taxon is assessed over the gradient of contamination and a benchmark level of response (such as 20 per cent or 50 per cent reduction in abundance) is evaluated. For each organism type, the concentration at which the threshold response is observed is documented, and the resulting concentrations are rank ordered. A hazardous concentration \( (H_{C_p}) \) is then derived by choosing the interpolated COC concentration that matches the target percentage \( (p) \) of all species observed to be affected. This approach requires that the statistical power to detect the threshold level of response is considered, and as such is best suited to studies with a large number of sampling stations and a wide gradient of COC concentrations. Due to the high data demands for this approach, it is recommended only for advanced stages of risk assessment, and is less suitable where habitat variations are large relative to variations in contamination levels.

4.5 Uncertainty Factors and Extrapolation

It is common practice in ERAs to collect effects information on an indicator organism or endpoint, and extrapolate the findings to the organisms of interest at a contaminated site. This is true for both literature-based evaluations (e.g., derivations of TRVs from historical studies) and site-specific analyses (e.g., use of laboratory test species to represent potential responses in a broader array of local species).

Following historical practice in human health risk assessment, extrapolations among species and endpoints have been conducted by applying various factors, known as application, assessment, safety, or uncertainty factors (Chapman et al. 1998). These factors are intended to compensate for uncertainty in the effects analysis, and are applied in order that risks are not underestimated.

In past risk assessments, uncertainty factors have commonly been applied to address several types of extrapolation in ERA, including:

- extrapolation from test species to wild species
- extrapolation from short-term to long-term exposures
- extrapolation from a significant biological effect to an insignificant magnitude or probability of effect.

### Key Concept

Uncertainty factors can be useful in developing conservative screening thresholds (such as generic environmental quality guidelines) but are not recommended for deriving effects thresholds used in detailed risk assessments.
Some jurisdictions have advocated the use of prescribed uncertainty factors. Forbes and Calow (2002) summarize the commonly applied uncertainty factors in Europe and the United States, although they note that these factors do not preclude the use of professional judgment.

This guidance does not advocate the application of uncertainty factors in establishing TRVs. Although uncertainty factors can be useful in developing conservative screening thresholds (such as generic environmental quality guidelines), their value diminishes greatly in quantitative risk evaluation. The major disadvantages of uncertainty factors include the following:

- **Bias**: The application of uncertainty factors is uni-directional, serving to increase risk estimates without considering that the uncertainty may apply in both directions.

- **Compounding conservatism**: Applying multiple uncertainty factors can result in predicted TRVs that are unrealistically low.

- **Lack of transparency**: Applying uncertainty factors buries the uncertainty such that the risk estimate is altered, but without a clear indication of the confidence (or lack thereof) in the numerical value.

- **Incompatibility with newer methods**: The application of arbitrary uncertainty factors is poorly aligned with the application of methods (e.g., SSDs, concentration-response analysis, effect-size approaches) that are preferred for quantitative TRV development.

## 5 RISK CHARACTERIZATION

Risk characterization is the process of estimating the probability, magnitude and extent of adverse ecological impacts based on the information obtained from the exposure and hazard assessments. Risk characterization also involves discussing the “strengths, limitations and uncertainties arising from the data and models used to provide conclusions” (CCME 1996-a). Risk characterization is the stage where the various study components are integrated and interpreted in terms of overall significance for ecological risk. Risk characterization also translates complex scientific information into a format that is useful for risk managers, by conveying the ecological consequences of the risk estimates along with the associated uncertainties.

The risk characterization merges the findings of the exposure assessment and hazard assessment for each line of evidence, and integrates findings across multiple lines of evidence. As such, risk characterization techniques encompass all methods used to analyze and interpret the relationships between measures of exposure and measures of effect.

Provided that the problem formulation has been well designed, many aspects of risk characterization should be contemplated *a priori*, and integration of exposure and effect should be seamless and relatively mechanical. However, risk characterization entails more than simply
merging exposure and effects information. Rather, it conveys the process by which numerous study results are evaluated to accomplish the following core objectives:

- **Synthesize** results from multiple measurements into a conclusion for each individual line of evidence, and synthesize conclusions from multiple lines of evidence into an overall conclusion regarding ecological risks.

- Provide a concluding narrative that presents conclusions in a clear and unambiguous manner. Where possible, conclusions are stated in plain language, emphasizing clarity, such that risk assessment output can be used effectively by site managers in their decision-making process.

- Evaluate the uncertainty in the conclusions, either qualitatively or quantitatively.

- Revisit the core questions and objectives of the study (which may have been framed as one or more study hypotheses), and provide conclusions in terms of these risk management objectives.

The objectives summarized in the above bullets are less mechanical, and sometimes require the application of professional judgment. In preparing a risk characterization, the practitioner should consider the broad assessment goals (see Section 2.2.1.1) to ensure that the content effectively answers the questions and addresses the hypotheses of interest.

### 5.1 Overview of Steps

The process of risk characterization includes the following steps, upon which the organization of the chapter is based:

- **Step 1: Conduct relevance check**: Following review of the data, a relevance check will determine whether any deviations occurred during field or lab studies that could affect the relevance of the data for supporting the line of evidence for which the data will be used. This step also provides an opportunity to identify adjustments that may be required to maintain the usefulness of the data for effective risk characterization (Section 5.2).

- **Step 2: Interpret and evaluate each line of evidence**: Select appropriate methods to evaluate and interpret the information generated during the risk assessment (Section 5.3).

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33 The role of professional judgment in ERA is contentious. Application of professional judgment in interpreting or synthesizing technical information requires the practitioner to present a rationale that is transparent, clear, consistent and reasonable, and should never be applied to circumvent or obscure sound decision making (i.e., distortion or selective analysis of results to accommodate a desired outcome). The role of professional judgment is explored further in Step 4 of the risk characterization process.
- **Step 3: Prepare compiled data summary:** Present a summary of the data for each line of evidence before applying detailed analyses (Section 5.4).

- **Step 4: Conduct WOE procedure:** Integrate results of the multiple lines of evidence, using a WOE framework established during problem formulation. Importantly, the WOE procedure is interlinked with Steps 5 to 8 below, and therefore Steps 4 to 8 are often implemented concurrently (Section 5.5).

- **Step 5: Evaluate ERA uncertainties:** Consider the uncertainties that affect the interpretation or reliability of each line of evidence (Section 5.6).

- **Step 6: Consider extrapolation and interpolation:** Assess the degree to which risk conclusions drawn from a limited number of analyses can be expected to reliably translate to other conditions at the site (Section 5.7).

- **Step 7: Develop site-specific remediation standards (optional):** Develop numerical standards in site media that will be used to distinguish action levels for substances of concern (Section 5.8).

- **Step 8: Summarize risk conclusions:** Prepare a risk summary that characterizes risk in terms of potential magnitude of response and other key attributes (e.g., likelihood [probability], spatial extent, temporal extent, level[s] of organization potentially affected, causality, and other aspects of ecological relevance) (Section 5.9).

- **Step 9: Conduct follow-up actions:** Prepare clear recommendations and articulation of next steps for site closure, approvals, regulatory liaison, and so on (Section 5.10).

Importantly, the steps in risk characterization do not require any particular level of detail. For simple sites or sites where estimated risks are negligible, risk characterization does not need to be overly cumbersome, whereas for complex sites more detail and rigour will usually be warranted.

### 5.2 Step 1: Conduct Relevance Check

As described previously, several aspects of the risk characterization are planned in the problem formulation stage, including the SAP. Strategic considerations described in Section 2.3.4 therefore influence the way the risk characterization is conducted. Planning during the problem formulation stage must anticipate the important linkages between exposure and effects during risk characterization. Consistent with a philosophy of “beginning with the end in mind,” the SAP should be designed and implemented in a way that facilitates the effective integration of effects and exposure information. For this reason, strategic considerations will not be new at this stage, but rather should be revisited in light of the findings of the hazard assessment and exposure assessment for each line of evidence.
5.2.1 Revisit the Overall Assessment Needs

Before conducting risk characterization, the risk practitioner should revisit the key risk assessment questions posed in the problem formulation, and address the following issues in light of the data that have been collected:

- Confirm that the measures or techniques selected during the problem formulation remain the most effective and appropriate for addressing key risk assessment needs (such as evaluating causation). Where the relevance and value of some measurement endpoints can be assessed in advance, others require retrospective examination.

- Assess whether the analyses proposed in the problem formulation remain applicable to the assessment of testable hypotheses.\(^{34}\) If studies could not be implemented as planned, or data quality considerations confound the application of the original methods, identify a modified approach that best meets the risk assessment needs, and explain the rationale for the modifications.

- Confirm that presentation methods defined in the problem formulation remain applicable and will provide output in a format useful to the risk manager for making decisions (i.e., data obtained are sufficient for the application of the proposed methods).

During the risk characterization, the practitioner should revisit these issues before selecting specific analysis techniques, and before investing significant effort to process and synthesize data. If the data collections were substantially compromised, it may be necessary to resample or add study components before proceeding with risk characterization. Although this decision may delay the project, it is better than preparing a risk assessment deliverable that does not properly address information needs for risk management.

5.2.2 Make Appropriate Modifications

When risk characterization begins, the risk assessor has completed the principal investigations, quality controls and a preliminary assessment of individual measurement endpoints. Often, a number of factors have diverged from the original plan created during problem formulation, so the analysis plan needs revising. Accordingly, Step 1 of risk characterization serves as a relevance check to determine the consequences of such modifications, and to make appropriate adjustments if data collections did not work out as planned. For example:

- If the problem formulation specified a gradient design in which effects measures were intended to be related to gradients in contamination, but the investigation failed to capture

\(^{34}\) This does not assume that classical hypothesis testing will be the only means of data analysis. Rather, the practitioner hypothesizes that certain effects may occur and attempts to determine whether or to what extent the evidence indicates effects.
a useful gradient in contamination, it may not be possible to implement the statistical analyses contemplated in the problem formulation.

- If analysis of analytical data reveals data quality issues (e.g., negative control failure, interference effects or protocol deviations), use of the data should be reconsidered (e.g., the data may be given less weight than originally envisioned, or eliminated entirely in the case of severe data quality failures).

- If the assumptions underlying statistical analyses are not satisfied (e.g., data distributions), then alternative methods of analysis may be needed.

- If community studies indicate significant variation in substrate or habitat type that confound analysis of contaminant-related effects, the strength of evidence from such studies may be lower than expected. In such cases, alternative statistical models may be useful in differentiating contaminant-related effects. If not, more weight may be given to other lines of evidence, or a different experimental design that controls for confounding variables may be appropriate.

- If the field data reveal new receptor groups or new exposure pathways that were not contemplated beforehand, additional analyses will be needed that were not considered in the problem formulation.

Importantly, any modifications to the analysis plan should be considered based on whether measures and techniques planned in the problem formulation delivered usable results. Modifications should not be made because the presence, magnitude or type of environmental response (e.g., presence of toxicity, patterns of community structure) differed from what the practitioner suspected. Where the risk assessor proposes significant changes to the analysis approach, they must document the deviations from what was expected and provide a supporting rationale for any changes.

Modifications to the risk characterization may entail changes to specific methods to facilitate a meaningful analysis. For example, it may be necessary to consider data transformations if underlying assumptions of statistical methods (e.g., normality, stable variances, lack of high influence values [outliers]) are not met. In other cases, it may be possible to proceed with the original analyses, but with explicit acknowledgement of the reduced value of the analysis. For example, a benthic community study found to be confounded by mechanical disturbances of substrate may require a reduced weighting in the risk characterization (based on lower statistical power than was originally contemplated). In a terrestrial setting, a similar situation may arise where human disturbance of the landscape confounds the application of idealized sampling strategies intended to evaluate a soil contamination gradient.

**Key Concept**

It is appropriate to modify the risk characterization approach based on constraints to acquisition of the data as originally planned, or if new information or methods have become available since the time the problem formulation was developed.

It is not appropriate to modify a risk characterization approach simply because the results are not desirable or are unexpected.
In the broadest terms, Step 1 incorporates important learning from the data collection stage and fine tunes the data analysis methods contemplated at the problem formulation stage (and only as needed). The practitioner should not make arbitrary changes to analysis, but rather link any required adjustments to the study goals and DQOs.

5.3 Step 2: Interpret and Evaluate Each Line of Evidence
After the measurement endpoints have been selected (during problem formulation) and applied, the practitioner must apply tools to interpret the findings. These interpretations must be consistent with the informational needs of the risk manager, as outlined in the problem formulation. A proper problem formulation should have already identified how the data will be analyzed in order to support risk characterization (see Table 2-7 as an example). The use of the data in the WOE procedure (see Step 4) should guide how information is summarized for individual lines of evidence.

The following subsections summarize some of the common tools available to interpret individual lines of evidence. This is not intended to be a comprehensive list or a recommendation for the universal use of any specific tool. Furthermore, the tools are not meant to describe discrete options and are not mutually exclusive.

All the methods described in this section are tools that are applied to interpret the results from individual lines of evidence. The purpose of this section is to provide a discussion of the common procedures in their application, and to summarize the advantages and limitations of each. The methods are organized as follows:

- **HQs and other quotient methods**: Simple ratios of point estimates for both exposure and effects.

- **Concentration-response relationships**: Using the mathematical relationship between site-specific exposure and response level to understand site-specific responses.

- **Adjustment to reference or background condition**: Standardizing endpoint data to provide information on relative responses rather than absolute responses only.

- **Gradients**: Patterns of responses over space (distance and direction) or over gradients in contamination.

- **Multivariate techniques**: Interpretations of complex data sets through consideration of multiple factors simultaneously.

### Key Concept
- The selection of specific methods is context-specific and cannot be prescribed. However, the following generic guidance applies:
  - It is desirable to retain available information (i.e., hazard quotients should not be applied when concentration-response profiles are readily available and reduce uncertainty).
  - Given a choice between two methods of equal value for evaluating an assessment endpoint or reducing uncertainty, the simpler method is preferred.

- Understanding of risk is improved by examining a measurement endpoint from multiple perspectives (i.e., multiple lines of evidence developed from a site-specific endpoint).
• **Probabilistic methods:** Replacement of point estimates with distributions to provide more information on the range and likelihood of potential outcomes.

Note that some of the above techniques involve replacing point estimates with a more robust analysis of the available data. It is common to begin with point estimates during screening-level risk assessments, beginning with a hazard quotient approach. However, as hazard quotients tend to incorporate conservative assumptions in the face of uncertainty, further evaluation is often needed following a screening assessment. In these situations, using methods that make greater use of the range of exposure and effects information is encouraged. It is acceptable to proceed in a sequential (tiered) manner through a range of methods that replace conservative point estimates with ranges of values or distributions.

### 5.3.1 Hazard Quotients and Hazard Indices

**Hazard quotient** (HQ) and **hazard index** (HI) are commonly used terms in risk assessment. However, other terms with similar definitions may be employed within the jurisdiction. For example, in Québec, the terms *quotient de danger* (hazard quotient) or *indice de danger* (hazard index) refer to theoretical risks, whereas the terms *quotient de risque* (risk quotient) and *indice de risque* (risk index) are used for sites with existing contamination to indicate existing potential risk.

The simplest tool for evaluating a line of evidence is a HQ, which is the ratio between the exposure measure (numerator term) and a corresponding effect-based threshold (denominator term). HQs are widely applied, particularly in screening assessments, due to the ease of application and the prevalence of point-estimate values for both exposure and effects. The HQ has particular value as a screening tool, which may be all that is required in some risk assessments. However, where HQs are calculated, care must be taken not to infer more information from the ratio than is warranted, and to consider the effect of uncertainty in both the numerator and denominator (Section 5.3.1.1).

The exposure term for an HQ can be derived from many sources, including (see Section 3 for details):

- a measured concentration in an environmental medium (e.g., mg/kg zinc in soil, mg/L selenium in water)
- a simulated concentration in abiotic environmental media or organism tissues using a model (which can range from a simple partitioning model to a complex mechanistic environmental fate and bioaccumulation model)
- a modeled dose to an organism (mg/kg-day) from a food chain or trophic transfer model.

The threshold effects term can also be derived from numerous sources, including (see Section 4 for details):

#### Key Concept

A HQ is a ratio between an exposure term (dose or concentrations) and a response term:

\[
HQ = \frac{\text{Exposure}}{\text{Threshold Effect Level}}
\]
• an environmental quality guideline for abiotic media (soil, sediment, water, groundwater, etc.)
• a threshold value gleaned from a compendium of toxicological summaries
• a threshold value obtained from an independent literature review
• a threshold value from an SSD analysis
• a site-specific threshold developed from interpreting the results of a toxicity or community study conducted over a range of exposure levels at the site of interest
• a meta-analysis of multiple sources of effects information (e.g., compilation of results from multiple studies that may cover a range of endpoints and species).

HQs may be applied for any of the four major categories of evidence. In practice, the most common HQs are derived for chemistry measurements in abiotic media (e.g., comparisons to soil, water or sediment quality guidelines), bioaccumulation endpoints (e.g., screening against TRGs) and dose-based wildlife assessments (e.g., dividing the estimated dose derived from a food-web model by a toxicity reference value [TRV]). However, an HQ can also be calculated for site-specific toxicology or community studies; threshold effects benchmarks can be calculated from concentration-response curves developed from site data\(^{35}\) and then used for application to other stations or samples for which only chemistry data are available. Section 4 discusses the HQ denominator further.

5.3.1.1 Common Errors in Application

Although easy to derive, HQs are often misinterpreted (Allard et al. 2010). The most common error is to incorrectly assume that an HQ is directly proportional to the magnitude of risk. HQs do not contain information about the specific probability that an adverse effect will occur, nor do they convey the magnitude of a potential adverse effect. Instead, a typical HQ is calculated using conservative assumptions, in which case the ratio indicates only whether existence of adverse effects is either possible (HQ > 1) or unlikely (HQ < 1).\(^{36}\) In ERA practice, there is broad agreement that HQ \(\leq 1\) indicates negligible risk for the specified endpoint, because the HQ is usually calculated on the basis of conservative assumptions.

Another common error is to assume that HQs can be scaled across different COCs to provide reliable rankings of contaminant risk (Allard et al. 2010). However, as quotient methods are only as reliable as the values in the numerator and denominator (with associated uncertainty), the degree of hazard cannot be directly compared. The derivation methods for different COCs can result in large differences in conservatism that are masked by presentation of simple ratios. Similarly, separate HQ values for the same COC cannot be linearly scaled to risk, because the intercept, slope

\(^{35}\) Concentration-response profiles are discussed further in Section 5.3.2.

\(^{36}\) In some cases, the threshold HQ is adjusted downward from 1.0 to 0.1, 0.2 or other values, to compensate for lack of data for background exposure. In general, these approaches are arbitrary and should be avoided (similar to arbitrary safety factors); instead, the uncertainty in total exposure estimates should be addressed explicitly.
and shape of the dose-response relationship is not reflected in the point estimate HQ (e.g., an HQ of 4 for APEC 1 cannot be assumed to be twice the risk of an HQ of 2 for APEC 2). Reliable comparisons can be made only through detailed understanding of the underlying concentration-response relationships, safety (application) factors and uncertainties, none of which are conveyed by an HQ.

Although a very large HQ suggests a greater “risk” than an HQ slightly greater than 1, it is not possible to draw conclusions about relative risk based on differences in HQs (e.g., HQs from 1 to 10 indicate moderate risk, while HQs above 10 indicate high risk). In addition, it is not true that minor changes in the HQ provide a meaningful differentiation (Allard et al. 2010; Ritter et al. 2002). For this reason, including excessive significant figures implies a level of certainty and precision that is not actually present; most HQs can be rounded to one or two significant figures.

5.3.1.2 Interpreting HQs
Because point estimates are applied in HQ derivation, the underlying uncertainty, bias and variability in the data are masked. Therefore, interpreting HQs requires explicitly considering the selection of the numerator and denominator terms by asking the following questions:

- Are the terms central tendencies or conservative estimates?
- If conservative estimates are applied, are they based on “worst-case” assumptions (such as use of maximum observed concentrations and the most sensitive species)?
- Is the effects term based on a NOAEL, LOAEL or a threshold effect size, and was the threshold response level bounded in the study design used to derive the threshold?
- Were application factors (margins of safety) applied to the estimates?
- Were thresholds derived from a broad range of studies and endpoints, or from limited data?
- Were exposures assessed through detailed profiling over space and time, or from isolated measurements?
- Were exposures estimated using uncertain models with high inherent uncertainty?

**Key Concept**
A HQ is only as reliable as the information used to parameterize the numerator and denominator. As such there is no universal system for interpreting the magnitude of an HQ (beyond comparison to 1.0), and different types of HQs are not directly comparable.

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37 In some cases, quasi-probabilistic HQs that account for some uncertainties in the numerator or denominator may be calculated. This approach is rarely applied, but may be appropriate if reviewers or regulators prefer to evaluate the site using only a quotient. Whereas it is always important to convey the types of uncertainty considered in a probabilistic assessment, it is particularly important for cases when probabilistic assessment is limited to the effects term or the exposure term.
In a screening-level assessment, the standard approach is to apply conservative measures in both the numerator (upper-bound estimate of exposure) and the denominator (lower-bound conservative guideline).

As a general rule, applying safety factors (application factors) when calculating HQs is discouraged, as discussed in Section 4.5. Chapman et al. (1998) and Forbes and Calow (2002) discuss the pitfalls of assigning arbitrary or default safety factors in ERAs. Depiction of uncertainty in HQs is better handled through a separate uncertainty analysis that conveys the plausible range of risk estimates using different assumptions for exposure and effects parameters. This may be done probabilistically or through a bounding analysis.

5.3.1.3 Linking HQs to Spatial Units

One issue in applying HQs is how they incorporate spatial variations in exposure levels. The procedures vary depending on the characteristics of the receptor under evaluation:

- For receptors with large home ranges, a single HQ can be calculated for the entire site.\(^{38}\) This entails use of an exposure metric such as the arithmetic mean or the 95 per cent UCLM for all the measured values for each medium, or the maximum measured concentration (Gilbert 1987). The degree of conservatism in the resulting HQ will depend on the metric used, the number of samples and the variability among the samples.

- For sessile receptors or those with small home ranges, spatially distinct risk quotients can also be calculated depending on the spatial definition of the local population, and the probability of exceeding an HQ of a given magnitude can be computed. This technique is generally applied when the single HQ method (screening assessment) yields a value above 1.0 and where the single HQ method is considered to be over-conservative.

Because these refinements still rely on the HQ as the underlying tool for evaluating risks, their primary use for highly contaminated sites may be to identify areas where more detailed evaluation of risks is warranted.

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\(^{38}\) If a wide-ranging receptor of concern has specific habitat preferences that discourage use of portions of the site, the procedure described here can be modified by adjusting the exposure metric (i.e., exclusion of data from non-relevant habitats).
5.3.1.4 Hazard Indices and Multiple Substance HQs

The hazard index (HI) is a simple metric used to aggregate hazard from multiple substances. The HI is the sum of the individual HQs for substances that have the same mechanism of toxic action. The implicit assumption in HI calculation is that risks from multiple substances are additive when the mechanism of toxic action is similar. Because different pollutants may cause similar adverse health effects, it may be appropriate to combine HQs associated with different substances.

As with the HQ, aggregate exposures below a HI of 1.0 will likely not result in adverse responses. However, the HI cannot be translated to a probability that adverse effects will occur and is not likely to be proportional to risk.

Combining values through summation (HI approach) using existing toxicological data is not well supported for most substances. There are two main reasons why hazard indexing is discouraged for most substances:

- Individual HQs are derived conservatively, and summation of individual HQs compounds this conservatism.
- Summation of risks is appropriate only for contaminants that act via the same mechanism of action. Most contaminants exhibit different toxicological mechanisms, so the scientific basis for calculating HIs for mixtures of contaminants is weak for most receptors.

Where the mechanism of action is known, and relative toxicity of related substances can be quantified, approaches are available to integrate the effects of groups of related contaminants. These approaches apply to select groups of contaminants that are known or strongly believed to exert toxicity through a single mode of toxic action. For example, non-polar organic contaminants commonly exert direct toxicity via narcosis, a reversible state caused by non-specific interaction of lipophilic molecules with biological membranes (Escher and Hermens 2002). As a result, some guidelines consider the cumulative effect of chemicals that act via this mechanism (Di Toro and McGrath 2000; Di Toro et al. 2000). Furthermore, for some hydrophobic chlorinated organic substances believed to act via the aryl hydrocarbon (Ah) receptor (e.g., dioxins, furans and a subset of PCB congeners), TEQ systems have been developed to simultaneously account for the relevant congeners once normalized to their receptor binding affinities. It is acceptable to apply these equivalence systems when calculating HQs, but

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39 Note: CCME has developed other indices to evaluate the potential effects of multiple contaminants. For example, the CCME water quality index (CCME 2001-b) evaluates water quality based on the number of contaminants exceeding CEQGs as well as the magnitude and frequency of those exceedances. The CCME water quality index scores water quality on a scale from 0 to 100 and categorizes scores from poor to excellent.
it is not acceptable to add HQs developed using different systems (e.g., one should not add a total PCB HQ to a PCB TEQ HQ, or add a total dry weight PAH HQ to either an HQ derived using the narcosis model or to HQs calculated for individual PAHs). Where such systems exist, they are preferable to application of HIs, as the latter may not account for the mechanistic understanding of contaminant potencies in mixtures.40

Where multiple contaminants are considered simultaneously, several assumptions may apply to derivation of the effects threshold, including:

• Concentrations of substances in the mixture are treated additively, with no assessment of relative toxic potential (e.g., total PAH threshold in sediment that does not discriminate among individual PAHs in the mixture).

• Concentrations of substances in the mixture are treated additively, and adjusted for relative potency (e.g., TEQ systems for narcotic effects of PAHs in porewater, or for dioxins/furans through \(\text{Ah} \) receptor binding affinity).

• Concentrations of substances in the mixture are treated additively, but with a bioavailability correction prior to screening (e.g., molar difference between acid volatile sulphides and sum of simultaneously extractable metals).

However, most COCs do not have established methods for assessing the synergistic or antagonistic effects of interactions with other substances.

5.3.2 Using Concentration-Response Relationships

Concentration-response relationships are typically derived as part of a hazard assessment, and the general methods of analysis are first described in the problem formulation. However, their use is almost always tailored to the data, which means that details regarding analysis of concentration-response relationships are often part of risk characterization. For example, a concentration-response model that fits the data well at low concentrations but not at high concentrations may be acceptable if measured or estimated concentrations are low. Concentration-response relationships are presented here rather than in Section 4 because of the emphasis on their use in practice.

A concentration-response relationship provides an assessment of the statistical relationship between an exposure term and a response term. Rather than provide a single threshold to describe the chemical potency of a COC, a concentration-response relationship describes the relationship between exposure and response over a range of exposure levels and effect sizes.

Definition:

Concentration-response relationship: A mathematical assessment of how an exposure term relates to the observation of a biological or toxicological effect.

40 Note that individual substances may have established putative effect levels that are based on empirical association (co-occurrence assessment) rather than relative potency established by mechanistic assessment. In the former case, summation of HQs is inappropriate.
Figure 5-1: Graphic example of a concentration-response relationship

The above graphic depicts an example of a concentration-response relationship for a single experiment. In the example, the y-axis represents the response measure, which could be survival, growth, reproduction, or any other toxicological or biological measure at the organism, population or community level. The x-axis displays a range of exposure conditions under which the experiment was performed (in this example, seven evenly spaced treatment levels, with multiple replicates for each treatment, and variance for each treatment indicated by the error bars).

5.3.2.1 Advantages
The information contained in Figure 5-1 is substantially greater than what is conveyed through use of a point estimate, such as the NOAEL or LOAEL, for several reasons, including:

- The response magnitude is defined at multiple exposure concentrations. For example, whereas a 50 per cent response occurs at approximately 40 mg/kg, the dashed line shows that a 25 per cent response occurs at approximately 34 mg/kg.
- The response curve illustrates the steepness of the response profile (total inhibition of response occurs within a factor of 3 of the concentration showing negligible response).
- The variability among replicates is depicted, providing an indication of the variation of the response (part of the uncertainty).

The preferred procedure is to perform a direct estimate of effects and risks using concentration-response analysis. The key difference is that a quotient compares exposure to a point estimate for effects information (e.g., a TRV), whereas a true risk estimate explicitly evaluates the mathematical relationship between site-specific exposure and response level across a range of
relevant exposures (e.g., mathematical evaluation of response magnitude versus chemical concentration or dose).

In the above example, the 25 per cent effect level (34 mg/kg) could be used to derive a TRV. Alternatively, a more stringent response magnitude, such as the 10 per cent effect level (25 mg/kg) could be applied. By plotting measured or estimated exposure on the curve, the estimated response can be understood for exposures greater than or less than these exposure values. Without the curve, it is not possible to understand the magnitude of response associated with the measured or estimated exposure.

5.3.2.2 Disadvantages
Although concentration-response models are conceptually attractive, there have limitations, including:

- In natural systems, a clear relationship between exposure and response is rarely observed (whether linear, sigmoidal or other shape). Frequently, the relationship is an “interrupted” concentration response in which one or more treatments do not follow a smooth pattern.41
- Data limitations (such as limited exposure levels or lack of information specific to the species of interest) restrict the application of the method.
- Other factors can confound the relationship between exposure and response. In the example of Figure 5-1, it is plausible that the response was caused by a factor that covaried with the COC concentration, such as soil pH or mean particle size.
- Concentration-response curves are challenging (and expensive) to derive on a site-specific basis, due to the number of treatments and replicates required to achieve confidence in the relationship.
- Numerous mathematical functions are available to quantify the relationship, and selecting the appropriate function can be challenging, especially when it is used to extrapolate beyond the range of measured exposures. Formal methods of model selection based on Akaike’s information criterion (AIC) and other criteria are available and should be considered (Burnham and Anderson 2002).
- Where concentration-response data are derived from the literature, results must be transferable to the context of the site. For example, if the example relationship was based on well-drained soils, but the site consisted of bog-like conditions, the relationship implied by the curve may be inapplicable to the site context. Similar considerations apply to other chemical-specific factors (e.g., metal speciation, modifying factors such as dissolved organic matter) and also to biological factors (e.g., representativeness of surrogate organism, physiological tolerance of local organisms).

41 An idealized concentration-response relationship is shown in Figure 5-1 to facilitate understanding of the approach.
Many of these disadvantages stem from data limitations. Because data limitations also affect our ability to derive point-estimate TRVs and HQs, practitioners should not be discouraged from exploring concentration-response relationships simply because data are limited. Those same data limitations will carry large uncertainties regardless of what methods are used for hazard assessment and risk characterization.

5.3.2.3 Application
The “true” risk estimate (based on concentration-response profiling) is what CCME (1996-a) envisioned as “detailed” risk characterization. This is consistent with the knowledge that quotient-based methods do not provide estimates of risk because they cannot characterize the probability and magnitude of effects. In some cases, quotient-based approaches have been applied even beyond screening-level risk assessments, due to several factors, including limited data for understanding concentration-response relationships. However, in many cases data are adequate for supporting concentration-response analysis, and practitioners should aim to analyze concentration-response data explicitly whenever possible for detailed risk assessments. For some measurement endpoints (e.g., aquatic toxicity tests conducted using dilution series), characterization of the concentration-response relationship is a natural outcome of the test results.

In addition to their use for risk estimation, site-specific concentration-response data have another use: establishing site-specific correlation or insight into causality (causality and its role in risk characterization is explored in detail in Section 5.5). Site-specific correlative approaches evaluate the association between contaminant concentrations and levels of response; they include formal statistical association methods and qualitative evaluations. Levels of response can be derived from any measurement endpoint used in a risk assessment, from a toxicity test endpoint (e.g., growth or reproduction in a lab bioassay) to a direct community measure (e.g., total organisms or total taxa measured in a benthic invertebrate sample).

Regardless of how concentration-response data are used, quantitative models that relate responses to any predictors should be appropriate for the data. For example, dichotomous outcomes (e.g., survival in a toxicity test) should usually be evaluated using a generalized linear model (e.g., logistic regression) that assumes the correct (binomial) error structure for the data. In addition, models that are fit to grouped data (e.g., dilution series bioassay results from more than one sample station) should use methods that account for the structured nature of the data (Pinheiro and Bates 2000; Wheeler and Bailer 2009).

In short, advanced statistical methods beyond simple linear regression are often necessary, and can facilitate evaluation of concentration-response relationships while simultaneously explicitly considering the influence of categorical and continuous factors on the nature of the relationship.

Key Concept
Developing concentration-response curves requires understanding the underlying statistical assumptions. Practitioners should consult toxicity test protocols and/or a biostatistician when applying statistical models.
5.3.3 Using Reference or Background Condition

Many measurement endpoints pertaining to biological or toxicological parameters cannot be readily interpreted at face value (e.g., a species richness value of 12 has little meaning until placed in ecological context). Rather, they must be compared to a reference condition if a gradient design (Section 5.3.4) is not used or not feasible (see Section 2.3.5.1 for further discussion). Accordingly, an important tool for risk characterization is the control-impact design or other comparative approaches. The general experimental design is established during problem formulation. This section focuses on application and use of the data during risk characterization.

Several types of samples can be used to standardize site responses, including:

- **Negative controls**: Clean artificial substrate or test media used in the laboratory to evaluate test acceptability. These are not recommended for standardizing site responses, as lab conditions may not represent the environment relevant to the site.\(^2\)

- **Reference condition**: Media collected in the general vicinity of the site, but confirmed to be less contaminated relative to site media.

- **Background condition**: Media collected from the region at stations known to exhibit a lack of incremental contamination beyond naturally occurring concentrations.

For example, if a quotient approach is used to evaluate a particular line of evidence, it can be useful to compare the quotients derived for on-site conditions and compare them to quotients derived for a range of off-site conditions. Sometimes, the difference in risks between a site and a reference condition are as important to understand as the absolute magnitude of estimated risks, particularly when conservative assumptions are built into a risk assessment. For example, in some environments, natural mineralization can elevate regional background concentrations above screening values and generate false-positive HQ values for metals that exceed 1.0.

For assessments of risks to wildlife that are based on total dose, comparative approaches are particularly useful for identifying how various exposure media are contributing to the incremental risk on site compared to off site.

One particular application of a control-impact design, based on application of multivariate methods, is the reference condition approach (RCA; also called the reference envelope approach). This procedure can be applied to both toxicity and field community studies. The procedure has been proposed as an alternative approach to overcome limitations of reference and negative control

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\(^{2}\) Negative control media are primarily intended to evaluate the sensitivity of test organisms to handling and manipulation. As such, the substrates are often simplified or artificial (e.g., silica sand), unless the laboratory has adopted a natural substrate. Practitioners should consult with laboratories before testing and consider using additional clean controls better matched to site conditions.
samples. These limitations include differences in non-contaminant characteristics (substrate, habitat, etc.) and low statistical power when many samples are compared to a single control or reference. The RCA for benthic invertebrate sampling (Reynoldson et al. 1997) selects multiple reference sites from a reference database to serve as the control, while individual test sites provide the treatment, and applies multivariate ordination methods (such as non-metric multi-dimensional scaling) to distinguish patterns among samples. Confidence bands (ovals) around the data (Figure 5-2) indicate the degree of statistical similarity of test samples in relation to the suite of references. This approach is currently the basis of the Canadian Aquatic Biomonitoring Network (CABIN) (Reynoldson et al. 2006) and has been promoted in southern California for the interpretation of biological data (Surface Water Ambient Monitoring Program [SWAMP] 2009). This approach first range standardizes toxicological data, and then applies Euclidean distance (a multivariate similarity measure) as the distance coefficient.

A drawback of this method is the difficulty of collecting data on a large enough pool of reference sites to ensure the data is conclusive. In the reference approach, practitioners must consistently apply the selection criteria for reference areas. Also, an ecological relevance check is required to ensure that reference stations are appropriately matched to exposed stations in terms of key environmental variables (organic enrichment, substrate type, depth, etc.). Finally, the RCA as envisaged by CABIN also requires a considerable pool of reference sites that meet these criteria.

5.3.4 Gradient Designs

Given the challenges of determining reference conditions against which site conditions can be compared, experimental designs based on gradients should be considered whenever possible, and the specific design should be determined as part of problem formulation (see Section 2.3.4 for more discussion). For example, if there is a historical point source of contaminants, it may be useful to correlate response measures to distance from that point source. Alternatively, if contaminant concentrations are known, the gradient may simply be based on categorizing spatial units according to contaminant concentrations. If a gradient design is envisaged in a risk assessment, it should consider how to best align the sampling design with the fate and transport pathways. For example, identification of spatial gradients may consider the following:

- distance or direction from a known source
- historically observed gradients in contaminant concentrations and interaction with physical factors such as water depth, salinity and substrate type.

Key Concept

In examining potential gradients, practitioners may need to consider information other than raw COC concentrations, such as factors influencing bioavailability (e.g., organic carbon, coal particles, sulphides) or physical factors (e.g., habitat, substrate).
Figure 5-2: Depiction of the reference condition approach for invertebrate communities

Source: Adapted from Rosenberg et al. 1999.

Note: Invertebrate communities at test sites that fall within the 90% probability ellipse are considered equivalent to reference sites, sites within the 99% probability ellipse are possibly different, sites within the 99.9% probability ellipse are different, and sites outside the 99.9% probability ellipse are very different.

In applying the gradient approach, practitioners should provide representation of a wide range of exposure levels, ranging from exposures at or near the background condition to “worst-case” conditions found at the site. The greater the range in exposure concentrations, the better the ability to characterize a concentration-response relationship. If gradients are weak or poorly defined, additional uncertainty will be incorporated in the assessment of responses. Furthermore, if the range of exposure levels is small, natural variability may obscure a meaningful underlying relationship that would be revealed if there was greater variety of exposure conditions.
5.3.5 Multivariate Techniques

Multivariate statistical analysis refers to any of various statistical methods for analyzing more than two variables simultaneously. Assessing effects at a community or ecosystem level usually involves measuring a large number of abiotic and biotic variables. Assessing each variable individually or with many pairwise bivariate analyses can be cumbersome and difficult to interpret, and this method cannot detect patterns that emerge from the interactions of variables. Multivariate techniques can be used to summarize overall patterns from a large suite of variables (Bier 1999; EC 2002; Fairbrother and Bennett 2000; Sparks et al. 1999). Once the number of variables has been reduced, patterns in the data can be evaluated and compared to other data (e.g., if a chemistry data set is reduced to a couple of summary variables, those variables could be correlated to toxicity data using multiple regression or similar techniques).

While general multivariate techniques may be discussed during problem formulation, the details of analysis often cannot be specified until the data are evaluated. The appendices of SAB-CS (2008) provide an overview of the common multivariate statistical approaches and identify potential pitfalls with their application (Landis et al. [2011] also discuss potential pitfalls). See Sparks et al. (1999) for more information on specific techniques as they have been applied to risk assessment. Because of the complexity of multivariate approaches relative to univariate statistics, risk assessors should consult a qualified statistician with experience in biological or ecological investigations. Broad types of applications for multivariate techniques in risk characterization include ordination, clustering or discrimination, and investigating relationships between sets of variables (correspondence):

- **Ordination techniques** (e.g., principal components analyses) reduce a large set of variables into a smaller set of factors, each of which is a combination of variables that captures as much as possible of the information in the original variables. In this way, a multi-dimensional set of data can be reduced into a more interpretable form.
- **Clustering or discrimination techniques** identify natural groupings among sampling units (e.g., most-similar groups of sampling sites) and the parameters that contribute most to this similarity (e.g., abundances of certain species).
- **Correspondence analysis techniques** (e.g., canonical correspondence analyses [CCA]) identify the degree of covariance between sets of variables (e.g., concentrations of several chemicals versus abundances of several species). They also identify the variables within each set that contribute most to this covariance.

5.3.5.1 Advantages and Disadvantages

Multivariate methods are aimed primarily at data exploration and are usually used to reveal patterns that warrant more specific quantitative evaluation. They distill complex data sets down to a low number of dimensions (usually two or three) that capture the main sources of variation in
the data. Multivariate approaches are amenable to graphical presentation of results (e.g., cluster analysis dendrograms, ordination plots) that are often intuitive relative to a large stream of univariate plots (e.g., intercorrelation matrix). These advantages must be traded off against the following drawbacks:

- The results of multivariate analysis are complex and can be difficult to communicate. The underlying assumptions of the statistical procedures must be thoroughly evaluated.
- Multivariate methods are usually exploratory, and therefore cannot be defined in detail before data acquisition (e.g., one cannot define the number of required dimensions \(a\) \(priori\)).
- Environmental data are prone to violations of parametric statistical frameworks (e.g., normality of distributions, independence among inputs), requiring great care in application and interpretation, or use of non-parametric techniques.
- Output of some multivariate methods cannot easily be translated to decision rules for ecological significance. For example, the axes of a principal components analysis ordination do not have defined units, and therefore differences in any dimension are challenging to interpret in terms of environmental relevance.
- Some methods are sensitive to data constraints such as missing values and non-detected concentration data.
- The meaning of each axis must be evaluated using correlations with the individual inputs.

With respect to interpreting findings, a significant issue for risk characterization is how to score and weight the findings of ordination methods. The results of these techniques are not conducive to an IC\(_{20}\) or other effect size–based categorization. The output is useful for identifying effects (relative differences among stations, station groupings or relative to reference), but interpreting the ecological significance is more challenging. Determining whether the differences among stations are ecologically meaningful requires a two-stage evaluation:

- Analyze the factors and variables that caused the observed divergence in ordination (e.g., which taxa are more or less common at extremes of each non-metric multidimensional scaling [NMDS] axis).
- Assess the functional importance of these differences in terms of community health. This step requires professional judgment, as it entails discerning between observed differences that are not necessarily negative phenomena and differences that indicate degradation of the community.

One specific application of multivariate methods, the RCA, is elaborated in Section 5.3.3.
5.3.6 Probabilistic Methods

Probabilistic methods acknowledge that natural ecological features are not constants, but rather are variable and complex, and that our understanding of their properties is not complete. Probabilistic models describe the state of one or more random variables as a distribution of possible values rather than fixed values (point estimates). Using probabilistic methods, important biological, chemical, physical and environmental parameters are assumed to vary or are uncertain and therefore are specified using distributions.

Most ERAs are conducted using point estimates for exposure and effects parameters. This is acceptable for many assessments (e.g., preliminary assessments) because using point estimates with appropriate conservatism to account for uncertainty can effectively screen numerous pathways with relatively little effort. However, for residual risks it is sometimes difficult to ascertain the influence of compounding conservatism on the risk assessment. Additionally, there are some parameters for which it is difficult to incorporate conservatism because the degree to which a parameter is conservative depends on how it is applied.

For example, a specific dietary preference, such as consumption of fish by mink, can be increased to err on the side of overestimating exposures when applied in a forward modelling mode. This is because fish tend to have higher concentrations of contaminants relative to other food items. However, if the purpose of the risk assessment is to identify threshold concentrations in various dietary items, such that the total daily dose is no larger than the TRV, the situation is more complex. Specifically, the calculation of total blended ingestion rate would be biased toward the fish pathway, such that the sensitivity to changes in aquatic contamination would be exaggerated, whereas changes to terrestrial contamination would be understated. In this situation, intentional use of a high-end point estimate for dietary preferences could result in a management decision for soil (e.g., threshold soil concentration derivation) that is contrary to the objective of conservatism. Probabilistic methods can help to resolve such problems by representing parameters as a range of plausible values rather than relying upon the relevance of a point estimate.

Probabilistic methods can be used when applying the quotient method, or when investigating concentration-response information (e.g., estimates of actual risk). In the case of the quotient method, the result may be a probability distribution of quotients that allows estimation of the probability that HQ > 1.0. In the case of concentration-response information, the result of a probabilistic assessment may be a probability distribution of effect rates; integration of this distribution provides an estimate of the expected risk, rather than the maximum likelihood estimate of risk. Probabilistic risk assessment explicitly acknowledges the stochastic or uncertain nature of model parameters, and attempts to describe the effect of multiple and linked parameter distributions. Probabilistic methods can be applied separately during the exposure assessment or hazard assessment, but can also be applied during risk characterization. Section 5.6 explores some

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**Key Concept**

Probabilistic methods replace point estimates with distributions. These methods may simulate the effect of natural variations (stochasticity), uncertainty in knowledge (incertitude) or a combination of both.
details regarding probabilistic methods in the context of evaluating uncertainties. Additional guidance and references are summarized by Suter (2007).

5.4 Step 3: Prepare Compiled Data Summary

A simplified data summary is a relatively simple but effective risk characterization tool. The intent is simply to summarize the range of endpoint data (without any sophisticated interpretation), with results for multiple endpoints organized by sampling station, habitat type or management unit. The risk assessor (or a reviewer) can refer to this table during risk characterization. The compiled data provide useful reference material that may be lost in a complicated WOE process (Section 5.5). Table 5-1 provides an example of a simplified compiled data summary. The data (normalized to the reference conditions and guidelines) are placed into categories of response, with no additional interpretation provided.

Table 5-1: Example of a compiled data summary

<table>
<thead>
<tr>
<th>Station ID</th>
<th>Chemistry</th>
<th>Toxicity</th>
<th>Community</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Metals</td>
<td>PAHs</td>
<td>Amphipod</td>
</tr>
<tr>
<td></td>
<td>As</td>
<td>Cu</td>
<td>LPAH</td>
</tr>
<tr>
<td>NF-1</td>
<td>O</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>NF-2</td>
<td>2</td>
<td>3</td>
<td>2</td>
</tr>
<tr>
<td>FF-1</td>
<td>2</td>
<td>O</td>
<td>3</td>
</tr>
<tr>
<td>FF-2</td>
<td>3</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>FF-3</td>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>FF-4</td>
<td>O</td>
<td>O</td>
<td>2</td>
</tr>
</tbody>
</table>

Chemistry data: O indicates below sediment quality guideline; 2 indicates above sediment quality guideline, with number in symbol representing degree of exceedance.

Toxicity data: O indicates negligible to low effect size (below 20%); 2 indicates moderate effect size (20%–50%); 3 indicates high effect size (> 50%) (all relative to reference).

Benthic data: O indicates negligible to low effect size (below 20%); 2 indicates moderate effect size (20–50%); 3 indicates high effect size (> 50%) (all relative to reference).

Table 5-1 provides details on a sample-by-sample basis, but is simplified in two ways. First, it bins the raw data into categories rather than reporting actual effect sizes. Second, it presents only the information on magnitude of effects, not information on causality, uncertainty, ecological relevance or any other attribute that may be relevant for evaluating lines of evidence. This simplified table may be most appropriate for cases where data indicate minimal risks, or where the complexity of the response profile is low.
An alternative to Table 5-1, more applicable in cases where data show significant or complex indications of responses, is to present the absolute values of the endpoint responses (with numerical values and no categorization) and to also present raw information on evaluation of causality (in anticipation of supporting the WOE assessment outlined in Section 5.5). Table 5-2 shows an example format for a single line of evidence, soil invertebrate richness. A summary of this type attempts to present information at face value without complex interpretation. In this case, results are usually not presented on a sample-by-sample basis but for an entire site or portion of a site.

The challenge is to present a condensed version of the field results (for simplicity or review) without introducing excessive manipulation of the data or professional judgments. Data summaries may vary in scope, but their role in facilitating review by regulators and others should not be underestimated. The two formats presented here have advantages and disadvantages as described above. For complex ERAs, using both formats may be appropriate.

**Table 5-2: Example of a compiled data summary**

<table>
<thead>
<tr>
<th>Line of evidence</th>
<th>Magnitude</th>
<th>Uncertainty about magnitude</th>
<th>Evidence for causality</th>
<th>Uncertainty about causality</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Soil invertebrate community richness</strong></td>
<td>Average richness 15% lower compared to reference condition.</td>
<td>t-test not significant (p = 0.22) but sample size limited; reference condition based on only three sites.</td>
<td>Linear regression indicates richness weakly inversely correlated with soil zinc concentrations</td>
<td>Regression not significant (p = 0.48) and explains little of the variation (r² = 0.08). The best predictor of richness is soil moisture (marginally significant at p = 0.09).</td>
</tr>
</tbody>
</table>
5.5 Step 4: Conduct Weight of Evidence Procedure

The term weight of evidence (WOE) is defined here to mean any process used to aggregate information from different lines of scientific evidence to render a conclusion regarding the probability and magnitude of harm. This definition encompasses a range of practice, ranging from best professional judgment (BPJ) assessments to complex quantitative methods (see Figure 5-3).

Box 5.1: WOE procedure

| The default procedure recommended in this guidance involves the following steps: |
| 1. Summarize each line of evidence based on magnitude of effects (including spatial extent), evidence for causal relationships between contaminants and effects, and ecological relevance. The methods of scoring or ranking each of these attributes should be established in advance (as in Table 5-3). The final line of evidence summary tables should be organized by assessment endpoint. Table 5-4 (terrestrial) and Table 5-5 (aquatic) provide examples for typical lines of evidence. |
| 2. As part of the line of evidence summary, evaluate uncertainty regarding magnitude of effects and evidence for causality for each line of evidence. (Uncertainty is evaluated more broadly in Step 5 following the WOE procedure, but must also be evaluated here to characterize specific uncertainty regarding magnitude and causality for each line of evidence). |
| 3. For each assessment endpoint, make an integrated evaluation of findings for all lines of evidence, taking into account the degree of concordance among the various lines of evidence for that assessment endpoint (i.e., do the lines of evidence tell the same story?). The integrated evaluation should be based on a narrative rationale that clearly articulates how the overall evaluation was derived. |

This guidance prescribes a default WOE procedure (see Box 5.1) that will be applicable to most sites.

During the WOE step, the results for the individual lines of evidence obtained in Step 2 (and summarized in Step 3) are integrated. This provides a basic structure for all WOE assessments that provides a degree of consistency and transparency necessary for technical review of the document. The following subsections provide rationale for and details of the recommended default procedure for conducting WOE evaluations, as presented in Box 5.1.

5.5.1 Frame Purpose and Type of WOE

This guidance document provides a default WOE approach that builds upon the prescribed three-step WOE approach described in Box 5.1. The default WOE approach described below is likely to be applicable for most sites. Other WOE approaches have been described in the literature and may be used if better suited for specific sites or specific types of ERAs. Linkov et al. (2009) provide a simplified but useful summary of the range of WOE methods available (Figure 5-3). They note
that although all WOE methods may include both qualitative and quantitative considerations, the methods can be ordered by increasing degree of quantification along a continuum.

Figure 5-3: Classification of WOE approaches in risk assessment

<table>
<thead>
<tr>
<th>Method</th>
<th>Method description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Listing evidence</td>
<td>Presentation of individual lines of evidence without attempt at integration</td>
</tr>
<tr>
<td>Best professional judgment</td>
<td>Qualitative integration of multiple lines of evidence</td>
</tr>
<tr>
<td>Causal criteria</td>
<td>Criteria-based methodology for determining cause and effect relationships</td>
</tr>
<tr>
<td>Logic</td>
<td>Standardized evaluation of individual lines of evidence based on qualitative logic models</td>
</tr>
<tr>
<td>Scoring</td>
<td>Quantitative integration of multiple lines of evidence using simple weighting or ranking</td>
</tr>
<tr>
<td>Indexing</td>
<td>Integration of lines of evidence into a single measure based on empirical models</td>
</tr>
<tr>
<td>Quantification</td>
<td>Integrated assessment using formal decision analysis and statistical methods</td>
</tr>
</tbody>
</table>

Source: Based on Linkov et al. (2009).
An intermediate degree of quantification is likely to be appropriate for most cases (Suter and Cormier 2011), and therefore is recommended as a starting point for sites. The most qualitative approaches and the most quantitative approaches (e.g., the extremes along the continuum depicted in Figure 5-3) often may not be appropriate. Qualitative approaches do not provide a transparent system of reaching integrated conclusions, whereas quantitative approaches can be difficult for risk managers or stakeholders to understand due to computational complexity. This does not mean that such approaches may never be applied. Rather, it requires that rationales be provided where the “extremes” are chosen, and that consideration should be given to the potential weaknesses of these approaches during implementation.

The following broad principles should be incorporated into the default WOE procedure or any alternative procedure that is used:

- Lines of evidence should be integrated in the context of the assessment endpoints, the valued ecosystem components (VECs) and environmental protection goals. Specifically, the WOE needs to consider the level of organization of interest (individual, population or community) and explicitly address the linkage of the various lines of evidence to that level.
- The magnitudes of response observed for various measurement endpoints should be evaluated using rules that are as consistent as possible, such that various lines of evidence are compared using compatible decision criteria.
- The concurrence or divergence among outcomes of multiple measurement endpoints should be carefully evaluated.
- WOE determinations may be quantitative or qualitative, but should always be transparent.
- Professional judgment may be exercised, but a transparent analysis should be applied to elucidate the influence of professional judgment on the results.
- The degree of confidence in the conclusion for each endpoint is nearly as important as the conclusion itself.

Put more succinctly, when presenting the results of an assessment, the risk assessor should strive for the achievement of the following TCCR principles (US EPA 2000):

- transparency
- clarity
- consistency
- reasonableness.

These principles can be difficult to quantify but are important to any risk assessment. In some situations, such as ERAs conducted for FCSAP, these principles will become mandatory aspects of the WOE procedure.
5.5.2 Major Attributes Used to Evaluate Lines of Evidence

Applying WOE is based in part on considering attributes that are used to evaluate each line of evidence. The recommended default WOE procedure for federal sites considers the following attributes:\(^{43}\)

- **Magnitude of response and associated uncertainty**: This includes effect size, probability of occurrence, spatial scale and temporal scale.
- **Evidence for causality and associated uncertainty**: This is the observed response likely to be associated with site-related contaminants.
- **Ecological relevance**: To what extent does the line of evidence represent the assessment endpoint of interest?

Each of these attributes is discussed below. This list of attributes emphasizes the importance of both magnitude and causality, although to some extent they can be evaluated sequentially. If there is zero magnitude of response (i.e., no effects), there is no need to look for a cause. Conversely, if a large response is measured, evaluation of causality is of critical importance (for further discussion see FCSAP Module 4 [EC 2013]; Hull and Swanson [2006]; Landis et al. [2011]; Suter [2007]; Suter et al. [2010]).

For the default WOE procedure, uncertainty regarding the magnitude of response and evidence for causality is not specified as a stand-alone attribute per se, but is an integral component of the evaluation of each line of evidence. Uncertainty is a function of many factors, including the quality of the data, the ability of the line of evidence to detect effect sizes of interest, the degree to which responses are specific to the stressors of interest, and the spatial and temporal representativeness of the data. Several of these factors were listed in Section 2.3.4.2 in the context of selecting lines of evidence.\(^{44}\) A thorough WOE evaluation of uncertainty must consider these factors.

Importantly, ecological relevance and some of the factors driving uncertainty are considered not only during risk characterization, but also during problem formulation (see Section 2.3.4.2). Specifically, these considerations may serve as criteria for selecting measurement endpoints and lines of evidence. For ecological relevance, judgments made during problem formulation should be carried through to risk characterization unchanged. For example, a practitioner may judge that a lab-based bioassay has only moderate ecological relevance. That judgment should be made during problem formulation and will not change based on results of the bioassay. In the case of uncertainty, some of the criteria listed in Section 2.3.4.2 for selection of lines of evidence (e.g., anticipated data quality) ultimately become contributors to uncertainty assessment during risk characterization.

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\(^{43}\) Based in part on consideration of available WOE frameworks, including those developed by Exponent (2010), Hull and Swanson (2006) and Menzie et al. (1996).

\(^{44}\) Several factors influencing uncertainty have been identified in other WOE frameworks as formalized attributes (e.g., Menzie et al. 1996).
5.5.2.1 Magnitude of Response
The magnitude of any observed responses is arguably the most important attribute of a line of evidence. Defining the meaning of “magnitude” is an important consideration, as the term can refer to a number of characteristics, including:

- effect size (change or difference in the response variable) relative to AELs or levels considered potentially ecologically relevant
- spatial scale of the change or difference
- temporal scale of the change or difference
- probability of harm suggested by the analysis.\(^\text{45}\)

Given the importance of spatial scale for most receptors of concern, it is usually appropriate to separate spatial scale from effect size so that the two types of information are communicated clearly. Specifying the characteristics of magnitude is a mandatory component of risk characterization; without this articulation, narrative conclusions such as “high risk” have no clear meaning. For example, if soil at a particular site is highly toxic (e.g., mortality > 50%), risks would be considered more significant if the entire site was toxic versus only one small portion of the site.

Determinations of magnitude may be qualitative or quantitative. If categorical assignments are used, it is best to constrain the number of categories to five (e.g., negligible, low, moderate, high, very high) or less and to define the terms (and decision rules for break points among categories) clearly.

5.5.2.2 Causality
An assessment of causality in an ERA attempts to identify the cause of observed effects, and attempts to distinguish between associations that are coincidental (or caused by external factors) and associations that are driven by specific contaminant influences.

Ideally, causality is evaluated systematically. For example, a toxicity identification evaluation (TIE) evaluates the relationship between a cause (e.g., adjustment to a sample treatment) and an effect (e.g., modification of toxicity response magnitude) by testing each potential causal agent one at a time. However, in the absence of this type of systematic approach, wholly empirical methods can be used to provide insight into causality (e.g., circumstantial evidence), provided that a defensible underlying explanation for the response is given. Causality is explored in detail in FCSAP Module 4 [EC 2013] and by Suter et al. (2010).

\[\text{Definition}\]

\textit{Causation} is the act or fact of causing, or the production of an effect by a cause. Causation differs from \textit{association} (correlation) in that the latter does not imply a mechanistic linkage between observations.

\(^{45}\) Depending on the risk assessment type, probability of harm may not be a pertinent consideration. In a retrospective condition assessment, the site conditions are already manifested. In contrast, a predictive risk assessment involving a population model may invoke probabilities of population decline or extinction.
5.5.2.3 Ecological Relevance
Ecological relevance is a key attribute of any line of evidence considered during problem formulation and risk characterization. Ecological relevance reflects how relevant the line of evidence is to the assessment endpoint that it is intended to address. For example, direct measures of a community (e.g., invertebrate abundance and diversity) are generally considered to be more ecologically relevant than laboratory bioassays. Thus, a direct community measure carries greater strength for this attribute than a laboratory-based measure. However, laboratory-based measures may be more precise and better able to detect responses, so they would score higher for other attributes. The ecological relevance of any line of evidence should be evaluated during problem formulation as one of the criteria for selecting lines of evidence (see Section 2.3.4).

5.5.2.4 Attribute Uncertainty
Uncertainty is an integral component of risk characterization. Although considering uncertainty is part of the WOE procedure, Step 5 of the risk characterization process is dedicated to this issue (see Section 5.6) to ensure that uncertainty is rigorously addressed.

Uncertainty is the culmination of many individual factors (see Section 2.3.4.2 and Menzie et al. 1996). Some important categories of uncertainties include:

- **Sensitivity and specificity**: Sensitivity refers to the ability of a line of evidence to reliably detect a change in an environmental response despite the presence of natural or analytical variability and uncertainty. Specificity refers to the extent to which data, media, species, environmental conditions and habitat types used in the study design reflect the site of interest (Exponent 2010).

- **Data quality**: Data quality is the extent to which data quality objectives (DQOs) and other recognized characteristics of high-quality studies are met. Lines of evidence that apply precise and standard methods with accepted quality assurance and quality control (QA/QC) procedures are more valued. Lines of evidence that use novel methods not yet accepted by the jurisdiction in question or imprecise data with unacceptable QA/QC will have higher uncertainty in application (Exponent 2010). In addition, studies designed with appropriate statistical power and robust study designs are more valued.

- **Representativeness**: Representativeness is the degree to which the spatial and temporal nature of the data collected reflects real potential exposure and effects. The representativeness attribute is strongest for studies that:
  - conduct synoptic (simultaneous) sampling of measurement endpoints
- repeat sampling over multiple seasons or environmental conditions
- describe natural spatial or temporal variation through replication and characterization of stochasticity (random error).

### 5.5.3 Scoring or Ranking Attributes

Results for individual attributes described in the previous section must be evaluated. Commensurate with an intermediate level of quantification in risk characterization, each attribute can be summarized using scores or ranks such as negligible, low, moderate and high, or using integer scores or continuous numerical scores. The scoring and ranking system should be defined in advance during problem formulation to facilitate transparency in interpretation of results. Table 5-3 shows examples for typical types of lines of evidence, using the attributes for magnitude, causality and ecological relevance, along with associated uncertainties. For cases where more resolution is needed, or less resolution will suffice, practitioners may provide rationale for alternative approaches. Importantly, the classification of attribute performance as “negligible” or “low” or “moderate” or “high” should be consistent with protection goals and AELs articulated during problem formulation.

Once attributes are scored, they must be considered simultaneously to support overall evaluation of several lines of evidence for an assessment endpoint. In other words, the relative importance of magnitude, causality and ecological relevance must be weighed. This can be done quantitatively by combining some or many attributes into a common metric (e.g., Exponent 2010) or qualitatively by leaving attributes in their own units (e.g., Hull and Swanson 2006). The default approach presented here is based on leaving the major attributes in their own units, to increase transparency. Examples for the results of scoring lines of evidence are provided in Table 5-4 for terrestrial cases and Table 5-5 for aquatic cases. These example summary tables include evaluation of uncertainty (Step 5 of risk characterization), and they also include an overall evaluation of risks for each assessment endpoint (the subject of the next section below). The results shown in Table 5-4 and Table 5-5 must be generated transparently, using criteria defined in advance during problem formulation (e.g., Table 5-3).

### 5.5.4 Integrated Evaluation by Assessment Endpoint

Once individual lines of evidence have been characterized, the findings must be evaluated separately for each assessment endpoint (e.g., risks to wildlife are not traded off against risks to invertebrates). The final columns of Table 5-4 and Table 5-5 contain short narrative summaries of the key rationale used to make judgments about risks for each assessment endpoint. That rationale is a succinct

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46 Trade-offs among valued environmental attributes may be considered later as part of risk management. In that context, other factors, including human health concerns, socio-economics, and legal and financial concerns, may ultimately influence site management.
summary that can be elaborated in the main text of an ERA, usually as part of the narrative summary of risk conclusions that is articulated in Step 7 (Section 5.8).

The most important element of integrating findings across multiple lines of evidence is coherence. Coherence can be defined as the degree to which components are logical and internally consistent. This does not mean that all components must provide the same response type. Rather, it means that lines of evidence should ideally tell a story that is logical and orderly.

Coherence assessment is an opportunity for the risk assessor to provide a unifying explanation for the responses observed, given the information on each line of evidence, the uncertainty in the lines of evidence, and the relevance of the lines of evidence to the assessment endpoint.

The coherence assessment is where the logic connecting the various line of evidence findings should be articulated. The risk assessor should articulate overall findings for the line of evidence with a narrative explaining how contradictory results are reconciled. Also, the risk assessor should consider and acknowledge information and associated lines of evidence that were not available and therefore could not be considered in the WOE procedure.
Table 5-3: Example criteria for scoring attributes for major types of lines of evidence

<table>
<thead>
<tr>
<th>Degree of contamination and effect size</th>
<th>Chemistry (water, soil, sediment, tissue)</th>
<th>Toxicity tests</th>
<th>Quantitative measures of plant or invertebrate community abundance, biomass, richness</th>
<th>Qualitative measures of presence/absence or relative abundance</th>
<th>Comparison of dose-based exposure to TRVs (if food chain models are used)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Rating</strong></td>
<td><strong>Type of line of evidence</strong></td>
<td><strong>MAGNITUDE</strong></td>
<td><strong>CAUSALITY</strong></td>
<td><strong>Uncertainty about magnitude</strong></td>
<td><strong>Comparison of dose-based exposure to TRVs (if food chain models are used)</strong></td>
</tr>
<tr>
<td>Negligible</td>
<td>Chemistry is simply characterized as &quot;above benchmarks&quot; (for water, soil, sediment) or &quot;elevated&quot; relative to reference or local gradient; differentiation on the basis of degree of contamination is not used.</td>
<td>Relative effect size &lt; 10%</td>
<td>Relative effect size &lt; 10%</td>
<td>Subjective evaluation based on combined consideration of HQs on site relative to reference, and for common species, likely population-level implications.</td>
<td></td>
</tr>
<tr>
<td>Low</td>
<td>Above standards/criteria/guidelines.</td>
<td>Relative effect size 10%–20%</td>
<td>Relative effect size 10%–20%</td>
<td>Subjective evaluation based on spatial patterns.</td>
<td></td>
</tr>
<tr>
<td>Moderate</td>
<td>Analysis for individual samples and groups of samples across portions of the site.</td>
<td>Relative effect size 20%–50%</td>
<td>Relative effect size 20%–50%</td>
<td>Analysis of spatial gradients over the areas where sampling occurs.</td>
<td></td>
</tr>
<tr>
<td>High</td>
<td>Analysis for individual samples and groups of samples across portions of the site.</td>
<td>Relative effect size &gt; 50%</td>
<td>Relative effect size &gt; 50%</td>
<td>Analysis on an area basis (probably the entire site or sampling area).</td>
<td></td>
</tr>
</tbody>
</table>

**MAGNITUDE**: Spatial scale for evaluation of magnitude

**CAUSALITY**: Evidence for causality

**Uncertainty about magnitude**: Subjective evaluation based on number of samples, quality and number of reference samples, and any other relevant considerations.

**Evidence for causality**: Qualitative or quantitative evaluation of potential link between contamination and a site-related source. For tissue chemistry, spatial concordance between tissue and other media is evaluated.

**Analysis for individual samples and groups of samples across portions of the site.**

Subjective evaluation based on combined consideration of study design, sample size, statistical significance, explanatory power. Rationale provided in each case.

Subjective evaluation based on combined consideration of study design, sample size, statistical significance, explanatory power. Rationale provided in each case.

Subjective evaluation based on combined consideration of spatial patterns with analysis of spatial gradients over the areas where sampling occurs. Analysis on an area basis (probably the entire site or sampling area).
<table>
<thead>
<tr>
<th>CAUSALITY</th>
<th>Uncertainty about causality</th>
<th>Rating</th>
<th>Chemistry (water, soil, sediment, tissue)</th>
<th>Toxicity tests</th>
<th>Quantitative measures of plant or invertebrate community abundance, biomass, richness</th>
<th>Qualitative measures of presence/absence or relative abundance</th>
<th>Comparison of dose-based exposure to TRVs (if food chain models are used)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low</td>
<td>Subjective evaluation based on combined consideration of study design, sample sizes, and understanding of site characterization. Rationale provided in each case.</td>
<td>Low</td>
<td>Subjective evaluation based on statistical significance, number of samples, number of controls and reference samples, extrapolation assumptions, and any other relevant considerations.</td>
<td>Subjective evaluation based on statistical significance, number of samples, number of controls and reference samples, extrapolation assumptions, and any other relevant considerations.</td>
<td>Subjective evaluation based on level of rigour in the measures used.</td>
<td>Subjective evaluation based on degree of concordance with chemistry data, sample sizes, etc.</td>
<td></td>
</tr>
<tr>
<td>Moderate</td>
<td>Tissue chemistry, when compared to tissue-based critical body residues.</td>
<td>Moderate</td>
<td>Endpoints for mortality, growth, reproduction.</td>
<td>Direct measures of plant and invertebrate communities such as abundance, biomass and richness typically have high ecological relevance.</td>
<td>Direct measures of presence/absence and abundance typically have high ecological relevance.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>High</td>
<td>Tissue chemistry, when compared to tissue-based critical body residues.</td>
<td>High</td>
<td>Endpoints for mortality, growth, reproduction.</td>
<td>Direct measures of plant and invertebrate communities such as abundance, biomass and richness typically have high ecological relevance.</td>
<td>Direct measures of presence/absence and abundance typically have high ecological relevance.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

1. EC$_x$ = effect concentration, with percent X of organisms affected.
<table>
<thead>
<tr>
<th>Assessment endpoint</th>
<th>Line of evidence group</th>
<th>Magnitude</th>
<th>Spatial scale</th>
<th>Uncertainty about magnitude</th>
<th>Evidence for causal relationship between exposure and effects</th>
<th>Uncertainty about causality</th>
<th>Ecological relevance</th>
<th>Overall assessment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plants</td>
<td>Soil chemistry</td>
<td>Above benchmarks</td>
<td>1,000 m²</td>
<td>Moderate</td>
<td>No evidence of links between benchmarks and site-specific effects to plants, because benchmarks for site-specific COCs are based on invertebrate data only.</td>
<td>High</td>
<td>Low</td>
<td>Low effects, high uncertainty. Soil and chemistry benchmarks for site-specific COCs are not based on plants but rather are based on invertebrates. The community survey indicates there are low effects, but the cause may be fungal infection rather than site-related COCs, and uncertainty is high.</td>
</tr>
<tr>
<td></td>
<td>Community survey</td>
<td>Low</td>
<td>n/a</td>
<td>High</td>
<td>No evidence of relationships between biomass/richness and soil chemistry. Leaf spots and shoot blights that are evident on a few species and are believed to be related to fungal infection, not contaminants.</td>
<td>High</td>
<td>High</td>
<td></td>
</tr>
<tr>
<td>Soil invertebrates</td>
<td>Soil chemistry</td>
<td>Above benchmarks</td>
<td>1,000 m²</td>
<td>Moderate</td>
<td>Weak evidence (from literature) of links between benchmarks and effects to soil invertebrates, but application to specific sites limited by variation in toxicity modifying factors.</td>
<td>High</td>
<td>Low</td>
<td>Low effects, moderate to high uncertainty. Although tissue concentrations of COCs in earthworms are elevated and there is some site-specific toxicity observed, the toxicity results are not correlated with COCs. Furthermore, the most ecologically relevant line of evidence (invertebrate abundance and richness) indicates no effects.</td>
</tr>
<tr>
<td></td>
<td>Earthworm tissue bioaccumulation</td>
<td>Moderate</td>
<td>300 m²</td>
<td>High</td>
<td>Weak evidence (from literature) that observed contaminant concentrations could be causing toxicity.</td>
<td>Moderate</td>
<td>Moderate</td>
<td></td>
</tr>
<tr>
<td>Assessment endpoint</td>
<td>Line of evidence group</td>
<td>Magnitude</td>
<td>Spatial scale</td>
<td>Uncertainty about magnitude</td>
<td>Evidence for causal relationship between exposure and effects</td>
<td>Uncertainty about causality</td>
<td>Ecological relevance</td>
<td>Overall assessment</td>
</tr>
<tr>
<td>----------------------</td>
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<td>-----------------------------</td>
<td>-------------------------------------------------</td>
<td>-----------------------------</td>
<td>---------------------</td>
<td>-------------------</td>
</tr>
<tr>
<td>Soil invertebrates</td>
<td>Diverse and abundant invertebrate community, and ecological function as food for wildlife</td>
<td>Earthworm (<em>Eisenia fetida</em>) survival in laboratory toxicity test</td>
<td>Low</td>
<td>30 m²</td>
<td>Moderate</td>
<td>No evidence of a concentration-response relationship. One sample yielded significant toxicity, but not at high contaminant concentration.</td>
<td>Moderate</td>
<td>Moderate</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Abundance and richness in quadrat sampling</td>
<td>Negligible</td>
<td>n/a</td>
<td>High</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Birds</td>
<td>Healthy and reproducing local population</td>
<td>Community survey</td>
<td>Negligible</td>
<td>n/a</td>
<td>High</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Food chain model (dose to reproducing females in breeding season)</td>
<td>Low</td>
<td>1,000 m²</td>
<td>Moderate</td>
<td>Literature-based dose-response relationship well established but highly variable among species</td>
<td>Moderate</td>
<td>Moderate</td>
</tr>
<tr>
<td>Mammals</td>
<td>Healthy and reproducing local population</td>
<td>Food chain model</td>
<td>Negligible</td>
<td>1,000 m²</td>
<td>Moderate</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

1 This table may be based on a more detailed summary table that provides raw results rather than summary results.
2 Decision rules for defining scores (e.g., negligible, low, moderate, high) need to be defined in advance for each of the attributes, during problem formulation. See Table 5-1 for a default example.
3 n/a: no need to evaluate causality where no effect exists.
Table 5-5: Example summary table of WOE by assessment endpoint for aquatic ecosystems¹,²

<table>
<thead>
<tr>
<th>Assessment endpoint</th>
<th>Line of evidence group</th>
<th>Magnitude</th>
<th>Spatial scale</th>
<th>Uncertainty about magnitude</th>
<th>Evidence for causal relationship between exposure and effects³</th>
<th>Uncertainty about causality</th>
<th>Ecological relevance</th>
<th>Overall assessment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Macrophytes</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ecological function as food for fish and wildlife</td>
<td>Sediment and surface water chemistry</td>
<td>Above benchmarks</td>
<td>100 m²</td>
<td>Moderate</td>
<td>No evidence of links between benchmarks and site-specific effects to macrophytes</td>
<td>High</td>
<td>Low</td>
<td>Negligible effects, high uncertainty. Sediment and surface water chemistry benchmarks are not based on macrophytes. The community survey indicates there are no effects, but uncertainty is high.</td>
</tr>
<tr>
<td></td>
<td>Community survey</td>
<td>Negligible</td>
<td>n/a</td>
<td>High</td>
<td>n/a</td>
<td>n/a</td>
<td>High</td>
<td></td>
</tr>
<tr>
<td>Benthos</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aquatic invertebrate community structure, and ecological function as food for fish and wildlife</td>
<td>Sediment and surface water chemistry</td>
<td>Above benchmarks</td>
<td>100 m²</td>
<td>Moderate</td>
<td>Weak evidence of links between benchmarks and site-specific effects to benthos</td>
<td>High</td>
<td>Low</td>
<td>Moderate effects, moderate uncertainty. Three of the four effects-based measures show moderate effects, with varying evidence for causal relationships to site contaminants.</td>
</tr>
<tr>
<td></td>
<td>Amphipod toxicity test: survival</td>
<td>Moderate</td>
<td>100 m²</td>
<td>High</td>
<td>Weak evidence that growth related to COCs (based on regressions)</td>
<td>Low</td>
<td>Moderate</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Amphipod toxicity test: growth</td>
<td>Low</td>
<td>30 m²</td>
<td>High</td>
<td>Weak evidence that richness related to COCs (based on regressions)</td>
<td>Moderate</td>
<td>High</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Abundance as total organisms</td>
<td>Moderate</td>
<td>30 m²</td>
<td>Moderate</td>
<td>No evidence of relationship between abundance and contamination</td>
<td>Moderate</td>
<td>High</td>
<td>Moderate effects, moderate uncertainty. Three of the four effects-based measures show moderate effects, with varying evidence for causal relationships to site contaminants.</td>
</tr>
<tr>
<td></td>
<td>Richness as total taxa</td>
<td>Moderate</td>
<td>30 m²</td>
<td>Moderate</td>
<td>Weak evidence that richness related to COCs (based on regressions)</td>
<td>Moderate</td>
<td>High</td>
<td></td>
</tr>
<tr>
<td>Assessment endpoint</td>
<td>Line of evidence group</td>
<td>Magnitude</td>
<td>Uncertainty about magnitude</td>
<td>Evidence for causal relationship between exposure and effects</td>
<td>Uncertainty about causality</td>
<td>Ecological relevance</td>
<td>Overall assessment</td>
<td></td>
</tr>
<tr>
<td>---------------------</td>
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<td>-------------------------------------------------------------</td>
<td>-----------------------------</td>
<td>---------------------</td>
<td>-------------------</td>
<td></td>
</tr>
<tr>
<td>Fish Abundance and viability of local fish populations</td>
<td>Surface water quality</td>
<td>Above benchmarks</td>
<td>n/a</td>
<td>Moderate</td>
<td>Weak evidence of links between benchmarks and actual effects to fish</td>
<td>High</td>
<td>Moderate</td>
<td>Negligible to low effects, moderate uncertainty. Data do not indicate effects on fish directly, but there is high uncertainty. Some effects on food sources may occur, but the spatial scale is limited and population-level impacts are unlikely.</td>
</tr>
<tr>
<td></td>
<td>Relative abundance</td>
<td>Negligible</td>
<td>n/a</td>
<td>High</td>
<td>n/a</td>
<td>n/a</td>
<td>High</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Abundance and diversity of benthos as food</td>
<td>Moderate</td>
<td>30 m²</td>
<td>Moderate</td>
<td>No evidence that abundance of benthos is affected, but weak evidence for richness</td>
<td>n/a</td>
<td>High</td>
<td></td>
</tr>
<tr>
<td>Wildlife Abundance and viability of local bird, mammal and amphibian populations</td>
<td>Food chain model</td>
<td>Negligible</td>
<td>n/a</td>
<td>Moderate</td>
<td>n/a</td>
<td>n/a</td>
<td>Moderate</td>
<td></td>
</tr>
</tbody>
</table>

1 This table may be based on a more detailed summary table that provides raw results rather than summary results.
2 Decision rules for defining scores (e.g., negligible, low, moderate, high) need to be defined in advance for each of the attributes, during problem formulation. See Table 5-1 for a default example.
3 n/a: no need to evaluate causality where no effect exists.
For many ERAs, the diversity in measurement tools may yield divergent results, and trade-offs among contradictory lines of evidence will need to be made to derive an overall evaluation of risks for each assessment endpoint. In making judgments in this regard, practitioners should consider the following:

- Lines of evidence that are highly ecologically relevant should be given more emphasis when making trade-offs among lines of evidence, provided that uncertainties are comparable.
- If there is negligible magnitude of response and low uncertainty, there is no need to consider causality. However, if magnitude is high and/or uncertainty is great, causality becomes more important.
- If there is no evidence for causality and low uncertainty in the causality assessment, then observed responses are not related to site contaminants.

Redundancy is an important consideration when evaluating multiple lines of evidence. If any of the four major categories of evidence (see Section 2.3.4.1) is missing, it is effectively assigned zero weight, whereas multiple measures within a major line of evidence can result in double-counting (or more) of redundant or strongly correlated information. For example, there may be two different measures of soil invertebrate diversity, but no information at all on soil toxicity. To address this problem, the overall WOE evaluation for an assessment endpoint should take redundancy into account by acknowledging overlap in metrics and explaining in the narrative rationale how the redundant lines of evidence were considered. For highly complex sites with many lines of evidence, more formal methods may be appropriate, such as combining redundant lines of evidence first, before integrating across all lines relevant to an assessment endpoint.

Best professional judgement (BPJ) plays a significant role in the default WOE procedure during the integrated evaluation of each assessment endpoint. Particularly for cases where individual lines of evidence provided contradictory results, the narrative summary must provide rationale, using professional judgment, as to how the WOE conclusions were derived. This is the primary role of BPJ. In contrast, the use of BPJ is more limited in the analysis of individual lines of evidence, because lines of evidence are evaluated based on criteria that are defined in advance during problem formulation.

The role of professional judgment is not limited to the specific default WOE procedure recommended in this guidance. Even when more formal quantitative methods are used to combine results of multiple lines of evidence, professional judgment is used to define how trade-offs are made among contradictory lines of evidence. Nevertheless, although BPJ is a necessary and important part of WOE (Chapman et al. 2002), there are pitfalls of reliance on BPJ, including:

- challenges with demonstrating that determinations are reasonable
- lack of consistency in risk conclusions reached by different practitioners when faced with similar input (i.e., repeatability issue)
- potential for abuse by practitioners seeking to find a predetermined outcome
• unintended bias resulting from perception of results according to an established paradigm, rather than objective evaluation of all possible explanations.

Despite these challenges, much of the problem can be resolved through proper articulation of “good practice” in application of BPJ. Wandall (2004) argues that proper application of professional judgment in risk assessment requires that risk assessors are aware of what underlying values they are relying on, the values are justifiable and transparency is ensured. This requirement for transparency is the foundation of properly applied professional judgment and translates into the following guiding principles: 47

• All assumptions and decisions must be supported with a rationale, especially for those instances where education and training were used as the basis for the professional judgment.

• Declarative and unqualified conclusions such as “The risk assessment proved that there are no adverse effects” should be avoided. Instead, conclusions should reflect where professional judgment was applied in the evaluation, for example: “The risk assessment, based on our professional judgment of ABC data, and subject to assumptions XYZ, found no evidence of adverse effects.”

5.6 Step 5: Evaluate ERA Uncertainties

There are numerous sources of uncertainty and variability in ERA. These uncertainties fall in multiple categories (see Box 2.1). Uncertainties must be evaluated in order to determine the level of confidence associated with risk estimates and to determine to what extent additional work is warranted to reduce uncertainties.

Importantly, the level of detail and rigour needed to address uncertainty will vary depending on the complexity of the ERA and the results. If estimated risks are either extremely low or extremely high, it may be easy to demonstrate that uncertainty is unlikely to change that conclusion. On the other hand, more rigorous evaluation of uncertainty is usually warranted when estimated risks are in the range that may or may not be acceptable.

Many aspects of uncertainty can be integrated directly into WOE summary tables, as shown in Table 5-4 and Table 5-5 and discussed in Section 5.5.2. However, uncertainty evaluation extends beyond the assessment of uncertainties for individual attributes and endpoints. Therefore, this section is identified as a separate step from WOE (even though uncertainties are evaluated during the WOE procedure).

Addressing uncertainties requires that the practitioner:

47 Further discussion of BPJ in ERA and WOE evaluation can be found in Bay et al. (2007), Lee and Jones-Lee (2002) and WDNR (2009).
• Identify uncertainties in the risk assessment, and distinguish them from elements of the risk assessment where there is reasonable certainty.

• Evaluate the implications of uncertainties. For instance, could risk conclusions change if uncertainties were reduced, and how likely is it that risk management decisions may change?

• If warranted, explicitly integrate uncertainties into risk characterization methods (e.g., using probabilistic methods).

• If warranted, determine the potential value of reducing uncertainty through follow-up investigations. For instance, to what extent would additional work increase accuracy and precision of risk estimates and lead to a more informed risk management decision?

5.6.1 Identifying Uncertainties
The first step in addressing uncertainties in ERA is to differentiate factors and conclusions that are known with reasonable certainty from those that are uncertain. Specific uncertainties may apply to any data, parameters, models or assumptions used in the risk assessment. The various sources of data and information related to characterizing exposure and effects (see Section 3 and Section 4) may all be subject to uncertainty to varying degrees. For uncertainties that can be quantified using data, basic plots (e.g., box plots) and descriptive statistics can be used to characterize the uncertainty in the data (e.g., minimum, maximum, median, mean, variance).

5.6.2 Evaluating the Implications of Uncertainties
Uncertainties are important because of their potential implications for risk estimates and ultimately for risk management decisions. The implications of specific uncertainties are most easily evaluated using sensitivity analysis to test how risk estimates change according to various “what-if” scenarios for each quantity. Sensitivity may be tested using the minimum and maximum possible values for a given quantity, or any other metrics (e.g., the 5th and 95th percentiles). For example, a HQ could be estimated using the minimum and maximum measured COC concentration in food items as a bounding analysis. If the HQ does not differ appreciably between the two scenarios (e.g., if it was well below 1 in both cases), the risk assessor may conclude that uncertainty related to the tissue concentration is negligible. In contrast, if the HQ changes from less than 1 to greater than 1, the uncertainty in the tissue concentration may need to be explored further.

Each of parameters used to estimate risks can be varied, independently or at the same time, to generate a range of “what-if” scenarios. Sensitivity analyses are useful for understanding which uncertainties have the most potential influence on risk estimates. The cumulative effect of multiple uncertainties can be understood to some extent using these methods. However, simultaneous consideration of the cumulative effect of multiple uncertainties is better addressed using probabilistic methods, as outlined below.
5.6.3 Integrating Uncertainties into Risk Characterization

The cumulative influence of uncertainties is best understood using probabilistic methods. As discussed earlier in Section 5.3.6, probabilistic methods are useful for characterizing risks because they provide accuracy and realism that is not captured when data and parameters are represented with point estimates. In the context of evaluating uncertainty, the key benefit of a probabilistic assessment is facilitating understanding of the cumulative effects of multiple uncertainties on risk estimates.

What is a probabilistic assessment? Probabilistic methods are distinguished from deterministic methods in that exposure is characterized not as a point estimate but as a probability distribution (or frequency distribution) of possible estimates, based on the use of distributions to characterize some or all of the uncertain input quantities. For example, all of the equations in a food chain model could be based on distributions rather than point estimates for each input parameter.

When should probabilistic methods be used? Risk assessors should consider developing probabilistic models whenever more accurate estimates of risk could be important from a risk management perspective, or to simply evaluate the cumulative effects of multiple uncertainties. Consistent with the iterative approach to ERA (Section 1.6), if a deterministic (point estimate) risk assessment based on conservative assumptions shows that risks are acceptable, then the increased accuracy provided by a probabilistic model is not warranted. However, if risks are identified using deterministic methods, probabilistic methods should be considered.

How to implement probabilistic methods: Uncertainties are usually modelled using numerical simulation techniques such as the Monte Carlo simulation.48 A simulation model may be run a few thousand times; each realization or trial involves random selection of a value for each uncertain quantity (according to a probability or frequency distribution). In the case of a wildlife food chain model, the output from each simulation trial might be, for example, an HQ or an estimate of expected mortality.

Although commercial software packages have made implementing simulation techniques much simpler, some of the design elements require careful consideration. First, the risk assessor must decide whether the simulation model will deal with variability among individuals in a population, or only with incertitude49 (e.g., uncertainty regarding an average individual), or with both (Hoffman and Hammonds 1994). Simultaneous consideration of individual variability and incertitude may warrant a two-dimensional simulation. Conversely, a simpler model may suffice

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48 Analytical methods and Taylor series approximation methods of propagating uncertainties are reviewed in Cullen and Frey (1999).

49 Incertitude is uncertainty caused by incomplete descriptions of a mechanism or process and other limitations of scientific knowledge. The term is used here to distinguish this aspect of uncertainty from natural variation and other types of uncertainty. In statistical terms, for a parameter such as body weight, variability among individuals might be characterized with a standard deviation, whereas incertitude about the mean body weight might be characterized with a standard error.
in many cases for ERA, provided that results are interpreted correctly. Second, any correlations among uncertain variables should be accounted for in simulation models, otherwise the estimated probability distribution of risks will be too wide and may be skewed. In reality, many ecological parameters are highly inter-correlated (e.g., feeding rate and growth rate of a species, or feeding rates of several species that are all a function of temperature). There are ways to account for these correlations in simulations (Haas 1999), but they require additional information about the form of the correlation. Even where inter-correlation structures are available, there is still uncertainty in the structure of the model itself, and it is difficult to determine the quantitative effect of the inability of our models to exactly represent natural processes.

Risks represented as a probability or frequency distribution are informative, but the risk assessor must communicate the information in a way that risk managers and stakeholders can easily interpret. For example, it may be useful at the risk characterization stage to report particular statistics, such as the probability that an average individual would exceed a particular effects threshold. Further guidance on probabilistic exposure methods is provided by Cullen and Frey (1999), Suter et al. (2000) and US EPA (1997-a, 1997-b, 2001).

**Data requirements for probabilistic models:** Although any model is best when data are plentiful, risk assessors should not shy away from probabilistic analyses in cases where data are sparse. In general, if a probabilistic analysis is appropriate for an ERA, the advantages of implementation will outweigh the disadvantages created by data limitations, provided that limitations are explicitly described. Methods exist for using limited data to construct probability distributions (Cullen and Frey 1999; Morgan and Henrion 1990), and simple distributions (e.g., uniform, discrete, triangular) can be used in data-poor cases. In addition, sensitivity analyses usually reveal that many uncertain quantities have little impact on the cumulative uncertainty, such that precise characterization of uncertainties is not always critical for all quantities.

### 5.6.4 Determining the Value of Reducing Uncertainty: When to Refine Risk Estimates

If preliminary risk estimates indicate the potential for adverse effects, the underlying conservative assumptions and uncertainties should be critically evaluated using approaches outlined above (e.g., sensitivity analysis). The practitioner (and client) must either decide to further refine the exposure or hazard assessments to reflect site-specific conditions, or conclude that risk is unacceptable or unresolvable and that remediation or other risk management options should be considered. A matrix based on varying levels of estimated risk and uncertainty (based on Pearsons and Hopley 1999) can be a useful way to conceptualize interpretation of uncertainties:

<table>
<thead>
<tr>
<th></th>
<th>Low magnitude of risk</th>
<th>High magnitude of risk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low uncertainty in risk estimate</td>
<td>Low precaution</td>
<td>Moderate precaution</td>
</tr>
<tr>
<td>High uncertainty in risk estimate</td>
<td>Moderate precaution</td>
<td>High precaution</td>
</tr>
</tbody>
</table>
Refining risk estimates for the “high” category of precaution is recommended. The “moderate” category of precaution may also indicate a need to reduce uncertainty as necessary to support management actions. This refinement may involve one or more of the following strategies:

- Reduce parameter uncertainty by gathering additional data. Supplemental data collection should be targeted to deal with the underlying cause of the parameter uncertainty (e.g., address spatial coverage, improve analytical detection limits, collect bioavailability information, evaluate cause and effect mechanisms).

- Reduce structural (model) uncertainty by adopting a more appropriate model and any additional data needed to support that model.\(^5\) Risk assessment should be an iterative process where new data may require reassessment of previous approaches or conclusions. This iterative process allows risk assessment to be a dynamic process well suited to ecological study, and does not indicate a failure of the initial screening risk estimate.

- Provide risk managers with multiple risk scenarios for consideration as a series of risk estimates with different assumptions and descriptions of uncertainty.

Several other strategies are often employed; however, they do not directly reduce parameter or model uncertainty. For example:

- Professional judgment is often used to fill in gaps in model structure. This may reduce uncertainty, but it may not, and there is no objective way to know. Conservative assumptions are often used as part of this strategy; although they do not reduce uncertainty, they ensure that the majority of the uncertainty errs on the side of caution. The challenge in using conservative assumptions lies in balancing conservatism and ecological realism relative to site-management needs.

- Increase the number and types of lines of evidence considered in a WOE approach. This strategy does not reduce the uncertainty in any single line of evidence, but does reduce overall uncertainty in the conclusions of the risk assessment because the limitations of one line of evidence are frequently balanced by the strengths of another.

### 5.7 Step 6: Consider Extrapolation and Interpolation

This aspect of uncertainty in risk assessments warrants its own step in the risk characterization process because it addresses transferability of ERA findings over time, space or alternate site-use scenarios. This is particularly relevant to site managers because management decisions may require confidence that risk narratives remain applicable even if some of the underlying assumptions change. For example, where a site is divested or otherwise changes in ownership or land use, repeating the entire ERA process is not desirable.

\(^5\) Additional model complexity may not reduce uncertainty, and often increases uncertainty. The benefits of additional model complexity should be evaluated on a case-by-case basis.
By design, risk assessments focus resources on a narrow subset of potential receptors, spatial locations and measurement endpoints. In concentrating on a narrow and focussed set of risk hypotheses, there is a danger of “not seeing the forest for the trees.” Therefore, near the end of the risk assessment, it is prudent to conduct a reality check to assess how representative the risk assessment is expected to be in terms of the broad site-management goals.

Conceptually, the extrapolation and interpolation assessment entails broadening the scope of the risk conclusions from the detailed findings (e.g., specific risk estimates for representative organisms and exposure scenarios) to the broadly defined assessment endpoints. Due to practical constraints, ERAs are limited in the spatial and temporal domains they consider, and in the degree to which they explicitly evaluate combinations of chemical, physical and biological components. The extrapolation and interpolation assessment serves as a reality check for the relevance of the study results to the VECs, and provides context for the overall findings.

Some specific issues to be addressed at this stage include:

- **Can results for one receptor be extended to other species at the site?** For example, if a mallard duck was selected as an receptor of concern to represent a VEC, can we assert that risks to other dabbling ducks, other waterfowl or other omnivorous birds in general are expected to be lower than those for the mallard? In some cases, the receptor of concern is selected based on its presumed sensitivity to relevant COCs and pathways of exposure. However, in other cases, other considerations may dictate receptor of concern selection (e.g., data availability, standardized methods for assessment). In these cases, the risk assessor should qualitatively evaluate the degree of protectiveness afforded other species not rigorously evaluated in the risk assessment.

- **Are thresholds for individual COCs protective of the entire contaminant mixture?** Where a site-specific standard has been developed for an individual substance of concern, and that substance serves as a surrogate for other COCs, there is an implicit assumption that the other COCs will not increase relative to the individual (indicator or surrogate) substance.

- **Are the study conclusions dependent on an assumption of fixed site use, or would the results also apply to site redevelopment or restoration?**

- **Can conclusions or quantitative relationships based on limited sample sizes be extended to other spatial units, habitats, depths or physical conditions?** The underlying assumption is that exposure-effect relationships observed at sampled areas will remain applicable when extended to other unsampled portions of the site. However, if the unsampled areas are substantially different in terms of factors that may influence COC bioavailability, or represent habitat conditions not evaluated in the risk assessment, there is uncertainty in extrapolating study findings. The specific issue of deriving site-specific standards or benchmarks, which implicitly assumes transferability of quantitative relationships, is considered further in Step 7.

The risk assessor should specify constraints or caveats to the extension of study findings across space, time, habitat type or biological assemblage. Note that the requirement for extrapolating to
new conditions (or predicting future responses) is closely linked to the assessment objectives identified during the problem formulation.

5.8 **Step 7: Develop Site-Specific Remediation Standards (Optional)**

Where significant ecological effects are observed over some or all of a contaminated site, it may be appropriate to develop site-specific remediation standards. These values are often also referred to as site-specific target levels and represent concentrations in environmental media that, once achieved, will meet the environmental protection goals for the site. This step is listed as optional because formally developing site-specific remediation standards may not be required, depending on the type of assessment, jurisdictional requirements and the risk management needs. For example, if risk characterization is conducted using a parcel-based or spatially explicit evaluation of risks (e.g., grid cells evaluated individually for acceptability of risks), then development of numerical target levels for specific substances might not be required.

CCME (1996-a) provides a framework for developing site-specific environmental remediation objectives. Under the framework, where a risk-based approach is applied, risk assessment procedures can be used to establish remediation objectives on a site-specific basis, as discussed in the following subsections.

5.8.1 **Choosing Appropriate Site Media**

Most risk assessments evaluate more than one site medium (e.g., soil, sediment, tissue, surface water, groundwater, porewater). In many cases, one exposure medium can be identified as the “driver” (i.e., dominate the magnitude of risk estimates) by strongly influencing the environmental exposures. The practitioner should ensure that the choice of a medium for development of a site-specific standard sufficiently addresses the risk pathways of relevance and does not leave other important pathways unaddressed. For example, if a wildlife risk assessment determined that metals uptake through soil-based pathways and drinking water were both important, it would be necessary to either develop standards for both pathways or to develop standards for one pathway with explicit acknowledgement that the other pathway remains.

Another consideration is the degree to which the site media can effectively be used to develop a remediation or monitoring plan. Soil and sediment are commonly applied media because they represent sinks for contaminants, are relatively immobile and can easily be sampled. In contrast, tissues or organisms are not commonly used, because the organisms may be mobile, availability of tissues may be seasonal and monitoring of post-remediation results may not be practical.

5.8.2 **Identifying Appropriate Contaminants**

The contaminants that are the dominant sources of risk must be identified. This may be a simple decision, or quite complex, depending on the nature of the contaminant mixtures and relative risks estimated for each COC. Some important considerations are:
• **Cumulative risks**: For related substances, a surrogate compound or integrated value may be useful. For example, total PAH may be used as an exposure measure if the composition of component PAHs across the site is stable.

• **Practical considerations**: The parameter adopted should be relatively easy to measure. For example, certain parameters may be difficult to measure or quantify based on lab detection limitations or limitations in separation from a complex mixture, and may warrant adoption of other, simpler measurements.

• **Degree of causation**: The identified contaminant should have a strong correspondence to environmental response, and ideally have strong evidence of causation. Where multiple COCs are present and causation has not been determined, developing a site-specific standard requires an assumption that the indicator COC is an effective surrogate for the effects of the entire mixture.

### 5.8.3 Contamination Pathways

In developing a site-specific standard, it is important to consider the pathways by which risk occurs and the assumptions required for a standards-based remediation to be effective. For example:

- Will the site be recontaminated by either on-site or off-site contributions?

- Are residual concentrations likely to attenuate over time or increase through chemical reaction?

### 5.8.4 Spatial Scale

Applying a site-specific standard requires considering the spatial domain relevant to the receptors. For mobile receptors, weighted averaging of exposures can be incorporated in the development of standards. For sessile receptors, the spatial scale at which monitoring of risks will be conducted needs to be addressed (e.g., depth of soil or sediment, resolution of lateral COPC characterization).

The scale of relevance will strongly influence the methods used to apply the standard. For a sessile receptor, the standard may be a “not to exceed” threshold, whereas an averaging procedure could be applied to migratory organisms. For wildlife, an area-based average is often applied, depending on the home range of the receptor relative to the size of the site.

### 5.8.5 Modifying Factors

Where site conditions are variable, it may be appropriate to adjust site-specific standards on a location-by-location basis. This would account for bioavailability or toxicity differences that could be relevant across small spatial scales. For example, values of soil or sediment organic carbon content may be variable, and adjustment to account for bioavailability differences may be appropriate if the site-specific standard was developed on a dry-weight basis. Alternatively, if the risk assessment data were amenable, the standard could be developed on an organic carbon-normalized basis. Other modifying factors include pH and salinity in aqueous samples.
5.8.6 Approval and Application
Regulatory review is typically required for any site-specific standard and may entail consideration of:

- management checks for consistency with law or policy considerations
- public consultation
- socio-economic factors
- technical constraints.

Furthermore, removal or remediation actions defined using site-specific standards typically require a clear linkage to a risk management plan, including long-term monitoring. For this reason, site-specific remedial standards are often developed in parallel with the risk management process, as described in Step 9 below.

5.8.7 Methods for Developing Site-Specific Remediation Standards
Site-specific standards rely on underlying concentration-response relationships. In some situations, it is possible to directly adopt a TRV developed in the hazard assessment stage. However, additional data synthesis or modelling is often required to develop a site-specific standard, particularly once the considerations discussed in Section 5.8.1 are taken into account. For example:

- Converting tissue-based TRVs to soil or sediment media may require bioaccumulation models or equations for back-calculation purposes.
- Concentration-response relationships from multiple lines of evidence may need to be synthesized (simplified) to yield a single threshold for management purposes.

Developing site-specific standards may be complicated by the numerous COPCs and the range of responses observed, but typically involves the following steps:

1. Identify a level of harm considered acceptable based on the risk characterization findings. This could be quantitative (e.g., soil concentration associated with a HQ of 1.0 for a wildlife species) or qualitative (e.g., low risk as determined from a sediment-quality WOE assessment).
2. Plot the degree of harm (response) versus COPC concentration, either graphically or using a mathematical relationship (such as regression analysis).
3. Resolve the uncertainty associated with an impact relationship between response and the exposure measure. For example, determine whether it is acceptable to have a “smoothed” target concentration considered protective of a receptor even if an individual station exhibited a significant ecological response.
4. Convert the target concentration to the desired units, scale and media of interest (as outlined in Section 5.8.1). The target concentration must be clearly defined in terms of spatial application (e.g., spatially weighted threshold, or a maximum not to be exceeded at any
location), the parameter details (e.g., dry weight sediment versus organic carbon normalized, fillet tissue versus whole body), and the conditions or assumptions required for applicability of the target concentration.

5.9 Step 8: Summarize Risk Conclusions
Following the technical application of a risk assessment, it is important to summarize results in a manner that is clear, accurate, concise and meaningful to the risk manager. A risk narrative is often provided for this purpose. This risk narrative may be combined with the summary of estimated risks for each assessment endpoint that concluded at the end of the WOE procedure (Step 4, Section 5.5), or it may be presented separately. Risk assessors must provide an opinion regarding their results generated with respect to confidence, uncertainty and significance of impacts. As described by Exponent (2010):

A full narrative is analogous to writing the results and discussion sections of scientific papers and is intended to help other reviewers or risk managers understand how the risk assessor reached their conclusions based on the evidence in hand. The narrative can be used to help reach agreements, identify disagreements, and identify aspects of the risk assessment that require additional clarity. (p. 22)

Although the WOE procedure has already articulated findings for each assessment endpoint based on magnitude of effects (including spatial and temporal scales), evidence for causality, ecological relevance and uncertainty for individual lines of evidence, the risk narrative should integrate that information into a form that is useful for decision-makers. Specific goals of the risk narrative may be to:

- present in lay language the key rationales used to draw overall conclusions for each assessment endpoint during the WOE procedure
- summarize overall confidence in the specific findings, in light of the ecological relevance of the various lines of evidence and the strength of evidence implicating site-related contaminants as the cause of any observed effects
- summarize confidence that overall the risk assessment methods are relevant and that the findings can be extrapolated to the general conditions at the site (both now and under foreseeable future conditions)
- summarize the extent to which key uncertainties may affect risk conclusions, and whether further work to refine those uncertainties may be warranted
- clarify the spatial and temporal scales at which effects are observed, or provide separate summaries of risk conclusions for different spatial or temporal units
- summarize the potential for cumulative impacts of site-related contaminants and other stressors.
5.10 Step 9: Conduct Follow-up Actions

The final step in risk characterization is to link the study findings to the risk management process. Risk communication is an important aspect of the overall risk management process, and therefore it is helpful to frame the path forward at the conclusion of the risk assessment process. This may entail a summary of recommendations and a clear articulation of next steps for site closure, approvals, regulatory liaison, and so on. Details may not be included in the ERA if risk management considerations are addressed as a separate deliverable.

Depending on the outcome, and provided that the scope of the ERA includes recommendations for next steps, recommendations for site management may include:

- **No further action required**: The rationale for the decision should be succinctly summarized.

- **Additional investigation or risk assessment required**: If the residual uncertainty in the risk assessment is large, a decision could be made to refine the assumptions and reduce uncertainties. Where iteration is contemplated, the advantages and limitations of follow-up studies should be assessed.

- **Risk management strategies required**: No physical actions are deemed necessary, but management activities may still be required (e.g., administrative controls, monitoring program).

- **Remediation required**: Considerations for conceptual remedial design may be articulated, often to site-specific standards that were developed during the risk assessment.

- **Provisions to protect existing VECs during the remedial phase required**: This may occur where remediation is recommended.

The evaluation of potential follow-up actions should reconsider the overall assessment goals in light of the conclusions of the ERA. In some cases, as part of an adaptive management approach, the focus for management may shift to one of the other quadrants of the overall assessment framework (Section 2.2.1.1). If monitoring has begun, the results of monitoring must be assessed to determine what to do next. If a past management action has not resulted in expected environmental improvements, then assessing causation may become more important. Alternatively, if environmental improvements have been substantial, the requirements for long-term monitoring may be re-evaluated. Step 9 provides an opportunity for risk managers to conduct a check of the site-management recommendations against the broad site-management goals, adjust the course of the investigation as appropriate, and update the conceptual model of the site to reflect recent information.
REFERENCES

Note to Reader

Links to websites and online references are accurate at the time of publication.

For an overview of additional, general references regarding ERA information refer to Section 1.9.


EC 2010-b. FCSAP Ecological Risk Assessment Guidance - Module 2 : Selection or Development of Site-Specific Toxicity Reference Values. EC, Ottawa.


