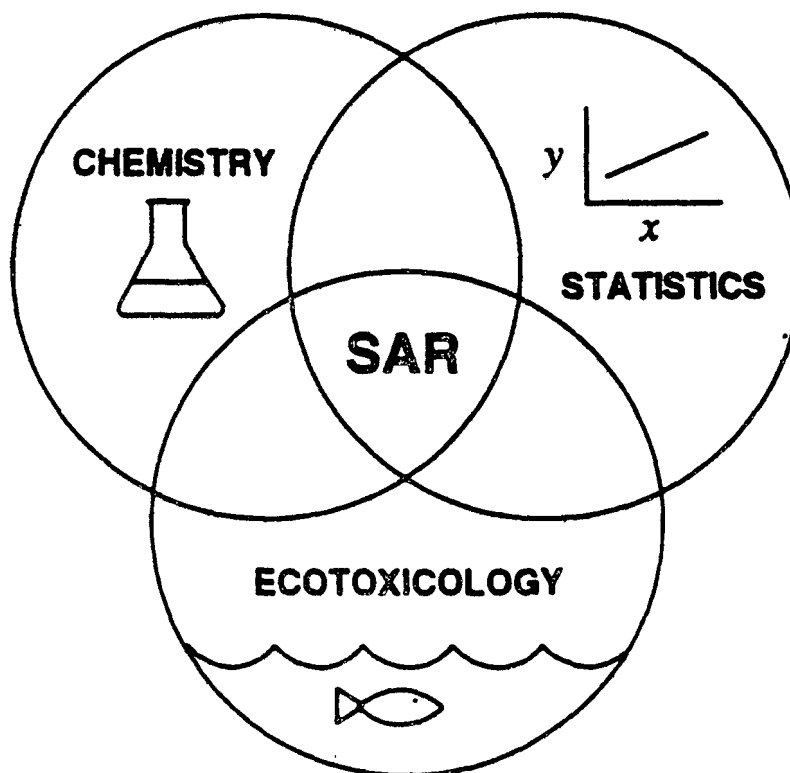




# Estimating Toxicity of Industrial Chemicals to Aquatic Organisms Using Structure Activity Relationships

## 2nd Edition

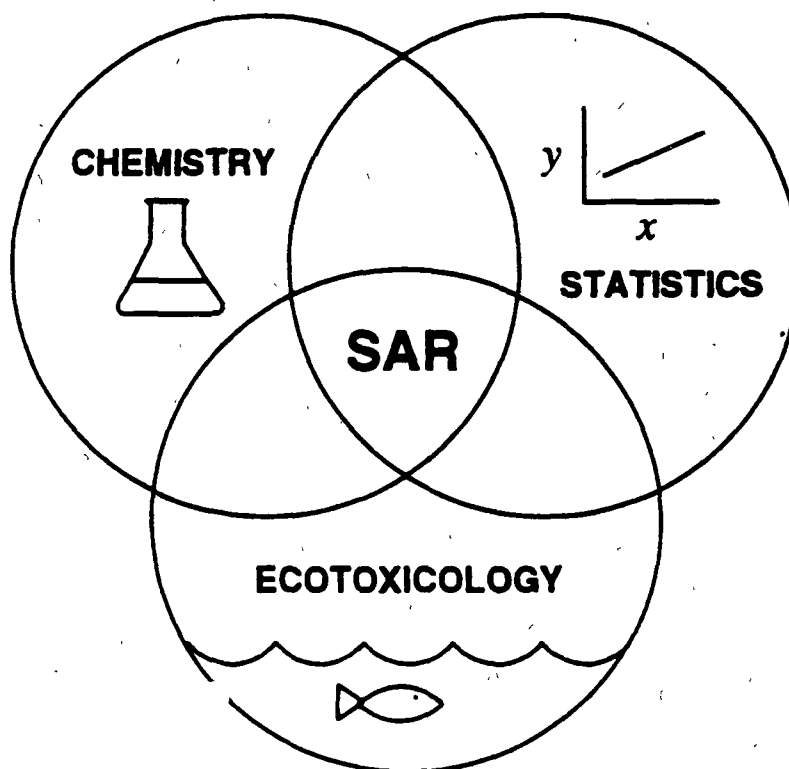


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# Estimating Toxicity of Industrial Chemicals to Aquatic Organisms Using Structure Activity Relationships

## 2nd Edition



**ESTIMATING TOXICITY OF INDUSTRIAL CHEMICALS  
TO AQUATIC ORGANISMS USING  
STRUCTURE-ACTIVITY RELATIONSHIPS**

**Second Edition**

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**August, 1996**

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## FORWARD to the SECOND EDITION

As discussed in the FORWARD to the first edition of **Estimating Toxicity of Industrial Chemicals to Aquatic Organisms Using Structure-Activity Relationships**, the development of predictive ecotoxicology models for industrial chemicals creates challenges that are unique compared to those faced in drug or agrichemical design. Under the requirements of the Toxic Substances Control Act there is, however, no choice but to face these challenges and provide the means to assess the ecological risks of new and existing compounds.

Since releasing their first edition in 1988, the scientists within the Environmental Effects Branch of the Office of Pollution Prevention and Toxics have continued to develop property-activity correlations that are relevant for industrial chemicals found in commerce. In publish the second edition of **Estimating Toxicity of Industrial Chemicals to Aquatic Organisms Using Structure-Activity Relationships**, the contributors have once again demonstrated their commitment to share the results of these efforts with the scientific and regulatory community. This second edition contains over 70 additional property-activity correlations and is a companion document to ECOSAR, which is a computerized version of the relationships developed by the contributors. Through this on-going contribution to the world-wide 'structure-activity relationship knowledge base', the scientist in the Environmental Effects Branch are also providing the means to identify first-order uncertainties in the development and use of these predictive models. The influence of the first and second edition on the application of structure-activity relationships and the course of future research in are of environmental toxicology can not be minimized. Of particular interest is the continuing to develop models for chronic effects and to establish objective techniques whereby compounds can be assigned to specific relationships.

Again, I congratulate the contributors to this document for their dedication in implementing structure-activity relationships in ecological risk assessments and for fostering the exchange of information that is essential for the advancement of the field.

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Duluth, Minnesota  
April, 1995

## INTRODUCTION

For many years, the manufacturers of pharmaceuticals, pesticides, and dyes have used the relationship between chemical structure and a specific effect to search for new chemicals. These relationships are called structure-activity relationships (SARs). Under Section 5 of the Toxic Substances Control Act of 1976, EPA must review and evaluate all new chemicals before they enter commerce. The Environmental Effects Branch (EEB) of OPPT has been responsible for the assessment and evaluation of these new chemicals and for identifying those chemicals of greatest concern for environmental hazard. Since 1976, of all chemicals submitted to EPA under Section 5 of TSCA, fewer than 5% of the Premanufacture Notices have contained toxicity data pertaining to terrestrial and aquatic organisms. To meet its regulatory mandate, EEB began using SARs in 1979 to estimate the toxicity of chemicals in the absence of test data.

The application of SARs in the field of environmental toxicology is relatively new. Some of the early research work began in the 1960's. During the 1970's, many investigators began examining the relationships among chemical properties and the toxicity to aquatic and terrestrial organisms. Among the leaders in this area was the U.S. EPA Environmental Research Laboratory at Duluth (ERL-Duluth) who pioneered research in the development and application of SARs to environmental toxicology. In the mid-1970's they developed and later published the SAR for predicting the bioconcentration of neutral organic compounds in fish based upon the octanol/water partition coefficient. In 1979, they initiated a long-term research program to develop SARs for industrial organic chemicals. Between 1981 and 1983, EEB staff evaluated and adopted 13 of these SARs for use in predicting toxicity to fish, aquatic invertebrates, and green algae. To date, the scientists at ERL-Duluth have measured the toxicity of over 800 compounds. From this research, they have developed SARs for at least a dozen classes of compounds to both freshwater and marine fish. Recently, emphasis at ERL-Duluth has shifted toward SARs for chronic toxicity with numerous chronic values now being published.

The octanol/water partition coefficient ( $K_{ow}$ ) has been the major attribute used by most investigators to correlate structure and toxic effect. The most frequently used relationship is the logarithm of the  $K_{ow}$  versus the logarithm of the median toxicity ( $LC_{50}$  and  $EC_{50}$ ) value. To date the major focus has been centered around the class of industrial organic chemicals known as neutral organics. These compounds are non-ionizable, non-reactive and neutral with respect to charge, however, SARs have been developed for other classes of chemicals and new ones continue to be derived as data become available.

This manual is intended to accompany an SAR program, called ECOSAR, that has been developed by EEB for use on a personal computer. ECOSAR is menu-driven and contains on-line help, including a User's Guide. ECOSAR includes all of the chemical classes and SARs contained in this manual. Most toxicity values (in mg/L) are based on  $\log K_{ow}$  and molecular weight information supplied by the user, although as discussed below, some SARs require other physical data, such as number of ethoxylates or percent amine nitrogen. ECOSAR may be obtained from the sources listed at the end of the Introduction.

### Chemical Classes

This manual presents information for deriving toxicity values for four primary classes of chemicals:

- (1) Neutral organics that are nonreactive and nonionizable;
- (2) Organics that are reactive and ionizable and that exhibit excess toxicity in addition to narcosis;
- (3) Surface-active organic compounds such as surfactants and polycationic polymers, and
- (4) Inorganic compounds including organometallics.

## FORWARD

The world of the scientist in a regulatory agency which is responsible for chemical safety is quite different from that in physical organic chemistry, toxicology, or drug design. Nothing highlights the difference quite like the scientists who implement the EPA mandate to review industrial chemicals for health and environmental effects. More conventional research scientists work in data-rich areas where chemical models are comparatively precise. Only under the Toxic Substances Control Act (TSCA), implemented by the Office of Toxic Substances, can we find the responsibility to evaluate the broad spectrum of chemical safety with little or no data on either new or existing chemicals.

These scientists responded to the EPA responsibilities by adapting approaches used in drug design and chemistry to predict the environmental behavior and toxicology of chemicals from their structure rather than extensive test data. The Office of Research and Development has enjoyed a ten-year partnership with the Office of Toxic Substances in developing quantitative structure-activity relationships to estimate the bioaccumulation potential, the persistence, and the toxicity of chemicals in the environment. Much developmental work remains to be done in efforts to more thoroughly evaluate chronic effects of long term exposure; nonetheless, many relationships are adequate to assist regulatory scientists in making judgements concerning the risks of chemicals.

As we continue to improve our understanding of relationships between chemical structure and effects, the Environmental Effects Branch realized the value of this technology to other scientists in EPA Regions and states. Their initiative to summarize the state-of-the-art in this document and make it available to others is another example of the futuristic planning of this group. The predictive power of the methods included in this document varies with the available data and complexity of the toxicity mechanisms. However, the predictive power will continue to increase over the next decade as new chemical models are formed. I congratulate the contributors to this document for the increasing effort to formulate structure-activity relationships from scant data, and for their desire to share this work with others in the scientific and regulatory community.

Gilman D. Veith  
Director  
Environmental Research Laboratory  
U.S. Environmental Protection Agency  
Duluth, Minnesota  
Jun, 1988

Neutral organic compounds that are nonelectrolytic and nonreactive act as anesthetics or narcotics. This class of compounds includes alcohols, ketones, ethers, alkyl halides, aryl halides, aromatic hydrocarbons, aliphatic hydrocarbons, many cyanates, sulfides, and disulfides

Organic compounds with a more specific mode of toxicity may contain reactive functional groups such as electrophilic moieties. These compounds are more toxic than would be predicted by using an SAR for a narcotic compound. Chemicals which exhibit excess toxicity include acrylates, methacrylates, aldehydes, anilines, beta-diketones (linear forms), benzotriazoles, esters, phenols, aziridines, and epoxides. A separate SAR has been developed for each of these classes.

Surface-active chemicals may act on the respiratory membranes of aquatic organisms. These chemicals consist primarily of surfactants that can be absorbed through respiratory membranes and charged polymers that cannot be absorbed. SARs have been developed for anionic surfactants such as linear alkyl benzene sulfonates, nonionic surfactants such as alcohol ethoxylates and cationic surfactants, such as ethoxylated beta-amine surfactants (ethomeen) and linear N-alkyl quaternary ammonium compounds. The SARs for surfactants are parabolic, i.e., toxicity is related to the size of the hydrophobic component in a parabolic manner when the size of the hydrophilic component remains constant. The size of the hydrophobic component, usually a linear alkyl carbon chain, can be estimated by simply counting the number of carbons in the hydrophobic alkyl chain. Maximum toxicity occurs when there are approximately 16 or 17 carbons in the linear alkyl chain. Toxicity for the nonionic surfactants is also affected by the number of ethoxylate units and the size of the hydrophobe and the number of ethoxy groups must be known to use the SAR.

Polycationic polymers include those with primary, secondary, and tertiary amines and/or quaternary ammoniums, phosphoniums, and sulfoniums. The molecular descriptor used to predict toxicity for these polymers is equivalent charge density as determined from chemical structure, i.e., percent amine-nitrogen, number of cationic charges per 1000 units of molecular weight, or cation equivalent weight. These polymers must be water soluble or self-dispersing or both.

Quantitative SARs have not been developed for inorganic compounds. However, in lieu of such equations, water quality criteria values have been used to predict their toxicity. Water quality criteria have been developed for several metals. These criteria are usually indicative of the lowest concentration that is believed to be protective of aquatic life in the receiving water. Consequently, most criteria are expressed only for acute or chronic toxicity to freshwater or marine organisms in general. SAR equations will eventually be developed for organometallics based on their  $K_{ow}$  values

Some chemical classes do not have quantitative SARs. These include polyanionic polymers, cationic dyes, and most classes of pesticides. Two classes of polyanionic polymers are known to be toxic to aquatic organisms; polyaromatic sulfonic acids are moderately toxic to aquatic organisms; and polycarboxylic acids are moderately toxic only to green algae. However, the high molecular weight of these polymers indicate that they will not be absorbed through the surface membranes of these organisms and their toxicity is the result of their surface activity and is not correlated with their anionic charge density. Cationic dyes can be absorbed and are known to be highly toxic to aquatic organisms. During acute exposure, the toxicity of these dyes is believed to be mostly the result of their activity on the surface membrane while chronic exposure also results in systemic toxicity. Dyes with delocalized cationic charges may be more toxic, followed by dyes with four localized charges, then three localized charges, etc. Most commercial dyes contain impurities which may, in part, be responsible for some of the toxic effects seen in these dyes. Acid dyes are moderately toxic only to green algae which results more from shading of the algae by the dye rather than from direct toxic effects. Data on which to validate this assumption are lacking in most PMN submissions.

## How SARs Are Developed

Work sheets were developed to provide pertinent information about each SAR, especially the mathematical procedures for calculating toxicity values based on molecular weight and  $K_{ow}$ . Data to develop new SARs are entered in a spreadsheet that allows the SAR equations to be calculated based on a measured toxicity values (in mmoles/L) and an estimated  $K_{ow}$ . Using these estimated values, regression equations can be developed for a class of chemicals, e.g. neutral organics, acrylates, anionic surfactants, etc. Toxicity values for new chemicals may then be calculated by inserting the estimated  $K_{ow}$  into the regression equation and correcting the resultant values for the molecular weight of the compound.

As discussed above, the mode of toxic action for most neutral organics appears to be narcosis; however, some organic chemicals have a more specific mode of toxicity with comparable  $K_{ow}$ s and molecular weights. For these chemicals, the toxicity is also related to the  $K_{ow}$  and as the  $K_{ow}$  decreases (i.e., as the chemicals become more water soluble), the amount of excess toxicity compared to neutral organic compounds increases. Consequently, at some higher  $K_{ow}$  the toxicity of the compound is not significantly different from the toxicity of the equivalent neutral organic. For organic chemicals which have excess toxicity and for which are data poor, e.g., amino anilines, a neutral organic data point may be used in addition to the measured toxicity value to give a regression equation. These are the chemicals that have a N=2 entry under statistics but show only one chemical in the list of chemicals used to develop the SAR. The second point is a neutral organic  $K_{ow}$  value. In addition, for some lists of chemicals used to develop the SAR, a single chemical is listed more than once. This is because the chemical has been tested more than once. Each toxicity value is included for the chemical if it provides a reliable data point, i.e., if a second study confirmed a previously derived toxicity value.

To date, over 100 SARs have been developed for over 40 classes of organic chemicals (see Table 1). These chemical classes include neutral organics, surfactants, polymers, and other organic compounds. Most of the SARs are for acute toxicity to fish or daphnids; however, acute and chronic SARs have been developed for other organisms. Some classes, such as acid chlorides, only have one SAR (e.g., fish 96-hour  $LC_{50}$ ), while for other classes such as neutral organics more than 10 SARs have been developed ranging from acute and chronic SARs for fish to a 14-day  $LC_{50}$  for earthworms in artificial soil. New SARs will be added as data become available. This manual will be periodically updated to reflect the additions.

## Selecting an Appropriate SAR

Selecting the appropriate SAR for a new chemical is based on a variety of chemical-specific information. This information includes the exact chemical structure, chemical class, predicted  $K_{ow}$ , molecular weight of the compound, physical state, water solubility, number of carbons or ethoxylates or both, and percent amine nitrogen or number of cationic charges or both, per 1000 molecular weight. The most important factor for deriving a SAR is the chemical class as SARs are chemical class specific. An alphabetical listing of chemical classes and appropriate SARs to use for each is included at the conclusion of this section.

To estimate the toxicity to aquatic organisms of neutral, nonreactive, non-ionizable organics and organics that exhibit excess toxicity, the  $K_{ow}$  and molecular weight are required. The value for the  $K_{ow}$  should be obtained from estimated values using the computer program CLOGP, Version 3.3. The range of  $K_{ow}$  values are valid to estimate the toxicity is SAR specific and is given for each SAR in a chemical class. In general, when the  $\log K_{ow}$  is less than or equal to 5.0, valid predictions can be obtained for estimating acute toxicity to aquatic organisms from neutral organic compounds. If the  $\log K_{ow}$  is greater than 5.0, the decreased solubility of a compound will result in no effects in a saturated solution during a 96-hour test and a longer exposure duration should be used to determine the  $LC_{50}$ . For chronic exposures, the applicable  $\log K_{ow}$  may be extended up to 8.0. If the  $\log K_{ow}$  of the compound exceeds 8.0, no adverse effects are

Table 1. Existing SARs

SAR Class	Acute Toxicity			Chronic Toxicity			Other
	Fish	Daphnid	Algae	Fish	Daphnid	Algae	
Acid chlorides	X						
Acrylates	X	X					
Acrylates, methacrylates	X						
Alcohols, propargyl	X						
Aldehydes	X	X	X	X		X	
Amines, aliphatic	X	X	X			X	
Anilines	X	X		X	X	X	
Anilines, amino, meta or 1,3-substituted	X	X	X				
Anilines, amino, ortho or 1,2-substituted	X	X	X				
Anilines, amino, para or 1,4-substituted	X	X	X		X		
Anilines, dinitroanilines	X	X		X			
Aziridines	X	X	X				
Benzenes, dinitro	X	X		X	X		
Benzotriazoles	X	X	X				
Carbamates							X
Carbamates, dithio	See	SAR	Title	Page			
Crown Ethers	See	SAR	Title	Page			
Diazoniums, aromatic	X						
Epoxides, monoepoxides	X	X					
Epoxides, diepoxides	X	X					
Esters	X	X	X			X	
Esters, monoesters, aliphatic				X			
Esters, diesters, aliphatic				X			
Esters, phosphate	X						
Esters, phthalate	X	X			X		
Hydrazines	X	X	X				
Hydrazines, semicarbazide, alkyl substituted			X				
Hydrazines, semicarbazides, aryl, meta/para substituted			X				
Hydrazines, semicarbazides, aryl, ortho substituted			X				
Imides	X						
Ketones, diketones, aliphatic	X	X			X	X	

SAR Class	Acute Toxicity			Chronic Toxicity			Other
	Fish	Daphnid	Algae	Fish	Daphnid	Algae	
Malononitriles	X						
Neutral organics	X	X	X	X	X	X	X
Peroxy acids	X	X					
Phenols	X	X	X	X	X	X	
Phenols, dinitrophenols	X	X		X	X		
Polymers, polycationic	X	X	X				
Surfactants, anionic	X	X	X	X	X	X	
Surfactants, cationic, quaternary ammonium, monoalkyl	X	X					X
Surfactants, cationic, quaternary ammonium, dialkyl	X	X	X	X	X	X	
Surfactants, ethomeen	X	X	X				
Surfactants, nonionic	X	X					
Thiazolinones, iso	X	X	X			X	
Thiols (mercaptans)	X	X					
Triazines, substituted	X	X					
Ureas, substituted			X				

for neutral organic compounds in saturated solutions even with long-term exposures. Other chemical classes have other upper limits for  $K_{ow}$ . For examples, the maximum log  $K_{ow}$  for aldehydes is 6.0 and 7.0 for phenols.

#### Using SARs

All SARs contain an equation that predicts the aquatic toxicity of a chemical. Most of the SARs require the user to know the predicted log of the octanol water partition coefficient ( $K_{ow}$ ). When this number is entered into the equation, a toxicity values in millimoles/L (mM/L) is derived. The molecular weight of the subject compound is required to convert the SAR estimates from millimoles/L to mg/L. The ECOSAR program does this automatically, however, manual estimates require that conversions be made. For example, the equation for predicting the fish 96-hour  $LC_{50}$  values for neutral organics is:

$$\text{Log } LC_{50} = 1.75 - 0.94 \log K_{ow}$$

Using 1,1'-biphenyl (CASRN [92-52-4] as a representative chemical, the estimated log  $K_{ow}$  for this compound is 4.0, to give a log  $LC_{50}$  of -2.01. Taking the antilog of -2.01, gives an  $LC_{50}$  value of 0.009 mM/L. However, to express the toxicity of the 1,1'-biphenyl as mg/L, the toxicity must be multiplied by the molecular weight of the compound which is 154.20, to give a final toxicity value of 1.5 mg/L. Conversions from mM/L to mg/L are not necessary for compounds and equations (e.g., surfactants, polymers) that do not use  $K_{ow}$  as the input parameter for toxicity.

Molecular weight is also used to determine the absorption cutoff limit for aquatic organisms. As the molecular weight of a chemical increases above 600, passive absorption through respiratory membranes decreases significantly. Therefore, for chemicals with molecular weights above 1000, it has been assumed that such absorption is negligible. For surface active chemicals such as cationic polymers, molecular weight is not limiting because the toxic effect is not due to absorption, for example, some polycationic polymers with molecular weights in excess of 1,000,000 are highly toxic to aquatic organisms.

An important aspect of determining the toxicity of a compound is knowing the water solubility. The water solubility of a compound can be compared with the SAR toxicity value derived for that compound. If the toxicity value is significantly greater than the measured or predicted maximum water solubility, then an effect is not expected to occur in a saturated solution. In addition, a determination of the physical state (liquid, solid, or gas) of the compound is helpful in selecting an SAR. SARs currently used by EEB were developed using toxicity data on chemicals that are liquids at room temperature (25 °C). If an organic chemical is a solid at room temperature, then the melting point should be known because of the effect it has on water solubility, i.e., assuming  $K_{ow}$  is constant, the higher the melting point of a neutral organic chemical, the lower its water solubility. For other chemicals such as surfactants, water dispersibility is used, however, for practical purposes, water solubility and dispersibility are considered to be synonymous.

To determine the toxicity of a surfactant, it is necessary to know the number of carbon atoms in the alkyl chain for anionic surfactants or the number of ethoxylate units in the compound if it is an cationic (ethomeen) or nonionic surfactant. For cationic quaternary ammonium surfactants, the toxicity is based on the average length of a linear carbon chain, if the chain length is between 10 and 24 carbons long. The surfactant SARs developed by EEB are based on surfactants where the hydrophobic component is composed of a single linear chain of carbons and/or chains of ethoxylate units. Surfactants that have complex hydrophobic components are assessed by calculating the  $K_{ow}$  of the complex hydrophobic component alone and determining which aliphatic alkyl (carbon) chain has an equivalent  $K_{ow}$ . Toxicity predictions are based on this equivalent chemical structure. See the SAR for cationic dialkyl quaternary ammonium surfactants for more details on these calculations.

For polycationic polymers, it is necessary to calculate the percent amine nitrogen and/or number of cationic charges per 1000 molecular weight.

For inorganic and organometallic compounds, only the molecular weight of the compound is used for calculating the toxicity value. Acute and/or chronic toxicity values will be expressed in mg/L, and further conversions and/or calculations are not necessary.

#### Reliability of SARs

As may be seen by reviewing the chemicals used to derive the individual SARs in this manual, some chemical classes have a greater number of chemicals with accompanying toxicity values than do others. For example, the neutral organic 96-hour fish  $LC_{50}$  SAR was based on toxicity values for over 60 chemicals, whereas, the fish 96-hour  $LC_{50}$  SAR for propargyl alcohols was based on only one toxicity value. In the cases where there is only one toxicity value for a chemical class, the SAR is based on the line drawn between the one toxicity value and the maximum toxicity value of a neutral organic compound. Obviously SARs developed using only one or two toxicity values taken from the literature or premanufacture notices may not have the same reliability as an SAR developed from a larger toxicity database, however, on a regulatory basis this is the best estimate that can be scientifically achieved.

To determine how reliable the SARs in this manual are, Nabholz et al (1993) conducted a validation study which compared the predicated toxicity values of chemicals with their measured toxicity values. Several chemical classes were included in the study: neutral organics, organic chemicals which show

excess toxicity compared with neutral organics of a similar structure, anionic surfactants, cationic surfactants, polycationic polymers, cationic dyes, acid dyes, polyanionic monomers which are strong chelators of nutrient elements, and compounds which undergo hydrolysis (e.g., acid chlorides and alkyloxysilanes). In all, test data from 462 chemicals were used in the validation study. SARs for acute and chronic toxicity for fish, daphnids, and green algae were reviewed. Validation was expressed as a ratio, i.e., predicted toxicity:measured toxicity. A ratio of 1.0 would indicate that the predictions were perfectly accurate, a ratio of less than 1.0 would indicate an over-prediction of toxicity, and a ratio of more than 1.0 would indicate that SARs were under predicting the toxicity of the chemicals. The results of the study indicated that the algal chronic effect was most accurately predicted (ratio 1.07) while the fish chronic value was the least reliable (ratio 0.24). The fish 96-hour  $LC_{50}$  ratio was 0.64, the daphnid 48-hour  $LC_{50}$  was 0.79, and the algae 96-hour  $EC_{50}$  was 0.81. Work on validating the SARs is continuously ongoing in EEB.

## **FURTHER DISCUSSION NEEDED FOR USE OF NEAREST ANALOG; WHY GASES DON'T HAVE SAR; AND RADIONUCLIDES**

### Sources for ECOSAR

ECOSAR: Computer Program and User's Guide for Estimating the Ecotoxicity of Industrial Chemicals Based on Structure Activity Relationships (Publication Number EPA-748-R-93-002) is available from the following sources:

- National Center for Environmental Publications and Information  
U.S. Environmental Protection Agency  
26 West Martin Luther King Drive  
Cincinnati, OH 45268  
(513) 569-7562
- National Technical Information Service  
U.S. Department of Commerce  
5285 Port Royal Road  
Springfield, VA 22161  
(703) 487-4650

### References

Clements, RG, Nabholz, JV, Johnson, DW, Zeeman, M. 1993. The Use and Application of QSARs in the Office of Toxic Substances for Ecological Hazard Assessment of New Chemicals. In: Landis, WG, Hughes, JS, and Lewis, MA, eds. Environmental Toxicology and Risk Assessment, ASTM STP 1179. Philadelphia, PA: American Society for Testing and Materials pp. 56-64.

Nabholz, JV, Clements, RG, Zeeman, MG, Osborn, KC, Wedge, R. 1993. Validation of Structure Activity Relationships Used by the USEPA's Office of Pollution Prevention and Toxics for the Environmental Hazard Assessment of Industrial Chemicals. In: Gorsuch, JW, Dwyer, FJ, Ingersoll, CG, and LaPoint, TW, eds. Environmental Toxicology and Risk Assessment 2nd Vol. ASTM STP 1216 Philadelphia, PA: American Society for Testing and Materials. pp. 571-590

Nabholz, JV, Miller, P, Zeeman, M. 1993. Environmental Risk Assessment of New Chemicals Under the Toxic Substances Control Act (TSCA) Section Five. In: Landis, WG, Hughes, JS, and Lewis, MA, eds. Environmental Toxicology and Risk Assessment, ASTM STP 1179. Philadelphia, PA: American Society for Testing and Materials pp. 40-55



## CHEMICAL CLASSES AND APPLICABLE SARs

### Chemical Class

### SAR to Use

ACETATES  
ACETYLENIC CARBAMATES  
ACID CHLORIDES  
ACID DYES with ONE ACID

ACID DYES with TWO ACIDS

ACID DYES with THREE ACIDS

Use SAR for **ESTERS**  
No SAR available, excess toxicity  
Use SAR for **ACID CHLORIDES**  
Some are moderately toxic to fish and daphnids, others are not; No SAR available, Use nearest analog  
Some are moderately toxic to fish and daphnids, others are not; No SAR available, Use nearest analog  
Only moderately toxic to green algae due to the indirect effect of shading; shading inhibits growth due to the colored water; Use nearest analog based on chemical structure, color, and intensity of color.

ACRYLAMIDES and SUBSTITUTED  
ACRYLAMIDES

ACRYLATES (log Kow <5.0)  
ACRYLATES (log Kow >5.0)  
ACRYLATES, METHACRYLATES  
ACTINIUM  
ALCOHOLS  
ALCOHOLS, PROPARGYL  
ALDEHYDES  
ALDEHYDES, VINYL

Excess toxicity, Use toxicity data for acrylamides with MW adjustment  
Use SAR for **ACRYLATES**  
Use SAR for **NEUTRAL ORGANICS**  
Use SAR for **ACRYLATES, METHACRYLATES**  
No SAR available  
Use SAR for **NEUTRAL ORGANICS**  
Use SAR for **ALCOHOLS PROPARGYL**  
Use SAR for **ALDEHYDES, R-C(=O)-H,**  
No SAR available; some exhibit excess toxicity, e.g., acrolein,  
Use SAR for **AMINES, ALIPHATIC**  
Use SAR for **ESTERS, DI, ALIPHATIC**  
Use SAR for **KETONES, DI, ALIPHATIC**

ALIPHATIC AMINES  
ALIPHATIC DIESTERS  
ALIPHATIC DIKETONES, LINEAR  
ALIPHATIC HYDROCARBON,  
 $\alpha$ -HYDROXY- $\beta$ -NITRO SUBSTITUTED  
or ALIPHATIC HYDROCARBON,  
1-HYDROXY-2-NITRO SUBSTITUTED

Excess toxicity towards algae, e.g., tris(hydroxymethyl)nitromethane  
Use SAR for **ESTERS**  
Use SAR for **NEUTRAL ORGANICS** Straight chain or cycloalkane,  
Use SAR for **NEUTRAL ORGANICS**  
Use SAR for **NEUTRAL ORGANICS**  
Use SAR for **NEUTRAL ORGANICS**  
Use SAR for **ANILINES**  
Use SAR for **SURFACTANTS, ANIONIC**  
No SAR available  
Use SAR for **NEUTRAL ORGANICS**  
Use SAR for **SURFACTANTS, ETHOMEEN**  
Use SAR for **SURFACTANTS, ANIONIC**  
  
Use SAR for **MALONONITRILES**  
Use SAR for **ESTERS , DI, ALIPHATIC**

ALIPHATIC MONOESTERS  
ALIPHATIC HYDROCARBONS

ALKANES, CYCLO  
ALKANES, STRAIGHT & BRANCHED  
ALKENES  
ALKYLANILINES  
ALKYL BENZENE SULFONATES  
ALKYL ESTERS OF CARBAMIC ACID  
ALKYL HALIDES  
ALKYL-NITROGEN-ETHOXYLATES  
ALKYL SULFONATES  
ALKYL SULFONATES AND CARBOXYLIC ACID  
ALLYL CYANIDES  
ALLYL DIESTERS

ALLYL ESTER  
ALLYL HALOGENS

ALKYNES  
ALLYL, NITRILES  
ALUMINUM  
AMERICIUM  
AMIDES, VINYL

AMINES, ALIPHATIC, PRIMARY

AMINES, ALIPHATIC, SECONDARY

AMINES, ALIPHATIC, TERTIARY

AMINES, ALIPHATIC,  
QUATERNARY, SURFACTANT

AMINES, ALIPHATIC,  
QUATERNARY, NOT A SURFACTANT

AMINES, SCHIFF BASES  
AMINES, AROMATIC  
AMINO-PHENOLS  
AMINOTRIAZOLES  
AMPHOTERIC DYES

ANILINES  
ANILINES, ALKYL  
ANILINES AR-NH<sub>2</sub> with  
N-substitutions  
ANILINES, AMINO, META,  
OR 1,3-SUBSTITUTED

ANILINES, AMINO, ORTHO,  
OR 1,2-SUBSTITUTED  
**SUBSTITUTED**

ANILINES, AMINO, PARA,  
OR 1,4-SUBSTITUTED  
**SUBSTITUTED**

ANILINES, DINITRO  
ANILINES, MONOHYDROXY  
ANILINES, POLYNITRO  
ANTIMONY  
used  
ARGON  
AROMATIC DIAZONIUMS  
ARSENIC(III)

Use SAR for **ESTERS**

No SAR available ALLYL CHLORIDES show excess  
toxicity, ALLYL BROMIDES are even more toxic  
Use SAR for **NEUTRAL ORGANICS**

Use SAR for ALUMINUM

No SAR available

No SAR available. Excess toxicity, Use toxicity data for  
arylamides with MW adjustment

Use SAR for **AMINES, ALIPHATIC** when log Kow < 7.0,

Use nearest analog when log Kow > 7.0

Use SAR for **AMINES, ALIPHATIC** when log Kow < 7.0,

Use nearest analog when log Kow > 7.0

Use SAR for **AMINES, ALIPHATIC** when log Kow < 7.0,

Use nearest analog when log Kow > 7.0

Use SAR for **SURFACTANTS, CATIONIC,**  
QUATERNARY AMMONIUM, MONOALKYL

Calculate Kow for the tertiary amine and Use SAR for  
**AMINES, ALIPHATIC** when log Kow < 7.0, nearest  
analog when log Kow > 7.0; or Use nearest analog  
method

No SAR available

Use SAR for **ANILINES**

Use SAR for **ANILINES**

If charges are balanced, low toxicity towards fish and  
daphnids, and shading only towards algae; if more  
cationic than anionic, see **CATIONIC DYES**; and if more  
anionic than cationic, see **ACID DYES**

Use SAR for **ANILINES**

Use SAR for **ANILINES**

Use SAR for **NEUTRAL ORGANICS**

Use SAR for **ANILINES, AMINO, META, OR 1,3-**  
**SUBSTITUTED**

Use SAR for **ANILINES, AMINO, ORTHO, OR 1,2-**

Use SAR for **ANILINES, AMINO, PARA OR 1,4-**

Use SAR for **ANILINES, DINITRO**

Use SAR for **ANILINES**

Use SAR for **ANILINES, DINITRO**

No SAR available; however water quality criteria may be

Gas; No SAR available

Use SAR for **DIAZONIUMS, AROMATIC**

Use SAR for **ARSENIC**

ARYL HALIDES  
 ASTATINE  
 AZIRIDINES  
 AZO DYES  
 BARIUM  
 BENZENE, DINITRO  
 BENZENEAMINES  
 BENZOATES  
 BENZOTRIAZOLES  
 BENZOTRIAZOLES with free -NH  
 BENZOTRIAZOLES with  
   N-alkyl substitution  
 BENZOTRIAZOLES with  
   N-thiol substitution  
 BENZOYL PEROXIDES  
  
 BERKELIUM  
 BERYLLIUM  
 BIPHENYLS, POLYBROMINATED  
 BISMUTH  
 BORON  
 BROMINE  
 CADMIUM  
 CALCIUM  
 CALIFORNIUM  
 CAPROLACTAMS  
 CARBAMATES  
 CARBAMATES, BIS(ETHYL)-  
   JOINED AT -NRN- BY ALKYL  
   OR ARYL GROUPS  
 CARBAMATES, ETHYL, N-ALKYL  
   OR ARYL SUBSTITUTED  
 CARBAMATES, BIS OR TRIS,  
   ESTERIFIED ON A SINGLE  
   PHENYL RING  
 CARBAMATES, THIO  
 CARBON  
 CARBOXYLIC ACIDS  
 CATIONIC DYES  
  
 CERIUM  
 CESIUM  
 CHLORINATED HYDROCARBONS  
 CHLORINE  
 CHLOROANILINES  
 CHLOROFLUOROCARBONS (CFCs)  
 CHROMIUM  
 CHROMIUM(III)  
 CHROMIUM(VI)  
 COBALT  
 COPPER

Use SAR for **NEUTRAL ORGANICS**  
 No SAR available  
 Use SAR for **AZIRIDINES**,  
 No SAR available; see DYES  
 No SAR available  
 Use SAR for **BENZENES, DINITRO**  
 Use SAR for **ANILINES**  
 Use SAR for **ESTERS**  
 Use SAR for **BENZOTRIAZOLES**  
 Use SAR for **BENZOTRIAZOLES**, has excess toxicity,  
  
 Use SAR for **NEUTRAL ORGANICS**  
  
 No SAR available  
 Use SAR for **PEROXY ACIDS**,  $RC(=O)OOC(=O)R$ ,  
 excess toxicity,  
 No SAR available  
 Use SAR for **BERYLLIUM**  
 Use SAR for **NEUTRAL ORGANICS**  
 No SAR available  
 Use SAR for **BORON**  
 No SAR available  
 Use SAR for **CADMIUM**  
 No SAR available  
 No SAR available  
 Use SAR for **NEUTRAL ORGANICS**  
 No SAR available  
  
 No SAR available  
  
 No SAR available  
  
 No SAR available  
 No SAR available  
 No SAR available  
  
 No SAR available, Use nearest analog, MWs can be  
 over 1000  
 No SAR available  
 Use SAR for **CESIUM**  
 Use SAR for **NEUTRAL ORGANICS**  
 Use SAR for **CHLORINE**  
 Use SAR for **ANILINES**  
 Use SAR for **NEUTRAL ORGANICS**  
 Use SAR for **CHROMIUM**  
 Use SAR for **CHROMIUM**  
 Use SAR for **CHROMIUM**  
 Use SAR for **COBALT**  
 Use SAR for **COPPER**

CROWN ETHERS  
CURIUM  
CYANIDE, VINYL

CYANATES

CYCLIC DIKETONES  
CYCLOALKANES  
CYCLODIENE  
DIAMINES, PHENYLENE  
(META OR 1,3 SUBSTITUTED)

DIAMINES, PHENYLENE  
(ORTHO OR 1,2-SUBSTITUTED)  
**SUBSTITUTED**

DIAMINES, PHENYLENE  
(PARA OR 1,4-SUBSTITUTED)  
**SUBSTITUTED**

DIAZONIUMS, ALIPHATIC

DIAZONIUMS, AROMATIC  
DICARBOXYLIC ALIPHATIC ESTERS  
DIEPOXIDES  
DIESTER, ALLYL

DIESTERS, AROMATIC OR  
ALIPHATIC/AROMATIC  
DIKETONES,  $\alpha,\gamma$ -diketone or  
1,3-diketones, linear  
pentanediols, excess toxicity  
DIKETONES, 1,3-diketones, cyclic  
DINITROANILINES  
DINITROBENZENES  
DINITROPHENOLS  
DIPHENOLS  
DISPERSE DYES  
DISULFIDES  
DISULFIDE, PHENYL  
DITHIOCARBAMATES  
DITHIOCARBAMATES, POLY  
DYES

DYSPROSIUM  
ERBIUM

See SAR for **CROWN ETHERS**

No SAR available

No SAR available,  $R-C=C-C\equiv N$ , e.g., acrylonitrile,  
fumaronitrile, have excess toxicity,

No SAR available, (NCO-R) or (R-OCN), excess toxicity,  
Use nearest analog

Use SAR for **NEUTRAL ORGANICS**

Use SAR for **NEUTRAL ORGANICS**

Use SAR for **NEUTRAL ORGANICS**

Use SAR for **ANILINES, AMINO, META, OR 1,3-  
SUBSTITUTED**

Use SAR for **ANILINES, AMINO, ORTHO OR 1,2-**

Use SAR for **ANILINES, AMINO, PARA OR 1,4-**

No SAR available ( $R-N\equiv N-A$ ), (very explosive and are  
used as synthesizing agents)

Use SAR for **DIAZONIUM, AROMATIC**( $AR-N\equiv N-AR$ ),

Use SAR for **ESTERS**

Use SAR for **EPOXIDES, DI**

No SAR available  $R-C-C=C-C-(O-C(=O)-C-R)-O-C(=O)-$   
 $C-R$  excess toxicity e.g., 2-propene-1,1-diol, diacetate,  
1000X more toxic than an equivalent NEUTRAL  
ORGANIC,

Use SAR for **ESTERS, PHTHALATE**

Use SAR for **KETONES, DI, ALIPHATIC**; e.g., 2,4-

Use SAR for **NEUTRAL ORGANICS**

Use SAR for **ANILINES, DINITRO**

Use SAR for **BENZENES, DINITRO**

Use SAR for **PHENOLS, DINITRO**

Use SAR for **PHENOLS**

Use SAR for **NEUTRAL ORGANICS**

Use SAR for **NEUTRAL ORGANICS**

No SAR available, excess toxicity,

See SAR for **CARBAMATES, DITHIO**

No SAR available, excess toxicity,

see ACID DYES if ANIONIC DYES; see CATIONIC DYES  
if cationic; see NEUTRAL DYES if neutral, and see  
AMPHOTERIC DYES if both cationic and anionic; MWs  
can be over 1000 for CATIONIC DYES, AMPHOTERIC  
DYES, and ACID DYES; MWs of NEUTRAL DYES have  
to be less than 1000 for toxicity towards fish and  
daphnids; toxicity to green algae is based on color and  
intensity of color, and is an indirect effect

No SAR available

No SAR available

EPOXIDES, AZIRIDINES  
 EPOXIDES, DIEPOXIDES  
 EPOXIDES, MONOEPOXIDES  
 EPOXIDES, POLYEPOXIDES  
 ESTERS (log Kow <5.0)  
 ESTERS (log Kow >5.0)  
 ESTERS  
 ESTER, ALLYL  
  
 ESTERS,  $\alpha$ -HALO-  
  
 ESTERS, DICARBOXYLIC, ALIPHATIC  
 ESTERS, DIESTERS, ALIPHATIC  
 ESTERS, METHANESULFONATES  
 ESTERS, PHOSPHATE  
 ESTERS, PHOSPHINOTHIOIC ACID,  
 TRISUBSTITUTED  
  
 ESTERS, PHOSPHINOTHIOIC ACID,  
 DISUBSTITUTED-FREE ACID  
  
 ESTERS, PHOSPHINOTHIOIC ACID,  
 MONOSUBSTITUTED-FREE DIACID  
  
 ESTERS, PHOSPHOROTHIOIC,  
 MONOESTER  
  
 ESTERS, PHOSPHOROTHIOIC,  
 MONOSUBSTITUTED ESTER  
  
 ESTERS, PHOSPHOROTHIOIC,  
 DISUBSTITUTED ESTER  
  
 ESTERS, PHOSPHOROTHIOIC,  
 TRIESTER  
 ESTERS, PHOSPHOROTHIOIC,  
 TRISUBSTITUTED  
 ESTERS, PHTHALATE  
 ESTERS, POLY  
 ESTERS, PROPARGYLIC  
 ESTERS, SULFONATE  
 ESTERS, TRIALKYL PHOSPHATE  
 ESTERS, VINYL  
 EINSTEINIUM  
 ETHERS  
 ETHOXYLATES, ALKYL  
 EUROPIUM  
 FATTY ACIDS  
 FERMIUM

Use SAR for **AZIRIDINES**  
 Use SAR for **EPOXIDES**, DI  
 Use SAR for **EPOXIDES**, MONO  
 Use SAR for **EPOXIDES**, DI  
 Use SAR for **ESTERS**  
 Use SAR for **NEUTRAL ORGANICS**  
 Use SAR for **ESTERS**, RC(=O)OR,  
 Use SAR for **ESTERS**, R-C=C-C-O-C(=O)-C-R, excess  
 toxicity,  
 No SAR available, C-O-C(=O)-C-X, excess toxicity,  
 BROMIDES are more toxic than CHLORIDES  
 Use SAR for **ESTERS**  
 Use SAR for **ESTERS**, DI,ALIPHATIC  
 Use SAR for **ESTERS**  
 Use SAR for **ESTERS**, PHOSPHATE  
  
 No SAR Available, R-O-P(=S)(O-R)R, pesticide, Use  
 nearest analog  
  
 Use SAR for **SURFACTANTS**, **ANIONIC** if alkyl chains  
 are long; if alkyl chains are short, use nearest analog  
 (R-O-P(=S)(OH)R)  
  
 Use SAR for **SURFACTANTS**, **ANIONIC** if alkyl chains  
 are long; if alkyl chains are short, Use nearest analog  
 (HO-P(=S)(OH)R)  
  
**WE NEED A DESCRIPTION FOR THIS; USES BOTH  
 ANIONIC SURFACTANT AND DIESTER SARs**  
  
 Use SAR for **SURFACTANTS**, **ANIONIC** if alkyl chain is  
 long; if alkyl chain is short, Use nearest analog (R-O-  
 P(=S)(OH)OH)  
  
 Use SAR for **SURFACTANTS**, **ANIONIC**, if alkyl chain is  
 long, if alkyl chain is short, Use nearest analog  
  
 Use SAR for **ESTERS**, **PHOSPHATE**  
  
 Use SAR for **ESTERS**, **PHOSPHATE**  
 Use SAR for **ESTERS**, **PHTHALATE**  
 Use SAR for **ESTERS**  
 No SAR available, have excess toxicity  
 Use SAR for **ESTERS**  
 Use SAR for **ESTERS**, **PHOSPHATE**  
 No SAR available, excess toxicity  
 No SAR available  
 Use SAR for **NEUTRAL ORGANICS**  
 Use SAR for **SURFACTANTS**, **NONIONIC**  
 No SAR available  
 Use SAR for **SURFACTANTS**, **ANIONIC**  
 No SAR available

FLUORINE  
 FRANCIUM  
 GADOLINIUM  
 GALLIUM  
 GERMANIUM  
 GOLD  
 GUANIDINE  
 HAFNIUM  
 HALIDES, ALKYL  
 HALIDES, ARYL  
 HALOGENATED FLUOROCARBONS  
 HELIUM  
 HFCs  
 HOLMIUM  
 HYDRAZIDES  
 HYDRAZINES  
 HYDRAZINES, CARBOXYLIC  
 (FREE) ACID SUBSTITUTION

HYDRAZINES, SEMICARBAZIDES, ARYL,  
 META/PARA SUBSTITUTED  
**SUBSTITUTED**

HYDRAZINES, SEMICARBAZIDES, ARYL,  
 ORTHO SUBSTITUTED  
**SUBSTITUTED**

HYDRAZONES

HYDROCARBONS, AROMATIC

HYDROCARBONS, AROMATIC,  
 HALOGENATED

HYDROCARBONS, ALIPHATIC,  
 HALOGENATED

HYDROGEN

HYDROQUINONES or PARA-HYDROXY

PHENOL

IMIDES

INDIUM

INDOLES, HALOGENATED

IODINE

IRIDIUM

IRON

ISOCYANATES, MONO- AND DI-  
 ISOCYANATES (R-NCO) and  
 ISOTHIOCYANATES

ISOTHIAZOLINONES

KETONES,  $\alpha$ -HALO-

KETONES, MONO

KETONES, DIKETONES, ALIPHATIC

KRYPTON

LANTHANUM

LAWRENCIUM

LEAD

No SAR available

No SAR available

No SAR available

No SAR available

Use SAR for **GERMANIUM**

Use SAR for **GOLD**

Use SAR for **AMINES, ALIPHATIC**

No SAR available

Use SAR for **NEUTRAL ORGANICS**

Use SAR for **NEUTRAL ORGANICS**

Use SAR for **NEUTRAL ORGANICS**

No SAR available

Use SAR for **NEUTRAL ORGANICS**

No SAR available

Use SAR for **HYDRAZINES**

Use SAR for **HYDRAZINES**

No SAR available, about 10 times less toxic than  
 HYDRAZINES

Use SAR for **SEMICARBAZIDES, ARYL, META/PARA**

Use SAR for **SEMICARBAZIDES, ARYL, ORTHO**

Use SAR for **HYDRAZINES**

Use SAR for **NEUTRAL ORGANICS**

Use SAR for **NEUTRAL ORGANICS**

Use SAR for **NEUTRAL ORGANICS**

No SAR available; toxicity is based on pH \*\*\*\*\*

No SAR available, excess toxicity

Use SAR for **IMIDES**

No SAR available

Use SAR for **NEUTRAL ORGANICS**

No SAR available

No SAR available

Use SAR for **IRON**

No SAR available, excess toxicity if very water soluble.

Use nearest analog

Use SAR for **THIAZOLINOES, ISO**

No SAR available, excess toxicity

Use SAR for **NEUTRAL ORGANICS**

Use SAR for **KETONES, DI Aliphatic**

No SAR available

Use SAR for **LANTHANUM**

No SAR available

Use SAR for **LEAD**

LINALOOLS  
 LINEAR ALKYL BENZENES  
 LINEAR ALKYL BENZENE  
 SULFONATES  
 LINEAR ALKYL SULFONATES  
 LITHIUM  
 LUTETIUM  
 MAGNESIUM  
 MALEIMIDES  
 MALONONITRILES  
 MANGANESE  
 MENDELEVIUM  
 MERCAPTANS/THIOLS  
 MERCAPTOTRIAZOLES:  
 MERCURY  
 METHACRYLAMIDES  
 SUBSTITUTED ACRYLAMIDES  
 METHACRYLATES (log Kow <5.0)  
 METHACRYLATES (log Kow >5.0)  
 METHANESULFONATES  
 MOLYBDENUM  
 MONOEPOXIDES  
 NEON  
 NEUTRAL DYES  
 NEUTRAL ORGANICS  
 NEODYMIUM  
 NEPTUNIUM  
 NICKEL  
 NIOBIUM  
 NITRILES  
 NITRILES, ALLYL  
 NITRILES, VINYL  
 NITROBENZENES,  
 DINITROBENZENES  
 NITROGEN  
 NITROSO COMPOUNDS  
 NOBELIUM  
 OSMIUM  
 OXYGEN  
 PALLADIUM  
 PEROXY ACIDS  
 PHENOLS  
 PHENOLS, AMINO  
 PHENOLS, DI  
 PHENOLS, DINITRO  
 PHENOLS, HALOGENATED  
 PHENOL, META-HYDROXY  
 PHENOL, ORTHO-HYDROXY  
 PHENOL, PARA-HYDROXY or  
 HYDROQUINONE  
 PHENOLS, POLY

Use SAR for **NEUTRAL ORGANICS**  
 Use SAR for **SURFACTANTS, ANIONIC**  
 Use SAR for **SURFACTANTS, ANIONIC**  
 Use SAR for **SURFACTANTS, ANIONIC**  
 No SAR available  
 No SAR available  
 No SAR available  
 Use SAR for **IMIDES**  
 Use SAR for **MALONONITRILES**  
 No SAR available  
 No SAR available  
 Use SAR for **THIOLS**; (R-SH)  
 No SAR available , excess toxicity  
 Use SAR for **MERCURY**  
 No SAR available, less toxic than ACRYLAMIDES and  
 Use SAR for **METHACRYLATES**  
 Use SAR for **NEUTRAL ORGANICS**  
 Use SAR for **ESTERS**  
 Use SAR for **MOLYBDENUM**  
 Use SAR for **EPOXIDES, MONO**  
 No SAR available  
 Use SAR for **NEUTRAL ORGANICS**  
 Use SAR for **NEUTRAL ORGANICS**  
 No SAR available  
 No SAR available  
 Use SAR for **NICKEL**  
 No SAR available  
 Use SAR for **NEUTRAL ORGANICS**  
 Use SAR for **MALONONITRILES**  
 Use SAR for **MALONONITRILES**  
 Use SAR for **BENZENES, DINITRO**  
 No SAR available  
 No SAR available, excess toxicity  
 No SAR available  
 No SAR available  
 No SAR available  
 No SAR available  
 Use SAR for **PEROXY ACIDS**  
 Use SAR for **PHENOLS**  
 Use SAR for **ANILINES**  
 Use SAR for **PHENOLS**  
 Use SAR for **PHENOLS, DINITRO**  
 Use SAR for **PHENOLS**  
 Use SAR for **PHENOLS**  
 No SAR available, CATECHOL, 16 times excess fish  
 acute toxicity  
 No SAR available, 1400 times excess fish acute toxicity  
 Use SAR for **PHENOLS**

PHENOLS, SUBSTITUTED  
PHENYLENEDIAMINES  
PHOSPHINOTHIOIC ACID ESTERS,  
DISUBSTITUTED FREE ACID  
PHOSPHINOTHIOIC ACID ESTERS,  
MONOSUBSTITUTED FREE ACID  
PHOSPHITES  
PHOSPHONIUM

PHOSPHOROTHIOIC ESTERS,  
DIESTER  
PHOSPHOROTHIOIC ESTERS,  
MONOESTER  
PHOSPHORUS  
PLATINUM  
PLUTONIUM  
POLONIUM  
POLYANIONIC MONOMERS

POLYAROMATIC HYDROCARBONS  
POLYBROMINATED BIPHENYLS  
POLYCATIONIC POLYMERS  
POLYEPOXIDES

POLYISOCYANATES

POLYMERS, POLYNONIONIC  
POLYMERS, POLYANIONIC,  
POLY(CARBOXYLIC ACID)  
POLYMERS, POLYANIONIC,  
POLY(ACRYLIC ACID)  
POLYMERS, POLYANIONIC,  
POLY(METHACRYLIC ACID)  
POLYMERS, POLYANIONIC,  
POLY(AROMATIC SULFONIC ACID)  
POLYMERS, POLYANIONIC,  
POLY(ALIPHATIC SULFONIC ACID)  
POLYMERS, POLYCATIONIC  
POLYMERS, POLYAMINE  
POLYMERS, POLYQUATERNARY  
AMMONIUM  
POLYMERS, POLYPHOSPHONIUM  
POLYMERS, POLYSULFONIUM  
POLYNUCLEAR AROMATICS  
POLYSULFIDES

Use SAR for **PHENOLS**  
Use SAR for **ANILINES, AMINO \*\*\*\*\***

Use SAR for **SURFACTANTS, ANIONIC**

Use SAR for **SURFACTANTS, ANIONIC**  
No SAR available, excess toxicity  
Use SAR for **SURFACTANTS, CATIONIC,**  
QUATERNARY AMMONIUM if a surfactant; if not a  
surfactant use nearest analog: SULFONIUM or  
QUATERNARY AMMONIUM analogs are acceptable

Use SAR for **SURFACTANTS, ANIONIC**

Use SAR for **SURFACTANTS, ANIONIC**  
Use SAR for **PHOSPHORUS**  
Use SAR for **PLATINUM**  
No SAR available  
No SAR available  
No SAR available, monomers with two or more acid  
groups and which act like organic acid chelators, Use  
nearest analog  
Use SAR for **NEUTRAL ORGANICS**  
Use SAR for **NEUTRAL ORGANICS**  
Use SAR for **POLYMERS, POLYCATIONIC**  
Use SAR for **EPOXIDES, DI,**

No SAR available, if water solubility is 13 mg/L or less,  
then no effects at saturation; these chemicals will  
polymerize: one NCO will hydrolyze to the amine and  
the amine will react with another NCO to form a  
urethane; a crosslinked polymer will be formed  
No SAR available, low environmental hazard.

No SAR available, Use nearest analog  
No SAR available, Use nearest analog

No SAR available, Use nearest analog

No SAR available, Use nearest analog

No SAR available, Use nearest analog  
Use SAR for **POLYMERS, POLYCATIONIC**  
Use SAR for **POLYMERS, POLYCATIONIC**

Use SAR for **POLYMERS, POLYCATIONIC**  
Use SAR for **POLYMERS, POLYCATIONIC**  
Use SAR for **POLYMERS, POLYCATIONIC**  
Use SAR for **NEUTRAL ORGANICS**  
Use SAR for **NEUTRAL ORGANICS**

POTASSIUM	No SAR available
PRASEODYMIUM	No SAR available
PROMETHIUM	No SAR available
PROPARGYL ALCOHOLS	Use SAR for <b>ALCOHOLS, PROPARGYL</b>
PROPARGYL CARBAMATES	No SAR available, excess toxicity
PROPARGYLIC ESTERS	No SAR available, excess toxicity
PROPARGYL HALIDE	No SAR available, excess toxicity, PROPARGYL
	BROMIDE more toxic than PROPARGYL CHLORIDE
PROTACTINIUM	No SAR available
QUATERNARY AMMONIUM SURFACTANTS, DIALKYL	Use SAR for <b>SURFACTANTS, CATIONIC,</b>
QUATERNARY AMMONIUM, DIALKYL	
QUATERNARY AMMONIUM SURFACTANTS, MONOALKYL	Use SAR for <b>SURFACTANTS, CATIONIC,</b>
QUATERNARY AMMONIUM, MONOALKYL	
QUINONES	No SAR available, para-benzoquinone, 5500X excess
toxicity to fish	
RADIUM	No SAR available
RADON	No SAR available
RHENIUM	No SAR available
RHODIUM	No SAR available
RUBIDIUM	No SAR available
RUTHENIUM	No SAR available
SAMARIUM	No SAR available
SCANDIUM	No SAR available
SCHIFF BASES	Use SAR for SCHIFF BASES a subclass of AMINES with excess toxicity; (R-N=C-R)
	Use SAR for SELENIUM
SELENIUM	
SEMICARBAZIDES, ALKYL SUBSTITUTED	Use SAR for <b>SEMICARBAZIDES, ALKYL SUBSTITUTED</b>
SEMICARBAZIDES, ARYL META/PARA SUBSTITUTED	Use SAR for <b>SEMICARBAZIDES, ARYL, META/PARA</b>
SUBSTITUTED	
SEMICARBAZIDES ARYL ORTHO SUBSTITUTED	Use SAR for <b>SEMICARBAZIDES, ARYL, ORTHO</b>
SUBSTITUTED	
SEMICARBAZIDES	Use SAR for <b>HYDRAZINES</b>
SEMICARBAZONES	Use SAR for <b>HYDRAZINES</b>
SILANES, ALKOXY RSi(OR)(OR)(OR) and CHLOROSILANES	reactive with water (hydrolyses) and generally shows low toxicity towards fish, moderate toxicity towards daphnids, and high toxicity towards green algae; the hydrolysis products (silic acids and silanols) probably overchelate nutrient elements and inhibit the growth of algae, all SARs for silanes have to be based on Kows which have C substituted for Si
SILICON	No SAR available
SILVER	Use SAR for <b>SILVER</b>
SODIUM	No SAR available
STRONTIUM	No SAR available
SULFIDES	Use SAR for <b>NEUTRAL ORGANICS</b>

SULFIDES (C-S-C), DISULFIDES  
(C-S-S-C), and POLYSULFIDES  
SULFONATES, ALKYL BENZENE  
SULFONATES, ALKYL

SULFONATES, METHANE  
SULFONIUM

SULFUR  
SULFONATES, LINEAR ALKYL  
SULFONYL CHLORIDES  
SURFACTANTS, ALCOHOL  
ETHOXYLATE  
SURFACTANTS, ALKYL  
ETHOXYLATE  
SURFACTANTS, AMPHOTERIC  
SURFACTANTS, ANIONIC  
SURFACTANTS, ANIONIC,  
CARBOXYLIC ACID  
SURFACTANTS, ANIONIC,  
ALKYL-BENZENE-SULFONATE  
SURFACTANTS, ANIONIC,  
ALKYL-SULFONATE

SURFACTANTS, ANIONIC,  
PHOSPHATE

SURFACTANTS, ANIONIC,  
ALKYL-ETHOXYLATE-SULFONATE

SURFACTANTS, ANIONIC,  
ALKYL-(SULFONATE and  
CARBOXYLIC ACID)

SURFACTANTS, ANIONIC,  
TWEEN-TYPE  
SURFACTANTS, CATIONIC,  
ALKYL-NITROGEN-ETHOXYLATES  
ETHOMEEN  
SURFACTANTS, CATIONIC,  
QUATERNARY AMMONIUM  
DIALKYL

SURFACTANTS, CATIONIC,  
QUATERNARY AMMONIUM,

Use SAR for **NEUTRAL ORGANICS**  
Use SAR for **SURFACTANTS, ANIONIC**  
Use SAR for **SURFACTANTS, ANIONIC**

Use SAR for **ESTERS**  
Use SAR for **SURFACTANTS, CATIONIC**,  
QUATERNARY AMMONIUM, if a surfactant; if not a  
surfactant, Use nearest analog: PHOSPHONIUM or  
QUATERNARY AMMONIUM analogs are acceptable.  
No SAR available  
Use SAR for **SURFACTANTS, ANIONIC**  
No SAR available, excess toxicity (RS(=O)(=O)Cl)

Use SAR for **SURFACTANTS, NONIONIC**

Use SAR for **SURFACTANTS, NONIONIC**  
Use SAR for **SURFACTANTS, ANIONIC**  
Use SAR for **SURFACTANTS, ANIONIC**

No SAR available

Use SAR for **SURFACTANTS, ANIONIC**

Use SAR for **SURFACTANTS, ANIONIC**, calculate Kow  
of alkyl, convert to equivalent alkyl-benzene based on  
equivalent Kow and use SAR for **SURFACTANTS,**  
**ANIONIC**

Use SAR for **SURFACTANTS, ANIONIC**

Use SAR for **SURFACTANTS, ANIONIC** to predict  
toxicity of alkyl-sulfonate and then adjust toxicity  
depending on number of ethoxylates

No SAR available; predict toxicity of alkyl-sulfonate and  
divide effective concentration by 10 times

No SAR available, Use nearest analog

Use SAR for **SURFACTANTS, ETHOMEEN**  
Use SAR for **SURFACTANTS, ETHOMEEN**

Use SAR for **SURFACTANTS, CATIONIC**,  
QUATERNARY AMMONIUM, DIALKYL, with two large  
alkyl chains

MONOALKYL

Use SAR for **SURFACTANTS, CATIONIC, QUATERNARY AMMONIUM, MONOALKYL**, with one large alkyl chain

SURFACTANTS, CATIONIC,  
QUATERNARY AMMONIUM,  
N-ETHOXYLATED

Use SAR for **SURFACTANTS, CATIONIC QUATERNARY AMMONIUM, MONOALKYL**, if ethoxy groups are less than five. If ethoxylates are greater than five, Use SAR for **SURFACTANTS, CATIONIC, QUATERNARY AMMONIUM MONOALKYL** and then reduce toxicity due to the presence of the ethoxylates through the use of the SAR for **SURFACTANTS, NONIONIC**.

SURFACTANTS, CATIONIC,  
QUATERNARY AMMONIUM  
TRIALKYL

Use SAR for **SURFACTANTS, CATIONIC, QUATERNARY AMMONIUM, MONOALKYL**, three large alkyls,

SURFACTANTS, CATIONIC,  
QUATERNARY AMMONIUM,  
TETRAALKYL

Use SAR for **SURFACTANTS, CATIONIC, QUATERNARY AMMONIUM, MONOALKYL**, four large alkyls

SURFACTANTS, NONIONIC  
SURFACTANTS, ETHOMEEN  
SURFACTANTS, LINEAR ALKYL  
BENZENE SULFONATES  
SURFACTANTS, NONIONIC  
SURFACTANT, NONIONIC,  
ALKYL-ETHOXYLATES  
SURFACTANT, NONIONIC,  
ALKYL-ETHOXYLATE-ALKYL  
SURFACTANT, NONIONIC,  
TWEEN-TYPE  
SULFONATES, LINEAR ALKYL  
BENZENE  
TANTALUM  
TECHNIUM  
TELLURIUM  
TERBIUM  
TERPENES  
THALLIUM  
THIAZOLINONES, ISO  
THIOLS (MERCAPTANS)  
THIOHYDRAZIDES  
THIOSEMICARBAZIDES  
THIOSEMICARBAZONES  
THORIUM  
THULIUM  
TIN

Use SAR for **SURFACTANTS, ETHOMEEN**

Use SAR for **SURFACTANTS, ANIONIC**  
Use SAR for **SURFACTANTS, NONIONIC**

Use SAR for **SURFACTANTS, NONIONIC**

No SAR available, use nearest analog

No SAR available, Use nearest analog

Use SAR for **SURFACTANTS, ANIONIC**

No SAR available

No SAR available

No SAR available

No SAR available

Use SAR for **NEUTRAL ORGANICS**

Use SAR for **THALLIUM**

Use SAR for **THIAZOLINONES, ISO**

Use SAR for **THIOLS**

Use SAR for **HYDRAZINES**

Use SAR for **HYDRAZINES**

Use SAR for **HYDRAZINES**

No SAR available

No SAR available

No SAR available for inorganic tins or organotins, Use nearest analog

Use SAR for **TITANIUM**

TITANIUM

TRIAZIDES, BENZO, N-ALKYL  
SUBSTITUTED  
TRIAZINES, SUBSTITUTED  
TRIAZOLES  
TRIAZOLES, AMINO  
toxicity,  
TRIAZOLES, BENZO  
TUNGSTEN  
VANADIUM  
URANIUM  
UREAS, CYCLIC  
UREAS, SUBSTITUTED

VINYL AMIDES  
VINYL ESTERS  
VINYL NITROS

VINYL SULFONE

XENON  
YTTERBIUM  
YTTRIUM  
ZINC  
ZIRCONIUM

Use SAR for **NEUTRAL ORGANICS**  
See SAR for **TRIAZINES, SUBSTITUTED**  
No SAR available, excess toxicity, Use nearest analogs  
Use SAR for **NEUTRAL ORGANICS**, herbicide, excess

Use SAR for BENZOTRIAZOLES  
Use SAR for TUNGSTEN  
Use SAR for VANADIUM  
No SAR available

Use SAR for **NEUTRAL ORGANICS**  
Use SAR for UREAS, SUBSTITUTED for green algae; to  
predict toxicity to fish and aquatic invertebrates, Use  
SAR for **NEUTRAL ORGANICS**

No SAR available  
No SAR available  
No SAR available,  $\alpha$ -nitro-styrene, excess toxicity, (R-  
 $C \equiv C = N(=O)(=O)$ )  
No SAR available, e.g., divinyl sulfone, excess toxicity,  
( $C = C - S(=O)(=O) - R$ )  
No SAR available  
No SAR available  
No SAR available  
Use SAR for **ZINC**  
Use SAR for **ZIRCONIUM**

**CHEMICAL CLASSES  
AND THEIR  
STRUCTURE ACTIVITY RELATIONSHIPS**



**SAR**

**ACID CHLORIDES**

**Organism:** Fish  
**Duration:** 96-h  
**Endpoint:** LC50 (Mortality)  
**Equation:**  $\text{Log LC50 (mM/L)} = 0.565 - 0.613 \log K_{ow}$   
**Statistics:**  $N = 3; R^2 = 1.0$   
**Maximum log  $K_{ow}$ :** 8.0  
**Maximum MW:** 1000.0  
**Application:** This SAR may be used to estimate the toxicity of acid chlorides.  
**Limitations:** If the log  $K_{ow}$  value is greater than 8.0, or if the compound is solid and the LC50 exceeds the water solubility.  
**References:** Curtis MW, Copeland TL, and Ward CH. 1978. Aquatic toxicity of substances proposed for spill prevention regulation. Proc. Natl. Conf. Control of Hazardous Material Spills, Miami Beach, FL. p. 93-103.  
 Curtis MW and Ward CH. 1981. Aquatic toxicity of forty industrial chemicals: testing in support of hazardous substance spill prevention regulation. J. Hydrol. 51:359-367.

LIST OF ACID CHLORIDES USED TO DEVELOP THE FISH 96-h LC50 SAR

CHEMICAL	96-h LC50 (mg/L)	Log $K_{ow}$	Ref
Benzoyl chloride	34.7	1.9	C1
Benzoyl chloride	34.1	1.9	C2

C1 = Curtis et al (1978)  
 C2 = Curtis et al (1981)

**ACID CHLORIDES**  
**9/1993**

9/1993

**SAR**                      **ALCOHOLS, PROPARGYL**

**Organism:** Fish  
**Duration:** 96-h  
**Endpoint:** LC50 (Mortality)  
  
**Equation:**  $\text{Log 96-h LC50 (mM/L)} = 0.056 - 0.511 \log K_{ow}$   
  
**Statistics:**  $N = 2, R^2 = 1.0$   
  
**Maximum log  $K_{ow}$ :** 5.0  
**Maximum MW:** 1000.0  
  
**Application:** This SAR may be used to estimate toxicity for propargyl alcohols.  
  
**Limitations:** If the log  $K_{ow}$  value is greater than 5.0, or if the compound is solid and the LC50 exceeds the water solubility, use SAR with longer duration.  
  
**References:** United States Environmental Protection Agency (USEPA). 1991. OTS PMN ECOTOX. Washington, DC: Office of Toxic Substances, USEPA

## LIST OF PROPARGYL ALCOHOLS USED TO DEVELOP THE FISH 96-h LC50 SAR.

CHEMICAL	96-h LC50 (mg/L)	Log $K_{ow}$	Ref.
Chemical identity CBI	310.0	-0.4	EPA

EPA = USEPA (1991); chemical identity is Confidential Business Information under TSCA.

**ALCOHOL, PROPARGYL**  
9/1993

## ACRYLATES

7/1988

### SAR

### ACRYLATES

**Organism:**

Fish

**Duration:**

96-h

**Endpoint:**

LC50 (Mortality)

**Equation:**

$\text{Log LC50 (mM/L)} = -1.46 - 0.18 \log K_{ow}$

**Statistics:**

$N = 10, R^2 = 0.627$

**Maximum  $K_{ow}$ :**

5.0

**Maximum MW:**

1000.0

**Application:**

This SAR may be used to estimate the toxicity of acrylates and polyacrylates. Allyl acrylate is expected to be about 30 times more toxic than predicted by this SAR.

**Limitations:**

**References:**

Nabholz JV and Platz RD. 1987. Environmental effects of acrylates and methacrylates. I. Category Program Support Document - Generic SNUR and II. Generic Environmental Hazard Assessment (Addendum to Standard Review of PMN 87-930/931). Washington, DC: Environmental Effects Branch, Health and Environmental Review Division (TS-796), Office of Toxic Substances, United States Environmental Protection Agency 20460-0001

United States Environmental Protection Agency (USEPA). 1991. OTS PMN ECOTOX Washington, DC. Office of Toxic Substances, USEPA

**ACRYLATES**

7/1988

## LIST OF ACRYLATES USED TO DEVELOP THE FISH 96-h LC50 SAR.

CHEMICAL	96-h LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
2-Hydroxyethyl acrylate	4.8	-0.058	EPA
2-Hydroxypropyl acrylate	3.61	0.251	EPA
2-Hydroxypropyl acrylate	3.26	0.251	EPA
2-Hydroxypropyl acrylate	3.10	0.251	EPA
Chemical identity CBI	13.0	1.6	EPA
Isobutyl acrylate	2.110	2.204	EPA
Isobutyl acrylate	2.090	2.204	EPA
Cyclohexyl acrylate	1.48	2.778	EPA
Hexyl acrylate	1.14	3.392	EPA
Hexyl acrylate	1.09	3.392	EPA
Lauryl acrylate	*	6.566	EPA

\* No mortalities within 96 hours at saturation

EPA = USEPA (1991); chemical identity is Confidential Business Information under TSCA.

## SAR

## ACRYLATES

Organism: Daphnid

Duration: 48-h

Endpoint: LC50

Equation:  $\text{Log LC50 (mM/L)} = 0.009 - 0.511 \log K_{ow}$ Statistics:  $N = 2; R^2 = 1.0$ Maximum  $K_{ow}$ : 5.0

Maximum MW: 1000.0

Application: This SAR may be used to estimate toxicity for acrylates.

## Limitations:

References: Beach SA. 1990. Acute toxicity of isooctyl acrylate to Daphnia magna  
 St Paul, MN: 3M Environmental Laboratory, 3M Company; Toxicity  
 Test Report.

United States Environmental Protection Agency (USEPA). 1991. OTS  
 PMN ECOTOX Washington, DC: USEPA, Office of Toxic Substances.

## LIST OF ACRYLATES USED TO DEVELOP THE DAPHNID LC50 SAR.

CHEMICAL	48-h LC50 (mg/L)	Log $K_{ow}$	Ref.
Chemical identity CBI	59.0	0.78	EPA
Isooctyl acrylate	1.2	4.3	B

B = Beach (1990)

EPA = USEPA (1991); Chemical identity is Confidential Business Information under TSCA.

ACRYLATES  
7/1988

<b>SAR</b>	<b>ACRYLATES</b>
<b>Organism:</b>	Green Algae
<b>Duration:</b>	96-h
<b>Endpoint:</b>	EC50 (Growth)
<b>Equation:</b>	$\text{Log EC50 (mM/L)} = -1.02 - 0.49 \log K_{ow}$
<b>Statistics:</b>	$N = 3; R^2 = 0.91$
<b>Maximum log <math>K_{ow}</math>:</b>	6.4
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This SAR may be used to estimate toxicity for acrylates.
<b>Limitations:</b>	If the log $K_{ow}$ value is greater than 6.4, or if the compound is solid and the EC50 exceeds the water solubility, use SAR with longer exposure.
<b>References:</b>	United States Environmental Protection Agency (USEPA). 1991. OTS PMN ECOTOX. Washington, DC: USEPA, Office of Toxic Substances.

## LIST OF ACRYLATES USED TO DEVELOP THE SAR.

CHEMICAL	96-h EC50 (mg/L)	Log $K_{ow}$	Ref.
Chemical identity CBI	2.2	0.78	EPA
Chemical identity CBI	18.5	1.6	EPA

EPA = U.S. EPA (1991); Chemical identities are Confidential Business Information under TSCA.

## ACRYLATES

7/1988

## SAR

## ACRYLATES

<b>Organism:</b>	Fish
<b>Duration:</b>	32-d
<b>Endpoint:</b>	Chronic Value (Survival/Growth)
<b>Equation:</b>	$\text{Log ChV (mM/L)} = -1.99 - 0.526 \log K_{ow}$
<b>Statistics:</b>	$N = 2; R^2 = 1.0$
<b>Maximum log <math>K_{ow}</math>:</b>	8.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This SAR may be used to estimate toxicity for acrylates
<b>Limitations:</b>	If the ChV is greater than water solubility or the log $K_{ow}$ is greater than 8.0, no effects expected at saturation.
<b>References:</b>	United States Environmental Protection Agency (USEPA). 1991. Fish Chronic Toxicity Data Base. Duluth, MN: Environmental Research Laboratory (ERL), Office of Research and Development, USEPA, 6201 Congdon Boulevard, 55804; contact C.L. Russom (218) 720-5500.

## LIST OF ACRYLATES USED TO DEVELOP THE FISH CHRONIC VALUE (ChV) SAR.

CHEMICAL	96-h LC50 (mg/L)	Log $K_{ow}$	Ref.
2-Hydroxyethyl acrylate	1.33	-0.1	D

D = USEPA (1991)

**ACRYLATES**  
7/1988

**SAR****ACRYLATES, METHACRYLATES****Organism:** Fish**Duration:** 96-h**Endpoint:** LC50 (Mortality)**Equation:**  $\text{Log LC50 (mM/L)} = 0.552 - 0.715 \log K_{ow}$ **Statistics:**  $N = 19; R^2 = 0.774$ **Maximum log  $K_{ow}$ :** 5.0**Maximum MW:** 1000.0**Application:** This SAR may be used to estimate the toxicity of methacrylates and polyacrylates. Allyl methacrylate is about 35 times more toxic than predicted by this SAR.**Limitations:** If the log  $K_{ow}$  value is greater than 5.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.**References:** Nabholz JV and Platz RD. 1987. Environmental effects of acrylates and methacrylates. I. Category Program Support Document - Generic SNUR and II. Generic Environmental Hazard Assessment (Addendum to Standard Review of PMN 87-930/931). Washington, DC: Environmental Effects Branch, Health and Environmental Review Division (TS-796), Office of Toxic Substances, United States Environmental Protection Agency 20460-0001.

United States Environmental Protection Agency (USEPA). 1991. OTS PMN ECOTOX Washington, DC: Office of Toxic Substances, USEPA

**ACRYLATES, METHACRYLATES**  
9/1993

LIST OF METHACRYLATES USED TO DEVELOP THE FISH 96-h LC50 SAR.

CHEMICAL	96-h LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
Methylene chloride	322 895	1.25	Z
2-Hydroxyethyl methacrylate	227.0	0.251	EPA
Methylmethacrylate	151.0	1.056	EPA
Tetrahydrofurfuryl	34.7	1.297	EPA
2-Ethoxyethyl methacrylate	27.7	1.402	EPA
3-(Trimethoxysilyl)propyl	175.0	1.464	EPA
Allyl methacrylate	0.99	1.570	EPA
Chemical identity CBI	34.0	1.774	EPA
Chemical identity CBI	32.0	1.774	EPA
Isopropyl methacrylate	38.0	1.894	EPA
Benzyl methacrylate	4.67	2.824	EPA

EPA = USEPA (1991); chemical identity is Confidential Business Information under TSCA.

**SAR**

**ALDEHYDES**

**Organism:**

Fish

**Duration:**

96-h

**Endpoint:**

LC50 (Mortality)

**Equation:**

$\text{Log LC50 (mM/L)} = -0.449 \log K_{ow} - 0.314$

**Statistics:**

$N = 54; R^2 = 0.527$

**Maximum log  $K_{ow}$ :**

5.0

**Maximum MW:**

1000.0

**Application:**

This SAR may be used to estimate the toxicity of aldehydes. Acrolein is about 1400 times more toxic than predicted by this SAR.

**Limitations:**

**References:**

Brooke LT, Call DJ, Geiger DL, and Northcott CE. 1984. Acute toxicity of organic chemicals to fathead minnows (Pimephales promelas). Volume I. Center for Lake Superior Environmental Studies, University of Wisconsin - Superior. Superior, Wisconsin.

Geiger DL, Northcott CE, Call DJ, and Brooke LT. 1985. Acute toxicity of organic chemicals to fathead minnows (Pimephales promelas). Volume II. Center for Lake Superior Environmental Studies, University of Wisconsin - Superior. Superior, Wisconsin.

Geiger DL, Poirier SH, Brooke LT, and Call DJ. 1986. Acute toxicity of organic chemicals to fathead minnows (Pimephales promelas). Volume III. Center for Lake Superior Environmental Studies, University of Wisconsin - Superior. Superior, Wisconsin.

United States Environmental Protection Agency (USEPA). 1991. Fish acute toxicity database. Duluth, MN: Environmental Research Laboratory (ERL), Office of Research and Development, USEPA. 6201 Congdon Blvd, 55804; contact C L. Russom (218) 720-5500

# ALDEHYDES

7/1988

## LIST OF ALDEHYDES USED TO DEVELOP THE FISH 96-h LC50 SAR.

CHEMICAL	96-h LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
Ethanal	30.800	-0.22	EPA
Butanal #1	19.000	0.88	EPA
Butanal #2	16.000	0.88	EPA
Butanal #3	13.400	0.88	EPA
2-Methylbutyraldehyde	9.970	1.14	EPA
Vanillin #2	123.000	1.21	EPA
Vanillin #1	57.000	1.21	EPA
Isovaleraldehyde	3.250	1.23	EPA
Valeraldehyde #1	12.400	1.36	EPA
Valeraldehyde #2	13.400	1.37	EPA
o-Vanillin #1	2.600	1.37	EPA
o-Vanillin #2	2.200	1.37	EPA
2,4,5-Trimethoxybenzaldehyde	49.500	1.38	EPA
Benzaldehyde #2	12.800	1.48	EPA
Benzaldehyde #1	7.610	1.48	EPA
4-Nitrobenzaldehyde	10.100	1.50	EPA
5-Hydroxy-2-nitrobenzaldehyde	41.900	1.65	EPA
2-Methylvaleraldehyde	18.800	1.67	EPA
2,4-Dihydroxybenzaldehyde	13.100	1.71	EPA
o-Nitrobenzaldehyde #1	12.500	1.74	EPA
o-Nitrobenzaldehyde #2	16.600	1.74	EPA
o-Fluorobenzaldehyde	1.350	1.76	EPA
Hexanal #1	22.000	1.78	EPA
Hexanal #2	14.000	1.78	EPA
p-Dimethylaminobenzaldehyde	45.700	1.81	EPA
Salicylaldehyde	2.300	1.81	EPA
3-Ethoxy-4-hydroxybenzaldehyde	87.600	1.88	EPA
5-Bromo-2-nitrovanillin	73.300	1.88	EPA
2,4-Dimethoxybenzaldehyde	20.100	1.91	EPA
2,3-Dimethylvaleraldehyde	16.000	2.07	EPA
5-Bromovanillin	59.700	2.09	EPA
4-Chlorobenzaldehyde	2.200	2.10	EPA
o-Tolualdehyde	52.900	2.26	EPA
2-Chloro-5-nitrobenzaldehyde #1	3.800	2.28	EPA
2-Chloro-5-nitrobenzaldehyde #2	3.950	2.28	EPA
p-Ethoxybenzaldehyde	28.100	2.31	EPA
4,6-Dimethoxy-2-hydroxybenzaldehyde	2.680	2.33	EPA
Pentafluorobenzaldehyde	1.100	2.45	EPA
α,α,α-Trifluoro-m-tolualdehyde #3	1.130	2.47	EPA
α,α,α-Trifluoro-m-tolualdehyde #2	0.760	2.47	EPA
α,α,α-Trifluoro-m-tolualdehyde #1	0.920	2.47	EPA

## ALDEHYDES

7/1988

Continued

CHEMICAL	96-h LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
2-Chloro-6-fluorobenzaldehyde	9.410	2.54	EPA
4-(Diethylamino)benzaldehyde	23.900	2.94	EPA
5-Chlorosalicylaldehyde	0.770	3.00	EPA
p-Isopropyl benzaldehyde	6.620	3.07	EPA
2,4-Dichlorobenzaldehyde	1.800	3.11	EPA
5-Bromosalicylaldehyde	1.300	3.15	EPA
4-(Diethylamino)salicylaldehyde	5.360	3.34	EPA
3,5-Dibromosalicylaldehyde	0.850	3.83	EPA
p-Phenoxybenzaldehyde	4.600	3.96	EPA
4-(Hexyloxy)-m-anisaldehyde	2.670	3.99	EPA
3-(3,4-Dichlorophenoxy) benzaldehyde	0.300	5.49	EPA
3-(4-Tert-butylphenoxy) benzaldehyde	0.370	5.93	EPA
Tetradecanal	*	6.12	EPA

\* No effects at saturation.

EPA = USEPA (1991)

**ALDEHYDES**  
7/1988

**SAR**

**ALDEHYDES**

**Organism:** Daphnid  
**Duration:** 48-h  
**Endpoint:** LC50 (Mortality)  
**Equation:**  $\text{Log 48-h LC50 (mM/L)} = -0.059 - 0.607 \log K_{ow}$   
**Statistics:**  $N = 4, R^2 = 1.0$   
**Maximum log  $K_{ow}$ :** 6.0  
**Maximum MW:** 1000.0  
**Application:** This SAR may be used to estimate toxicity for aldehydes.  
**Limitations:**  
**References:** Sloof W, Canton JH, and Hermens JLM. 1983 Comparison of the susceptibility of 22 freshwater species to 15 chemical compounds. I (Sub)Acute toxicity tests. Aquatic Toxicology 4:113-128.

LIST OF ALDEHYDES USED TO DEVELOP THE DAPHNID 48-h LC50 SAR.

CHEMICAL	48-h LC50 (mg/L)	Log $K_{ow}$	Ref
Salicylaldehyde	5.4	2.1	S
Salicylaldehyde	5.5	2.1	S
Salicylaldehyde	5.8	2.1	S

S = Sloof et al (1983)

## ALDEHYDES

9/1993

**SAR**

**ALDEHYDES**

**Organism:**

Green Algae

**Duration:**

96-h

**Endpoint:**

EC50 (Growth)

**Equation:**

Use green algae 96-h EC50 SAR developed for neutral organics.

**Maximum log  $K_{ow}$ :**

6.4

**Maximum MW:**

1000.0

**Application:**

The green algae 96-h SAR for neutral organics may be used to estimate toxicity for aldehydes.

**Limitations:**

**References:**

See references for neutral organics.

**ALDEHYDES**  
9/1993

**SAR**

**ALDEHYDES**

**Organism:** Fish  
**Duration:** 32-d  
**Endpoint:** Chronic Value (Survival/Growth)  
**Equation:**  $\text{Log ChV} = -0.81 - 0.68 \log K_{ow}$   
**Statistics:**  $N = 3; R^2 = 0.97$   
**Maximum log  $K_{ow}$ :** 8.0  
**Maximum MW:** 1000.0  
**Application:** This SAR may be used to estimate toxicity for aldehydes.  
**Limitations:** If the log Kow is greater than 8.0, or if the ChV exceeds the water solubility, no effects expected at saturation.  
**References:** United States Environmental Protection Agency (USEPA). 1991. Fish Chronic Toxicity Data Base. Duluth, MN: Environmental Research Laboratory (ERL), Office of Research and Development, USEPA, 6201 Congdon Boulevard, 55804; contact C.L. Russom (218) 720-5500.

LIST OF ALDEHYDES USED TO DEVELOP THE FISH 32-d Chronic Value (Survival/Growth) SAR.

CHEMICAL	32-d ChV (mg/L)	Log $K_{ow}$	Ref.
o-Tolualdehyde	1.61	2.1	D
$\alpha, \alpha, \alpha$ -Trifluoro-m-tolualdehyde	0.19	2.6	D

D = USEPA (1991)

**ALDEHYDES**  
9/1993

**SAR**

**ALDEHYDES**

**Organism:**

Green Algae

**Duration:**

96-h

**Endpoint:**

Chronic Value (Growth)

**Equation:**

Use the equation for the green algae chronic value SAR developed for neutral organics.

**Maximum log  $K_{ow}$ :**

8.0

**Maximum MW:**

1000.0

**Application:**

The green algae chronic value SAR for neutral organics may be used to estimate toxicity for aldehydes.

**Limitations:**

If the log  $K_{ow}$  is greater than 8.0, or if the ChV exceeds the water solubility, no effects expected at saturation

**References:**

Sloof W, Canton JH, and Hermens JLM. 1983 Comparison of the susceptibility of 22 freshwater species to 15 chemical compounds. I (Sub)Acute toxicity tests. Aquatic Toxicology 4:113-128

**ALDEHYDES**  
9/1993

**SAR****AMINES, ALIPHATIC****Organism:**

Fish

**Duration:**

96-h

**Endpoint:**

LC50 (Mortality)

**Equation:** $\text{Log 96-h LC50 (mM/L)} = 0.72 - 0.64 \log K_{ow}$ **Statistics:** $N = 55; R^2 = 0.82$ **Maximum log  $K_{ow}$ :**

6.0

**Maximum MW:**

1000.0

**Application:**

This SAR may be used to estimate the toxicity of aliphatic amines.

**Limitations:**If the log  $K_{ow}$  value is greater than 6.0, no effects expected in a saturated solution.**References:**

Bridie AL, Wolff CJM, and Winter M. 1979. The acute toxicity of some petrochemicals to goldfish. *Water Research* 13:623-626

Brooke LT, Call DJ, Geiger DL, and Northcott CE (eds). 1984. Acute toxicities of organic chemicals to fathead minnows (Pimephales promelas). Superior, WI: Center for Lake Superior Environmental Studies, University of Wisconsin-Superior. Volume I.

Calamari D, DaGasso R, Galassi S, Provini A, and Vighi M. 1980. Biodegradation and toxicity of selected amines on aquatic organisms. *Chemosphere* 9:753-762.

Geiger DL, Piorier SH, Brooke LT, and Call DJ (eds). 1986. Acute toxicities of organic chemicals to fathead minnows (Pimephales promelas). Superior, WI: Center for Lake Superior Environmental Studies, University of Wisconsin-Superior. Volume III.

Geiger DL, Call DJ, and Brooke LT (eds). 1988. Acute toxicities of organic chemicals to fathead minnows (Pimephales promelas). Superior, WI: Center for Lake Superior Environmental Studies, University of Wisconsin-Superior. Volume IV.

Platz RD and Nabholz JV. 1990. Generic environmental hazard assessment of aliphatic amines. Washington, DC. Environmental Effects Branch, Health and Environmental Review Division (TS-796), Office of Toxic Substances, United States Environmental Protection Agency. Unpublished manuscript

## **AMINES, ALIPHATIC**

9/1993

United States Environmental Protection Agency (USEPA). 1990. Summary of structure-activity data files: University of Wisconsin - Superior (UWS) and Environmental Research Laboratory, Duluth, MN (ERL-D) research team. Computer printout from Environmental Effects Branch, HERD, USEPA, Washington, DC.

United States Environmental Protection Agency (USEPA). 1991. OTS PMN ECOTOX. Washington, DC: Office of Toxic Substances; United States Environmental Protection Agency.

LIST OF ALIPHATIC AMINES USED TO DEVELOP THE FISH 96-h LC50 SAR.

CHEMICAL	96-h LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
Triethanolamine	1180.000	-1.59	EPA1
1,3-Diaminopropane	1190.000	-1.49	BR
Diethanolamine	47100.000	-1.46	EPA1
Ethanolamine	2070.000	-1.30	EPA1
Ethylenediamine	220.000	-1.22	EPA1
1,2-Diaminopropane	1010.000	-0.91	BR
Morpholine	380.000	-0.72	C
2-Methoxyethylamine	524.000	-0.67	BR
Dimethylamine	118.000	-0.52	C
2-(Ethylamino)ethanol	1480.000	-0.46	BR
Allylamine	27.000	-0.15	B
Ethylamine	227.000	-0.14	EPA1
N-(3-Methoxypropyl)-3,4,5- trimethoxybenzylamine	136.000	0.09	EPA1
5-Diethylamino-2-pentanone	336.000	0.35	G1
Propylamine	308.000	0.39	BR
N,N-Diethylethanolamine	1780.000	0.40	G1
Diallylamine	20.000	0.51	B
Diethylamine	855.000	0.54	BR
Diethylamine	182.000	0.54	C
tert-Butylamine	270.000	0.57	C
3-Dimethylaminopropyl chloride hydrochloride	133.000	0.66	G1
sec-Butylamine	275.000	0.70	EPA1
2-(Diisopropylamino)ethanol	201.000	0.86	G2
n-Butylamine	268.000	0.92	G1
Benzylamine	102.000	1.09	EPA1
1,2-Dimethylpropylamine	284.000	1.10	G1
Diisopropylamine	196.000	1.16	C
2,2-Dimethyl-1-propylamine	475.000	1.19	G1
1,8-Diamino-p-menthane	65.300	1.23	G2
Tripropargylamine	296.000	1.26	G1
Cyclohexylamine	90.000	1.37	C
N,N-bis(2,2-Diethoxyethyl) methylamine	634.000	1.39	G1
N,N-bis(2,2-Diethoxyethyl) methylamine	637.000	1.39	G1
Amylamine	177.000	1.45	G1
3,3-Dimethylbutylamine	602.000	1.72	G2
Chemical Identity CBI	778.000	1.93	EPA
N,N-Dimethylbenzylamine	37.800	1.98	EPA1
Hexylamine	56.600	1.98	G1

**AMINES, ALIPHATIC**  
9/1993

Continued

CHEMICAL	96-h LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
1-Adamantylamine	25.000	2.00	G1
N-Ethylbenzylamine	57.100	2.04	EPA1
tert-Octylamine	24.600	2.43	G2
Heptylamine	21.800	2.51	BR
Dibutylamine	37.000	2.66	C
Tripropylamine	50.900	2.82	G1
1-Methylheptylamine	5.110	2.82	BR
1-Methylheptylamine	5.280	2.82	BR
N,N-Diethylcyclohexylamine	21.400	2.98	G2
Octylamine	5.190	3.04	G2
Nonylamine	2.160	3.57	EPA1
Chemical identity CBI	2.800	4.10	EPA
Decylamine	1.030	4.10	EPA1
Undecylamine	0.210	4.63	EPA1
Dihexylamine	0.780	4.77	BR
Dodecylamine	0.103	5.16	EPA1
Tridecylamine	0.065	5.68	EPA1

EPA = USEPA (1991); Chemical identity is Confidential Business Information under TSCA.

EPA1 = USEPA (1990)

BR = Brooke et al (1984)

B = Bridie et al (1979)

C = Calamari et al (1980)

G1 = Geiger et al (1986)

G2 = Geiger et al (1988)

## SAR

## AMINES, ALIPHATIC

Organism:

Daphnids

Duration:

48-h

Endpoint:

LC50 (Mortality)

Equation:

 $\text{Log 48-h LC50 (mM/L)} = -0.524 - 0.584 \log K_{ow}$ 

Statistics:

 $N = 10; R^2 = 0.78$ Maximum log  $K_{ow}$ :

5.0

Maximum MW:

1000.0

Application:

This SAR may be used to estimate the toxicity of aliphatic amines.

Limitations:

If the log  $K_{ow}$  value is greater than 5.0, no effects expected in a saturated solution.

References:

Cowgill UM, Takahashi IT, and Applegath SL. 1985. A comparison of the effect of four benchmark chemicals on Daphnia magna and Ceriodaphnia dubia/affinis tested at two different temperatures.

Gersich FM, Milazzo DP, and Voos-Esquivel C. 1988. An invertebrate life-cycle study of the toxicity of Daphnia magna Straus. Mammalian and Environmental Toxicology Research Laboratory. Dow Chemical Company Study ID: ES-DR-0065-5425-6.

LeBlanc GA. 1980 Acute toxicity of priority pollutants to water flea (Daphnia magna). Bulletin of Environmental Contamination and Toxicology 24:684-691.

Platz RD and Nabholz JV. 1990. Generic environmental hazard assessment of aliphatic amines. Washington, DC: Environmental Effects Branch, Health and Environmental Review Division (TS-796), Office of Toxic Substances, United States Environmental Protection Agency. Unpublished manuscript.

United States Environmental Protection Agency (USEPA). 1991. OTS PMN ECOTOX Washington, DC: Office of Toxic Substances, United States Environmental Protection Agency

Van Leeuwen CJ, Maas-Diepeveen JL, Niebeek G, Vergouw WHA, Griffioen PS, and Luijken MW. 1985. Aquatic toxicological aspects of dithiocarbamates and related compounds I. Short-term toxicity tests. Aquatic Toxicology 7:145-164

## AMINES, ALIPHATIC

### LIST OF ALIPHATIC AMINES USED TO DEVELOP THE DAPHNID 48-h LC50 SAR

CHEMICAL	48-h LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
Diethanolamine	131.000	-1.46	C
Diethanolamine	55.000	-1.46	L
Ethylenediamine	26 500	-1.22	VL
Chemical identity CBI	1760.000	-0.90	EPA
Dimethylamine	50.000	-0.52	VL
Chemical identity CBI	4.300	0.44	EPA
Diethylamine	56.000	0.54	VL
Chemical identity CBI	15 000	1.03	EPA
Chemical identity CBI	3.800	2.74	EPA
2-(Decylthio)ethylamine hydrochloride	0.033	4.85	G

C = Cowgill et al (1985)

EPA = USEPA (1991); Chemical identity is Confidential Business Information under TSCA

G = Gersich et al (1988)

L = LeBlanc (1980)

VL = Van Leeuwen et al (1985)

## SAR AMINES, ALIPHATIC

**Organism:** Green Algae  
**Duration:** 96-h  
**Endpoint:** EC50 (Growth)

**Equation:**  $\text{Log 96-h EC50 (mM/L)} = -0.548 - 0.434 \log K_{ow}$

**Statistics:**  $N = 14; R^2 = 0.74$

**Maximum log  $K_{ow}$ :** 7.0  
**Maximum MW:** 1000.0

**Application:** This SAR may be used to estimate toxicity for aliphatic amines.

**Limitations:** If the log  $K_{ow}$  value is greater than 7.0, no effects expected in a saturated solution.

**References:** Calamari D, DaGasso R, Galassi S, Provini A, and Vighi M. 1980. Biodegradation and toxicity of selected amines on aquatic organisms. Chemosphere 9:753-762.

Platz RD and Nabholz JV. 1990. Generic environmental hazard assessment of aliphatic amines. Washington, DC: Environmental Effects Branch, Health and Environmental Review Division (TS-796), Office of Toxic Substances, United States Environmental Protection Agency. Unpublished manuscript.

United States Environmental Protection Agency (USEPA). 1990. Summary of structure-activity data files: University of Wisconsin - Superior (UWS) and Environmental Research Laboratory, Duluth, MN (ERL-D) research team. Computer printout from Environmental Effects Branch, HERD, USEPA, Washington, DC.

United States Environmental Protection Agency (USEPA). 1991. OTS PMN ECOTOX. Washington, DC: Office of Toxic Substances, United States Environmental Protection Agency.

Van Leeuwen CJ, Maas-Diepeveen JL, Niebeek G, Vergouw WHA, Griffioen PS, and Luijken MW. 1985. Aquatic toxicological aspects of dithiocarbamates and related compounds I. Short-term toxicity tests Aquatic Toxicology 7:145-164.

## AMINES, ALIPHATIC

### LIST OF ALIPHATIC AMINES USED TO DEVELOP THE ALGAL 96-h EC50 SAR.

CHEMICAL	96-h EC50 (mg/L)	Log K <sub>ow</sub>	Ref.
Ethylenediamine	61.000	-1.22	VL
Morpholine	28.000	-0.72	C
Dimethylamine	30.000	-0.52	VL
Dimethylamine	9.000	-0.52	C
Diethylamine	20.000	0.54	C
Diethylamine	56.000	0.54	VL
tert-Butylamine	16.000	0.57	C
Chemical identity CBI	1.800	1.03	EPA
Diisopropylamine	20.000	1.16	C
Cyclohexylamine	20.000	1.37	C
Dibutylamine	19.000	2.66	C
Chemical identity CBI	1.040	2.74	EPA
Octylamine	0.220	3.04	EPA1
Chemical identity CBI	0.130	6.85	EPA

C = Calamari et al (1980)

EPA1 = USEPA (1990)

EPA = USEPA (1991); Chemical identity is Confidential Business Information under TSCA.

VL = Van Leeuwen et al (1985)

## AMINES, ALIPHATIC

### SAR

### AMINES, ALIPHATIC

**Organism:**

Green Algae

**Duration:**

**Endpoint:**

Chronic Value (Growth)

**Equation:**

$\text{Log ChV (mM/L)} = -1.399 - 0.334 \log K_{ow}$

**Statistics:**

$N = 11, R^2 = 0.61$

**Maximum log  $K_{ow}$ :**

7.0

**Maximum MW:**

1000.0

**Application:**

This SAR may be used to estimate toxicity for aliphatic amines.

**Limitations:**

If the log  $K_{ow}$  value is greater than 7.0, no effects expected at saturation.

**References:**

Calamari D, DaGasso R, Galassi S, Provini A, and Vighi M. 1980. Biodegradation and toxicity of selected amines on aquatic organisms. Chemosphere 9 753-762.

Platz RD and Nabholz JV. 1990. Generic environmental hazard assessment of aliphatic amines. Washington, DC: Environmental Effects Branch, Health and Environmental Review Division (TS-796), Office of Toxic Substances, United States Environmental Protection Agency. Unpublished manuscript.

United States Environmental Protection Agency (USEPA). 1989. Report on alga toxicity tests on selected OTS chemicals. Unpublished preliminary draft. Corvallis Environmental Research Laboratory. Corvallis, OR: United States Environmental Protection Agency.

United States Environmental Protection Agency (USEPA). 1991. OTS PMN ECOTOX. Washington, DC: Office of Toxic Substances, United States Environmental Protection Agency.

## AMINES, ALIPHATIC

### LIST OF ALIPHATIC AMINES USED TO DEVELOP THE ALGAL ChV SAR.

CHEMICAL	ChV (mg/L)	Log K <sub>ow</sub>	Ref.
Morpholine	1.000	-0.72	C
Dimethylamine	2.000	-0.52	C
Diethylamine	2.000	0.54	C
tert-Butylamine	2.000	0.57	C
Chemical identity CBI	0.110	1.03	EPA2
Diisopropylamine	5.000	1.16	C
Cyclohexylamine	5.000	1.37	C
Dibutylamine	2.500	2.66	C
Chemical identity CBI	0.410	2.74	EPA2
Octylamine	0.650	3.04	EPA1
Chemical identity CBI	0.050	6.85	EPA2

C = Calamari et al (1980)

EPA1 = USEPA (1989)

EPA2 = USEPA (1991); Chemical identity is Confidential Business Information under TSCA.

**SAR**

**ANILINES**

**Organism:** Fish  
**Duration:** 96-h  
**Endpoint:** LC50 (Mortality)

**Equation:**  $\text{Log 96-h LC50 (mM/L)} = 0.956 - 0.739 \log K_{ow}$

**Statistics:**  $N = 20; R^2 = 0.882$

**Maximum log  $K_{ow}$ :** 7.0  
**Maximum MW:** 1000.0

**Application:** This SAR may be used to estimate toxicity for anilines.

**Limitations:** Di- and tri-nitroanilines are more toxic than predicted; a fish 96-h LC50 SAR has been developed for dinitroanilines.

2,3,5,6-Tetrachloroaniline is 19 times more toxic than predicted by this SAR. Tetrabromoaniline may be more toxic than predicted by this SAR as well.

N-substituted anilines are less toxic than predicted by this SAR; for these compounds use the neutral organics fish 96-h LC50 SAR.

If the log  $K_{ow}$  value is greater than 7.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.

**References:** Veith GD and Broderius SJ 1987. Structure-toxicity relationships for industrial chemicals causing type (II) narcosis syndrome. In: Kaiser KLE (ed.). QSAR in Environmental Toxicology-II Boston, MA: D. Reidel Pub. Co., pp 385-391.

**ANILINES**

7/1988

## LIST OF ANILINES USED TO DEVELOP THE FISH 96-h LC50 SAR.

CHEMICAL	96-h LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
ANILINES USED IN CALCULATION OF THIS SAR			
aniline	134.0	0.9	VB
4-nitroaniline	125.0	1.3	VB
4-toluidine	149.0	1.4	VB
4-chloroaniline	32.5	1.8	VB
4-ethylaniline	73.0	2.0	VB
pentafluoroaniline	37.1	2.2	VB
2-chloro-4-nitroaniline	20.2	2.2	VB
4-bromoaniline	47.5	2.3	VB
4-ethoxy-2-nitroaniline	26.0	2.5	VB
$\alpha,\alpha,\alpha$ -4-tetrafluoro- 2-toluidine	29.6	2.6	VB
$\alpha,\alpha,\alpha$ -4-tetrafluoro- 3-toluidine	30.1	2.6	VB
3,4-dichloroaniline	7.6	2.7	VB
3-benzyloxyaniline	9.14	2.8	VB
4-butylaniline	10.2	3.2	VB
2,3,6-trichloroaniline	3.64	3.3	VB
4-hexyloxyaniline	3.2	3.7	VB
2,6-diisopropylaniline	15.3	4.1	VB
4-octylaniline	0.120	5.3	VB
4-decylaniline	0.062	6.3	VB
4-dodecyl aniline	*	7.4	VB
ANILINES WITH EXCESS TOXICITY			
2,3,5,6-tetrachloroaniline	0.270	4.1	VB

\* No fish mortality in saturated solutions.

VB = Veith and Broderius (1987)

**SAR**

**ANILINES**

**Organism:**

Daphnid

**Duration:**

48-h

**Endpoint:**

LC50 (Mortality)

**Equation:**

$\text{Log 48-h LC50 (mM/L)} = -1.623 - 0.271 \log K_{ow}$

**Statistics:**

$N = 24; R^2 = 0.24$

**Maximum log  $K_{ow}$ :**

7.0

**Maximum MW:**

1000.0

**Application:**

This SAR may be used to estimate toxicity for anilines

**Limitations:**

Di- and tri-nitroanilines are more toxic than predicted by this SAR; a daphnid 48-h LC50 SAR has been developed for dinitroanilines.

Tetrachloro- and tetrabromo-aniline may be 20 times more toxic than predicted by this SAR.

N-substituted anilines are less toxic than predicted by this SAR; for these compounds use the neutral organics daphnid 48-h LC50 SAR

If the log  $K_{ow}$  value is greater than 7.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.

**References:**

Canton JH and Adema DMM. 1978. Reproducibility of short-term and reproduction toxicity experiments with Daphnia magna and comparison of the sensitivity of Daphnia magna with Daphnia pulex and Daphnia cucullata in short-term experiments. *Hydrobiologia* 2:135-140.

Kuhn R, Pattard M, Pernak K-D and Winter A. 1989. Results of the harmful effects of selected water pollutants (anilines, phenols, aliphatic compounds) to Daphnia magna. *Water Research* 23:495-499.

Sloof W, Canton JH, and Hermens JLM. 1983. Comparison of the susceptibility of 22 freshwater species to 15 chemical compounds. I. (Sub)Acute toxicity tests. *Aquatic Toxicology* 4:113-128.

# ANILINES

9/1993

## LIST OF ANILINES USED TO DEVELOP THE DAPHNID 48-h LC50 SAR.

CHEMICAL	48-h LC50 (mg/L)	K <sub>ow</sub>	Log	Ref.
p-aminophenol	0.240	0.2	K	
m-aminophenol	1.1	0.2	K	
aniline	0.640	0.6	S	
benzidine (dianiline)	0.600	1.6	K	
4-aminoacetophenone	5.0	0.9	K	
aniline	0.300	0.9	K	
aniline	0.100	0.9	CA	
aniline	0.680	0.9	CA	
p-methoxyaniline	1.9	1.0	K	
2-amino-4-methoxyphenol	3.0	1.3	K	
5-chloro-2,4-dimethoxyaniline	1.62	1.8	K	
p-chloroaniline	0.310	1.9	K	
m-chloroaniline	0.350	1.9	K	
o-chloroaniline	1.8	1.9	K	
p-ethylaniline	2.0	2.1	K	
o-bromoaniline	3.0	2.1	K	
o-ethylaniline	14.0	2.1	K	
2,4-dimethylaniline	9.9	2.2	K	
3-trifluoromethylaniline	2.7	2.3	K	
4-chloro-2-nitroaniline	3.2	2.6	K	
3-chloro-4-methylaniline	0.620	2.6	K	
2,6-dichloroaniline	1.4	2.8	K	
2,4-dichloroaniline	2.7	2.8	K	

K = Kuhn et al (1989)

S = Sloof et al (1983)

CA = Canton and Adema (1978)

**SAR**

**ANILINES**

**Organism:** Fish  
**Duration:** 32-d  
**Endpoint:** Chronic Value (Survival/Growth)

**Equation:**  $\text{Log ChV (mM/L)} = -1.516 - 0.625 \log K_{ow}$

**Statistics:**  $N = 11; R^2 = 0.66$

**Maximum log  $K_{ow}$ :** 8.0  
**Maximum MW:** 1000.0

**Application:** This SAR may be used to estimate toxicity for anilines.

**Limitations:** N-substituted anilines are less toxic than predicted by this SAR; for these compounds use the neutral organics fish ChV SAR.

If the log  $K_{ow}$  is greater than 8.0, or if the compound is solid and the ChV exceeds the water solubility, no effects expected at saturation.

**References:**

Bresch H, Beck H, Ehlermann D, Schlaszus H and Urbanek M. 1990. A long-term toxicity test comprising reproduction and growth of zebrafish with 4-chloroaniline. Archives of Environmental Contamination and Chemistry 19:419-427.

Call DJ, Poirier SH, Knuth ML, Harting SL and Lindbery CA. 1987. Toxicity of 3,4-dichloroaniline to fathead minnow, Pimephales promelas, in acute and early life-stage exposures. Bulletin of Environmental Contamination and Toxicology 38:352-358.

United States Environmental Protection Agency (USEPA). 1990. Rainbow trout early life stage toxicity test with 2,6-dichloro-4-nitrobenzeneamine. TSCA Section 4 Test Report. Washington, DC: Office of Toxic Substances, USEPA

United States Environmental Protection Agency (USEPA). 1991. Fish Chronic Toxicity Data Base. Duluth, MN: Environmental Research Laboratory (ERL), Office of Research and Development, USEPA, 6201 Congdon Boulevard, 55804; contact C.L. Russom (218) 720-5500.

Van Leeuwen CJ, Adema DMM and Hermens J. 1990. Quantitative structure-activity relationships for fish early life stage toxicity. Aquatic Toxicology 16:321-334.

**ANILINES**

9/1993

**LIST OF ANILINES USED TO DEVELOP THE FISH CHRONIC VALUE (ChV) SAR.**

CHEMICAL	ChV (mg/L)	Log K <sub>ow</sub>	Ref.
aniline	1.8	0.9	VL
aniline	0.569	0.9	D
4-chloroaniline	0.400	1.8	B
3-chloroaniline	1.0	1.9	VL
3,4-dichloroaniline	0.020	2.7	C
3,4-dichloroaniline	0.006	2.7	C
3,5-dichloroaniline	0.320	2.9	VL
2,6-dichloro-4- nitroaniline	0.016	3.0	EPA
2,4,5-trichloroaniline	0.056	3.7	VL
2,3,4,5-tetrachloroaniline	0.032	4.6	VL
pentachloroaniline	0.010	5.1	VL

EPA = USEPA (1990)

C = Call et al (1987)

D = USEPA (1991)

VL = Van Leeuwen et al (1990)

B = Bresch et al (1990)

**SAR**

**ANILINES**

**Organism:** Daphnid  
**Duration:** 16-d  
**Endpoint:** Chronic Value (Survival/Reproduction)

**Equation:**  $\text{Log ChV (mM/L)} = -3.12 - 0.36 \log K_{ow}$

**Statistics:**  $N = 3;; R^2 = 0.98$

**Maximum log  $K_{ow}$ :** 9.0  
**Maximum MW:** 1000.0

**Application:** This SAR may be used to estimate toxicity for anilines.

**Limitations:** N-substituted anilines are less toxic than predicted by this SAR; for these compounds use the daphnid ChV SAR for neutral organics.

If the log  $K_{ow}$  value is greater than 9.0, or if the compound is solid and the ChV exceeds the water solubility, no effects expected at saturation.

**References:** United States Environmental Protection Agency (USEPA). 1990. Daphnid Chronic Toxicity Tests with aniline and 2-chloroaniline. TSCA Sec. 4 Test Reports. Washington, DC: U S. Environmental Protection Agency, Office of Toxic Substances.

LIST OF ANILINES USED TO DEVELOP THE DAPHNID CHRONIC VALUE (ChV) SAR.

CHEMICAL	ChV (mg/L)	Log $K_{ow}$	Ref.
aniline	0.021	0.9	EPA
2-chloroaniline	0.034	1.9	EPA

EPA = USEPA (1990)

**ANILINES**  
9/1993

**SAR**

**ANILINES**

**Organism:** Green Algae  
**Duration:**  
**Endpoint:** Chronic Value (Growth)  
**Equation:**  $\text{Log ChV (mM/L)} = -0.411 - 0.588 \log K_{ow}$   
**Statistics:**  $N = 5; R^2 = 1.0$   
**Maximum log  $K_{ow}$ :** 9.0  
**Maximum MW:** 1000.0  
**Application:** This equation may be used to estimate toxicity for anilines.  
**Limitations:** N-substituted anilines are less toxic than predicted by this SAR; for these compounds use the neutral organics green algae ChV SAR.  
 If the log Kow value is greater than 9.0, or if the compound is solid and the ChV exceeds the water solubility, no effects expected at saturation.  
**References:** Sloof W, Canton JH, and Hermens JLM. 1983. Comparison of the susceptibility of 22 freshwater species to 15 chemical compounds. I. (Sub)Acute toxicity tests. Aquatic Toxicology 4:113-128.

LIST OF ANILINES USED TO DEVELOP THE GREEN ALGAE CHRONIC VALUE (ChV) SAR.

CHEMICAL	Log ChV (mg/L)	Log $K_{ow}$	Ref
aniline	11.0	0.9	S
aniline	8.0	0.9	S
aniline	16.0	0.9	S
aniline	10.0	0.9	S

S = Slooff et al (1987)

**ANILINES**

9/1993

**SAR**

**ANILINES**

**Organism:** Fish  
**Duration:** 14-d  
**Endpoint:** LC50 (Mortality)  
**Equation:**  $\text{Log LC50 (mM/L)} = 1.02 - 0.988 \log K_{ow}$   
**Statistics:**  $N = 17; R^2 = 0.783$

**Maximum log  $K_{ow}$ :** 5.0  
**Maximum MW:** 1000.0

**Application:** This SAR may be used to estimate toxicity for the following classes of compounds:

1. Anilines
2. Chloroanilines
3. Alkylanilines

**Limitations:** If the log  $K_{ow}$  value is greater than 5.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.

**References:** Hermans J, Leeuwangh P, and Musch A. 1984 Quantitative structure-activity relationships and mixture toxicity studies of chloro- and alkylanilines at an acute lethal toxicity level to the guppy, Poecilia reticulata. Ecotoxicology and Environmental Safety 8:388-394.

# ANILINES

9/1993

## LIST OF ANILINES USED TO DEVELOP THE FISH 14-d LC50 SAR.

CHEMICAL	Log LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
Aniline	125.0	1.03	H
2-Methylaniline	81.3	1.54	H
3-Methylaniline	36.3	1.54	H
4-Methylaniline	10.7	1.54	H
2-Chloroaniline	6.2	1.76	H
3-Chloroaniline	13.4	1.76	H
4-Chloroaniline	26.0	1.76	H
2-Ethylaniline	74.7	2.07	H
3-Ethylaniline	27.1	2.07	H
4-Ethylaniline	29.1	2.07	H
2,5-Dichloroaniline	1.65	2.42	H
2,4-Dichloroaniline	6.3	2.42	H
3,5-Dichloroaniline	3.9	2.42	H
3,4-Dichloroaniline	6.3	2.42	H
2,3,4-Trichloroaniline	1.4	3.17	H
2,4,5-Trichloroaniline	2.0	3.17	H
2,3,4,5-Tetrachloroaniline	0.36	3.92	H

H = Hermans et al (1984)

**AMINO ANILINES, META OR 1,3 SUBSTITUTED**  
9/1993

**SAR**

**ANILINES, AMINO, META OR 1,3-SUBSTITUTED**

**Organism:** Fish  
**Duration:** 96-h  
**Endpoint:** LC50 (Mortality)

**Equation:**  $\text{Log 96-h LC50 (mM/L)} = 0.978 - 0.740 \log K_{ow}$

**Statistics:**  $N = 2; R^2 = 1.0$

**Maximum log  $K_{ow}$ :** 7.0  
**Maximum MW:** 1000.0

**Application:** This equation may be used to estimate toxicity for meta or 1,3 substituted amino anilines.

**Limitations:** If the log  $K_{ow}$  value is greater than 7.0, no effects expected at saturation. duration

**References:** United States Environmental Protection Agency (USEPA). 1991. OTS TSCA Section 4 database. Washington, DC: USEPA, Office of Toxic Substances.

LIST OF META SUBSTITUTED AMINO ANILINES USED TO DEVELOP THE FISH 96-h LC50 SAR.

CHEMICAL	96-h LC50 (mg/L)	Log $K_{ow}$	Ref.
m-Phenylenediamine	1618	-0.3	EPA

EPA = USEPA (1991)

## ANILINES, AMINO, META OR 1,3-SUBSTITUTED

### SAR ANILINES, AMINO, META OR 1,3-SUBSTITUTED

**Organism:** Daphnid  
**Duration:** 48-h  
**Endpoint:** LC50 (Mortality)

**Equation:**  $\text{Log 48-h LC50 (mM/L)} = -1.44 - 0.466 \log K_{ow}$

**Statistics:**  $N = 2; R^2 = 1.0$

**Maximum log  $K_{ow}$ :** 7.0  
**Maximum MW:** 1000.0

**Application:** This equation may be used to estimate toxicity for meta or 1,3 substituted amino anilines.

**Limitations:** If the log  $K_{ow}$  value is greater than 7.0, no effects expected at saturation.

**References:** United States Environmental Protection Agency (USEPA). 1991. OTS TSCA Section 4 database. Washington, DC: USEPA, Office of Toxic Substances

#### LIST OF META SUBSTITUTED AMINO ANILINES USED TO DEVELOP THE DAPHNID 48-h LC50 SAR.

CHEMICAL	48-h LC50 (mg/L)	Log $K_{ow}$	Ref.
m-Phenylenediamine	5.9	-0.3	EPA

EPA = USEPA (1991)

**ANILINES, AMINO, META OR 1,3-SUBSTITUTED**

## ANILINES, AMINO, META OR 1,3-SUBSTITUTED

### SAR

### ANILINES, AMINO, META OR 1,3-SUBSTITUTED

**Organism:** Green Algae

**Duration:** 96-h

**Endpoint:** EC50

**Equation:**  $\text{Log 96-h EC50 (mM/L)} = -1.8 - 0.333 \log K_{ow}$

**Statistics:**  $N = 2, R^2 = 1.0$

**Maximum log  $K_{ow}$ :** 6.0

**Maximum MW:** 1000.0

**Application:** This equation may be used to estimate toxicity for meta or 1,3 substituted amino anilines.

**Limitations:** If the log  $K_{ow}$  value is greater than 6.0, no effects expected at saturation.

**References:** United States Environmental Protection Agency (USEPA). 1991. OTS TSCA Section 4 database. Washington, DC: USEPA, Office of Toxic Substances.

### LIST OF META SUBSTITUTED AMINO ANILINES USED TO DEVELOP THE GREEN ALGAE 96-h EC50 SAR.

CHEMICAL	96-h EC50 (mg/L)	Log $K_{ow}$	Ref.
m-Phenylenediamine	2.4	-0.3	EPA

EPA = USEPA (1991)



**ANILINES, AMINO, META OR 1,3-SUBSTITUTED**

9/1993

**SAR****ANILINES, AMINO, META OR 1,3-SUBSTITUTED**

**Organism:** Daphnid  
**Duration:** 16-d  
**Endpoint:** Chronic Value

**Equation:**  $\text{Log ChV (mM/L)} = -3.29 - 0.301 \log K_{ow}$

**Statistics:**  $N = 2; R^2 = 1.0$

**Maximum log  $K_{ow}$ :** 8.0  
**Maximum MW:** 1000.0

**Application:** This equation may be used to estimate toxicity for meta or 1,3 substituted amino anilines.

**Limitations:** If the log  $K_{ow}$  value is greater than 8.0, or if the compound is solid and the ChV exceeds the water solubility, no effects expected at saturation.

**References:** United States Environmental Protection Agency (USEPA). 1991. OTS TSCA Section 4 database. Washington, DC. USEPA, Office of Toxic Substances.

**LIST OF META SUBSTITUTED AMINO ANILINES USED TO DEVELOP THE DAPHNID ChV SAR.**

CHEMICAL	ChV (mg/L)	Log $K_{ow}$	Ref.
m-Phenylenediamine	0.070	-0.3	EPA

EPA = USEPA (1991)



**ANILINES, AMINO, ORTHO OR 1,2-SUBSTITUTED**  
9/1993

**SAR**

**ANILINES, AMINO, ORTHO OR 1,2-SUBSTITUTED**

<b>Organism:</b>	Fish
<b>Duration:</b>	96-h
<b>Endpoint:</b>	LC50 (Mortality)
<b>Equation:</b>	$\text{Log 96-h LC50 (mM/L)} = -0.547 - 0.522 \log K_{ow}$
<b>Statistics:</b>	$N = 2; R^2 = 1.0$
<b>Maximum log <math>K_{ow}</math>:</b>	7.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This equation may be used to estimate toxicity for ortho or 1,2 substituted amino anilines
<b>Limitations:</b>	If the log $K_{ow}$ value is greater than 7.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.
<b>References:</b>	United States Environmental Protection Agency (USEPA). 1991. OTS, TSCA Section 4 database. Washington, DC: USEPA, Office of Toxic Substances.

**LIST OF ORTHO SUBSTITUTED AMINO ANILINES USED TO DEVELOP THE FISH 96-h LC50 SAR**

CHEMICAL	96-h LC50 (mg/L)	Log $K_{ow}$	Ref.
o-Phenylenediamine	44	-0.3	EPA

EPA = USEPA (1991)



**ANILINES, AMINO, ORTHO OR 1,2-SUBSTITUTED**

9/1993

**SAR****ANILINES, AMINO, ORTHO OR 1,2-SUBSTITUTED****Organism:**

Daphnid

**Duration:**

48-h

**Endpoint:**

LC50 (Mortality)

**Equation:**
$$\text{Log 48-h LC50 (mM/L)} = -2.21 - 0.356 \log K_{ow}$$
**Statistics:**
$$N = 2, R^2 = 1.0$$
**Maximum log  $K_{ow}$ :**

7.0

**Maximum MW:**

1000.0

**Application:**

This equation may be used to estimate toxicity for ortho or 1,2 substituted amino anilines.

**Limitations:**

If the log  $K_{ow}$  value is greater than 7.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation

**References:**

United States Environmental Protection Agency (USEPA). 1991 OTS TSCA Section 4 database. Washington, DC: USEPA, Office of Toxic Substances.

LIST OF ORTHO SUBSTITUTED AMINO ANILINES USED TO DEVELOP  
THE DAPHNID 48-h LC50 SAR.

CHEMICAL	48-h LC50 (mg/L)	Log $K_{ow}$	Ref.
o-Phenylenediamine	0.880	-0.3	EPA

EPA = USEPA (1991)



**ANILINES, AMINO, ORTHO OR 1,2-SUBSTITUTED**

9/1993

**SAR****ANILINES, AMINO, ORTHO OR 1,2-SUBSTITUTED****Organism:** Green Algae**Duration:** 96-h**Endpoint:** EC50**Equation:**  $\text{Log 96-h EC50 (mM/L)} = -2.848 - 0.159 \log K_{ow}$ **Statistics:**  $N = 2; R^2 = 1.0$ **Maximum log  $K_{ow}$ :** 6.0**Maximum MW:** 1000.0**Application:** This equation may be used to estimate toxicity for ortho or 1,2 substituted amino anilines.**Limitations:** If the log  $K_{ow}$  value is greater than 6.0, or if the compound is solid and the EC50 exceeds the water solubility, no effects expected at saturation.**References:** United States Environmental Protection Agency (USEPA). 1991. OTS TSCA Section 4 database. Washington, DC: USEPA, Office of Toxic Substances.**LIST OF ORTHO SUBSTITUTED AMINO ANILINES USED TO DEVELOP  
THE GREEN ALGAE 96-h EC50 SAR.**

CHEMICAL	96-h EC50 (mg/L)	Log $K_{ow}$	Ref.
o-Phenylenediamine	0.160	-0.3	EPA

EPA = USEPA (1991)



**ANILINES, AMINO, PARA OR 1,4-SUBSTITUTED**

9/1993

**SAR****ANILINES, AMINO, PARA OR 1,4-SUBSTITUTED**

<b>Organism:</b>	Fish
<b>Duration:</b>	96-h
<b>Endpoint:</b>	LC50 (Mortality)
<b>Equation:</b>	$\text{Log 96-h LC50 (mM/L)} = -3.337 - 0.123 \log K_{ow}$
<b>Statistics:</b>	$N = 2, R^2 = 1.0$
<b>Maximum log <math>K_{ow}</math>:</b>	7.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This equation may be used to estimate toxicity for para or 1,4 substituted amino anilines
<b>Limitations:</b>	If the log $K_{ow}$ value is greater than 7.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.
<b>References:</b>	United States Environmental Protection Agency (USEPA). 1991. OTS TSCA Section 4 database. Washington, DC: USEPA, Office of Toxic Substances

**LIST OF PARA SUBSTITUTED AMINO ANILINES USED TO DEVELOP THE FISH 96-h LC50 SAR.**

CHEMICAL	96-h LC50 (mg/L)	Log $K_{ow}$	Ref.
p-Phenylenediamine	0.060	-0.3	EPA

EPA = USEPA (1991)

**ANILINES, AMINO, PARA OR 1,4-SUBSTITUTED**

9/1993

**ANILINES, AMINO, PARA OR 1,4-SUBSTITUTED**  
9/1993

<b>SAR</b>	<b>ANILINES, AMINO, PARA OR 1,4-SUBSTITUTED</b>
<b>Organism:</b>	Daphnid
<b>Duration:</b>	48-h
<b>Endpoint:</b>	LC50 (Mortality)
<b>Equation:</b>	$\text{Log 48-h LC50 (mg/L)} = -2.686 - 0.288 \log K_{ow}$
<b>Statistics:</b>	$N = 2, R^2 = 1.0$
<b>Maximum log <math>K_{ow}</math>:</b>	7.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This equation may be used to estimate toxicity for para or 1,4 substituted amino anilines.
<b>Limitations:</b>	If the log $K_{ow}$ is greater than 7.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.
<b>References:</b>	United States Environmental Protection Agency (USEPA). 1991. OTS TSCA Section 4 database. Washington, DC: USEPA, Office of Toxic Substances.

**LIST OF PARA SUBSTITUTED AMINO ANILINES USED TO DEVELOP THE DAPHNID 48-h LC50 SAR.**

CHEMICAL	48-h LC50 (mg/L)	Log $K_{ow}$	Ref.
p-Phenylenediamine	0.280	-0.3	EPA

EPA = USEPA (1991)

**ANILINES, AMINO, PARA OR 1,4-SUBSTITUTED**

9/1993

**ANILINES, AMINO, PARA OR 1,4-SUBSTITUTED**

9/1993

**SAR****ANILINES, AMINO, PARA OR 1,4-SUBSTITUTED****Organism:** Green Algae**Duration:** 96-h**Endpoint:** EC50**Equation:**  $\text{Log 96-h EC50 (mM/L)} = -2.657 - 0.190 \log K_{ow}$ **Statistics:**  $N = 2, R^2 = 1.0$ **Maximum log  $K_{ow}$ :** 6.0**Maximum MW:** 1000.0**Application:** This equation may be used to estimate toxicity for para or 1,4 substituted amino anilines.**Limitations:** If the log  $K_{ow}$  value is greater than 6.0, or if the compound is solid and the EC50 exceeds the water solubility, no effects expected at saturation.**References:** United States Environmental Protection Agency (USEPA). 1991. OTS TSCA Section 4 database. Washington, DC. USEPA, Office of Toxic Substances.**LIST OF PARA SUBSTITUTED AMINO ANILINES USED TO DEVELOP  
THE GREEN ALGAE 96-h EC50 SAR.**

CHEMICAL	96-h EC50 (mg/L)	Log $K_{ow}$	Ref
p-Phenylenediamine	0.28	-0.3	EPA

EPA = USEPA (1991)

**SAR**

**ANILINES, DINITRO**

**Organism:** Fish  
**Duration:** 96-h  
**Endpoint:** LC50 (Mortality)  
**Equation:**  $\text{Log 96-h LC50 (mM/L)} = -0.027 - 0.596 \log K_{ow}$   
**Statistics:**  $N = 2; R^2 = 1.0$   
**Maximum log  $K_{ow}$ :** 7.0  
**Maximum MW:** 1000.0  
**Application:** This SAR may be used to estimate toxicity for dinitroanilines and other polynitroanilines.  
**Limitations:** If the log  $K_{ow}$  value is greater than 7.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.  
**References:** Veith GD and Broderius SJ 1987. Structure-toxicity relationships for industrial chemicals causing type (II) narcosis syndrome. In: Kaiser KLE (ed ). QSAR in Environmental Toxicology-II. Boston, MA: D. Reidel Pub. Co., pp. 385-391.

LIST OF DINITROANILINES USED TO DEVELOP THE FISH 96-h LC50 SAR

CHEMICAL	96-hour LC50 (mg/L)	Log $K_{ow}$	Ref.
2,4-dinitroaniline	15.5	1.8	VB

VB = Veith and Broderius (1987)

**ANILINES, DINITRO**  
9/1993

**SAR**

**ANILINES, DINITRO**

**Organism:** Daphnid  
**Duration:** 48-h  
**Endpoint:** LC50 (Mortality)

**Equation:**  $\log 48\text{-h LC50 (mM/L)} = -0.296 - 0.558 \log K_{ow}$

**Statistics:**  $N = 2; R^2 = 1.0$

**Maximum log  $K_{ow}$ :** 7.0  
**Maximum MW:** 1000.0

**Application:** This SAR may be used to estimate toxicity for dinitroanilines and other polynitroanilines.

**Limitations:** If the log  $K_{ow}$  value is greater than 7.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.

**References:** Kuhn R, Pattard M, Pernak K-D, and Winter A 1989 Results of the harmful effects of selected water pollutants (anilines, phenols, aliphatic compounds) to Daphnia magna. Water Research 23:495-499.

LIST OF DINITROANILINES USED TO DEVELOP THE DAPHNID 48-h LC50.

CHEMICAL	48-h LC50 (mg/L)	Log $K_{ow}$	Ref.
2,4-dinitroaniline	9.6	1.8	K

Kuhn = Kuhn et al (1989)

**ANILINES, DINITRO**  
9/1993

## SAR ANILINES, DINITRO

<b>Organism:</b>	Fish
<b>Duration:</b>	32-d
<b>Endpoint:</b>	Chronic Value (Survival/Growth)

**Equation:**  $\text{Log ChV (mM/L)} = -0.91 - 0.661 \log K_{ow}$

**Statistics:** N = 2;  $R^2$  = 1.0

Maximum log K<sub>ow</sub>: 8.0  
Maximum MW: 1000.0

**Application:** This SAR may be used to estimate toxicity for dinitroanilines and other polynitroanilines.

**Limitations:** If the log  $K_{ow}$  value is greater than 8.0, or if the compound is solid and the ChV exceeds the water solubility, no effects expected at saturation.

**References:** United States Environmental Protection Agency (USEPA). 1991. Fish Chronic Toxicity Data Base. Duluth, MN: Environmental Research Laboratory (ERL), Office of Research and Development, USEPA, 6201 Congdon Boulevard, 55804; contact C.L. Russom (218) 720-5500.

LIST OF DINITROANILINES USED TO DEVELOP THE FISH CHRONIC (ChV)-SAR.

CHEMICAL	ChV (mg/L)	Log K <sub>ow</sub>	Ref.
2,4-dinitroaniline	1.41	1.8	D

D = USEPA (1991)

**ANILINES, DINITRO**  
9/1993

**SAR**

**AZIRIDINES**

<b>Organism:</b>	Fish
<b>Duration:</b>	Acute
<b>Endpoint:</b>	LC50 (Mortality)
<b>Equation:</b>	$\text{Log LC50 (mM/L)} = -1.65 - 0.364 \log K_{ow}$
<b>Statistics:</b>	$N = 2; R^2 = 1.0$
<b>Maximum log <math>K_{ow}</math>:</b>	7.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This equation may be used to estimate toxicity for aziridines.
<b>Limitations:</b>	If the log $K_{ow}$ value is greater than 7.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation
<b>References:</b>	Juhnke I and Luedemann D. 1978. Results of the investigation of 200 chemical compounds for acute toxicity with the Golden Orfe test. Z.F. Wasser-Und Abwasser-Forschung 11 161-164. Translation by SCITRAN (Scientific Translation Service), Santa Barbara, CA 93108.

LIST OF AZIRIDINES USED TO DEVELOP THE FISH ACUTE LC50 SAR.

CHEMICAL	LC50 (mg/L)	Log $K_{ow}$	Ref.
Aziridine	2.4	-1.1	J

J = Juhnke and Luedemann (1978)

## AZIRIDINES

9/1993

**SAR**

**AZIRIDINES**

<b>Organism:</b>	Daphnid
<b>Duration:</b>	48-h
<b>Endpoint:</b>	LC50 (Mortality)
<b>Equation:</b>	$\text{Log 48-h LC50 (mg/L)} = -1.062 - 0.52 \log K_{ow}$
<b>Statistics:</b>	$N = 2; R^2 = 1.0$
<b>Maximum log <math>K_{ow}</math>:</b>	7.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This equation may be used to estimate toxicity for aziridines.
<b>Limitations:</b>	If the log $K_{ow}$ value is greater than 7.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.
<b>References:</b>	Bringmann G and Kuhn R. 1977. Results of the damaging effect of water pollutants on <u>Daphnia magna</u> . Z. Wasser Abwasser Forsch. 10(5):161-166

LIST OF AZIRIDINES USED TO DEVELOP THE DAPHNID 48-h LC50 SAR.

CHEMICAL	LC50 (mg/L)	Log $K_{ow}$	Ref
Aziridine	14.0	-1.1	B

B = Bringmann and Kuhn (1977)

**AZIRIDINES**  
9/1993

<b>SAR</b>	<b>AZIRIDINES</b>
<b>Organism:</b>	Green Algae
<b>Duration:</b>	7-d
<b>Endpoint:</b>	Chronic Value
<b>Equation:</b>	$\text{Log ChV (mM/L)} = -2.4 - 0.33 \log K_{ow}$
<b>Statistics:</b>	$N = 2; R^2 = 1.0$
<b>Maximum log <math>K_{ow}</math>:</b>	8.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This equation may be used to estimate toxicity for aziridines
<b>Limitations:</b>	If the log $K_{ow}$ value is greater than 8.0, or if the compound is solid and the ChV exceeds the water solubility, no effects expected at saturation
<b>References:</b>	Bringmann G and Kuhn R 1980. Comparison of the toxicity thresholds of water pollutants to bacteria, algae, and protozoa in the cell multiplication inhibition test. Water Research 14(3).231-241.

LIST OF AZIRIDINES USED TO DEVELOP THE GREEN ALGAE CHV SAR.

CHEMICAL	ChV (mg/L)	Log $K_{ow}$	Ref.
Aziridine	0.370	-1.1	B

B = Bringmann and Kuhn (1980)

**AZIRIDINES**  
9/1993

**SAR****BENZENES, DINITRO**

<b>Organism:</b>	Fish
<b>Duration:</b>	96-h
<b>Endpoint:</b>	LC50 (Mortality)
<b>Equation:</b>	$\text{Log 96-h LC50 (mM/L)} = -1.867 - 0.333 \log K_{ow}$
<b>Statistics:</b>	$N = 2; R^2 = 1.0$
<b>Maximum log <math>K_{ow}</math>:</b>	7.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This SAR may be used to estimate toxicity for dinitrobenzenes and other polynitrobenzenes
<b>Limitations:</b>	If the log $K_{ow}$ value is greater than 7.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation
<b>References:</b>	<p>Veith GD and Broderius SJ. 1987. Structure-toxicity relationships for industrial chemicals causing type (II) narcosis syndrome. In: Kaiser KLE (ed.). QSAR in Environmental Toxicology-II. Boston, MA: D. Reidel Pub. Co., pp. 385-391.</p> <p>United States Environmental Protection Agency (USEPA). 1991. OTS PMN ECOTOX. Washington, DC: USEPA, Office of Toxic Substances.</p>

## LIST OF DINITROBENZENES USED TO DEVELOP THE FISH 96-h LC50 SAR:

CHEMICAL	96-h LC50 (mg/L)	Log $K_{ow}$	Ref
1,3-dinitrobenzene	0.71	1.5	VB
Chemical identity CBI	0.013	3.2	EPA

VB = Veith and Broderius (1987)

EPA = USEPA (1991), chemical identity is Confidential Business Information under TSCA.

**BENZENES, DINITRO**  
9/1993

**SAR**

**BENZENES, DINITRO**

<b>Organism:</b>	Daphnid
<b>Duration:</b>	48-h
<b>Endpoint:</b>	LC50 (Mortality)
<b>Equation:</b>	$\text{Log 48-h LC50 (mM/L)} = -0.325 - 0.634 \log K_{ow}$
<b>Statistics:</b>	$N = 3; R^2 = 0.86$
<b>Maximum log <math>K_{ow}</math>:</b>	7.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This SAR may be used to estimate toxicity for dinitrobenzenes or other polynitrobenzenes
<b>Limitations:</b>	If the log $K_{ow}$ value is greater than 7.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.
<b>References:</b>	<p>Hermens J, Canton J, Janssen P, and DeJong R. 1984. Quantitative structure-activity relationships and toxicity studies of mixtures of chemicals with anaesthetic potency: Acute lethal and sublethal toxicity to <u>Daphnia magna</u>. <u>Aquatic Toxicology</u> 5:143-154</p> <p>LeBlanc. 1980. Acute toxicity of priority pollutants to water flea (<u>Daphnia magna</u>). <u>Bulletin of Environmental Contamination and Toxicology</u>. 24: 684-691.</p> <p>United States Environmental Protection Agency (USEPA). 1991. OTS PMN ECOTOX. Washington, DC: USEPA, Office of Toxic Substances.</p>

LIST OF DINITROBENZENES USED TO DEVELOP THE DAPHNID 48-h LC50 SAR

CHEMICAL	48-h LC50 (mg/L)	Log $K_{ow}$	Ref.
1,3-dinitrobenzene	43.0	1.6	H
2,3-dinitrotoluene	0.66	2.0	LB
Chemical identity CBI	0.012	3.2	EPA

LB = LeBlanc (1987)

H = Hermens et al (1984)

EPA = USEPA (1991), chemical identities are Confidential Business Information under TSCA.

**BENZENES, DINITRO**  
9/1993

<b>SAR</b>	<b>BENZENES, DINITRO</b>
<b>Organism:</b>	Fish
<b>Duration:</b>	32-d
<b>Endpoint:</b>	Chronic Value (Survival/Growth)
<b>Equation:</b>	$\text{Log ChV (mM/L)} = -3.0 - 0.40 \log K_{ow}$
<b>Statistics:</b>	$N = 2; R^2 = 1.0$
<b>Maximum log <math>K_{ow}</math>:</b>	8.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This SAR may be used to estimate toxicity for dinitrobenzenes or other polynitrobenzenes.
<b>Limitations:</b>	If the log $K_{ow}$ value is greater than 8.0, or if the compound is solid and the ChV exceeds the water solubility, no effects are expected at saturation.
<b>References:</b>	United States Environmental Protection Agency (USEPA) 1991. Fish Chronic Toxicity Data Base. Duluth, MN. Environmental Research Laboratory (ERL), Office of Research and Development, USEPA, 6201 Congdon Boulevard, 55804; contact C.L. Russom (218) 720-5500.

LIST OF DINITROBENZENES USED TO DEVELOP THE FISH CHRONIC VALUE (ChV) SAR.

CHEMICAL	ChV (mg/L)	Log $K_{ow}$	Ref.
1,3-dichloro-4,6-dinitro benzene	0.023	2.5	D

D = USEPA (1991)

**BENZENES, DINITRO**  
9/1993

<b>SAR</b>	<b>BENZENES, DINITRO</b>
<b>Organism:</b>	Daphnid
<b>Duration:</b>	16-d
<b>Endpoint:</b>	Chronic Value (Survival/Reproduction)
<b>Equation:</b>	$\text{Log ChV (mM/L)} = -0.7 - 0.625 \log K_{ow}$
<b>Statistics:</b>	$N = 2; R^2 = 1.0$
<b>Maximum log <math>K_{ow}</math>:</b>	8.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This SAR may be used to estimate toxicity for dinitrobenzenes or other polynitrobenzenes.
<b>Limitations:</b>	If the log $K_{ow}$ value is greater than 8.0, or is the compound is solid and the ChV exceeds the water solubility, no effects are expected at saturation.
<b>References:</b>	Hermens J, Canton H, Janssen P, and DeJong R. 1984. Quantitative structure-activity relationships and toxicity studies of mixtures of chemicals with anaesthetic potency: Acute lethal and sublethal toxicity to <u>Daphnia magna</u> Aquatic Toxicology 5.143-154.

LIST OF DINITROBENZENES USED TO DEVELOP THE DAPHNID CHRONIC VALUE (ChV) SAR.

CHEMICAL	ChV (mg/L)	Log $K_{ow}$	Ref.
1,3-dinitrotoluene	3.2	1.6	H

H = Hermens et al (1984)

**BENZENES, DINITRO**  
9/1993

<b>SAR</b>	<b>BENZOTRIZOLES</b>
<b>Organism:</b>	Fish
<b>Duration:</b>	96-h
<b>Endpoint:</b>	LC50 (Mortality)
<b>Equation:</b>	$\text{Log LC50 (mM/L)} = 0.366 - 0.587 \log K_{ow}$
<b>Statistics:</b>	$N = 2; R^2 = 1.00$
<b>Maximum log <math>K_{ow}</math>:</b>	5.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	<p>This SAR may be used to estimate the toxicity of substituted benzotriazoles with substitution on the 5th position. Toxicity estimates for substituted benzotriazoles with substitutions on the triazole ring may not be valid with this SAR.</p> <p>This SAR may be used for substituted benzotriazoles with substitutions on the 3rd, 4th or 6th positions (other benzo positions).</p>
<b>Limitations:</b>	If the log $K_{ow}$ value is greater than 5.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.
<b>References:</b>	<p>Nabholz JV. 1987. Generic review of various benzotriazoles. Washington, DC: Environmental Effects Branch, Health and Environmental Review Division (TS-796), Office of Toxic Substances, United States Environmental Protection Agency.</p>

LIST OF BENZOTRIAZOLES USED TO DEVELOP THE FISH 96-h LC50 SAR.

CHEMICAL	LC50 (mg/L)	Log $K_{ow}$	Ref.
Benzotriazole	39.0	1.45	N
5-Butylbenzotriazole	2.8	3.68	N

N = Nabholz (1987)

**BENZOTRIAZOLES**  
7/1988

**SAR**

**BENZOTRIAZOLES**

**Organism:** Daphnid  
**Duration:** 48-h  
**Endpoint:** LC50 (Mortality)

**Equation:** To determine the acute toxicity of benzotriazoles to daphnids use the neutral organic daphnid 48-h LC50 SAR.

**Statistics:**

**Maximum log  $K_{ow}$ :** 5.0  
**Maximum MW:** 1000.0

**Application:** The neutral organic SAR may be used to estimate the toxicity of substituted benzotriazoles with substitution on the 5th position, log  $K_{ow}$  values of less than 5.0, and molecular weights less than 1000. Toxicity estimates for substituted benzotriazoles with substitutions on the triazole ring may not be valid with this SAR.

This SAR may be used for substituted benzotriazoles with substitutions on the 3rd, 4th or 6th positions (other benzo positions).

**Limitations:** If the log  $K_{ow}$  value is greater than 5.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.

**References:** Nabholz JV. 1987. Generic review of various benzotriazoles. Washington, DC: Environmental Effects Branch, Health and Environmental Review Division (TS-796), Office of Toxic Substances, United States Environmental Protection Agency.

LIST OF BENZOTRIAZOLES USED TO DEVELOP THE DAPHNID 48-h LC50 SAR.

CHEMICAL	LC50 (mg/L)	Log $K_{ow}$	Ref.
Benzotriazole	141.6	1.45	N
5-Butylbenzotriazole	10.7	3.68	N

N = Nabholz (1987)

## **BENZOTRIAZOLES**

7/1988

**SAR**

**BENZOTRIAZOLES**

**Organism:** Green Algae  
**Duration:** 96-h  
**Endpoint:** EC50 and EC10 (Growth)

**Equation:**  $\text{Log EC50 (mM/L)} = 0.061 - 0.573 \log K_{ow}$

The 96-h EC10 may be determined by:

$$\text{EC10} = \text{EC50}/8$$

**Statistics:**  $N = 2; R^2 = 1.00$

**Maximum log  $K_{ow}$ :** 8.0  
**Maximum MW:** 1000.0

**Application:** This SAR may be used to estimate the toxicity of substituted benzotriazoles with substitution on the 5th position. Toxicity estimates for substituted benzotriazoles with substitutions on the triazole ring may not be valid with this SAR.

This SAR may be used for substituted benzotriazoles with substitutions on the 3rd, 4th or 6th positions (other benzo positions).

**Limitations:** If the log  $K_{ow}$  value is greater than 8.0, or if the compound is solid and the EC50 or EC10 exceeds the water solubility, no effects expected at saturation.

**References:** Nabholz JV. 1987. Generic review of various benzotriazoles. Washington, DC: Environmental Effects Branch, Health and Environmental Review Division (TS-796), Office of Toxic Substances, United States Environmental Protection Agency.

LIST OF BENZOTRIAZOLES USED TO DEVELOP THE GREEN ALGAE 96-h EC50 AND EC10 SARs

CHEMICAL	EC50 (mg/L)	EC10 (mg/L)	Log $K_{ow}$	Ref.
Benzotriazole	15.4	1.75	1.45	N
5-Butylbenzotriazole	1.18	0.16	3.68	N

N = Nabholz (1987)

## **BENZOTRIAZOLES**

7/1988

**SAR**

**CARBAMATES**

**Organism:**

Sea Urchin

**Duration:**

48-h

**Endpoint:**

NEC (Early Development)

**Equation:**

$\text{Log NEC (mM/L)} = 0.51 - 0.72 \log K_{ow}$

**Statistics:**

$N = 35; R^2 = 0.62$

**Maximum log  $K_{ow}$ :**

4.5

**Maximum MW:**

1000.0

**Application:**

This SAR may be used to estimate toxicity for carbamates and the following classes of carbamates:

1. Alkyl esters of carbamic acid
2. N-alkyl or aryl substitutes on ethyl carbamate
3. Bis(ethylcarbamates) joined at -NRN- by alkyl or aryl groups
4. Bis- and tris- carbamates esterified on a single phenyl ring
5. Thiocarbamates

This SAR may be used for other similar substituted carbamates with log  $K_{ow}$  values less than 4.5 and molecular weights less than 1000.

**Limitations:**

The following classes of carbamates are more toxic than predicted by this SAR:

1. Meta-phenylene bis(ethyl carbamates) - 200 X
2. N-methyl-ortho phenyl biscarbamates - 1000 X
3. N-methyl-para phenyl biscarbamates - 400 X
4. N,N-dimethyl-1,2,3-phenyl triscarbamates - 400 X

If the log  $K_{ow}$  value is greater than 4.5, or if the compound is solid and the NEC exceeds the water solubility, no effects expected at saturation

**References:**

Cornman I. 1950. Inhibition of sea-urchin egg cleavage by a series of substituted carbamates. Journal of the National Cancer Institute 50:1123-1138.

# CARBAMATES

7/1988

## LIST OF CARBAMATES USED TO DEVELOP THE SEA URCHIN 48-h NEC SAR.

CHEMICAL	48-h NEC (mg/L)	Log K <sub>ow</sub>	Ref.
CARBAMATES USED FOR THIS SAR			
Methyl carbamate	2000.0	-0.70	C
Ethyl carbamate	999.0	-0.18	C
1,2-Hydrazine di(ethylcarboxylate)	2000.0	-0.11	C
1,2-Hydrazine di(ethylcarboxylate)	1000.0	-0.11	C
N-methyl-ethylcarbamate	10.3	0.37	C
N,N-dimethyl-ethylcarbamate	994.0	0.42	C
Propylene bis(ethylcarbamate)	998.0	0.58	C
1,4-Phenylene bis(N,N-dimethyl carbamate	501.0	0.88	C
1,4-Phenylene bis(N,N-dimethyl carbamate	101.0	0.88	C
N-ethyl-ethylcarbamate	99.0	0.90	C
Ethylidene bis(ethylcarbamate)	100.0	0.97	C
Ethylene bis(ethylcarbamate)	100.0	0.98	C
Tetramethylene bis(ethylcarbamate)	998.0	0.58	C
N-isopropyl-ethylcarbamate	9.2	1.21	C
3-Methylbutyl carbamate	10.5	1.28	C
Cyclohexyl carbamate	1.43	1.33	C
N,N-propyl-ethylcarbamate	97.0	1.43	C
N,N-diethyl-ethylcarbamate	95.7	1.48	C
N,N-cyclopentamethylene- ethylcarbamate	39.1	1.61	C
N,N-diethyl ethylcarbamodithioate	<41.0	1.68	C
N,N-butyl-ethylcarbamate	8.7	1.96	C
1,3-Phenylene bis(N,N-dimethyl carbamate	<39.0	2.09	C
N,N-di-isopropyl-ethylcarbamate	92.0	2.09	C
N-ethyl ethylcarbamothioate	<39.0	2.09	C
Para-xylene bis(ethylcarbamate)	19.6	2.14	C
Hexamethylene bis(ethylcarbamate)	502.0	2.16	C
Hexamethylene bis(ethylcarbamate)	99.0	2.16	C
N-phenyl-ethylcarbamate	1.0	2.29	C
N-cyclohexyl-ethylcarbamate	1.7	2.40	C
Ortho-phenylene bis (ethylcarbamate)	20.2	2.44	C
N,N-di-n-propyl-ethylcarbamate	9.4	2.53	C
N,N-di-n-butyl-ethylcarbamate	10.0	3.59	C
N,N-diphenyl-ethylcarbamate	9.6	NC	C
N-decyl carbamate	1.0	4.06	C
N-n-octyl-ethylcarbamate	1.0	4.07	C
N-2-fluorene-ethylcarbamate	<0.10	4.34	C
2,7-fluorene-bis(ethylcarbamate)	*	4.52	C

CONTINUED.

CHEMICAL	48-h NEC (mg/L)	Log K <sub>ow</sub>	Ref.
CARBAMATES USED FOR THIS SAR			
n-Dodecyl carbamate	*	5.12	C
N-n-decyl-ethylcarbamate	*	5.13	C
CARBAMATES WITH EXCESSIVE TOXICITY			
1,2-Phenylene bis(N-methyl carbamate	0.9	-0.11	C
1,3-Phenylene bis(N-methyl carbamate	0.9	-0.11	C
1,4-Phenylene bis(n-methyl carbamate	0.9	0.45	C
1,2,3-Phenylene tris(N,N-dimethyl carbamate	10.2	2.44	C

\* No effects in a saturated solution.

C = Cornman (1950)

**CARBAMATES, DITHIO**  
**9/1993**

**SAR**

**CARBAMATES, DIOTHIO**

Includes N,N-dialkyldithiocarbamates and ethylenebisdithiocarbamates and their metal salts which include but are not limited to Zn, Na, Fe, Mn, Cu, Pb, Hg, Ag, and Se. The SARs for the dithiocarbamates and their degradation products are sigmoidal with acute and chronic toxicity increasing with increasing Kow. The sigmoidal relationship between Kow and toxicity is very poor statistically. Consequently, toxicity predictions must be made using either the closest analog or averaging data for the two closest analogs which bracket the dithiocarbamate under question.

**CARBAMATES, DITHIO**  
**9/1993**

**SAR**

**CROWN ETHERS**

Use SAR for **NEUTRAL ORGANICS** for fish and daphnids; some should show excess toxicity toward green algae due to over-chelation of nutrient elements; each crown ether chelates a different element; the type of element chelated by a crown ether has to be matched up with a nutrient element needed by algae, e.g, Fe, Ca, Mg. There are no test data to show that crown ethers do in fact overchelate nutrient elements in the algal toxicity test. Conclusions about crown ethers are based on extrapolations of theory.

**CROWN ETHERS**  
**9/1993**

**SAR DIAZONIUMS, AROMATIC**

<b>Organism:</b>	Fish
<b>Duration:</b>	96-h
<b>Endpoint:</b>	LC50 (Mortality)
<b>Equation:</b>	$\text{Log 96-h LC50 (mM/L)} = -2.456 - 0.331 \log K_{ow}$
<b>Statistics:</b>	$N = 3; R^2 = 0.98$
<b>Maximum log <math>K_{ow}</math>:</b>	8.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This equation may be used to estimate toxicity for aromatic diazoniums
<b>Limitations:</b>	If the log $K_{ow}$ value is greater than 8.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation
<b>References:</b>	United States Environmental Protection Agency (USEPA). 1991. OTS TSCA 8(e) database Washington, DC: USEPA, Office of Toxic Substances

## LIST OF AROMATIC DIAZONIUMS USED TO DEVELOP THE FISH 96-h LC50 SAR

CHEMICAL	96-h LC50 (mg/L)	Log $K_{ow}$	Ref
4-Dimethylamino benzene diazonium	0.150	2.1	EPA
4-Dimethylamino benzene diazonium	0.330	2.1	EPA

EPA = USEPA (1991).

## DIAZONIUMS, AROMATIC

9/1993

**SAR**

**EPOXIDES, MONO**

**Organism:**

Fish

**Duration:**

96-h

**Endpoint:**

LC50 (Mortality)

**Equation:**

$\text{Log 96-h LC50 (mM/L)} = -0.290 - 0.382 \log K_{ow}$

**Statistics:**

$N = 4; R^2 = 0.92$

**Maximum log  $K_{ow}$ :**

5.0

**Maximum MW:**

1000.0

**Application:**

This equation may be used to estimate toxicity for monoepoxides.

**Limitations:**

If the log  $K_{ow}$  value is greater than 5.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.

Monoepoxides which are significantly more toxic than predicted by this SAR, based on the fish 14-d LC50 SAR, are:

epichlorohydrin, and  
epibromohydrin.

Endrin has an excess toxicity of over 33,000 times the value predicted by this SAR. Diepoxides are significantly more toxic than predicted by this SAR and a SAR for diepoxides has been developed.

**References:**

Bridie AL, Wolff CJM, and Winter M. 1979. The acute toxicity of some petrochemicals to goldfish. *Water Research*, 13: 623-626.

Conway RA, Waggy GT, Spiegel MH, and Berglund RL. 1983. Environmental fate and effects of ethylene oxide. *Environmental Science and Technology* 17:107-112.

Leach JM and Thakore AN. 1975. Isolation and identification of constituents toxic to juvenile rainbow trout (*Salmo gairdneri*) in caustic extraction effluents from kraft pulpmill bleach plants. *Journal of the Fisheries Research Board of Canada*, 32: 1249.

United States Environmental Protection Agency (USEPA). 1986. *Water Quality Criteria for 1986*. Washington, DC: USEPA

**EPOXIDES, MONO**  
9/1993

LIST OF MONOEPOXIDES USED TO DEVELOP THE FISH 96-h LC50 SAR.

CHEMICAL	96-h LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
MONOEPOXIDES USED IN CALCULATION OF THE SAR			
Ethylene oxide	84.0	-0.8	C
Allyl glycidyl ether	30.0	-0.33	B
Phenyl glycidyl ether	43.0	1.12	B
9,10-Epoxystearic acid	1.5	5.14	LT
MONOEPOXIDES HAVING EXCESS TOXICITY			
Endrin	0.000410	2.9	W

B = Bridie et al. (1979)

C = Conway et al (1983)

LT = Leach and Thakore (1975)

W = USEPA (1986); water quality criteria document

**SAR**

**EPOXIDES, MONO**

**Organism:**  
**Duration:**  
**Endpoint:**

Fish  
14-d  
LC50 (Mortality)

**Equation:**

$\text{Log 14-d LC50 (mM/L)} = -0.49506 - 0.34618 \log K_{ow}$

**Statistics:**

$N = 9; R^2 = 0.87$

**Maximum log  $K_{ow}$ :**  
**Maximum MW:**

5.0  
1000.0

**Application:**

This equation may be used to estimate toxicity for monoepoxides.

**Limitations:**

If the log  $K_{ow}$  value is greater than 5.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation

Monoepoxides which are significantly more toxic than predicted by this SAR are:

epichlorohydrin, 53 X excess toxicity; and  
epibromohydrin, 57 X excess toxicity.

Diepoxides are significantly more toxic than predicted by this SAR and a SAR for diepoxides has been developed.

**References:**

Deneer JW, Sinnige TL, Seinen W, and Hermens JLM. 1988. A quantitative structure-activity relationship for the acute toxicity of some epoxy compounds to the guppy. *Aquatic Toxicology* 13:195-204.

**EPOXIDES, MONO**

9/1993

**LIST OF MONOEPOXIDES USED TO DEVELOP THE FISH 14-d LC50 SAR.**

CHEMICAL	14-d LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
MONOEPOXIDES USED IN CALCULATION OF THE SAR			
Glycidol	50.0	-1.46	D
Propylene oxide	31.9	-0.27	D
1,2-Epoxybutane	32.9	0.26	D
Styrene oxide	7.07	0.73	D
1,2-Epoxyhexane	18.6	1.31	D
1,2-Epoxyoctane	10.4	2.37	D
1,2-Epoxydecane	3.26	3.43	D
1,2-Epoxydodecane	1.11	4.49	D
1,2-Epoxyhexadecane	*	6.60	D
MONOEPOXIDES HAVING EXCESS TOXICITY			
Epichlorohydrin	0.651	-0.21	D
Epibromohydrin	0.807	-0.07	D

\* No fish mortality in saturated solutions.

D = Deneer et al (1988)

<b>SAR</b>	<b>EPOXIDES, MONO</b>
<b>Organism:</b>	Daphnid
<b>Duration:</b>	48-h
<b>Endpoint:</b>	LC50 (Mortality)
<b>Equation:</b>	$\text{Log 48-h LC50 (mM/L)} = 0.036 - 0.567 \log K_{ow}$
<b>Statistics:</b>	$N = 2; R^2 = 1.0$
<b>Maximum log <math>K_{ow}</math>:</b>	5.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This equation may be used to estimate toxicity for monoepoxides.
<b>Limitations:</b>	If the log $K_{ow}$ value is greater than 5.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.
<b>References:</b>	Conway RA, Waggy GT, Speigel MH, and Berglund RL. 1983. Environmental fate and effects of ethylene oxide. Environmental Science and Technology 17:107-112.

LIST OF MONOEPOXIDES USED TO DEVELOP THE DAPHNID 48-h LC50 SAR.

CHEMICAL	48-h LC50 (mg/L)	Log $K_{ow}$	Ref.
Ethylene oxide	137.0	-0.8	C

C = Conway et al. (1983)

**EPOXIDES, MONO**  
9/1993

<b>SAR</b>	<b>EPOXIDES, DI</b>
<b>Organism:</b>	Fish
<b>Duration:</b>	96-h
<b>Endpoint:</b>	LC50 (Mortality)
<b>Equation:</b>	$\text{Log 96-h LC50 (mM/L)} = -1.184 - 0.263 \log K_{ow}$
<b>Statistics:</b>	$N = 2; R^2 = 1.0$
<b>Maximum log <math>K_{ow}</math>:</b>	5.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This equation may be used to estimate toxicity for diepoxides and other polyepoxides.
<b>Limitations:</b>	If the log $K_{ow}$ value is greater than 5.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.
<b>References:</b>	<p>Bailey RE and Rhinehart WL. 1976. Evaluation of D.E.R. 331, diglycidyl ether of bisphenol-A, in the aquatic environment. R&amp;D Report D0004653. Midland, MI: The Dow Chemical Company.</p> <p>United States Environmental Protection Agency (USEPA). 1991. OTS PMN ECOTOX. Washington, DC: USEPA, Office of Toxic Substances.</p>

LIST OF DIEPOXIDES USED TO DEVELOP THE FISH 96-h LC50 SAR.

CHEMICAL	96-h LC50 (mg/L)	Log $K_{ow}$	Ref.
Diglycidyl ether of bisphenol A	3.1	3.1	B
Chemical identity CBI	*	7.1	EPA

\* No fish mortality in saturated solutions.

B = Bailey and Rhinehart (1976)

EPA = USEPA (1991); chemical identity is Confidential Business Information under TSCA.

**EPOXIDES, DI**  
9/1993

<b>SAR</b>	<b>EPOXIDES, DI</b>
<b>Organism:</b>	Fish
<b>Duration:</b>	14-d
<b>Endpoint:</b>	LC50 (Mortality)
<b>Equation:</b>	$\text{Log 14-d LC50 (mM/L)} = -1.5692 - 0.1216 \log K_{ow}$
<b>Statistics:</b>	$N = 3; R^2 = 0.83$
<b>Maximum log <math>K_{ow}</math>:</b>	5.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This equation may be used to estimate toxicity for diepoxides and other polyepoxides.
<b>Limitations:</b>	If the log $K_{ow}$ value is greater than 5.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.
<b>References:</b>	Deneer JW, Sinnige TL, Seinen W and Hermens JLM. 1988. A quantitative structure-activity relationship for the acute toxicity of some epoxy compounds to the guppy. <i>Aquatic Toxicology</i> 13:195-204.

LIST OF DIEPOXIDES USED TO DEVELOP THE FISH 14-d LC50 SAR.

CHEMICAL	14-d LC50 (mg/L)	Log $K_{ow}$	Ref.
1,3-Butadiene diepoxide	2.66	-1.84	D
1,2,7,8-Diepoxyoctane	6.64	-0.18	D

D = Deneer et al (1988)

**EPOXIDES, DI**  
9/1993

<b>SAR</b>	<b>EPOXIDES, DI</b>
<b>Organism:</b>	Daphnid
<b>Duration:</b>	48-h
<b>Endpoint:</b>	LC50 (Mortality)
<b>Equation:</b>	$\text{Log 48-h LC50 (mM/L)} = -2.093 - 0.1474 \log K_{ow}$
<b>Statistics:</b>	$N = 2; R^2 = 1.0$
<b>Maximum log <math>K_{ow}</math>:</b>	5.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This equation may be used to estimate toxicity for diepoxides and other polyepoxides.
<b>Limitations:</b>	If the log $K_{ow}$ value is greater than 5.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.
<b>References:</b>	Bailey RE and Rhinehart WL. 1976. Evaluation of D.E.R. 331, diglycidyl ether of bisphenol-A, in the aquatic environment. R&D Report D0004653. Midland, MI: The Dow Chemical Company.

LIST OF DIEPOXIDES USED TO DEVELOP THE DAPHNID 48-h LC50 SAR.

CHEMICAL	48-h LC50 (mg/L)	Log $K_{ow}$	Ref.
Diglycidyl ether of bisphenol A	0.95	3.1	B

B = Bailey and Rhinehart (1976)

EPOXIDES, DI  
9/1993

**SAR**

**ESTERS**

**Organism:**

Fish

**Duration:**

96-h

**Endpoint:**

LC50 (Mortality)

**Equation:**

$\text{Log LC50 (mM/L)} = -0.535 \log K_{ow} + 0.25$

**Statistics:**

$N = 29, R^2 = 0.828$

**Maximum log  $K_{ow}$ :**

5.0

**Maximum MW:**

1000.0

**Application:**

This SAR may be used to estimate toxicity for the following esters:

1. Acetates
2. Benzoates
3. Dicarboxylic aliphatics
4. Phthalates derived from aliphatic alcohols and phenol.

**Limitations:**

If the log  $K_{ow}$  value is greater than 5.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.

**References:**

Veith GD, DeFoe D, and Knuth M. 1984. Structure-activity relationships for screening organic chemicals for potential ecotoxicity effects. Drug Metabolism Reviews 15(7):1295-1303.

# ESTERS

9/1993

## LIST OF ESTERS USED TO DEVELOP THE FISH 96-h LC50 SAR.

CHEMICAL	96-h LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
Methylene chloride	322.895	1.25	Z
Methyl acetate	320.0	0.18	V
Ethyl acetate	230.0	0.69	V
2-Ethoxyethyl acetate	42.2	0.71	V
Diethyl malonate	14.9	1.19	V
Ethyl-p-aminobenzoate	35.2	1.22	V
Propyl acetate	60.0	1.25	V
Methyl-2,4-dihydroxybenzoate	38.5	1.59	V
Butyl acetate	18.0	1.79	V
Diethyl adipate	19.3	1.80	V
Methyl-p-nitrobenzoate	23.6	2.10	V
Dimethyl-2-nitro-p-phthalate	6.52	2.28	V
Methyl-4-chloro-2-nitrobenzoate	27.2	2.35	V
Dimethyl-2-amino-p-phthalate	8.94	2.65	V
Diethyl-o-phthalate	30.0	2.69	V
Hexyl acetate	4.40	2.87	V
Ethyl hexanoate	8.90	2.87	V
Methyl-p-chlorobenzoate	10.9	3.15	V
Methyl-2,5-dichlorobenzoate	13.8	3.45	V
Ethyl salicylate	19.6	3.45	V
Dibutyl succinate	4.45	3.65	V
Dibutyl adipate	3.66	3.96	V
Diethyl sebacate	2.75	3.96	V
Di-n-butyl-o-phthalate	1.10	4.74	V
Di-n-butyl-m-phthalate	0.90	5.07	V
Diphenyl-1-phthalate	0.80	7.06	V
Di-2-ethylhexyl-o-phthalate	*	7.06	V
Di-n-octyl-o-phthalate	*	7.06	V
Di-n-octyl-m-phthalate	*	7.06	V
Di-n-octyl-p-phthalate	*	7.06	V

\* = No fish mortality in saturated solutions.

V = Veith et al (1984)

<b>SAR</b>	<b>ESTERS</b>
<b>Organism:</b>	Daphnid
<b>Duration:</b>	48-h
<b>Endpoint:</b>	LC50 (Mortality)
<b>Equation:</b>	To find the estimated acute toxicity of an ester, use the neutral organics daphnid 48-h LC50 SAR.
<b>Maximum log <math>K_{ow}</math>:</b>	5.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	The daphnid 48-h LC50 SAR for neutral organics may be used to estimate acute toxicity for esters. The neutral organic 48-h LC50 SAR for daphnids may be used for other esters; however, a separate SAR has been developed for phthalate esters.
<b>Limitations:</b>	If the log $K_{ow}$ value is greater than 5.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.
<b>References:</b>	Kuhn R, Pattard M, Pernack K-D, and Winter A. 1989. Results of the harmful effects of selected water pollutants (anilines, phenols, aliphatic compounds) to <u>Daphnia magna</u> . Water Research 23:495-499.  United States Environmental Protection Agency (USEPA). 1991. OTS PMN ECOTOX. Washington, DC: Office of Toxic Substances, USEPA.

LIST OF ESTERS USED TO DEVELOP THE DAPHNID 48-h LC50 SAR.

CHEMICAL	48-h LC50 (mg/L)	Log $K_{ow}$	Ref.
Methylene chloride	322.895	1.25	Z
Chloroacetic ethyl ester	1.6	?	K
Chemical identity CBI	3.32	3.7	EPA
Chemical identity CBI	*	4.4	EPA

\* = No daphnid mortality in saturated solutions.

EPA = USEPA (1991); chemical identities are Confidential Business Information under TSCA.

K = Kuhn et al (1989)

**ESTERS**  
7/1988

<b>SAR</b>	<b>ESTERS</b>
<b>Organism:</b>	Green Algae
<b>Duration:</b>	96-h
<b>Endpoint:</b>	EC50 (Growth)
<b>Equation:</b>	$\text{Log EC50 (mM/L)} = -0.881 - 0.519 \log K_{ow}$
<b>Statistics:</b>	$N = 2; R^2 = 1.0$
<b>Maximum log <math>K_{ow}</math>:</b>	6.4
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This SAR may be used to estimate toxicity for esters.
<b>Limitations:</b>	If the log $K_{ow}$ value is greater than 6.4, or if the compound is solid and the EC50 exceeds the water solubility, no effects expected at saturation
<b>References:</b>	United States Environmental Protection Agency (USEPA). 1991. OTS PMN ECOTOX. Washington, DC: Office of Toxic Substances, USEPA.

LIST OF ESTERS USED TO DEVELOP THE GREEN ALGAE 96-h EC50 SAR.

CHEMICAL	96-h EC50 (mg/L)	Log $K_{ow}$	Ref.
Chemical identity CBI	0.410	3.7	EPA

EPA = USEPA (1991), chemical identities are Confidential Business Information under TSCA.

**ESTERS**  
7/1988

**SAR**

**ESTERS**

**Organism:**

Green Algae

**Duration:**

16-d

**Endpoint:**

Chronic Value (Growth)

**Equation:**

$\text{Log ChV (mM/L)} = -1.01 - 0.51 \log K_{ow}$

**Statistics:**

$N = 2; R^2 = 1.0$

**Maximum log  $K_{ow}$ :**

8.0

**Maximum MW:**

1000.0

**Application:**

This SAR may be used to estimate toxicity for esters.

**Limitations:**

If the log  $K_{ow}$  value is greater than 8.0, or if the compound is solid and the ChV exceeds the water solubility, no effects expected at saturation.

**References:**

United States Environmental Protection Agency (USEPA). 1991. OTS PMN ECOTOX. Washington, DC: Office of Toxic Substances, USEPA.

LIST OF ESTERS USED TO DEVELOP THE GREEN ALGAE CHRONIC VALUE (ChV) SAR.

CHEMICAL	ChV (mg/L)	Log $K_{ow}$	Ref.
Chemical identity CBI	0.390	3.7	EPA

EPA = USEPA (1991); chemical identities are Confidential Business Information under TSCA.

**ESTERS**  
7/1988

**SAR**

**ESTERS, MONO**

**Organism:** Fish  
**Duration:** 32-d  
**Endpoint:** Chronic Value (Survival/Growth)  
**Equation:**  $\text{Log ChV (mM/L)} = 0.421 - 0.828 \log K_{ow}$   
**Statistics:**  $N = 2; R^2 = 1.0$   
**Maximum log  $K_{ow}$ :** 8.0  
**Maximum MW:** 1000.0  
**Application:** This SAR may be used to estimate toxicity for aliphatic monoesters.  
**Limitations:** If the log  $K_{ow}$  value is greater than 8.0, no effects expected at saturation.  
**References:** United States Environmental Protection Agency (USEPA). 1991. Fish Chronic Toxicity Data Base. Duluth, MN: Environmental Research Laboratory (ERL), Office of Research and Development, USEPA, 6201 Congdon Boulevard, 55804; contact C.L. Russom (218) 720-5500.

LIST OF ALIPHATIC MONOESTERS USED TO DEVELOP THE FISH CHRONIC VALUE (ChV) SAR.

CHEMICAL	ChV (mg/L)	Log $K_{ow}$	Ref
Methyl acetate	133.0	0.2	D

D = USEPA (1991)

**ESTERS, MONO, ALIPHATIC**  
9/1993

**SAR**

**ESTERS, DI**

**Organism:** Fish  
**Duration:** 32-d  
**Endpoint:** Chronic Value (Survival/Growth)  
**Equation:**  $\text{Log ChV (mM/L)} = -1.677 - 0.565 \log K_{ow}$   
**Statistics:**  $N = 3; R^2 = 1.0$   
**Maximum log  $K_{ow}$ :** 8.0  
**Maximum MW:** 1000.0  
**Application:** This SAR may be used to estimate toxicity for aliphatic diesters.  
**Limitations:** If the log  $K_{ow}$  value is greater than 8.0, or if the compound is solid and the ChV exceeds the water solubility, no effects expected at saturation.  
**References:** United States Environmental Protection Agency (USEPA). 1991. Fish Chronic Toxicity Data Base. Duluth, MN: Environmental Research Laboratory (ERL), Office of Research and Development, USEPA, 6201 Congdon Boulevard, 55804; contact C L. Russom (218) 720-5500.

LIST OF ALIPHATIC DIESTERS USED TO DEVELOP THE FISH CHRONIC VALUE (ChV) SAR

CHEMICAL	ChV (mg/L)	Log $K_{ow}$	Ref.
Diethyl malonate	0.759	1.1	D
Dibutyl fumarate	0.030	3.9	D

D = USEPA (1991)

**ESTERS, DI, ALIPHATIC**  
9/1993

**SAR****ESTERS, PHOSPHATE****Organism:**

Fish

**Duration:**

96-h

**Endpoint:**

LC50 (Mortality)

**Equation:**
$$\text{Log LC50 (mM/L)} = -0.0695 - 0.5178 \log K_{ow}$$
**Statistics:**
$$N = 16; R^2 = 0.595$$
**Maximum log  $K_{ow}$ :**

5.0

**Maximum MW:**

1000.0

**Application:**

This SAR may be used to estimate the toxicity of phosphate esters and other tri-alkyl-phenyl phosphate esters. This SAR may be used to estimate toxicity for the following classes of phosphate esters all of which are weak acetylcholinesterase inhibitors:

1. Tri-alkyl phosphate esters
2. Tri-phenyl phosphate esters
3. Halogenated tri-alkyl phosphate esters
4. Halogenated tri-phenyl phosphate esters

Some halogenated tri-alkylphosphate esters are significantly more toxic than predicted by this SAR as a result of their strong acetylcholinesterase and cholinesterase inhibition. These include:

1. 1,2-dibromoethyldiethyl phosphate ester - 400 X
2. 1,2-dichloroethyldiethyl phosphate ester -  $\approx 30$  X

**Limitations:**

If the log  $K_{ow}$  value is greater than 5.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.

**References:**

United States Environmental Protection Agency (USEPA). 1991. OTS PMN ECOTOX Washington, DC: Office of Toxic Substances, USEPA

**ESTERS, PHOSPHATE**

9/1993

**LIST OF PHOSPHATE ESTERS USED TO DEVELOP THE FISH 96-h LC50 SAR.**

CHEMICAL	96-h LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
Tris(betachloroethyl)	210.0	0.92	EPA
Tris(betachloroethyl)	90.0	0.92	EPA
Chemical identity CBI	21.0	1.80	EPA
Tris(dichloropropyl)	3.6	2.67	EPA
Tris(dichloropropyl)	5.1	2.67	EPA
Tris(2,3-dibromopropyl)	1.33	3.51	EPA
Tris(2,3-dibromopropyl)	1.45	3.51	EPA
Tributyl	11.0	3.53	EPA
Tributyl	8.18	3.53	EPA
Tributyl	8.8	3.53	EPA
Tributyl	9.6	3.53	EPA
Tributyl	11.8	3.53	EPA
Tributyl	11.4	3.53	EPA
Triphenyl	0.87	4.63	EPA
Triphenyl	0.70	4.63	EPA
Triphenyl	1.2	4.63	EPA

EPA = USEPA (1991)

<b>SAR</b>	<b>ESTERS, PHTHALATE</b>
<b>Organism:</b>	Fish
<b>Duration:</b>	96-h
<b>Endpoint:</b>	LC50 (Mortality)
<b>Equation:</b>	Use the ester fish 96-h SAR to determine the acute toxicity of a phthalate ester.
<b>Maximum log K<sub>ow</sub>:</b>	5.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	<p>The ester SAR may be used to estimate the toxicity of phthalate esters. The ester SAR is applicable for the following phthalate esters:</p> <ol style="list-style-type: none"> <li>1. Aliphatic diesters</li> <li>2. Aromatic diesters</li> <li>3. Aliphatic-aromatic diesters</li> <li>4. Phthalates, derived from aliphatic alcohols and phenol.</li> </ol>
<b>Limitations:</b>	If the log K <sub>ow</sub> value is greater than 5.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation SAR with longer exposure.
<b>References:</b>	Veith GD, DeFoe D, and Knuth M. 1984. Structure-activity relationships for screening organic chemicals for potential ecotoxicity effects. Drug Metabolism Reviews 15(7):1295-1303.

LIST OF PHTHALATE ESTERS USED TO DEVELOP THE FISH 96-h LC50 SAR.

CHEMICAL	96-h LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
Dimethyl-2-nitro-p-phthalate	6.52	2.28	V
Dimethyl-2-amino-p-phthalate	8.94	2.65	V
Diethyl-o-phthalate	30.0	2.69	V
Di-n-butyl-o-phthalate	1.10	4.74	V
Di-n-butyl-m-phthalate	0.90	4.74	V
Diphenyl-i-phthalate	0.80	5.07	V
Di-2-ethylhexyl-o-phthalate	*	7.06	V
Di-n-octyl-o-phthalate	*	7.06	V
Di-n-octyl-m-phthalate	*	7.06	V
Di-n-octyl-p-phthalate	*	7.06	V

## ESTERS, PHTHALATE

9/1993

\* No fish mortality in saturated solutions.

V = Veith et al (1984).

**SAR****ESTERS, PHTHALATE****Organism:**

Daphnid

**Duration:**

48-h

**Endpoint:**

LC50 (Mortality)

**Equation:**

Use the neutral organic daphnid 48-h SAR to determine the acute toxicity of a phthalate ester.

**Maximum log  $K_{ow}$ :**

5.0

**Maximum MW:**

1000.0

**Application:**

The neutral organic SAR may be used to estimate the toxicity of phthalate esters. The neutral organic SAR is applicable for the following phthalate esters:

1. Aliphatic diesters
2. Aromatic diesters
3. Aliphatic-aromatic diesters
4. Phthalates, derived from aliphatic alcohols and phenol.

**Limitations:**

If the log  $K_{ow}$  value is greater than 5.0, or if the compound is solid and the LC50 exceeds the water solubility, use SAR with longer exposure.

**References:**

Nabholz JV. 1987. The acute and chronic toxicity of dialkyl phthalate esters to daphnids. Interagency memorandum to "Whom It May Concern." Washington, DC: Environmental Effects Branch, Health and Environmental Review Division, Office of Toxic Substances, United States Environmental Protection Agency.

# ESTERS, PHTHALATE

9/1993

## LIST OF PHTHALATE ESTERS USED TO DEVELOP THE DAPHNID 48-h LC50 SAR.

CHEMICAL	48-h LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
Methylene chloride	322.895	1.25	Z
Dimethyl	>52.0	1.52	N
Diethyl	90.0	2.57	N
Di-n-butyl-ortho	3.4	4.69	N
Di-n-butyl-ortho	5.2	4.69	N
Butyl-benzyl	1.83	4.87	N
Butyl-benzyl	3.7	4.87	N
Butyl-benzyl	1.6	4.87	N
Butyl-benzyl	1.0	4.87	N
Butyl-benzyl	2.4	4.87	N
Butyl-benzyl	1.7	4.87	N
Diethyl	*	6.80	N
Butyl-2-ethylhexyl	*	7.93	N
Di-(n-hexyl, n-octyl, n-decyl)	*	8.57	N
Di-(2-ethylhexyl)	*	8.66	N
Di-(2-ethylhexyl)	*	8.66	N
Diisooctyl	*	8.66	N
Di-(n-octyl)	*	8.92	N
Di-(heptyl, nonyl, undecyl)	*	9.59	N
Diisononyl	*	9.72	N
Diisodecyl	*	10.78	N
Diisodecyl	*	10.78	N
Diundecyl	*	12.10	N
Ditridecyl	*	14.21	N

\* No daphnid mortality in saturated solutions.

N = Nabholz (1987)

**SAR**

**ESTERS, PHTHALATE**

<b>Organism:</b>	Daphnid
<b>Duration:</b>	21-d
<b>Endpoint:</b>	No Effect Concentration (NEC) (Reproduction)
<b>Equation:</b>	$\text{Log 21-d NEC (mM/L)} = 0.05 - 0.72 \log K_{ow}$
<b>Statistics:</b>	
<b>Maximum log <math>K_{ow}</math>:</b>	8.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	<p>The neutral organic 16-d NEC SAR may be used to estimate the toxicity of phthalate esters. The neutral organic SAR is applicable for the following phthalate esters:</p> <ol style="list-style-type: none"><li>1. Aliphatic diesters</li><li>2. Aromatic diesters</li><li>3. Aliphatic-aromatic diesters</li><li>4. Phthalates, derived from aliphatic alcohols and phenol.</li></ol>
<b>Limitations:</b>	<p>If the log <math>K_{ow}</math> value is greater than 8.0, or if the compound is solid and the NEC exceeds the water solubility, no effects expected at saturation.</p>
<b>References:</b>	<p>Nabholz JV. 1987. The acute and chronic toxicity of dialkyl phthalate esters to daphnids. Interagency memorandum to "Whom It May Concern." Washington, DC: Environmental Effects Branch, Health and Environmental Review Division, Office of Toxic Substances, United States Environmental Protection Agency.</p>

# ESTERS, PHTHALATE

9/1993

## LIST OF PHTHALATE ESTERS USED TO DEVELOP THE DAPHNID 21-d NEC SAR.

CHEMICAL	21-d NEC (mg/L)	Log K <sub>ow</sub>	Ref.
Methylene chloride	322.895	1.25	Z
Dimethyl	15.0	1.52	N
Diethyl	38.0	2.57	N
Di-n-butyl-ortho	1.0	4.69	N
Di-n-butyl-ortho	1.4	4.69	N
Di-n-butyl-ortho	1.5	4.69	N
Butyl-benzyl	0.63	4.87	N
Butyl-benzyl	0.44	4.87	N
Di-n-butyl-iso	0.15	5.53	N
Di-n-butyl-tere	0.32	5.53	N
Dihexyl	*	6.80	N
Butyl-2-ethylhexyl	*	7.93	N
Di-(n-hexyl, n-octyl, n-decyl)	*	8.57	N
Di-(2-ethylhexyl)	*	8.66	N
Di-(2-ethylhexyl)	*	8.66	N
Diisooctyl	*	8.66	N
Di-(n-octyl)	*	8.92	N
Di-(heptyl, nonyl, undecyl)	*	9.59	N
Diisononyl	*	9.72	N
Diisodecyl	*	10.78	N
Diisodecyl	*	10.78	N
Diundecyl	*	12.10	N
Ditridecyl	*	14.21	N

\* No daphnid systemic effects in saturated solutions.

N = Nabholz (1987).

**SAR**

**HYDRAZINES**

**Organism:** Fish  
**Duration:** 96-h  
**Endpoint:** LC50 (Mortality)  
**Equation:**  $\text{Log 96-h LC50 (mM/L)} = -1.53 - 0.438 \log K_{ow}$   
**Statistics:**  $N = 9; R^2 = 0.91$   
**Maximum log  $K_{ow}$ :** 5.0  
**Maximum MW:** 1000.0  
**Application:** This equation may be used to estimate toxicity for:

hydrazines  
 hydrazones  
 hydrazides  
 thiohydrazides  
 semicarbazides  
 thiosemicarbazides  
 semicarbazones  
 thiosemicarbazones

For hydrazines with missing fragment constants in CLOGP the following constants may be used:

1. missing fragment (-C(=S)-): -0.24
2. missing fragment (-NC(=O)N-N): -3.13
3. missing fragment (C=NNC(=O)N): -3.39.

**Limitations:**

Hydrazines which are 10 times less toxic than predicted by this SAR are those hydrazines which have a carboxylic acid substitution:

butanedioic acid mono-(2,2'-dimethylhydrazide).

If the log  $K_{ow}$  value is greater than 5.0 and less than 6.6, use the neutral organics fish 14-d LC50 SAR; and if the log  $K_{ow}$  value is equal to or greater than 6.6, use the neutral organics fish ChV SAR.

**References:**

Buccafusco RJ, Ells SJ, and LeBlanc GA 1981. Acute toxicity of priority pollutants to bluegill (*Lepomis macrochirus*). Bulletin of Environmental Contamination and Toxicology 26:446-452.

# HYDRAZINES

9/1993

Hammermeister D, Kahl M, and Broderius S. 1990. EEB/ERL-Duluth interaction on various join projects. Duluth, MN: Environmental Research Laboratory-Duluth, United States Environmental Protection Agency, 6201 Congdon Blvd., 55804, Unpublished memorandum to V. Nabolz.

Odenkirchen EW and Nabholz JV. 1989. Generic environmental hazard assessment of hydrazines and related compounds. Rockville, Maryland: Dynamac Corporation, 11140 Rockville Pike, 20852.

## LIST OF HYDRAZINES USED TO DEVELOP THE FISH 96-h LC50 SAR.

CHEMICAL	96-h LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
HYDRAZINES USED IN CALCULATION OF THE SAR			
Hydrazine	2.81	-1.37	H
Hydrazine	3.4	-1.37	ON
Monomethyl hydrazine	3.26	-1.06	ON
Monomethyl hydrazine	2.58	-1.06	ON
1,1-Dimethyl hydrazine	7.75	-1.50	H
1,1-Dimethyl hydrazine	10.0	-1.50	ON
1,1-Dimethyl hydrazine	26.5	-1.50	ON
Thiosemicarbazide	20.8	-2.4	ON
1,2-Diphenyl hydrazine	0.27	2.97	B
HYDRAZINES LESS TOXIC THAN PREDICTED			
Butanedioic acid mono- (2,2'-dimethylhydrazide)	423.0	-0.619	ON
Butanedioic acid mono- (2,2'-dimethylhydrazide)	149.0	-0.619	ON
HYDRAZINES NOT ACUTELY TOXIC AT SATURATION			
N-Acetyl-1,2-diphenylhydrazine	* (mp 164° C)	2.2	H

H = Hammermeister et al (1990)

ON = Odenkirchen and Nabholz (1989)

B = Buccafusco et al (1981)

**SAR****HYDRAZINES****Organism:**

Daphnid

**Duration:**

48-h

**Endpoint:**

LC50 (Mortality)

**Equation:** $\text{Log 48-h LC50 (mM/L)} = -1.2941 - 0.256 \log K_{ow}$ **Statistics:** $N = 4; R^2 = 0.46$ **Maximum log  $K_{ow}$ :**

5.0

**Maximum MW:**

1000.0

**Application:**

This equation may be used to estimate toxicity for:

hydrazines  
 hydrazones  
 hydrazides  
 thiohydrazides  
 semicarbazides  
 thiosemicarbazides  
 semicarbazones  
 thiosemicarbazones

For hydrazines with missing fragment constants in CLOGP the following constants may be used:

1. missing fragment (-C(=S)-): -0.24
2. missing fragment (-NC(=O)N-N): -3.13
3. missing fragment (C=NNC(=O)N): -3.39

**Limitations:**

Hydrazines which are significantly less toxic than predicted by this SAR are those hydrazines which have a carboxylic acid substitution:

butanedioic acid mono-(2,2'-dimethylhydrazide).

If the log  $K_{ow}$  value is greater than 5.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.

**References:**

LeBlanc. 1980. Acute toxicity of priority pollutants to water flea (*Daphnia magna*). Bulletin of Environmental Contamination and Toxicology. 24: 684-691.

Hammermeister D, Kahl M, and Broderius S. 1990. EEB/ERL-Duluth interaction on various joint projects. Duluth, MN: Environmental Research Laboratory-Duluth, United States Environmental Protection Agency, 6201 Congdon Blvd., 55804, Unpublished memorandum to V. Nabolz.

**HYDRAZINES**

9/1993

## LIST OF HYDRAZINES USED TO DEVELOP THE DAPHNID 48-h LC50 SAR.

CHEMICAL	48-h LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
HYDRAZINES USED IN CALCULATION OF THE SAR			
Hydrazine	0.280	-1.37	H
1,1-Dimethyl hydrazine	68.2	-1.50	H
1,2-Diphenyl hydrazine	4.1	2.97	L
HYDRAZINES NOT ACUTELY TOXIC AT SATURATION			
N-Acetyl-1,2-diphenylhydrazine	* (mp 164° C)	2.2	H

H = Hammermeister et al (1990)

L = LeBlanc (1980)

<b>SAR</b>	<b>HYDRAZINES</b>
<b>Organism:</b>	Green Algae
<b>Duration:</b>	144-h
<b>Endpoint:</b>	EC50 (Growth)
<b>Equation:</b>	$\text{Log 144-h EC50 (mM/L)} = -5.1725 - 0.0999 \text{ Log } K_{ow}$
<b>Statistics:</b>	$N = 3; R^2 = 0.3$
<b>Maximum log <math>K_{ow}</math>:</b>	8.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This equation may be used to estimate toxicity for hydrazines. SAR equations for other subclasses of hydrazines, i.e., alkylsemicarbazides and arylsemicarbazides, may be found elsewhere in this volume.
<b>Limitations:</b>	Hydrazines which are significantly less toxic than predicted by this SAR are those hydrazines which have a carboxylic acid substitution:  butanedioic acid mono-(2,2'-dimethylhydrazide).
<b>References:</b>	Odenkirchen EW and Nabholz JV. 1989. Generic environmental hazard assessment of hydrazines and related compounds. Rockville, Maryland: Dynamac Corporation, 11140 Rockville Pike, 20852.

LIST OF HYDRAZINES USED TO DEVELOP THE GREEN ALGAE 144-h EC50 SAR.

CHEMICAL	144-h EC50 (mg/L)	Log $K_{ow}$	Ref.
1,1-Dimethyl hydrazine	0.004100	-1.50	ON
Hydrazine	0.000041	-1.37	ON

ON = Odenkirchen and Nabholz (1989)

**HYDRAZINES**  
9/1993

## HYDRAZINES, SEMICARBAZIDES, ALKYL SUBSTITUTED

9/1993

### SAR

### HYDRAZINES, SEMICARBAZIDE, ALKYL SUBSTITUTED

**Organism:**

Green Algae

**Duration:**

6-h

**Endpoint:**

EC50 (Growth)

**Equation:**

(1) For  $\log K_{ow}$  less than -1.02:

$$\text{Log 6-h EC50 (mM/L)} = -2.1 - 0.521 \log K_{ow}$$

(2) For  $\log K_{ow}$  greater than -1.02:

$$\text{Log 6-h EC50 (mM/L)} = -0.89 + 0.625 \log K_{ow}$$

**Statistics:**

(1) For  $\log K_{ow}$  less than -1.02:  $N = 6$ ,  $R^2 = 0.75$ ;

(2) For  $\log K_{ow}$  greater than -1.02:  $N = 7$ ,  $R^2 = 0.86$

**Maximum  $\log K_{ow}$ :**

1.5

**Maximum MW:**

1000.0

**Application:**

This equation may be used to estimate toxicity for the following hydrazine classes with alkyl substitutions:

semicarbazides  
thiosemicarbazides  
semicarbazones  
thiosemicarbazones  
hydrazides  
thiohydrazides  
hydrazones

SAR equations for aryl substituted semicarbazides and hydrazines may be found elsewhere in this volume.

For semicarbazides with missing fragment constants in CLOGP the following constants may be used.

1. missing fragment (-C(=S)-): -0.24
2. missing fragment (-NC(=O)N-N): -3.13
3. missing fragment (C=NNC(=O)N): -3.39.

**Limitations:**

If the  $\log K_{ow}$  value is greater than 1.5, no effects expected at saturation

**References:**

Odenkirchen EW and Nabholz JV. 1989. Generic environmental hazard assessment of hydrazines and related compounds Rockville, Maryland: Dynamac Corporation, 11140 Rockville Pike, 20852.

# HYDRAZINES, SEMICARBAZIDES, ALKYL SUBSTITUTED

9/1993

## LIST OF ALKYL SUBSTITUTED SEMICARBAZIDES USED TO DEVELOP THE GREEN ALGAE 6-h EC50 SAR.

CHEMICAL	6-h EC50 (mg/L)	Log K <sub>ow</sub>	Ref
4-Methyl	9.9	-2.25	ON
4-Allyl	3.8	-1.74	ON
4-Ethyl	5.1	-1.73	ON
4,4-Dimethyl	4.2	-1.50	ON
4-Isopropyl	2.7	-1.42	ON
4-n-Propyl	2.5	-1.20	ON
4-t-Butyl	4.1	-1.02	ON
4-Isobutyl	3.9	-0.80	ON
4-Benzyl	12.8	-0.69	ON
4-n-Butyl	5.9	-0.67	ON
4,4-Diethyl	6.1	-0.44	ON
4-n-Pentyl	12.6	-0.14	ON
4-n-Hexyl	38.2	0.39	ON

ON = Odenkirchen and Nabholz (1989)

# HYDRAZINES, SEMICARBAZIDES, ARYL, ORTHO SUBSTITUTED

9/1993

## SAR

## HYDRAZINES, SEMICARBAZIDES, ARYL, ORTHO SUBSTITUTED

**Organism:**

Green Algae

**Duration:**

6-h

**Endpoint:**

EC50 (Growth)

**Equation:**

$\text{Log 6-h EC50 (mM/L)} = -0.88 - 0.563 \log K_{ow}$

**Statistics:**

$N = 7; R^2 = 0.98$

**Maximum log  $K_{ow}$ :**

8.0

**Maximum MW:**

1000.0

**Application:**

This equation may be used to estimate toxicity for the following arylsemicarbazides with ortho substituents on the aryl group:

thiosemicarbazides  
semicarbazones  
thiosemicarbazones  
hydrazides  
thiohydrazides  
hydrazones

SAR equations for arylsemicarbazides with meta and para substituents, alkylsemicarbazides, and hydrazines may be found elsewhere in this volume.

For semicarbazides with missing fragment constants in CLOGP the following constants may be used:

1. missing fragment (-C(=S)-): -0.24
2. missing fragment (-NC(=O)N-N): -3.13
3. missing fragment (C=NNC(=O)N): -3.39.

**Limitations:**

Arylsemicarbazides which are significantly more toxic than predicted by this SAR are:

4-(o-hydroxyphenyl)semicarbazide, 30X excess toxicity.

If the log  $K_{ow}$  is greater than 5.0, or if the compound is solid and the EC50 exceeds the water solubility, no effects expected at saturation

**References:**

Odenkirchen EW and Nabholz JV. 1989. Generic environmental hazard assessment of hydrazines and related compounds. Rockville, Maryland: Dynamac Corporation, 11140 Rockville Pike, 20852.

**HYDRAZINES, SEMICARBAZIDE, ARYL, ORTHO SUBSTITUTED**  
9/1993

**LIST OF ARYLSEMICARBAZIDES WITH ORTHO SUBSTITUENTS ON THE ARYL GROUP USED TO DEVELOP THE GREEN ALGAE 6-h EC50 SAR.**

CHEMICAL	6-h EC50 (mg/L)	Log K <sub>ow</sub>	Ref.
ARYLSEMICARBAZIDES USED IN CALCULATION OF THE SAR			
4-[o-Nitrophenyl]	194.6	-1.47	ON
4-[o-Carboxyphenyl]	176.6	-1.47	ON
4-[o-Methoxyphenyl]	116.2	-1.30	ON
4-[o-Methylphenyl]	52.0	-0.57	ON
4-[o-Chlorophenyl]	39.4	-0.50	ON
4-[m-Bromophenyl]	26.2	-0.35	ON
4-[o-Bromophenyl]	53.4	-0.35	ON
4-[2,5-Dichlorophenyl]	22.3	0.21	ON
ARYLSEMICARBAZIDES HAVING EXCESS TOXICITY			
4-[o-Hydroxyphenyl]	6.0	-1.88	ON

ON = Odenkirchen and Nabholz (1989)

**HYDRAZINES, SEMICARBAZIDE, ARYL, META/PARA SUBSTITUTED**  
9/1993

**SAR**

**HYDRAZINES, SEMICARBAZIDES, ARYL, META/PARA SUBSTITUTED**

**Organism:** Green Algae  
**Duration:** 6-h  
**Endpoint:** EC50 (Growth)

**Equation:**  $\text{Log 6-h EC50 (mM/L)} = -1.13 - 0.461 \log K_{ow}$

**Statistics:**  $N = 19; R^2 = 0.98$

**Maximum log  $K_{ow}$ :** 8.0  
**Maximum MW:** 1000.0

**Application:** This equation may be used to estimate toxicity for the following arylsemicarbazides with meta or para substituents on the aryl group:

thiosemicarbazides  
semicarbazones  
thiosemicarbazones  
hydrazides  
thiohydrazides  
hydrazones

For semicarbazides with missing fragment constants in CLOGP the following constants may be used:

1. missing fragment (-C(=S)-): -0.24
2. missing fragment (-NC(=O)N-N): -3.13
3. missing fragment (C=NNC(=O)N): -3.39

**Limitations:** SAR equations for arylsemicarbazides with ortho substituents, alkylsemicarbazides, and hydrazines may be found elsewhere in this volume. If the log  $K_{ow}$  value is greater than 8.0, or if the compound is solid and the EC50 exceeds the water solubility, no effects expected at saturation.

**References:** Odenkirchen EW and Nabholz JV. 1989. Generic environmental hazard assessment of hydrazines and related compounds. Rockville, Maryland: Dynamac Corporation, 11140 Rockville Pike, 20852.

**HYDRAZINES, SEMICARBAZIDE, ARYL, META/PARA SUBSTITUTED**  
9/1993,

**LIST OF ARYLSEMICARBAZIDES WITH META AND PARA SUBSTITUENTS ON THE ARYL GROUP  
USED TO DEVELOP THE GREEN ALGAE 6-h EC50 SAR.**

CHEMICAL	6-h EC50 (mg/L)	Log K <sub>ow</sub>	Ref.
4-[ <u>m</u> -Hydroxyphenyl]	144.8	-1.88	ON
4-[ <u>p</u> -Hydroxyphenyl]	170.0	-1.88	ON
4-[ <u>m</u> -Nitrophenyl]	109.5	-1.47	ON
4-[ <u>p</u> -Nitrophenyl]	98.0	-1.47	ON
4-[ <u>m</u> -Carboxyphenyl]	104.0	-1.47	ON
4-[ <u>p</u> -Carboxyphenyl]	92.7	-1.47	ON
4-[ <u>m</u> -Methoxyphenyl]	71.7	-1.30	ON
4-[ <u>p</u> -Methoxyphenyl]	70.0	-1.30	ON
4-Phenyl	42.5	-1.22	ON
4-[ <u>p</u> -Ethoxyphenyl]	38.7	-0.77	ON
4-[ <u>m</u> -Methylphenyl]	26.7	-0.57	ON
4-[ <u>p</u> -Methylphenyl]	24.9	-0.57	ON
4-[ <u>m</u> -Chlorophenyl]	22.7	-0.50	ON
4-[ <u>p</u> -Chlorophenyl]	22.2	-0.50	ON
4-[ <u>m</u> -Bromophenyl]	26.2	-0.35	ON
4-[ <u>p</u> -Bromophenyl]	22.3	-0.35	ON
4-[ <u>p</u> -Iodophenyl]	17.7	-0.09	ON
4-[3,4-Dichlorophenyl]	9.3	0.21	ON
4-[2,5-Dichlorophenyl]	22.3	0.21	ON

ON = Odenkirchen and Nabholz (1989)

**SAR**

**IMIDES**

**Organism:**

Fish

**Duration:**

96-h

**Endpoint:**

LC50 (Mortality)

**Equation:**

$\text{Log 96-h LC50 (mM/L)} = 1.256 - 0.76 \log K_{ow}$

**Statistics:**

$N = 4; R^2 = 0.98$

**Maximum log  $K_{ow}$ :**

5.0

**Maximum MW:**

1000.0

**Application:**

This SAR may be used to estimate toxicity for imides.

**Limitations:**

For imides with log  $K_{ow}$  values greater than 5.0, a test duration of greater than 96 hours may be required for proper expression of toxicity. Also, if the toxicity value obtained by the use of this equation exceeds the water solubility of the compound (measured or estimated), mortalities greater than 50% would not be expected in a saturated solution during an exposure period of 96 hours.

**References:**

Fukunaga K (ed). 1987. Japanese Pesticides Guide. Tokyo, Japan: Japan Plant Protection Association.

Worthing CR (ed). 1983. The Pesticide Manual. A World Compendium. 7th Ed. Croydon, Great Britain: British Crop Protection Council.

LIST OF IMIDES USED TO DEVELOP THE FISH 96-h LC50 SAR.

CHEMICAL	96-h LC50 (mg/L)	Log $K_{ow}$	Ref.
Methylene chloride	322.895	1.25	Z
Sumilex	*	2.2	F
Vinclozolin	32.5	2.8	W
Vinclozolin	52.5	2.8	W
Sparticide	5.5	3.7	F

\* = No fish mortality in saturated solutions.

F = Fukunaga (1987)

W = Worthing (1983)

**IMIDES**  
9/1993

**SAR****KETONES, DI, ALIPHATIC****Organism:**

Fish

**Duration:**

96-h

**Endpoint:**

LC50 (Mortality)

**Equation:** $\text{Log 96-h LC50 (mM/L)} = -0.151 - 0.433 \log K_{ow}$ **Statistics:** $N = 22; R^2 = 0.87$ **Maximum log  $K_{ow}$ :**

5.0

**Maximum MW:**

1000.0

**Application:**

This SAR may be used to estimate toxicity for aliphatic diketones.

**Limitations:**If the log  $K_{ow}$  value is greater than 5.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.**References:**

Brooke LT, Call DJ, Geiger DL, and Northcott CE. 1984. Acute toxicities of organic chemicals to fathead minnows (*Pimephales promelas*). Volume 1. Superior, WI: University of Wisconsin, Center for Lake Superior Environmental Studies. pp. 414.

Geiger DL, Northcott EC, Call DJ, and Brooke LT. 1985. Acute toxicities of organic chemicals to fathead minnows (*Pimephales promelas*). Volume 2. Superior, WI: University of Wisconsin, Center for Lake Superior Environmental Studies. pp. 326.

Juhnke I and Luedemann D. 1978. Results of the investigation of 200 chemical compounds for acute fish toxicity with the golden orfe test. Z. F. Wasser-Und-Abwasser-Forschung 11(5):161-164.

Nacci D, et al 1986. Comparative evaluation of three rapid marine toxicity tests: sea urchin early embryo growth test, sea urchin sperm cell toxicity test and microtox. Environmental Toxicology and Chemistry. 5 521-525.

Phipps GL and Holcombe GW. 1985. A method for aquatic multiple species toxicant testing: Acute toxicity of 10 chemicals to 5 vertebrates and 2 invertebrates. Environ Pollut. Ser. A Ecol. Biol. 38(2):141-157.

Thurston RV, Gilfoil TA, Meyn EL, Zajdel RK, Aoki, TL, and Veith GD. 1985. Comparative toxicity of ten organic chemicals to ten common aquatic species. Water Res. 19(9):1145-1155.  
 United States Environmental Protection Agency (USEPA). 1991. Fish Chronic Toxicity Data Base. Duluth, MN: Environmental Research Laboratory (ERL), Office of Research and Development, USEPA, 6201 Congdon Boulevard, 55804; contact C.L. Russom (218) 720-5500

KETONES, DI, ALIPHATIC  
9/1993

LIST OF ALIPHATIC DIKETONES USED TO DEVELOP THE FISH 96-h LC50 SAR.

CHEMICAL	96-h LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
Methylene chloride	322.895	1.25	Z
2,4-pentanedione	116.0	-0.5	J
2,4-pentanedione	104.0	-0.5	N
5,5-dimethyl-1,3-cyclohexanedione	11,500.0	0.5	EPA
2,4-pentanedione	175.0	-0.5	G
1-benzoyl acetone	1.1	1.0	EPA
2,4-pentanedione	104.0	-0.5	B
2,4-pentanedione	146.0	-0.5	J
2,4-pentanedione	155.0	-0.5	P
2,4-pentanedione	71.6	-0.5	P
2,4-pentanedione	107.0	-0.5	P
2,4-pentanedione	83.6	-0.5	P
2,4-pentanedione	74.3	-0.5	P
2,4-pentanedione	92.4	-0.5	T
2,4-pentanedione	71.7	-0.5	T
2,4-pentanedione	66.9	-0.5	T
2,4-pentanedione	60.1	-0.5	T
2,4-pentanedione	204.0	-0.5	T
2,4-pentanedione	151.0	-0.5	T
2,4-pentanedione	106.0	-0.5	T
2,4-pentanedione	121.0	-0.5	T
2,4-pentanedione	143.0	-0.5	T
2,4-pentanedione	141.0	-0.5	T

B = Brooke et al (1984)  
EPA = USEPA (1991)  
G = Geiger et al (1985)  
J = Juhnke and Luedemann (1978)  
N = Nacci et al (1986)  
P = Phipps and Holcombe (1985)  
T = Thurston et al (1985)

**SAR**

**KETONES, DI, ALIPHATIC**

**Organism:** Daphnid  
**Duration:** 48-h  
**Endpoint:** LC50

**Equation:**  $\text{Log 48-h LC50 (mM/L)} = -0.466 - 0.467 \log K_{ow}$

**Statistics:**  $N = 6; R^2 = 0.98$

**Maximum log  $K_{ow}$ :** 5.0  
**Maximum MW:** 1000.0

**Application:** This SAR may be used to estimate toxicity for aliphatic diketones.

**Limitations:** If the  $K_{ow}$  value is greater than 5.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.

**References:** Elnabarawy MT, Welter AN, and Robideau RR. 1986. Relative sensitivity of three daphnid species to selected organic and inorganic chemicals. Environ. Toxicol. Chem. 5(4):393-398.

Mount DI and Norberg TJ. 1984. A seven-day life-cycle cladoceran toxicity test. Environ. Toxicol. Chem. 3(3):425-434.

Nacci D, et al. 1986. Comparative evaluation of three rapid marine toxicity tests: sea urchin early embryo growth test, sea urchin sperm cell toxicity test and microtox. Environmental Toxicology and Chemistry. 5:521-525.

LIST OF ALIPHATIC DIKETONES USED TO DEVELOP THE DAPHNID 48-h LC50 SAR.

CHEMICAL	48-h LC50 (mg/L)	Log $K_{ow}$	Ref.
Methylene chloride	322 895	1.25	Z
2,4-pentanedione	47.6	-0.5	N
2,4-pentanedione	75.0	-0.5	E
2,4-pentanedione	75.0	-0.5	E
2,4-pentanedione	75.0	-0.5	E
2,4-pentanedione	35.4	-0.5	M

E = Elnabarawy et al (1986)  
M = Mount and Norberg (1984)

**KETONES, DI, ALIPHATIC**  
9/1993

N = Nacci et al (1986)

## KETONES, DI, ALIPHATIC

**Organism:** Daphnid  
**Duration:** 16-d  
**Endpoint:** ChV

**Equation:**  $\text{Log ChV (mM/L)} = -1.841 - 0.482 \log K_{ow}$

**Statistics:**  $N = 4; R^2 = 0.98$

Maximum log K<sub>ow</sub>: 8.0  
Maximum MW: 1000.0

**Application:** This SAR may be used to estimate toxicity for aliphatic diketones.

**Limitations:** If the log Kow value is greater than 8.0, or if the compound is solid and the ChV exceeds the water solubility, no effects expected at saturation.

**References:** Elnabarawy MT, Welter AN, and Robideau RR. 1986. Relative sensitivity of three daphnid species to selected organic and inorganic chemicals. Environ. Toxicol. Chem. 5(4):393-398.

LIST OF ALIPHATIC DIKETONES USED TO DEVELOP THE DAPHNID ChV SAR.

CHEMICAL	ChV (mg/L)	Log K <sub>ow</sub>	Ref.
Methylene chloride	322.895	1.25	Z
2,4-pentanedione	6.5	-0.5	E
2,4-pentanedione	2.6	-0.5	E
2,4-pentanedione	1.0	-0.5	E

E = Elnabarawy et al. (1986)

**KETONES, DI, ALIPHATIC**  
9/1993

<b>SAR</b>	<b>KETONES, DI, ALIPHATIC</b>
<b>Organism:</b>	Green Algae
<b>Duration:</b>	
<b>Endpoint:</b>	ChV
<b>Equation:</b>	$\text{Log ChV (mM/L)} = -1.806 - 0.412 \log K_{ow}$
<b>Statistics:</b>	$N = 2; R^2 = 1.0$
<b>Maximum log <math>K_{ow}</math>:</b>	8.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This SAR may be used to estimate toxicity for aliphatic diketones.
<b>Limitations:</b>	If the log Kow value is greater than 8.0, or if the compound is solid and the ChV exceeds the water solubility, no effects expected at saturation
<b>References:</b>	Bringmann G and Kuhn R. 1980. Comparison of the toxicity thresholds of water pollutants to bacteria, algae, and protozoa in the cell multiplication inhibition test. Water Res. 14(3):231-241.

LIST OF ALIPHATIC DIKETONES USED TO DEVELOP THE GREEN ALGAE ChV SAR.

CHEMICAL	ChV (mg/L)	Log $K_{ow}$	Ref.
2,4-pentanedione	2.7	-0.5	BK

BK = Bringmann and Kuhn (1980)

**KETONES, DI, ALIPHATIC**  
9/1993

**SAR**

**MALONONITRILES**

<b>Organism:</b>	Fish
<b>Duration:</b>	96-h
<b>Endpoint:</b>	LC50 (Mortality)
<b>Equation:</b>	$\text{Log 96-h LC50 (mM/L)} = -2.079 - 0.139 \log K_{ow}$
<b>Statistics:</b>	$N = 3; R^2 = 0.40$
<b>Maximum log <math>K_{ow}</math>:</b>	5.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This SAR may be used to estimate toxicity for malononitriles.
<b>Limitations:</b>	For malononitriles with log $K_{ow}$ values greater than 5.0, a test duration of greater than 96 hours may be required for proper expression of toxicity. Also, if the acute toxicity value obtained by the use of this equation exceeds the water solubility of the compound (measured or estimated), significant mortalities would not be expected in a saturated solution during an exposure period of 96 hours.
<b>References:</b>	Abram FSH and Wilson P. 1979. The acute toxicity of CS to rainbow trout. Water Research 13:631-635.

LIST OF MALONONITRILES USED TO DEVELOP THE FISH 96-H LC50 SAR.

CHEMICAL	96-h LC50 (mg/L)	Log $K_{ow}$	Ref.
Malononitrile	1.6	-1.2	A
o-Chlorobenzylidene malononitrile	0.22	1.8	A

A = Abram and Wilson (1979)

**MALONONITRILES**  
9/1993

## SAR

## NEUTRAL ORGANICS

Organism:

Fish

Duration:

96-h

Endpoint:

LC50 (Mortality)

Equation:

$$\text{Log LC50 (mM/L)} = 1.75 - 0.94 \log K_{ow}$$

Statistics:

$$N = 60; R^2 = 0.942$$

Maximum  $K_{ow}$ :

5.0

Maximum MW:

1000.0

Application:

Solvents, non-reactive, non-ionizable neutral organic compounds

1. Alcohols
2. Acetals
3. Ketones
4. Ethers
5. Alkyl halides
6. Aryl halides
7. Aromatic hydrocarbons
8. Halogenated aromatic hydrocarbons
9. Halogenated aliphatic hydrocarbons
10. Sulfides and di-sulfides

Limitations:

Use the fish 14-day LC50 for neutral organics with  $\log K_{ow}$  greater than 5 and less than 7. If the compound is and the LC50 is exceeds the water solubility, use SAR with longer exposure.

References:

Veith GD, Call DJ, and Brooke LT. 1983. Structure-toxicity relationships for the fathead minnow, Pimephales promelas: narcotic industrial chemicals. Canadian Journal of Fisheries and Aquatic Sciences 40:743-748.

# NEUTRAL ORGANICS

7/1988

## LIST OF NEUTRAL ORGANICS USED TO DEVELOP THE FISH 96-h LC50 SAR.

CHEMICAL	96-h LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
Triethylene glycol	69800	-1.17	V
2-Methyl-2,4-pentandiol	10700	-0.70	V
Methanol	28100	-0.66	V
Acetone	8120	-0.24	V
Ethanol	14200	-0.16	V
2-(2-Ethoxyethoxy)ethanol	26400	-0.08	V
2-Propanol	10400	0.14	V
2-Butanone	3200	0.28	V
3-Furanmethanol (static)	508	0.32	V
Tetrahydrofuran	2160	0.46	V
3-Methyl-2-butanone	864	0.62	V
2-Methyl-1-propanol	1430	0.74	V
Cyclohexanone	527	0.81	V
3-Pentanone	1540	0.84	V
1-Butanol	1730	0.88	V
3,3-Dimethyl-2-butanone	87	0.94	V
2',3',4'-Trimethoxyacetophenone	172	1.12	V
2-Phenoxyethanol	344	1.16	V
Cyclohexanol	704	1.23	V
4-Methyl-2-pentanone	505	1.25	V
t-Butylmethyl ether	706	1.30	V
Furan	61	1.34	V
2,2,2-Trichloroethanol	299	1.38	V
Diisopropyl ether	91.7	1.56	V
Acetophenone	162	1.66	V
5-Methyl-2-hexanone	159	1.79	V
1,3-Dichloroethane	118	1.79	V
p-Dimethoxybenzene	117	2.00	V
1-Fluoro-4-nitrobenzene	28.4	2.02	V
1-Hexanol	97.5	2.03	V
1,1,2-Trichloroethane	81.7	2.07	V
6-Methyl-5-heptene-2-one	85.7	2.13	V
2'-Hydroxy-4'-methoxyacetophenone	54.9	2.14	V
1,1,2,2-Tetrachloroethane	20.3	2.39	V
1,1,2-Trichloroethylene	44.1	2.42	V
2-Octanone	36	2.46	V
Tetrachloroethane	13.5	2.53	V
2,6-Dimethoxytoluene	20.5	2.67	V
5-Nonanone	31	3.00	V
2',4'-Dichloroacetophenone	11.7	3.02	V
1-Octanol	13.5	3.03	V
Di-n-butyl ether	32.5	3.08	V

Continued.

CHEMICAL	96-h LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
1,4-Dichlorobenzene	4.0	3.37	V
Benzophenone	15.3	3.38	V
1,3-Dichlorobenzene	7.8	3.38	V
1-Nonanol	5.7	3.53	V
2-Decanone	5.7	3.54	V
Pentachloroethane	7.3	3.64	V
2',3',4'-Trichloroacetophenone	2.0	3.73	V
p-Nitrophenyl phenyl ether	2.7	3.97	V
1-Decanol	2.4	4.03	V
Dipentyl ether	3.2	4.16	V
3,4-Dichlorotoluene	2.91	4.22	V
$\alpha,\alpha$ -2,6-Tetrachlorotoluene	0.97	4.24	V
Diphenyl ether	4.0	4.26	V
1,2,4-Trichlorobenzene	2.9	4.28	V
1-Undecanol	1.04	4.53	V
Hexachloroethane	1.5	4.62	V
1-Dodecanol	1.01	5.00	V
7-Tridecanone	*	5.16	V
1-Tridecanol	*	5.51	V
Pentachlorobenzene	*	5.71	V
1,2,3,4-Tetrachlorobenzene	1.1	5.71	V
Hexachlorobenzene	*	5.71	V
Diethyl ether	*	6.42	V

\* = No fish mortality in saturated solutions.

V = Veith et al. (1983)

## NEUTRAL ORGANICS

7/1988

**SAR****NEUTRAL ORGANICS**

**Organism:** Fish, Sheepshead Minnow (marine)  
**Duration:** 96-h  
**Endpoint:** LC50 (Mortality)

**Endpoint:**  $\text{Log LC50 (mM/L)} = 0.69 - 0.73 \log K_{ow}$

**Statistics:**  $N = 37; R^2 = 0.656$

**Maximum  $K_{ow}$ :** 5.0

**Maximum MW:** 1000.0

**Application:** Solvents, non-reactive, non-ionizable neutral organic compounds:

1. Alcohols
2. Acetals
3. Ketones
4. Ethers
5. Alkyl halides
6. Aryl halides
7. Aromatic hydrocarbons
8. Halogenated aromatic hydrocarbons
9. Halogenated aliphatic hydrocarbons
10. Sulfides and di-sulfides

**Limitations:** If the  $\log K_{ow}$  is greater than 5, or if the compound is solid and the LC50 exceeds the water solubility, use SAR with longer exposure.

**References:** Zarogian G, Heltshe JF, and Johnson M. 1985. Estimation of toxicity to marine species with structure activity models developed to estimate toxicity to freshwater fish. *Aquatic Toxicology* 6:251-270

# NEUTRAL ORGANICS

7/1988

## LIST OF NEUTRAL ORGANICS USED TO DEVELOP THE SHEEPSHEAD MINNOW 96-h LC50 SAR.

CHEMICAL	96-h LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
Methylene chloride	322.895	1.25	Z
Diethyl phthalate	29.979	1.40	Z
1,1-Dichloroethylene	249.174	1.48	Z
2,4-Dinitrophenol	28.515	1.53	Z
Dimethyl phthalate	57.310	1.61	Z
Nitrobenzene	58.924	1.83	Z
1,1,1,2,2,2-Hexachloroethane	1.380	1.91	Z
4-Nitrophenol	26.507	1.91	Z
1,3-Dichloropropene	1.759	1.98	Z
2,3-Dinitrotoluene	2.293	1.98	Z
2,4,6-Trinitrophenol	128.838	2.03	Z
Bromoform	17.893	2.30	Z
4-Chlorophenol	5.359	2.35	Z
1,1,2,2-Tetrachloroethane	11.883	2.39	Z
1,1,1-Trichloroethane	70.015	2.47	Z
1,1,1,2,2-Pentachloroethane	113.762	2.89	Z
Diazinon	1.457	3.14	Z
1,4-Dichlorobenzene	7.200	3.38	Z
1,2-Dichlorobenzene	9.491	3.40	Z
1,3-Dichlorobenzene	7.715	3.44	Z
2,4,5-Trichlorophenol	1.681	3.72	Z
Hexachlorobutadiene	0.545	3.74	Z
Chlorobenzene	9.804	3.79	Z
Disulfoton	0.739	3.81	Z
Lindane	0.801	3.89	Z
Dibenzofuran	1.761	4.10	Z
Diphenyl ether	2.350	4.21	Z
Dieldrin	0.010	4.31	Z
1,2,4-Trichlorobenzene	20.833	4.32	Z
1,2,3,5-Tetrachlorobenzene	3.666	4.46	Z
1,2,4,5-Tetrachlorobenzene	0.784	4.67	Z
Methoxychlor	0.049	4.68	Z
Chloropyrifos	0.881	4.82	Z
Heptachlor	0.004	5.44	Z
Kepone	0.693	6.08	Z
Fenvalerate	0.004	6.20	Z
Parmethrin	0.068	6.50	Z

Z = Zarogian et al. (1985)

**SAR**

**NEUTRAL ORGANICS**

**Organism:**

Fish

**Duration:**

14-day

**Endpoint:**

LC50 (Mortality)

**Equation:**

$$\text{Log LC50 (mM/L)} = 1.87 - 0.871 \log K_{ow}$$

**Statistics:**

$$N = 50; R^2 = 0.976$$

**Maximum  $K_{ow}$ :**

8.0

**Maximum MW:**

1000.0

**Application:**

Solvents, non-reactive, non-ionizable neutral organic compounds:

1. Aromatic hydrocarbons
2. Halogenated aromatic hydrocarbons
3. Halogenated aliphatic hydrocarbons
4. Alcohols
5. Ketones
6. Acetals
7. Ethers
8. Alkyl halides
9. Aryl halides
10. Sulfides and di-sulfides

Also applicable to reactive compounds (i.e., compounds which show excess toxicity) whose  $\log K_{ow}$  is greater than 5.0, such as:

1. Esters
2. Acrylates
3. Methacrylates
4. Substituted benzotriazoles

**Limitations:**

If the  $\log K_{ow}$  is greater than 8.0, or if the compound is solid and the LC50 exceeds the water solubility, use SAR with longer exposure.

**References:**

Konemann H. 1981. Quantitative structure-activity relationships in fish toxicity studies. Part 1: Relationship for 50 industrial pollutants. Toxicology 19(3):209-221.

# NEUTRAL ORGANICS

7/1988

## LIST OF NEUTRAL ORGANICS USED TO DEVELOP THE FISH 14-d LC50 SAR.

CHEMICAL	14-d LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
Ethanediol (ethyleneglycol)	49303.00	-1.35	K
Digol (diethyleneglycol)	61065.00	-1.30	K
Trigol (triethyleneglycol)	62601.00	-1.24	K
2-Methoxyethanol	17433.00	-0.74	K
Acetone	6368.00	-0.30	K
Ethanol	11051.00	-0.26	K
2-Ethoxyethanol	16399.00	-0.21	K
Propanol-2	7061.00	0.15	K
2-Isopropoxyethanol	5467.00	0.20	K
2-Methylpropanol-2	3547.00	0.77	K
2-Butoxyethanol	983.00	0.86	K
Diethylether	2137.00	0.88	K
Butyldigol	1148.00	0.91	K
Butyltrigol	197.00	0.97	K
Pentanol-3	989.00	1.21	K
Dichloromethane	294.00	1.61	K
1,3-Dichloropropane	83.80	1.71	K
1,2-Dichloroethane	106.00	1.76	K
2,2'-Dichlorodiethylether	54.40	1.81	K
1,1-Dichloroethane	202.00	1.92	K
Chloroform	102.00	2.02	K
Trans-1,4-dichloro-2-butene	39.50	2.11	K
Benzene	63.50	2.13	K
1,2-Dichloropropane	115.00	2.16	K
Trichloroethane	55.60	2.20	K
1-Chlorobutane	96.90	2.35	K
1,1,2-Trichloroethane	94.40	2.38	K
2,4-Dichloroaniline	11.70	2.42	K
1,1,1-Trichloroethane	133.00	2.49	K
Toluene	68.30	2.59	K
2,3-Dichloro-1-propane	11.10	2.60	K
1,2,3-Trichloropropane	41.60	2.63	K
1,5-Dichloropentane	11.20	2.77	K
Tetrachloromethane	67.10	2.79	K
Monochlorobenzene	19.10	2.81	K
α,α'-Dichloro-m-xylene	0.12	2.87	K
Tetrachloroethane	18.00	2.95	K
1,1,2,2-Tetrachloroethane	36.70	3.01	K
o-Xylene	35.20	3.09	K
m-Xylene	37.70	3.09	K
p-Xylene	35.20	3.09	K
Cyclohexane	84.20	3.18	K

Continued.

CHEMICAL	14-d LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
4-Chlorotoluene	5.90	3.31	K
3-Chlorotoluene	18.30	3.31	K
1,2-Dichlorobenzene	5.80	3.53	K
1,3-Dichlorobenzene	7.40	3.53	K
1,4-Dichlorobenzene	4.00	3.53	K
Pentachloroethane	15.00	3.58	K
2,4,α-Trichlorotoluene	0.24	3.87	K
2,4-Dichlorotoluene	4.60	3.98	K
3,4-Dichlorotoluene	5.10	3.98	K
1,2,3-Trichlorobenzene	2.30	4.20	K
1,2,4-Trichlorobenzene	2.40	4.20	K
1,3,5-Trichlorobenzene	3.30	4.20	K
Hexachlorobutadiene	0.39	4.63	K
2,4,5-Trichlorotoluene	1.70	4.72	K
1,2,4,5-Tetrachlorobenzene	0.30	4.94	K
1,2,3,5-Tetrachlorobenzene	0.80	4.94	K
1,2,3,4-Tetrachlorobenzene	0.80	4.94	K
Pentachlorobenzene	0.18	5.69	K
Hexachlorobenzene	0.32	6.44	K

K = Konemann (1981)

**NEUTRAL ORGANICS**  
7/1988

**SAR****NEUTRAL ORGANICS**

<b>Organism:</b>	Fish
<b>Duration:</b>	> 30 days
<b>Endpoint:</b>	Chronic Value (Survival/Growth; Early Life Stage)
<b>Equation:</b>	$\text{Log ChV (mM/L)} = 0.72 - 0.87 \log K_{ow}$
<b>Statistics:</b>	$N = 20; R^2 = 0.91$
<b>Maximum <math>K_{ow}</math>:</b>	7.9
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	Solvents, non-reactive, non-ionizable neutral organic compounds: <ol style="list-style-type: none"><li>1. Alcohols</li><li>2. Acetals</li><li>3. Ketones</li><li>4. Ethers</li><li>5. Alkyl halides</li><li>6. Aryl halides</li><li>7. Aromatic hydrocarbons</li><li>8. Halogenated aromatic hydrocarbons</li><li>9. Halogenated aliphatic hydrocarbons</li><li>10. Sulfides and di-sulfides</li></ol>
<b>Limitations:</b>	If the $\log K_{ow}$ is greater than 7.9 or if the compound is solid and the ChV exceeds the water solubility, no effects expected at saturation.
<b>References:</b>	United States Environmental Protection Agency (USEPA). 1991 Fish chronic toxicity data base. Duluth, MN: Environmental Research Laboratory (ERL), Office of Research and Development, USEPA, 6201 Congdon Blvd., 55804; contact C L. Russom (218) 720-5500.

## NEUTRAL ORGANICS

7/1988

**SAR**

**NEUTRAL ORGANICS**

**Organism:** Fish, Fathead minnow  
**Duration:** 28-d  
**Endpoint:** BCF (Bioconcentration Factor)  
**Equation:**  $\text{Log BCF} = 0.79 \log K_{ow} - 0.40$  (the BCF is without units)  
**Statistics:**  $N = 122$ ;  $R_2 = 0.927$

**Maximum  $K_{ow}$ :** 8.0  
**Maximum MW:** 1000.0

**Application:** Solvents, non-reactive, non-ionizable compounds:

1. Aromatic amines
2. Acetals
3. Cycloidiene
4. Ethers
5. Halogenated alkyl
6. Halogenated aromatic
7. Halogenated indoles
8. Halogenated phenols
9. Phosphate esters

**Limitations:** If  $\log K_{ow}$  is greater than 8.0, no significant BCF unless analog data can be found, e.g., PCBs.

**Reference:** Veith, GD, and Kosian, P. 1982. Estimating bioconcentration potential from octanol/water partition coefficients. IN: Physical Behavior of PCB's in the Great Lakes. MacKay, Paterson, Eisenreich, and Simmons, eds. Ann Arbor, MI: Ann Arbor Science.

# NEUTRAL ORGANICS

7/1988

## LIST OF CHEMICALS USED TO DEVELOP THE FISH BIOCONCENTRATION SAR.

CHEMICAL	Log BCF	Log K <sub>ow</sub>	Ref.
Lindane	2.67	3.85	VK
Atrazine	0.90	2.63	VK
Heptachlor	4.30	5.44	VK
2-Ethylhexiphthalate	2.93	4.20	VK
DASC-3	0.32	1.00	VK
DASC-4	0.32	1.00	VK
NTS-1	0.66	1.00	VK
BSB	0.32	1.00	VK
FWA-2-A	0.32	1.80	VK
FWA-3-A	0.32	1.48	VK
FWA-4-A	0.32	1.20	VK
Nitrobenzene	1.18	2.93	VK
p-Nitrophenol	1.88	1.91	VK
Naphthalene	2.63	3.59	VK
Chlorobenzene	2.65	3.79	VK
2,4,5-Trichlorophenol	3.28	3.72	VK
Endrin	3.66	4.56	VK
1,1,2,2-Tetrachloroethylene	2.06	2.88	VK
Hexachlorobenzene	4.37	6.18	VK
p-Biphenylphenyl ether	3.22	5.55	VK
Carbon tetrachloride	2.77	4.21	VK
p-Dichlorobenzene	1.72	2.64	VK
Biphenyl	2.81	3.38	VK
Chloropyrifos	2.67	4.82	VK
Endrin	3.17	4.56	VK
2,5,6-Trichloropyridinol	0.49	1.35	VK
Fluorene	3.11	4.38	VK
Dibenzofuran	3.13	4.12	VK
2-Chlorophenanthrene	3.63	5.16	VK
Phenanthrene	3.42	4.46	VK
2-Methylphenanthrene	3.48	4.86	VK
Heptachlor	3.98	5.44	VK
Heptachloroepoxide	4.16	5.40	VK
p,p'-DDE	4.71	5.69	VK
Pentachlorophenol	2.89	2.97	VK
Hexabromobiphenyl	6.39	4.26	VK
Methoxychlor	3.92	4.30	VK
Mirex	4.26	6.89	VK
Hexabromocyclododecane	4.26	5.81	VK
Heptachloronorborene	4.05	5.28	VK
Hexachloronorborene	3.81	5.28	VK
1,2-Dichlorobenzene	1.95	3.40	VK
1,3-Dichlorobenzene	1.82	3.44	VK

## NEUTRAL ORGANICS

7/1988

Continued.

CHEMICAL	Log BCF	Log K <sub>ow</sub>	Ref.
1,4-Dichlorobenzene	1.78	3.37	VK
1,2,3,5-Tetrachlorobenzene	3.26	4.46	VK
Pentachlorobenzene	3.53	4.94	VK
Carbon tetrachloride	1.48	2.73	VK
Chloroform	0.78	1.90	VK
1,2-Dichloroethane	0.30	1.45	VK
1,1,1-Trichloroethane	0.95	2.47	VK
1,1,2,2-Tetrachloroethane	0.90	2.39	VK
Pentachloroethane	1.83	3.21	VK
Hexachloroethane	2.14	3.93	VK
Bis(2-chloroethyl) ether	1.04	1.12	VK
1,1,2-Trichloroethylene	1.23	2.42	VK
Tetrachloroethylene	1.69	2.53	VK
Isophorone	0.84	1.67	VK
N-Nitrosophenylamine	2.34	3.13	VK
2-Chlorophenol	2.33	2.16	VK
2,4-Dimethylphenol	2.33	2.16	VK
Butylbenzylphthalate	2.89	4.05	VK
Dimethylphthalate	1.76	1.61	VK
Alkyl benzene sulfonate	2.02	1.59	VK
Alkyl benzene sulfonate	1.56	1.59	VK
Naphthalene	1.90	3.59	VK
2-Methylnaphthalene	2.28	3.84	VK
1-Methylnaphthalene	2.11	3.84	VK
Hexachlorocyclohexane	2.15	3.85	VK
Hexachlorocyclohexane	2.70	3.85	VK
Endrin	4.02	4.56	VK
Endrin	4.18	4.56	VK
Endrin	3.85	4.56	VK
Al254	4.60	6.47	VK
Al254	4.43	6.47	VK
1,4-Dichlorobenzene	1.96	3.37	VK
1,2,3-Trichlorobenzene	2.81	4.20	VK
1,3,5-Trichlorobenzene	2.85	4.20	VK
1,2,3,5-Tetrachlorobenzene	3.56	4.46	VK
Pentachlorobenzene	4.11	4.94	VK
Hexachlorobenzene	4.16	6.18	VK
Aroclor 1016	4.63	5.86	VK
Aroclor 1248	4.85	6.11	VK
Aroclor 1254	5.00	6.47	VK
Aroclor 1260	5.29	6.91	VK
Chlordane	4.58	6.00	VK
Octachlorostyrene	4.52	6.29	VK
p,p-DDT	4.47	5.75	VK

# NEUTRAL ORGANICS

7/1988

Continued.

CHEMICAL	Log BCF	Log K <sub>ow</sub>	Ref.
o,p-DDT	4.57	5.75	VK
Hexachlorobenzene	4.27	6.18	VK
1,2,4-Trichlorobenzene	3.32	4.23	VK
Lindane	2.26	3.85	VK
5-Bromoindole	1.15	2.97	VK
2,4,6-Tribromoanisol	2.94	4.48	VK
N-Phenyl-2-naphylamine	2.17	4.38	VK
Tricresyl phosphate	2.22	3.42	VK
Diphenyl amine	1.48	3.42	VK
Toluene	1.96	3.16	VK
1,1,2,2-Tetrachloroethylene	0.91	2.39	VK
Pentachloroethane	1.78	3.21	VK
Hexachloroethane	2.85	3.93	VK
1,3-Dichlorobenzene	1.99	3.44	VK
1,4-Dichlorobenzene	2.05	3.37	VK
1,2,4-Trichlorobenzene	2.60	4.52	VK
1,2,3,4-Tetrachlorobenzene	3.41	4.46	VK
Hexachlorobenzene	4.37	6.18	VK
Hexachloro-1,3-butadiene	3.84	5.10	VK
Acridine	2.10	3.30	VK
Toxaphene	3.64	5.28	VK
Toxaphene	3.59	5.28	VK
Pentachlorophenol	1.11	2.97	VK
Imidan	0.90	2.83	VK
Imidan	1.04	2.83	VK
Imidan	0.90	2.83	VK
Diazinon	1.56	1.92	VK
Diazinon	1.81	1.92	VK
Diazinon	1.24	1.92	VK
Endrin	3.21	4.56	VK
Acenaphthene	2.59	3.92	VK

VK = Veith and Kosian (1982)

SAR

NEUTRAL ORGANICS

**Organism:** Daphnid  
**Duration:** 48-h  
**Endpoint:** LC50 (Mortality)  
**Equation:**  $\text{Log LC50 (mM/L)} = 1.72 - 0.91 \log K_{ow}$   
**Statistics:**  $N = 19; R^2 = 0.992$   
**Maximum  $K_{ow}$ :** 5.0  
**Maximum MW:** 1000.0

**Application:** Solvents, non-reactive, non-ionizable compounds:

1. Aromatic hydrocarbons
2. Halogenated aromatic hydrocarbons
3. Halogenated aliphatic hydrocarbons
4. Alcohols
5. Ketones
6. Acetals
7. Ethers
8. Alkyl halides
9. Aryl halides
10. Sulfides and di-sulfides

Also be applied to some classes of reactive organic compounds which show excess toxicity to fish, such as:

1. Benzotriazoles
2. Phthalate esters
3. Esters

**LIMITATIONS:** If the  $\log K_{ow}$  is greater than 5.0 and less than 8.0, or if the compound is solid and the LC50 exceeds the water solubility, use SAR with longer exposure.

**References:** Hermans J, Canton H, Janssen P, and De Jong R. 1984. Quantitative structure-activity relationships and toxicity studies of mixtures of chemicals with anaesthetic potency: Acute lethal and sublethal toxicity to Daphnia magna Aquatic Toxicology 5.143-154.

**NEUTRAL ORGANICS**

7/1988

**LIST OF NEUTRAL ORGANICS USED TO DEVELOP THE DAPHNID 48-h LC50 SAR.**

CHEMICAL	48-h LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
Ethenediol	50452	-1.35	H
Acetone	6081	-0.30	H
Ethanol	5413	-0.26	H
2-Ethoxyethanol	7670	-0.21	H
Diethylether	1380	0.88	H
Benzene	56.6	2.13	H
1,2-Dichloropropane	45.0	2.16	H
1,1,2-Trichloroethene	20.8	2.20	H
Toluene	14.9	2.59	H
1,2,3-Trichloropropane	35.4	2.63	H
Monochlorobenzene	25.8	2.81	H
m-Xylene	14.3	3.09	H
4-Chlorotoluene	3.6	3.31	H
1,2-Dichlorobenzene	3.8	3.53	H
2,4-Dichlorotoluene	0.62	3.98	H
1,2,4-Trichlorobenzene	2.7	4.20	H
2,4,5-Trichlorobenzene	0.55	4.72	H
1,2,3,4-Tetrachlorobenzene	0.54	4.94	H
Pentachlorobenzene	0.12	5.69	H

H = Hermans et al. (1984)

**SAR**

**NEUTRAL ORGANICS**

**Organism:** Mysid shrimp  
**Duration:** 96-h  
**Endpoint:** LC50 (Mortality)  
**Equation:**  $\text{Log LC50 (mM/L)} = 1.83 - 1.25 \log K_{ow}$   
**Statistics:**  $N = 17; R^2 = 0.706$   
**Maximum  $K_{ow}$ :** 5.0  
**Maximum MW:** 1000.0  
**Application:** Solvents, non-reactive, non-ionizable compounds:

1. Alcohols
2. Acetals
3. Ketones
4. Ethers
5. Alkyl halides
6. Aryl halides
7. Aromatic hydrocarbons
8. Halogenated aromatic hydrocarbons
9. Halogenated aliphatic hydrocarbons
10. Sulfides and di-sulfides

**Limitations:** If the  $\log K_{ow}$  is greater than 5.0, or if the compound is solid and the LC50 exceeds the water solubility, use SAR with longer exposure.

**References:** Zaroogian G, Heltshe JF, and Johnson M. 1985. Estimation of toxicity to marine species with structure activity models developed to estimate toxicity to freshwater fish. *Aquatic Toxicology* 6 251-270.

**NEUTRAL ORGANICS**

7/1988

**LIST OF NEUTRAL ORGANICS USED TO DEVELOP THE MYSID SHRIMP 96-h LC50 SAR.**

CHEMICAL	96-h LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
Toluene	55.5198	2.21	Z
1,3-Dichloropropane	10.0702	2.28	Z
Tetrachloroethylene	9.9922	2.60	Z
Benthiocarb	0.3235	3.40	Z
Hexachlorobutadiene	0.0611	3.74	Z
Chlorobenzene	16.2699	3.79	Z
EPN	0.0032	3.85	Z
Lindane	0.0059	3.89	Z
Dieldrin	0.0050	4.31	Z
1,2,4-Trichlorobenzene	0.4454	4.32	Z
1,2,3,5-Tetrachlorobenzene	0.3344	4.46	Z
Acenaphthene	0.0250	4.49	Z
1,2,4,5-Tetrachlorobenzene	1.4596	4.67	Z
Pentachlorobenzene	0.1616	4.94	Z
Heptachlor	0.0030	5.44	Z
Leptophos	0.0033	6.08	Z
Fenvalerate	0.0001	6.20	Z

Z = Zaroogian et al. (1985)

**SAR**

**NEUTRAL ORGANICS**

**Organism:** Daphnid  
**Duration:** 16-d  
**Endpoint:** Chronic Value (EC50 Reproduction)

**Equation:**  $\text{Log ChV (mM/L)} = -0.72 \log K_{ow} + 0.05$

**Statistics:**  $N = 5; R^2 = 0.990$

**Maximum  $K_{ow}$ :** 8.0  
**Maximum MW:** 1000.0

**Application:** Solvents, non-reactive, non-ionizable compounds:

1. Aromatic hydrocarbons
2. Halogenated aromatic hydrocarbons
3. Halogenated aliphatic hydrocarbons
4. Alcohols
5. Ketones
6. Acetals
7. Ethers
8. Alkyl halides
9. Aryl halides
10. Sulfides and di-sulfides

This SAR can also be applied to some classes of reactive organic compounds which show excess toxicity to fish, such as:

1. Benzotriazoles
2. Phthalate esters
3. Esters

**Limitations:** If the  $\log K_{ow}$  is greater than 8.0, or if the compound is solid and the ChV exceeds the water solubility, no effects expected at saturation.

**References:** Hermans J, Canton H, Janssen P, and De Jong R. 1984. Quantitative structure-activity relationships and toxicity studies of mixtures of chemicals with anaesthetic potency: Acute lethal and sublethal toxicity to Daphnia magna. Aquatic Toxicology 5:143-154.

## NEUTRAL ORGANICS

7/1988

### LIST OF NEUTRAL ORGANICS USED TO DEVELOP THE DAPHNID 16-d EC50 SAR.

CHEMICAL	16-d EC50 (mg/L)	Log K <sub>ow</sub>	Ref.
Monochlorobenzene	25.8	2.81	H
4-Chlorotoluene	3.6	3.31	H
1,2,4-Trichlorobenzene	2.7	4.20	H
1,2,3,4-Tetrachlorobenzene	0.54	4.94	H
Pentachlorobenzene	0.23	5.96	H

H = Hermans et al (1984)

**SAR**

**NEUTRAL ORGANICS**

**Organism:** Green algae

**Duration:** 96-h

**Endpoint:** EC50 (Growth)

**Equation:**  $\text{Log 96-h EC50 (mM/L)} = 1.466 - 0.885 \log K_{ow}$

**Statistics:**  $N = 7; R^2 = 0.91$

**Maximum  $K_{ow}$ :** 6.4

**Maximum MW:** 1000.0

**Application:**

**Limitations:** If the  $\log K_{ow}$  is greater than 6.4, or if the compound is solid and the EC50 exceeds the water solubility, use SAR with longer exposure.

**References:**

Calamari D, Galassi S, Setti F, and Vighi M. 1983. Toxicity of selected chlorobenzenes to aquatic organisms. *Chemosphere* 12:253-262.

Galassi S and Vighi M. 1981. Testing toxicity of volatile substances with algae. *Chemosphere* 10:1123-1126.

United States Environmental Protection Agency (USEPA). 1991. OTS PMN ECOTOX. Washington, DC: USEPA, Office of Toxic Substances.

United States Environmental Protection Agency (USEPA). 1992. Aquatic toxicity database. Duluth, MN: USEPA, ERL - Duluth.

# NEUTRAL ORGANICS

7/1988

## LIST OF NEUTRAL ORGANICS USED TO DEVELOP THE GREEN ALGAE 96-h EC50 SAR.

CHEMICAL	96-h EC50 (mg/L)	Log K <sub>ow</sub>	Ref.
Polyether	315	1.9	EPA
Berizene	29	2.1	G
Isolinalool	14	2.4	EPA
Toluene	12.5	2.8	G
Chlorobenzene	12.5	2.9	C
trans-Anethole	4.24	3.3	D
Ethylbenzene	4.6	3.3	G
o-Xylene	4.7	3.4	G
m-Xylene	4.9	3.4	G
p-Xylene	3.2	3.4	G
1,2-Dichlorobenzene	2.2	3.6	C
1,4-Dichlorobenzene	0.57	3.6	C
Isopropylbenzene	2.6	3.7	G
n-Propylbenzene	1.8	3.8	G
1,2,3-Trichlorobenzene	0.22	4.3	C
1,2,4-Trichlorobenzene	0.37	4.3	C
Hexachlorobenzene	*	6.4	C

\* = No effects in saturated solutions

C = Calamari et al. (1983)

D = USEPA (1992).

EPA = USEPA (1991); chemical identity is Confidential Business Information under TSCA

G = Galassi and Vighi (1988)

**SAR**                      **NEUTRAL ORGANICS****Organism:** Green algae**Duration:****Endpoint:** Chronic Value (Growth)**Equation:**  $\text{Log ChV (mM/L)} = -0.036 - 0.634 \log K_{ow}$ **Statistics:**  $N = 7; R^2 = 0.99$ **Maximum  $K_{ow}$ :** 8.0**Maximum MW:** 1000.0**Applications:** May be applied to other neutral organics including aldehydes.**Limitations:** If the  $\log K_{ow}$  is greater than 8.0, or if the compound is solid and the ChV exceeds the water solubility, no effects expected at saturation.**References:** Calamari D, Galassi S, Setti F, and Vighi M. 1983. Toxicity of selected chlorobenzenes to aquatic organisms. Chemosphere 12:253-262.

Galassi S and Vighi M. 1981. Testing toxicity of volatile substances with algae. Chemosphere 10:1123-1126.

United States Environmental Protection Agency (USEPA). 1991. OTS PMN ECOTOX. Washington, DC. USEPA, Office of Toxic Substances.

United States Environmental Protection Agency (USEPA). 1992. Aquatic toxicity database. Duluth, MN: USEPA, ERL - Duluth.

## LIST OF NEUTRAL ORGANICS USED TO DEVELOP THE GREEN ALGAE ChV SAR.

CHEMICAL	ChV (mg/L)	Log $K_{ow}$	Ref
Polyether	15.9	1.9	EPA
Isolinalool	4.8	2.4	EPA
trans-Anethole	3.09	3.3	D
1,4-Dichlorobenzene	0.57	3.6	C
1,2,3-Trichlorobenzene	0.22	4.3	C
1,2,4-Trichlorobenzene	0.37	4.3	C
Hexachlorobenzene	0.027	6.4	C

## NEUTRAL ORGANICS

7/1988

C = Calamari et al. (1983)

D = USEPA (1992)

EPA = USEPA (1991)

**SAR**                      **NEUTRAL ORGANICS**

**Organism:** Earthworm

**Duration:** 14-d

**Endpoint:** LC50 (Mortality)

**Equation:**  $\text{Log 14-d LC50 (mM/L)} = 1.405 - 0.308 \log K_{ow}$

**Statistics:**  $N = 5; R^2 = 0.48$

**Maximum  $K_{ow}$ :** 5.0

**Maximum MW:** 1000.0

**Applications:** Neutral organics

**Limitations:** None

**References:** Neuhauser EF, Durkin PR, Malecki MR, and Anatra M. 1986. Comparative toxicity of ten organic chemicals to four earthworm species. Comp. Biochem. Physiol. 83C:197-200.

Neuhauser EF, Loehr RC, Malecki MR, Milligan DL, and Durkin PR. 1985 The toxicity of selected organic chemicals to the earthworm Eisenia fetida. Journal of Environmental Quality 14:383-388.

LIST OF NEUTRAL ORGANIC CHEMICALS USED TO DEVELOP THE EARTHWORM 14-d LC50 SAR.

CHEMICAL	14-d LC50 (mg/L)	Log $K_{ow}$	Ref.
2-chloroethylvinylether	740.0	1.0	N
nitrobenzene	319.0	1.9	N
1,2-dichloropropane	4240.0	2.0	N
fluorene	173.0	4.2	N
1,2,4-trichlorobenzene	197.0	4.3	N

N = Neuhauser et al. (1985, 1986)

## NEUTRAL ORGANICS

7/1988

**SAR****PEROXY ACIDS****Organism:**

Fish

**Duration:**

96-h

**Endpoint:**

LC50 (Mortality)

**Equation:** $\log 96\text{-h LC50 (mM/L)} = -2.6 \log K_{ow}$ **Statistics:** $N = 2; R^2 = 1.0$ **Maximum log  $K_{ow}$ :**

5.0

**Maximum MW:**

1000.0

**Application:**

This SAR may be used to estimate toxicity for peroxy acids.

**Limitations:**If the log  $K_{ow}$  value is greater than 5.0, or if the compound is solid and the LC50 exceeds the water solubility, use SAR with longer exposure.**References:**

United States Environmental Protection Agency (USEPA). 1991. OTS PMN ECOTOX. Washington, DC: Office of Toxic Substances, USEPA

## LIST OF PEROXY ACIDS USED TO DEVELOP THE FISH 96-h LC50 SAR.

CHEMICAL	96-h LC50 (mg/L)	Log $K_{ow}$	Ref.
Chemical identity CBI	0.750	2.6	EPA

EPA = USEPA (1991); chemical identity is Confidential Business Information under TSCA.

## PEROXY ACIDS

9/1993

**SAR**

**PEROXY ACIDS**

**Organism:** Daphnids  
**Duration:** 48-h  
**Endpoint:** LC50 (Mortality)  
**Equation:**  $\text{Log 48-h LC50 (mM/L)} = -0.717 - 0.417 \log K_{ow}$   
**Statistics:**  $N = 2; R^2 = 1.0$   
**Maximum log  $K_{ow}$ :** 5.0  
**Maximum MW:** 1000.0  
**Application:** This SAR may be used to estimate toxicity for peroxy acids.  
**Limitations:** If the log  $K_{ow}$  value is greater than 5.0, or if the compound is solid and the LC50 exceeds the water solubility, use SAR with longer exposure.  
**References:** United States Environmental Protection Agency (USEPA). 1991. OTS PMN ECOTOX. Washington, DC: Office of Toxic Substances, USEPA.

LIST OF PEROXY ACIDS USED TO DEVELOP THE DAPHNID 48-H SAR.

CHEMICAL	48-h LC50 (mg/L)	Log $K_{ow}$	Ref.
Chemical identity CBI	4.6	2.6	EPA

EPA = USEPA (1991); chemical identity is Confidential Business Information under TSCA.



## SAR

## PHENOLS

Organism: Fish  
Duration: 96-h  
Endpoint: LC50 (Mortality)

Equation:  $\text{Log 96-h LC50 (mM/L)} = 0.399 - 0.616 \log K_{ow}$

Statistics:  $N = 78; R^2 = 0.86$

Maximum  $\log K_{ow}$ : 7.0  
Maximum MW: 1000.0

Application: This equation may be used to estimate toxicity for phenols.

Limitations: Phenols which are significantly more toxic than predicted by this SAR are:

catechol with 16 x excess toxicity;  
hydroquinone with 1400 x excess toxicity; and  
p-benzoquinone with 5500 x excess toxicity.

If the  $\log K_{ow}$  value is greater than 7.0, or if the compound is solid and the LC50 exceeds the water solubility, use SAR with longer exposure.

## References:

- Alexander HC, Dill, DC, Smith LW, Guiney PD, and Dorn P. 1988. Bisphenol A: Acute aquatic toxicity. *Environ. Toxicol. Chem.* 7:19-26.
- Curtis MW and Ward CH. 1981. Aquatic toxicity of forty industrial chemicals: Testing in support of hazardous substance spill prevention regulation. *Journal of Hydrology* 51:359-367.
- DeGraeve GM, Geiger DL, Meyer JS, Bergman HL. 1980. Acute and embryo-larval toxicity of phenolic compounds to aquatic biota. *Arch. Environ. Contam. Toxicol.* 9:557-568.
- Holcombe GW, Phipps GL, Knuth M, and Felhaber T. 1984. The acute toxicity of selected substituted phenols, benzenes, and benzoic acid esters to fathead minnows, Pimephales promelas. *Environ. Pollution, Ser. A*, 35:367-381.
- Holcombe GW, Phipps GL, and Fiandt JT. 1982. Effects of phenol, 2,4-dimethylphenol, 2,4-dichlorophenol, and pentachlorophenol on embryo, larval, and early-juvenile fathead minnows (Pimephales promelas). *Arch. Environ. Contam. Toxicol.* 11:73-78.

## PHENOLS

9/1993

Konemann H, and Musch A. 1981. Quantitative structure-activity relationships in fish toxicity studies. Part 2: The influence of pH on the SAR of chlorophenols. *Toxicology* 19:223-228.

Marking LL, Howe GE, and Bills TD. 1991. Temperature and pH effects on acute and chronic toxicity of four chemicals to amphipods (Gammarus pseudolimnaeus) and rainbow trout (Oncorhynchus mykiss). EPA/600/X-90/286. Gulf Breeze, FL: Environmental Research Laboratory, Office of Research and Development, United States Environmental Protection Agency. August.

Saarikoski J and Viluksela M. 1982. Relation between physicochemical properties of phenols and their toxicity and accumulation in fish. *Ecotoxicology and Environmental Safety* 6:501-512.

United States Environmental Protection Agency (USEPA1). 1980. Ambient Water Quality Criteria for Phenol. EPA-440-5-80-066. Washington, DC: Criteria and Standards Division, Office of Water Regulations and Standards, USEPA.

United States Environmental Protection Agency (USEPA2). 1984. Dynamic 14-day acute toxicity of octylphenol to rainbow trout (Salmo gairdneri). TSCA Section 4(d). Document No. 40-8462075. Washington, DC: OTS Public Files, USEPA Fiche No. 0507489 (2).

United States Environmental Protection Agency (USEPA3). 1990. Section 8(e)908.

United States Environmental Protection Agency (USEPA4). 1991. OTS PMN-ECOTOX. Washington, DC: Office of Toxic Substances, USEPA.

Veith GD and Broderius SJ. 1987. Structure-toxicity relationships for industrial chemicals causing type (II) narcosis syndrome. IN: Kaiser KLE (ed.). *QSAR in Environmental Toxicology - II*. New York: D. Reidel Publishing Company. pp. 385-391.

LIST OF PHENOLS USED TO DEVELOP THE FISH 96-h LC50 SAR.

CHEMICAL	96-h LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
PHENOLS USED IN THE CALCULATION OF THIS SAR			
Resorcinol	60.0	0.8	C
Resorcinol	100.0	0.81	D
Phenol	44.5	1.5	EPA1
Phenol	36.3	1.5	EPA1
Phenol	36.0	1.5	EPA1
Phenol	34.9	1.5	EPA1
Phenol	26.0	1.5	EPA1
Phenol	19.0	1.5	EPA1
Phenol	16.7	1.5	EPA1
Phenol	16.4	1.5	EPA1
Phenol	10.2	1.5	EPA1
Phenol	8.9	1.5	D
Phenol	67.5	1.5	D
Phenol	29.8	1.5	K
Phenol	43.0	1.5	S
Phenol	37.0	1.5	V
3-Methoxyphenol	74.0	1.6	V
4-Methoxyphenol	110.0	1.6	V
4-Nitrophenol	14.2	1.9	S
3-Nitrophenol	11.8	1.9	S
4-Nitrophenol	41.0	1.9	H
4-Nitrophenol	6.9	1.9	M
3-Methylphenol	23.1	2.1	S
4-Methylphenol	16.5	2.1	V
p-Cresol	7.9	2.1	D
p-Cresol	28.6	2.1	D
o-Cresol	8.4	2.1	D
o-Cresol	18.2	2.1	D
m-Cresol	8.9	2.1	D
m-Cresol	55.9	2.1	D
2-Chlorophenol	11.2	2.2	K
2-Chlorophenol	13.8	2.2	S
2-Chlorophenol	9.4	2.2	V
2-Allylphenol	15.0	2.2	V
4-Chlorophenol	8.5	2.5	S
3-Chlorophenol	6.4	2.5	K
1-Naphthol	4.6	2.6	V
4-Ethylphenol	10.4	2.7	V
2,6-Dichlorophenol	7.8	2.8	S
2,4-Dimethylphenol	16.6	2.8	H
2-Chloro-4-methylphenol	35.9	2.9	V
2,4-Dichlorophenol	5.5	3.1	S

# PHENOLS

9/1993

CONTINUED.

CHEMICAL	96-h LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
PHENOLS USED IN CALCULATION OF THE SAR			
4,5-Dichloro-2-methoxy phenol	4.8	3.1	S
2,4-Dichlorophenol	4.2	3.1	K
3,4,5-Trichloro- 2,6-dimethoxyphenol	3.4	3.1	S
2,4-Dichlorophenol	7.75	3.1	H
4-Chloro-3-methylphenol	5.72	3.1	V
2,4-Dichlorophenol	7.75	3.1	V
4-Propylphenol	11.0	3.2	V
4-Phenylazophenol	1.17	3.2	V
3,5-Dichlorophenol	2.7	3.3	K
Bis(thiophenol)	1.5	3.4	EPA3
2,3,6-Trimethylphenol	0.390	3.4	S
2-Phenylphenol	6.15	3.4	V
4,4'-[oxybis(2,1-ethane diylthio)]bisphenol	1.5	3.4	EPA4
4-Tert-butylphenol	5.15	3.5	V
3,4,5-Trichloro- 2-methoxyphenol	2.1	3.6	S
2,4,6-Trichlorophenol	4.55	3.6	V
2,4,6-Trichlorophenol	2.3	3.6	S
2,3,6-Trichlorophenol	5.1	3.8	K
4-chloro-3,5-dimethyl phenol	3.4	3.8	S
4-Phenoxyphenol	4.96	3.8	V
Bisphenol A	4.6	3.8	A
2,3,5-Trichlorophenol	1.6	3.9	K
2,4,5-Trichlorophenol	1.2	3.9	S
3,4,5,6-Tetrachloro-2- hydroxyphenol	2.5	3.9	S
4-Tert-pentylphenol	2.59	4.0	V
2-Tert-butyl-4- methylphenol	2.1	4.1	S
2,3,5,6-Tetrachlorophenol	1.4	4.3	K
2,3,4,6-Tetrachlorophenol	1.1	4.3	S
2,3,4,5-Tetrachlorophenol	0.770	4.6	K
Pentachlorophenol	0.380	5.1	K
Pentachlorophenol	0.24	5.1	V
Pentachlorophenol	0.440	5.1	S
4-(Tert-octyl)phenol	0.250	5.3	EPA2
4-(Tert-octyl)phenol	0.210	5.3	EPA2
4-Nonylphenol	0.140	6.4	V

CONTINUED.

CHEMICAL	96-h LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
PHENOLS USED IN CALCULATION OF THE SAR			
Substituted benzophenone glyceride	*	8.0	EPA4
Hindered phenol	*	11.0	EPA4
PHENOLS HAVING EXCESS TOXICITY			
p-Benzoquinone	0.125	-0.3	D
p-Benzoquinone	0.045	-0.3	D
Hydroquinone	0.097	0.8	D
Hydroquinone	0.044	0.8	D
Catechol	8.9	0.81	D
Catechol	3.5	0.81	D

\* = No fish mortality in saturated solutions.

A = Alexander et al (1988)  
 C = Curtis and Ward (1981)  
 D = DeGraeve et al (1980)  
 H = Holcombe et al (1984, 1982)  
 K = Konemann and Musch (1981)  
 M = Marking et al (1991)  
 S = Saarikoski and Viluksela (1982)  
 EPA1 = USEPA (1980)  
 EPA2 = USEPA (1984)  
 EPA3 = USEPA (1990)  
 EPA4 = USEPA (1991)  
 V = Veith and Broderius (1987)

## PHENOLS

9/1993

**SAR**

**PHENOLS**

**Organism:**

Daphnid

**Duration:**

48-h

**Endpoint:**

LC50 (Mortality)

**Equation:**

$\text{Log 48-h LC50 (mM/L)} = -0.451 - 0.409 \log K_{ow}$

**Statistics:**

$N = 48; R^2 = 0.6$

**Maximum log  $K_{ow}$ :**

5.5

**Maximum MW:**

1000.0

**Application:**

This equation may be used to estimate toxicity for phenols.

**Limitations:**

Phenols which contain the following groups may have excess toxicity compared with the values predicted by this SAR:

1,2-di(OH) groups (e.g., catechol);  
1,4-di(OH) groups (e.g., hydroquinone); or  
1,4-di(=O) groups (e.g., benzoquinone).

If the log  $K_{ow}$  value is greater than 5.5, or if the compound is solid and the LC50 exceeds the water solubility, use SAR with longer duration.

For aminophenols, use the daphnid 48-h LC50 SAR for anilines.

**References:**

Alexander HC, Dill, DC, Smith LW, Guiney PD, and Dorn P. 1988. Bisphenol A: Acute aquatic toxicity. Environ. Toxicol. Chem. 7:19-26.

Kuhn R, Pattard M, Pernak, K-D, and Winter A. 1989. Results of the harmful effects of water pollutants to Daphnia magna in the 21 day reproduction test. Water Res. 23:501-510.

LeBlanc G. 1980. Acute toxicity of priority pollutants to water flea (Daphnia magna). Bull. Environm. Contam. Toxicol. 24:684-691.

United States Environmental Protection Agency (USEPA1). 1984. Dynamic 14-day acute toxicity of octylphenol to rainbow trout (Salmo gairdneri). TSCA Section 4(d). Document No. 40-8462075. Washington, DC: OTS Public Files, USEPA. Fiche No. 0507489 (2)

United States Environmental Protection Agency (USEPA2). 1991. OTS PMN ECOTOX Washington, DC: Office of Toxic Substances, USEPA.

# PHENOLS

9/1993

## LIST OF PHENOLS USED TO DEVELOP THE DAPHNID 48-h LC50 SAR.

CHEMICAL	48-h LC50 (mg/L)	Log K <sub>ow</sub>	Ref.
PHENOLS USED IN CALCULATION OF THE SAR			
3-Hydroxyphenylurea	93.0	0.2	K
4-Acetamidophenol	9.2	0.5	K
4-Hydroxybenzonitrile	15.0	1.6	K
4-Nitrophenol	22.0	1.9	L
2-Chlorophenol	2.6	2.2	L
4-Chlorophenol	4.1	2.5	L
2,4-Dimethylphenol	2.1	2.8	L
3-(Trifluoromethyl)phenol	11.0	2.9	K
2,4-Dichlorophenol	2.6	3.1	L
4-Chloro-6-methylphenol	0.290	3.1	L
o-Phenylphenol	1.5	3.4	K
2,4,6-Trichlorophenol	6.0	3.6	L
2,4-Dichloro-6-methyl phenol	0.430	3.7	L
4-Chloro-3,5-dimethyl phenol	4.5	3.8	K
Bisphenol A	10.2	3.8	A
2,4,5-Trichlorophenol	2.7	3.9	L
2,3,4,6-Tetrachlorophenol	0.290	4.3	L
2,3,5,6-Tetrachlorophenol	0.570	4.3	L
Pentachlorophenol	0.680	5.1	L
4-(Tert-octyl)phenol	0.270	5.3	EPA1
3,5-Di-tert-butylphenol	1.7	5.4	K
Substituted benzophenone glyceride	*	8.0	EPA2

\* = No daphnid mortalities in saturated solutions.

A = Alexander et al (1988)

EPA1 = USEPA1 (1984)

EPA2 = USEPA2 (1991)

K = Kuhn et al (1989)

L = LeBlanc (1980)

SAR

PHENOLS

Organism: Green Algae  
Duration: 96-h  
Endpoint: EC50 (Growth)

Equation: To find the estimated acute toxicity of a phenol, use the neutral organic green algae 96-h EC50 SAR.

Maximum log  $K_{ow}$ : 6.4  
Maximum MW: 1000.0

Application: The neutral organic green algae 96-h EC50 SAR may be used to estimate toxicity for phenols.

Limitations: Phenols which contain the following groups may have excess toxicity compared with the values predicted by this SAR:

1,2-di(OH) groups (e.g., catechol);  
1,4-di(OH) groups (e.g., hydroquinone); or  
1,4-di(=O) groups (e.g., benzoquinone).

If the log  $K_{ow}$  value is greater than 6.4, or if the compound is solid and the EC50 exceeds the water solubility use SAR with longer exposure.

For aminophenols, use the green algae chronic value SAR for anilines

References:

Alexander HC, Dill, DC, Smith LW, Guiney PD, and Dorn P. 1988. Bisphenol A: Acute aquatic toxicity. Environ. Toxicol. Chem. 7:19-26.

Kuhn R and Pattard M. 1990. Results of the harmful effects of water pollutants to green algae (*Scenedesmus subspicatus*) in the cell multiplication inhibition test. Water Res. 24:31-38.

United States Environmental Protection Agency (USEPA1). 1984. Dynamic 14-day acute toxicity of octylphenol to rainbow trout (*Salmo gairdneri*). TSCA Section 4(d). Document No. 40-8462075. Washington, DC: OTS Public Files, USEPA. Fiche No. 0507489 (2).

United States Environmental Protection Agency (USEPA2). 1991. OTS PMN ECOTOX. Washington, DC: Office of Toxic Substances, USEPA

# PHENOLS

9/1993

## LIST OF PHENOLS USED TO DEVELOP THE GREEN ALGAE 96-h EC50 SAR.

CHEMICAL	96-h EC50 (mg/L)	Log K <sub>ow</sub>	Ref.
PHENOLS USED IN CALCULATION OF THE SAR			
3,5-Dimethoxyphenol	110.0	1.4	K
4-Nitrophenol	26.0	1.9	K
p-Cresol	7.8	2.1	K
2-Chlorophenol	50.0	2.2	K
2-Bromophenol	60.0	2.4	K
2-Nitro-para-cresol	12.0	2.5	K
4-Chlorophenol	8.0	2.5	K
2,4-Dichlorophenol	11.5	3.1	K
4-Chloro-3-methylphenol	>10.0	3.1	K
2,4,6-Trimethylphenol	17.0	3.4	K
Bis(thiophenol)	0.740	3.4	EPA2
Bisphenol A	2.7	3.8	A
4-(Tert-octyl)phenol	1.6	5.3	EPA1
PHENOLS HAVING EXCESS TOXICITY			
2-Amino-4-methylphenol	4.6	1.3	K

A = Alexander et al (1988)

EPA1 = USEPA1 (1984)

EPA2 = USEPA2 (1991)

K = Kuhn and Pattard (1990)

**SAR**

**PHENOLS**

**Organism:**

Fish

**Duration:**

30-d

**Endpoint:**

Chronic Value

**Equation:**

$\text{Log ChV (mM/L)} = -0.401 - 0.652 \log K_{ow}$

**Statistics:**

$N = 20; R^2 = 0.94$

**Maximum log  $K_{ow}$ :**

8.0

**Maximum MW:**

1000.0

**Application:**

This equation may be used to estimate toxicity for phenols.

**Limitations:**

Phenols which contain the following groups may have excess toxicity compared with the values predicted by this SAR:

- 1,2-di(OH) groups (e.g., catechol);
- 1,4-di(OH) groups (e.g., hydroquinone); or
- 1,4-di(=O) groups (e.g., benzoquinone).

If the log  $K_{ow}$  value is greater than 8.0, or if the compound is solid and the ChV exceeds the water solubility, use SAR with longer exposure. A test duration of more than 30 days may result in a lower chronic toxicity; at 60 days the toxicity will be 20 x lower than predicted by this SAR for phenols with a log  $K_{ow}$  of 1.5 and 4 x lower for phenols with a log  $K_{ow}$  of 5.3. For an exposure period of 60 days, a separate SAR has been developed.

For aminophenols, use the fish ChV SAR for anilines.

**References:**

DeGraeve GM, Geiger DL, Meyer JS, and Bergman HL. 1980. Acute and embryo-larval toxicity of phenolic compounds to aquatic biota. Arch. Environ. Contam. Toxicol. 9:557-568.

Hedtke SF, West CW, Allen KN, Norberg-King TJ, and Mount DI. 1986. Toxicity of pentachlorophenol to aquatic organisms under naturally varying and controlled conditions. Environ. Toxicol. Chem. 5:531-542.

Holcombe GW, Phipps GL, and Fiandt JT. 1982. Effects of phenol, 2,4-dimethylphenol, 2,4-dichlorophenol, and pentachlorophenol on embryo, larval, and early-juvenile fathead minnows (*Pimephales promelas*). Arch Environ. Contam. Toxicol. 11:73-78.

Marking LL, Howe GE, and Bills TD. 1991. Temperature and pH effects on acute and chronic toxicity of four chemicals to amphipods (*Gammarus pseudolimnaeus*) and rainbow trout (*Oncorhynchus mykiss*).

## PHENOLS

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EPA/600/X-90/286. Gulf Breeze, FL: Environmental Research Laboratory, Office of Research and Development, United States Environmental Protection Agency. August.

Spehar RL, Nelson HP, Swanson MJ, and Renos JW. 1985. Pentachlorophenol toxicity to amphipods and fathead minnows at different test pH values. Environ. Toxicol. Chem. 4:389-397.

United States Environmental Protection Agency (USEPA1). 1980. Ambient Water Quality Criteria for Phenol. EPA-440-5-80-066. Washington, DC: Criteria and Standards Division, Office of Water Regulations and Standards, USEPA.

United States Environmental Protection Agency (USEPA2). 1984. Dynamic 14-day acute toxicity of octylphenol to rainbow trout (Salmo gairdneri). TSCA Section 4(d). Document No. 40-8462075. Washington, DC: OTS Public Files, USEPA. Fiche No. 0507489 (2).

United States Environmental Protection Agency (USEPA3). 1991. Fish Chronic Toxicity Data Base. Duluth, MN: Environmental Research Laboratory (ERL), Office of Research and Development, USEPA, 6201 Congdon Boulevard, 55804; contact C.L. Russom (218) 720-5500.

LIST OF PHENOLS USED TO DEVELOP THE FISH 30-d ChV SAR.

CHEMICAL	ChV (mg/L)	Log K <sub>ow</sub>	Ref.
PHENOLS USED IN CALCULATION OF THE SAR			
2,2'-Methylene bis (4-chlorophenol)	0.122	5.0	EPA3
4-Nitrophenol	3.38	1.9	M
Phenol	1.4	1.5	D
Phenol	2.56	1.5	EPA1
o-Cresol	1.8	2.1	EPA3
4-Nitrophenol	2.65	1.9	EPA3
2,4-Dimethylphenol	2.48	2.8	HO
2,4-Dimethylphenol	0.763	2.8	EPA3
p-Cresol	1.86	2.1	EPA3
Phenol	2.56	1.5	HO
2-Phenylphenol	1.22	3.4	EPA3
Pentachlorophenol	0.089	5.1	S
Pentachlorophenol	0.057	5.1	HO
Pentachlorophenol	0.040	5.1	S
Pentachlorophenol	0.144	5.1	HE
Pentachlorophenol	0.049	5.1	S
Pentachlorophenol	0.024	5.1	S
2,4,5-Trichlorophenol	0.232	3.9	EPA3
2,4-Dichlorophenol	0.365	3.1	HO
PHENOLS HAVING EXCESS TOXICITY			
Phenol	<0.200	1.5	D
4-(Tert-octyl)phenol	0.008	5.3	EPA2

D = DeGraeve et al (1980)  
EPA1 = USEPA (1980)  
EPA2 = USEPA (1984)  
EPA3 = USEPA (1991)  
HE = Hedtke et al (1986)  
HO = Holcombe et al (1982)  
M = Marking et al (1991)  
S = Spehar et al (1985)

**PHENOLS**  
9/1993

<b>SAR</b>	<b>PHENOLS</b>
<b>Organism:</b>	Fish
<b>Duration:</b>	60-d
<b>Endpoint:</b>	Chronic Value
<b>Equation:</b>	$\text{Log ChV (mM/L)} = -2.029 - 0.447 \log K_{ow}$
<b>Statistics:</b>	$N = 2; R^2 = 1.0$
<b>Maximum log <math>K_{ow}</math>:</b>	8.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This equation may be used to estimate toxicity for phenols.
<b>Limitations:</b>	If the log $K_{ow}$ value is greater than 8.0, or if the compound is solid and the ChV exceeds the water solubility, no effects expected at saturation.  For aminophenols, use the fish ChV SAR for anilines.
<b>References:</b>	DeGraeve GM, Geiger DL, Meyer JS, and Bergman HL. 1980. Acute and embryo-larval toxicity of phenolic compounds to aquatic biota. Arch. Environ. Contam. Toxicol. 9:557-568.  United States Environmental Protection Agency (USEPA). 1984. Dynamic 14-day acute toxicity of octylphenol to rainbow trout ( <i>Salmo gairdneri</i> ). TSCA Section 4(d). Document No. 40-8462075. Washington, DC: OTS Public Files, USEPA. Fiche No. 0507489 (2)

LIST OF PHENOLS USED TO DEVELOP THE FISH 60-d ChV SAR.

CHEMICAL	ChV (mg/L)	Log $K_{ow}$	Ref.
PHENOLS USED IN CALCULATION OF THE SAR			
Phenol	<0.200	1.5	D
4-(Tert-octyl)phenol	0.008	5.3	EPA

D = DeGraeve et al (1980)  
EPA = USEPA (1984)

**PHENOLS**  
9/1993

<b>SAR</b>	<b>PHENOLS</b>
<b>Organism:</b>	Daphnid
<b>Duration:</b>	
<b>Endpoint:</b>	Chronic Value
<b>Equation:</b>	$\text{Log ChV (mM/L)} = -0.573 - 0.614 \log K_{ow}$
<b>Statistics:</b>	$N = 12; R^2 = 0.92$
<b>Maximum log <math>K_{ow}</math>:</b>	8.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This equation may be used to estimate toxicity for phenols.
<b>Limitations:</b>	<p>Phenols which contain the following groups may have excess toxicity compared with the values predicted by this SAR:</p> <ul style="list-style-type: none"><li>1,2-di(OH) groups (e.g., catechol);</li><li>1,4-di(OH) groups (e.g., hydroquinone); or</li><li>1,4-di(=O) groups (e.g., benzoquinone).</li></ul> <p>3,5-Dimethoxyphenol has an excess toxicity of 18 x that predicted by this SAR.</p> <p>If the log <math>K_{ow}</math> value is greater than 8.0, or if the compound is solid and the ChV exceeds the water solubility, no effects expected at saturation.</p> <p>For aminophenols, use the daphnid ChV SAR for anilines.</p>
<b>References:</b>	<p>Kuhn R, Pattard M, Pernak, K-D, and Winter A. 1989. Results of the harmful effects of water pollutants to <u>Daphnia magna</u> in the 21 day reproduction test. Water Res. 23:501-510.</p> <p>Oris JT, Winner RW, and Moore MV. 1991. A four-day survival and reproduction toxicity test for <u>Ceriodaphnia dubia</u>. Environ. Toxicol. Chem. 10:217-224.</p> <p>United States Environmental Protection Agency (USEPA). 1984. Dynamic 14-day acute toxicity of octylphenol to rainbow trout (<u>Salmo gairdneri</u>). TSCA Section 4(d). Document No. 40-8462075. Washington, DC: OTS Public Files, USEPA. Fiche No. 0507489 (2).</p>

**PHENOLS**

9/1993

## LIST OF PHENOLS USED TO DEVELOP THE DAPHNID ChV SAR.

CHEMICAL	ChV (mg/L)	Log K <sub>ow</sub>	Ref.
PHENOLS USED IN CALCULATION OF THE SAR			
Phenol	4.9	1.5	O
4-Nitrophenol	1.8	1.9	K
4-Methylphenol	1.4	2.1	K
2-Chlorophenol	0.500	2.2	K
2-Bromophenol	1.5	2.4	K
4-Chlorophenol	0.840	2.5	K
2-Nitro-para-cresol	3.2	2.5	K
2,4-Dichlorophenol	0.290	3.1	K
4-Chloro-3-methylphenol	1.8	3.1	K
2,4,6-Trimethylphenol	0.160	3.4	K
4-(Tert-octyl)phenol	0.086	5.3	EPA
PHENOLS HAVING EXCESS TOXICITY			
2-Amino-4-methylphenol	0.400	1.3	K
3,5-Dimethoxyphenol	0.320	1.4	K

**SAR****PHENOLS****Organism:**

Green Algae

**Duration:****Endpoint:**

Chronic Value (Growth)

**Equation:**

To find the estimated chronic toxicity of a phenol, use the neutral organic green algae ChV SAR.

**Maximum log  $K_{ow}$ :**

8.0

**Maximum MW:**

1000.0

**Application:**

The neutral organic green algae ChV SAR may be used to estimate toxicity for phenols.

**Limitations:**

Phenols which contain the following groups may have excess toxicity compared with the values predicted by this SAR:

- 1,2-di(OH) groups (e.g., catechol);
- 1,4-di(OH) groups (e.g., hydroquinone); or
- 1,4-di(=O) groups (e.g., benzoquinone).

If the log  $K_{ow}$  is greater than 8.0, or if the compound is solid and the ChV exceeds the water solubility, no effects expected at saturation.

For aminophenols, use the aniline green algae ChV SAR.

**References:**

Kuhn R and Pattard M. 1990. Results of the harmful effects of water pollutants to green algae (*Scenedesmus subspicatus*) in the cell multiplication inhibition test. *Water Res.* 24:31-38.

Slooff W, Canton JH, and Hermens JLM. 1983. Comparison of the susceptibility of 22 freshwater species to 15 chemical compounds. I. (Sub)Acute toxicity tests. *Aquatic Toxicology* 4:113-128.

United States Environmental Protection Agency (USEPA1). 1984. Dynamic 14-day acute toxicity of octylphenol to rainbow trout (*Salmo gairdneri*). TSCA Section 4(d). Document No. 40-8462075. Washington, DC: OTS Public Files, USEPA. Fiche No. 0507489 (2).

United States Environmental Protection Agency (USEPA2). 1991. OTS PMN ECOTOX. Washington, DC: Office of Toxic Substances, USEPA.

**PHENOLS**  
9/1993

LIST OF PHENOLS USED TO DEVELOP THE GREEN ALGAE ChV SAR.

CHEMICAL	ChV (mg/L)	Log K <sub>ow</sub>	Ref.
PHENOLS USED IN CALCULATION OF THE SAR			
3,5-Dimethoxyphenol	40.0	1.4	K
4-Nitrophenol	2.1	1.9	K
o-Cresol	34.0	2.1	S
o-Cresol	11.0	2.1	S
o-Cresol	36.0	2.1	S
o-Cresol	65.0	2.1	S
p-Cresol	2.3	2.1	K
2-Chlorophenol	24.0	2.2	K
2-Bromophenol	28.0	2.4	K
2-Nitro-p-cresol	6.3	2.5	K
4-Chlorophenol	3.0	2.5	K
4-Chloro-3-methylphenol	5.2	3.1	K
2,4-Dichlorophenol	2.4	3.1	K
2,4,6-Trimethylphenol	5.8	3.4	K
Bis(thiophenol)	0.300	3.4	EPA2
4-(Tert-octyl)phenol	<0.860	5.3	EPA1
PHENOLS WITH EXCESS TOXICITY			
2-Amino-4-methylphenol	0.750	1.3	K

EPA1 = USEPA (1984)

EPA2 = USEPA (1991)

K = Kuhn and Pattard (1990)

S = Slooff et al (1983)

**SAR** **PHENOLS, DINITRO**

**Organism:** Fish  
**Duration:** 96-h  
**Endpoint:** LC50 (Mortality)

**Equation:**  $\text{Log 96-h LC50 (mM/L)} = -0.285 - 0.559 \log K_{ow}$

**Statistics:**  $N = 4; R^2 = 0.96$

**Maximum log  $K_{ow}$ :** 7.0  
**Maximum MW:** 1000.0

**Application:** This SAR may be used to estimate toxicity for dinitrophenols and other polynitrophenols.

**Limitations:** If the log  $K_{ow}$  value is greater than 7.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.

**References:** Veith GD and Broderius SJ. 1987. Structure-toxicity relationships for industrial chemicals causing type (II) narcosis syndrome. In: Kaiser KLE (ed.). QSAR in Environmental Toxicology-II. Boston, MA: D. Reidel Pub. Co, pp. 385-391.

LIST OF DINITROPHENOLS USED TO DEVELOP THE FISH 96-h LC50 SAR.

CHEMICAL	96-h LC50 (mg/L)	Log $K_{ow}$	Ref.
2,4-dinitrophenol	11.0	1.5	VB
4,6-dinitro-o-cresol	1.54	2.6	VB
2,4-dinitro-1-naphthol sodium	4.24	3.09	VB

VB = Veith and Broderius (1987)

**PHENOLS, DINITRO**

9/1993

<b>SAR</b>	<b>PHENOLS, DINITRO</b>
<b>Organism:</b>	Daphnid
<b>Duration:</b>	48-h
<b>Endpoint:</b>	LC50 (Mortality)
<b>Equation:</b>	$\text{Log 48-h LC50 (mM/L)} = 0.083 - 0.632 \log K_{ow}$
<b>Statistics:</b>	$N = 7; R^2 = 0.85$
<b>Maximum log <math>K_{ow}</math>:</b>	7.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This SAR may be used to estimate toxicity for dinitrophenols and other polynitrophenols
<b>Limitations:</b>	If the log $K_{ow}$ value is greater than 7.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.
<b>References:</b>	<p>Hermens J, Canton H, Janssen P, and DeJong R. 1984. Quantitative structure-activity relationships and toxicity studies of mixtures of chemicals with anaesthetic potency: Acute lethal and sublethal toxicity to <u>Daphnia magna</u>. Aquatic Toxicology 5:143-154.</p> <p>Kuhn R, Rattard M, Pernak K-D, and Winter A. 1989. Results of the harmful effects of selected water pollutants (anilines, phenols, aliphatic compounds) to <u>Daphnia magna</u>. Water Research 23:495-499.</p> <p>LeBlanc. 1980. Acute toxicity of priority pollutants to water flea (<u>Daphnia magna</u>). Bulletin of Environmental Contamination and Toxicology. 24: 684-691.</p>

LIST OF DINITROPHENOLS USED TO DEVELOP THE DAPHNID 48-h LC50 SAR

CHEMICAL	48-h LC50 (mg/L)	Log $K_{ow}$	Ref.
2,4,6-trinitrophenol	85.0	1.8	L
2,4,6-trinitrophenol	90.0	1.8	K
2,4-dinitrophenol	4.1	1.9	L
2,4-dinitro-6-methyl phenol	3.1	2.6	L
dinitro-o-cresol	3.3	2.6	H
2-methyl-4,6-dinitrophenol	2.7	2.6	K

ARSENIC(III)  
9/1993

**Organism:** Aquatic life (freshwater)  
**Duration:** Acute  
**Endpoint:** Lowest Observable Effect Concentration (LOEC)  
**Equation:**  $LOEC \text{ (mg/L)} = (0.130 \cdot MW)/9.012$   
**Application:** This equation may be used to estimate the acute toxicity of both organic and inorganic compounds containing beryllium.  
**Limitations:** Hardness has a substantial effect on acute toxicity.  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Regulations and Standards. EPA 440/5-86-001.

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**Organism:** Aquatic Life (freshwater)  
**Duration:** Chronic  
**Endpoint:** Lowest Observable Effect Concentration (LOEC)  
**Equation:**  $LOEC \text{ (mg/L)} = (0.0053 \cdot MW)/9.012$   
**Application:** This equation may be used to estimate the chronic toxicity of both organic and inorganic compounds containing beryllium.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Regulations and Standards. EPA 440/5-86-001.

**BERYLLIUM**  
9/1993

**Organism:** Fish (freshwater)  
**Duration:** 48-hour  
**Endpoint:** LC50

**Equation:**  $LC50 \text{ (mg/L)} = (315.0 \cdot MW)/10.81$

**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing boron.

**Limitations:** None

**References:** United States Environmental Protection Agency (USEPA). 1991. Hazard Profiles for Selected Heavy Metals. Washington, DC: Office of Pollution Prevention and Toxics, Health and Environmental Review Division, Environmental Effects Branch.

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**Organism:** Daphnid  
**Duration:** 48-hour  
**Endpoint:** LC50

**Equation:**  $LC50 \text{ (mg/L)} = (226.0 \cdot MW)/10.81$

**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing boron.

**Limitations:** None

**References:** United States Environmental Protection Agency (USEPA). 1991. Hazard Profiles for Selected Heavy Metals. Washington, DC: Office of Pollution Prevention and Toxics, Health and Environmental Review Division, Environmental Effects Branch.

**BORON**  
9/1993

**Organism:** Fish (freshwater)  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (0.05 \cdot \text{MW})/10.81$   
**Application:** This equation may be used to estimate the toxicity of both inorganic and organic compounds containing boron.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1991. Hazard Profiles for Selected Heavy Metals. Washington, DC: Office of Pollution Prevention and Toxics, Health and Environmental Review Division, Environmental Effects Branch.

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**Organism:** Daphnid  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (8.37 \cdot \text{MW})/10.81$   
**Application:** This equation may be used to estimate the toxicity of organic and inorganic compounds containing boron.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1991. Hazard Profiles for Selected Heavy Metals. Washington, DC: Office of Pollution Prevention and Toxics, Health and Environmental Review Division, Environmental Effects Branch.

## **PHENOLS, DINITRO**

9/1993

Kuhn = Kuhn et al (1989)

H = Hermens et al (1984)

L = LeBlanc (1980)

<b>SAR</b>	<b>PHENOLS, DINITRO</b>
<b>Organism:</b>	Fish
<b>Duration:</b>	32-d
<b>Endpoint:</b>	Chronic Value (Survival/Growth)
<b>Equation:</b>	$\text{Log ChV (mM/L)} = -1.78 - 0.552 \log K_{ow}$
<b>Statistics:</b>	$N = 4; R^2 = 1.0$
<b>Maximum log <math>K_{ow}</math>:</b>	8.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This SAR may be used to estimate toxicity for dinitrophenols and other polynitrophenols.
<b>Limitations:</b>	If the log $K_{ow}$ is greater than 8.0, or if the compound is solid and the ChV exceeds the water solubility, no effects expected at saturation.
<b>References:</b>	United States Environmental Protection Agency (USEPA). 1991. Fish Chronic Toxicity Data Base. Duluth, MN: Environmental Research Laboratory (ERL), Office of Research and Development, USEPA, 6201 Congdon Boulevard, 55804; contact C.L. Russom (218) 720-5500.

LIST OF DINITROPHENOLS USED TO DEVELOP THE FISH CHRONIC (ChV) SAR.

CHEMICAL	ChV (mg/L)	Log $K_{ow}$	Ref.
2,4-dinitrophenol	0.278	1.5	D
4,6-dinitro-o-cresol	0.171	2.3	D
2-(1-methylpropyl)- 4,6-dinitrophenol	0.027	3.7	D

D = USEPA (1991)

**PHENOLS, DINITRO**  
9/1993

<b>SAR</b>	<b>PHENOLS, DINITRO</b>
<b>Organism:</b>	Daphnid
<b>Duration:</b>	16-d
<b>Endpoint:</b>	Chronic Value (Survival/Reproduction)
<b>Equation:</b>	$\text{Log ChV (mM/L)} = -0.465 - 0.654 \log K_{ow}$
<b>Statistics:</b>	$N = 2; R^2 = 1.0$
<b>Maximum log <math>K_{ow}</math>:</b>	8.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This SAR may be used to estimate toxicity for dinitrophenols and other polynitrophenols.
<b>Limitations:</b>	If the log $K_{ow}$ is greater than 8.0, or if the compound is solid and the ChV exceeds the water solubility, no effects expected at saturation.
<b>References:</b>	Hermens J, Canton H, Janssen P, and DeJong R. 1984. Quantitative structure-activity relationships and toxicity studies of mixtures of chemicals with anaesthetic potency: Acute lethal and sublethal toxicity to <u>Daphnia magna</u> . Aquatic Toxicology 5:143-154.

LIST OF DINITROPHENOLS USED TO DEVELOP THE DAPHNID CHRONIC VALUE (ChV) SAR.

CHEMICAL	ChV (mg/L)	Log $K_{ow}$	Ref.
Dinitro-o-cresol	2.1	2.3	H

H = Hermens et al (1984)

PHENOLS, DINITRO

9/1993

**SAR****POLYMERS, POLYCATIONIC****Organism:**

Fish

**Duration:**

96-h

**Endpoint:**

LC50 (Mortality)

**Equation:**

Determine either the percent amine nitrogen or the number of positive charges per 1000 units of molecular weight and use the appropriate SAR:

1. If the percent amine nitrogen is less than 3.5:

$$\text{Log LC50 (mg/L)} = 1.3076 - 0.534 \times (\text{percent amine nitrogen})$$

If the percent amine nitrogen is greater than or equal to 3.5, then the fish 96-h LC50 is 0.27 mg/L.

2. If the number of positive charges per 1000 units of MW is less than 2.5:

$$\text{Log LC50 (mg/L)} = 1.3116 - 0.7606 \times (\text{number of positive charges per 1000 MW units})$$

If the number of positive charges per 1000 units of molecular weight is greater than or equal to 2.5, then the fish 96-h LC50 is 0.27 mg/L.

**Statistics:**

For the percent amine nitrogen SAR: (less than 3.5% amine nitrogen)  $N = 12$  and  $R^2 = 0.73$ , (greater than or equal to 3.5% amine nitrogen)  $N = 20$  and the standard deviation is plus or minus 0.18 logarithmic units; For the number of positive charges/1000 units MW SAR: (less than 2.5 charges/100 MW)  $N = 12$  and  $R^2 = 0.73$

**Minimum MW:**

1000.0

**Application:**

These SARs may be used for polycationic polymers which are highly water soluble or dispersible and contain nitrogen which may be protonated and/or quaternarized. These SARs may be used for polysulfoniums and polyphosphoniums which are dispersible.

**Limitations:**

Polycationic polymers which contain silicon may have limited water solubility or dispersibility. Polycationic polymers which contain anionic groups may be significantly less toxic than predicted by this SAR. For example, a polycationic polymer containing 4.7 percent amine nitrogen (or 3.4 cationic charges per 1000 molecular weight) and anionic groups with a cationic:anionic molar ratio of 1:1.1, will be about 24 times less toxic than predicted, i.e., fish 96-h LC50 is 6.6 mg/L.

# **POLYMERS, POLYCATIONIC**

9/1993

## **References:**

Nabholz JV. 1988. A structure-activity relationship for polycationic polymers. Washington, DC: Environmental Effects Branch, Health and Environmental Review Division (TS-796), Office of Toxic Substances, United States Environmental Protection Agency 20460-0001.

## **LIST OF COMPOUNDS USED TO DEVELOP THE POLYCATIONIC POLYMER FISH 96-h LC50 SAR.**

Percent Amine Nitrogen	Number of Positive Charges per 1000 Units of Mol. Weight	Average Molecular Weight (1000)	96-h LC50 (mg/L)
0.7	0.5	1.8	9.2
0.7	0.5	1.8	8.5
0.7	0.5	1.8	3.9
0.7	0.5	100.0	53.0
2.0	1.4	2500.0	0.97
2.0	1.4	2500.0	2.3
2.0	1.4	1100.0	0.64
2.0	1.4	1100.0	1.2
2.1	1.5	19000.0	0.84
3.0	2.1	100	0.94
3.4	2.4	*	0.6
3.4	2.4	*	0.3
6.0	4.3	>5.0	0.15
6.0	4.3	>5.0	0.16
6.0	4.3	>5.0	0.29
8.0	5.7	5.0	0.13
8.0	5.7	5.0	0.22
8.0	5.7	5.0	0.22
9.2	6.6	*	0.5
11.0	7.9	1.8	0.22
12.0	8.6	*	1.9
15.0	10.7	*	0.26
15.0	10.7	*	0.24
17.0	12.1	*	0.45
17.2	12.3	50.0	0.45
20.0	14.3	*	0.32
20.0	14.3	*	0.32
20.0	14.3	*	0.32
20.0	14.3	*	0.32
20.0	14.3	*	0.23
20.0	14.3	*	0.20

\* Unavailable at present.

**SAR** POLYMERS, POLYCATIONIC

**Organism:** Daphnid  
**Duration:** 48-h  
**Endpoint:** LC50 (Mortality)

**Equation:** The first SAR uses percent amine nitrogen to estimate toxicity while the second SAR uses the number of positive charges per 1000 unites of molecular weight. The toxicity increases rapidly from 0.1 to 2.3 percent amine nitrogen; thereafter, toxicity increases slowing with increasing charge density. The SAR equations used to estimate the acute toxicity are:

1.  $\text{Log LC50 (mg/L)} = 3.41 - 1.53 \times (\text{percent amine nitrogen})$
2.  $\text{Log LC50 (mg/L)} = 3.43 - 2.19 \times (\text{number of positive charges per 1000 MW units})$

**Maximum Value:** percent amine nitrogen SAR: 2.3% amine nitrogen;  
number of positive charges/1000 MW SAR: 1.6

**Minimum MW:** 1000.0

**Application:** This SAR may be used to estimate the toxicity of polycationic polymers which are highly water soluble or dispersible and contain a nitrogen which can be protonated and/or quaternarized. This SAR may be used for polysulfoniums and polyphosphoniums which are dispersible.

**Limitations:** Polycationic polymers which contain silicon may have limited water solubility or dispersibility.

Polycationic polymers which contain anionic groups may be significantly less toxic than predicted by this SAR. For example, a polycationic polymer containing 4.7 percent amine nitrogen (or 3.4 cationic charges per 1000 molecular weight) and anionic groups with a cationic:anionic molar ratio of 1:1.1, will be about 31 times less toxic than predicted, i.e., daphnid 48-h LC50 is 19.8 mg/L.

**References:** Nabholz JV. 1988. A structure-activity relationship for polycationic polymers. Washington, DC: Environmental Effects Branch, Health and Environmental Review Division (TS-796), Office of Toxic Substances, United States Environmental Protection Agency 20460-0001.

**POLYMERS, POLYCATIONIC**

0/1993

**LIST OF COMPOUNDS USED TO DEVELOP THE DAPHNID 48-h LC50 SAR.**

Percent Amine Nitrogen	Number of Positive Charges per 1000 Units of Mol. Weight	Average Molecular Weight	96-h LC50 (1000) (mg/L)
0.7	0.5	*	300.0
0.7	0.5	*	310.0
2.0	1.4	*	1.7
8.0	5.7	5.0	0.34
11.0	7.9	1.8	0.58
12.0	8.6	1.2	1.2
15.0	10.7	*	0.26
20.0	14.3	*	0.17

\* Unavailable at present.

**SAR**                      **POLYMERS, POLYCATIONIC****Organism:** Green Algae**Duration:** 96-h**Endpoint:** EC50 (Growth)

**Equation:** The algal 96-h EC50 can be estimated by dividing the equivalent fish 96-h LC50 estimate by 6. In addition, the algal 96-h no effect concentration (NEC; same as GMATC) can be estimated by dividing the algal 96-h EC50 by 2.5.

**Minimum MW:** 1000.0

**Application:** This SAR may be used to estimate the toxicity of polycationic polymers which are highly water soluble or dispersible and contain a nitrogen which can be protonated and/or quaternarized. This SAR may be used for polysulfoniums and polyphosphoniums which are dispersible.

**Limitations:** Polycationic polymers which contain silicon may have limited water solubility or dispersibility.

Polycationic polymers which contain anionic groups may be significantly less toxic than predicted by this SAR. For example, a polycationic polymer containing 4.7 percent amine nitrogen (or 3.4 cationic charges per 1000 molecular weight) and anionic groups with a cationic:anionic molar ratio of 1:1, will be about 30 times less toxic than predicted, i.e., the algal 96-h EC50 is 1.35 mg/L.

**References:** Nabholz JV. 1988. A structure-activity relationship for polycationic polymers. Washington, DC: Environmental Effects Branch, Health and Environmental Review Division (TS-796), Office of Toxic Substances, United States Environmental Protection Agency 20460-0001.

**POLYMERS, POLYCATIONIC**  
9/1993

**LIST OF COMPOUNDS USED TO DEVELOP THE GREEN ALGAE 96-h EC50 SAR.**

Percent Amine Nitrogen	Number of Positive Charges per 1000 Units of Mol. Weight	Average Molecular Weight (1000)	96-h EC50 (mg/L)	96-h NEC (mg/L)
0.7	0.5	*	300.0	0.88
8.0	5.7	5.0	0.16	*
11.0	7.9	1.8	0.07	0.034

\* Unavailable at present.

**SAR**                      **SURFACTANTS, ANIONIC**

**Organism:** Fish  
**Duration:** 96-h and 28-d  
**Endpoint:** LC50 and NEC

**Equation:** Determine the average length of the carbon chain to the nearest tenth and use the SAR equation:

$$\text{Log LC50 (mg/L)} = [(\text{avg no. of carbons} - 16)^2 - 10.643]/12.9346$$

The fish 28-d no effect concentrations (NEC, GMATC, or chronic value) can be estimated by dividing the estimated acute value derived above by 6.5.

**Statistics:**                       $N = 14; R^2 = 0.624$

**Maximum Value:** carbon chain length of 18 carbons  
**Minimum Value:** carbon chain length of 10 carbons

**Application:** This SAR may be used for the following classes of compounds:

1. Alkyl benzene sulfonates
2. Linear alkyl sulfonates (LAS)
3. Amphoteric surfactants with a sulfonate, phosphonate, or carboxylate terminus
4. Anionic surfactants terminated with phosphates
5. Anionic surfactants

**Limitations:** If the acute or chronic toxicity of linear alkyl benzene sulfonates which vary only in carbon chain length are to be estimated, then the weighted average of carbons in the alkyl chains (excluding the aromatic benzene ring) has to be determined.

**References:** Nabholz JV. 1985. Standard Environmental Hazard Assessment of PMNs 85-1156/1163. Intra-agency memorandum to O. Gutenson, Chemical Review and Evaluation Branch, Health and Environmental Review Division (TS-796), Office of Toxic Substances, United States Environmental Protection Agency, Washington, DC 20460-0001. August.

**SURFACTANTS, ANIONIC**  
9/1993

LIST OF ANIONIC SURFACTANTS USED TO DEVELOP THE FISH LC50 SAR.

Number of Fish Carbons	LC50 (mg/L)
10	21.2 - 47.5
11	11.6
12	1.18 - 6.5
13	1.11
14	0.25 - 0.42
16	0.087
18	0.38

**SAR****SURFACTANTS, ANIONIC****Organism:**

Daphnid

**Duration:**

48-h and 21-d NEC

**Endpoint:**

LC50 and NEC

**Equation:**

Determine the average length of the carbon chain to the nearest tenth and use the fish 96-h LC50 SAR equation:

$$\text{Log LC50 (mg/L)} = [(\text{ave. no. of carbons} - 16)^2 - 10.643]/12.9346$$

The daphnid 21-d no effect concentration (NEC, GMATC, or chronic value) can be estimated by dividing the estimated acute value derived above by 6.5.

**Statistics:**

$$N = 14; R^2 = 0.624$$

**Maximum Value:**

carbon chain length of 18 carbons

**Minimum Value:**

carbon chain length of 10 carbons

**Application:**

These SARs may be used for the following classes of compounds:

1. Alkyl benzene sulfonates
2. Alkyl sulfonates
3. Amphoteric surfactants with a sulfonate, phosphonate, or carboxylate terminus
4. Anionic surfactants terminated with phosphates
5. Anionic surfactants

**Limitations:**

If the acute or chronic toxicity of linear alkyl benzene sulfonates which vary only in carbon chain length are to be estimated, then the weighted average of carbons in the alkyl chains (excluding the aromatic benzene ring) have to be determined.

**References:**

Nabholz JV. 1985. Standard Environmental Hazard Assessment of PMNs 85-1156/1163. Intra-agency memorandum to O. Gutenson, Chemical Review and Evaluation Branch, Health and Environmental Review Division (TS-796), Office of Toxic Substances, United States Environmental Protection Agency, Washington, DC 20460-0001. August.

**SURFACTANTS, ANIONIC**  
9/1993

· LIST OF ANIONIC SURFACTANTS USED TO DEVELOP THE DAPHNID LC50 SAR.

Number of Carbons	Daphnid LC50 (mg/L)
10	29.55
11	21.15
12	5.88
13	2.63
14	0.68
16	0.11
18	0.12

<b>SAR</b>	<b>SURFACTANTS, ANIONIC</b>
<b>Organism:</b>	Green Algae
<b>Duration:</b>	96-h
<b>Endpoint:</b>	EC50 and NEC (Growth)
<b>Equation:</b>	<p>Determine the average length of the carbon chain to the nearest tenth and use the SAR equation:</p> $\text{Log EC50 (mg/L)} = [(\text{ave. no. of carbons} - 16)^2 - 42.466]/12.368$ <p>The green algae 96-h no effect concentration (NEC, GMATC, or chronic value) can be estimated by dividing the estimated EC50 value by 1.4.</p>
<b>Statistics:</b>	$N = 14; R^2 = 0.89$
<b>Maximum Value:</b>	carbon chain length of 18 carbons
<b>Minimum Value:</b>	carbon chain length of 10 carbons
<b>Maximum MW:</b>	
<b>Application:</b>	<p>These SARs may be used for the following classes of compounds:</p> <ol style="list-style-type: none"><li>1. Alkyl benzene sulfonates</li><li>2. Alkyl sulfonates</li><li>3. Amphoteric surfactants with a sulfonate, phosphonate, or carboxylate terminus</li><li>4. Anionic surfactants terminated with phosphates</li><li>5. Anionic surfactants</li></ol>
<b>Limitations:</b>	<p>If the toxicity of linear alkyl benzene sulfonates which vary only in carbon chain length are to be estimated, then the weighted average of carbons in the alkyl chains (excluding the aromatic benzene ring) have to be determined.</p>
<b>References:</b>	<p>Nabholz JV. 1987. Predicting the algal 96-h EC50 from the daphnid and fish SAR for LAS's. Intra-agency memorandum to "Whom It May Concern." Washington, DC: Office of Toxic Substances, United States Environmental Protection Agency, Washington, DC, 20460-0001.</p>

**SURFACTANTS, ANIONIC**  
**9/1993**

DATA FOR A C8 ANIONIC SURFACTANT USED TO DEVELOP THE GREEN ALGAE SAR.

Organism	EC50 (mg/L)	EC10 (mg/L)
Algae	12	8.5
Fish	366	
Daphnid	289	

## SURFACTANTS, CATIONIC, QUATERNARY AMMONIUM, MONOALKYL

9/1993

### SAR

### SURFACTANTS, CATIONIC, QUATERNARY AMMONIUM, MONOALKYL

**Organism:**

Fish

**Duration:**

Acute

**Endpoint:**

LC50 (Mortality)

**Equation:**

Determine the average number of carbons in the hydrophobic alkyl chain of the surfactant. If the average length of the carbon chain is between 16 and 24 carbons, use the SAR equation:

$$\text{Log LC50 (mg/L)} = -0.0918 + 0.023 (\text{average length of carbon chain})$$

If the length of the carbon chain is at least 10 but less than 16, use the SAR equation:

$$\text{Log LC50 (mg/L)} = 5.43 - 0.37 (\text{average length of carbon chain})$$

**Maximum Value:**

average carbon chain length of 24 carbons

**Minimum Value:**

average carbon chain length of 10 carbons

**Application:**

This SAR may be applied to monoalkyl (trimethyl) quaternary ammonium surfactants which are dispersible in water. This SAR may be used to estimate toxicity for:

1. monoalkyl cationic surfactants
2. monoalkyl phosphonium surfactants
3. monoalkyl sulfonium surfactants

**Limitations:**

This SAR may be used for monoalkyl quaternary ammonium surfactants where the anionic salt has less than 8 carbons in the alkyl chain. If the alkyl chain contains 8 or more carbons, the cationic surfactant and the anionic surfactant will form a strong ion pair. This ion pair will be much less soluble in water and consequently will be less toxic to fish.

**References:**

Nabholz JV. 1987. The SAR for monoalkyl (trimethyl) quaternary ammonium surfactants. Washington, DC: Environmental Effects Branch, Health and Environmental Review Division, Office of Toxic Substances, United States Environmental Protection Agency

Knauf W. 1973. Summary of the toxicity of surfactants to aquatic organisms. Tenside Detergents 5:251-255.

**SURFACTANTS, CATIONIC, QUATERNARY AMMONIUM, MONOALKYL**  
9/1993

**LIST OF MONOALKYL-TRIMETHYL-AMMONIUM CHLORIDE SURFACTANTS USED TO DEVELOP THE  
SAR FOR QUATERNARY AMMONIUM SURFACTANTS FOR FISH**

Number of Carbons	Species Tested	Acute LC50 (mg/L)
10	Golden orfe	68
12	Golden orfe	9.0
14	Golden orfe	2.1
16	Golden orfe	0.36
18	Golden orfe	0.41
21	Golden orfe	0.42

## SURFACTANTS, CATIONIC, QUATERNARY AMMONIUM, MONOALKYL

9/1993

### SAR

### SURFACTANTS, CATIONIC, QUATERNARY AMMONIUM, MONOALKYL

**Organism:** Daphnid  
**Duration:** Acute  
**Endpoint:** LC50 (Mortality)

**Equation:** Determine the average number of carbons in the hydrophobic alkyl chain of the surfactant. If the average length of the carbon chain is between 16 and 22 carbons, use the SAR equation:

$$\text{Log LC50 (mg/L)} = -1.64 + 0.115 (\text{average length of carbon chain})$$

If the length of the carbon chain is at least 10 but less than 16, use the SAR equation:

$$\text{Log LC50 (mg/L)} = 2.07 - 0.13 (\text{average length of carbon chain})$$

**Maximum Value:** average carbon chain length of 22 carbons  
**Minimum Value:** average carbon chain length of 10 carbons

**Application:** This SAR may be applied to monoalkyl (trimethyl) quaternary ammonium surfactants which are dispersible in water. This SAR may be used to estimate toxicity for:

1. monoalkyl cationic surfactants
2. monoalkyl phosphonium surfactants
3. monoalkyl sulfonium surfactants

**Limitations:** This SAR may be used for monoalkyl quaternary ammonium surfactants where the anionic salt has less than 8 carbons in the alkyl chain. If the alkyl chain contains 8 or more carbons, the cationic surfactant and the anionic surfactant will form a strong ion pair. This ion pair will be much less soluble in water and consequently will be less toxic to daphnids.

**References:** Nabholz JV. 1987. The SAR for monoalkyl (trimethyl) quaternary ammonium surfactants. Washington, DC: Environmental Effects Branch, Health and Environmental Review Division, Office of Toxic Substances, United States Environmental Protection Agency.

Knauf W. 1973. Summary of the toxicity of surfactants to aquatic organisms. Tenside Detergents 5:251-255.

**SURFACTANTS, CATIONIC, QUATERNARY AMMONIUM, MONOALKYL**  
9/1993

**LIST OF MONOALKYL-TRIMETHYL-AMMONIUM CHLORIDE SURFACTANTS USED TO DEVELOP THE  
SAR FOR QUATERNARY AMMONIUM SURFACTANTS FOR DAPHNIDS**

Number of Carbons	Species Tested	Acute LC50 (mg/L)
10	<u>Daphnia magna</u>	7.0
12	<u>Daphnia magna</u>	3.2
14	<u>Daphnia magna</u>	1.7
16	<u>Daphnia magna</u>	1.2
18	<u>Daphnia magna</u>	3.2
21	<u>Daphnia magna</u>	6.0

**SURFACTANTS, CATIONIC, QUATERNARY AMMONIUM, MONOALKYL**  
9/1993

<b>SAR</b>	<b>SURFACTANTS, CATIONIC, QUATERNARY AMMONIUM, MONOALKYL</b>
<b>Organism:</b>	Snail
<b>Duration:</b>	Acute
<b>Endpoint:</b>	LC50 (Mortality)
<b>Equation:</b>	<p>Determine the average number of carbons in the hydrophobic alkyl chain of the surfactant. If the average length of the carbon chain is between 16 and 22 carbons, use the SAR equation:</p> $\text{Log LC50 (mg/L)} = -1.56 + 0.087 (\text{average length of carbon chain})$ <p>If the length of the carbon chain is at least 10 but less than 16, use the SAR equation:</p> $\text{Log LC50 (mg/L)} = 5.74 - 0.37 (\text{average length of carbon chain})$
<b>Maximum Value:</b>	carbon chain length of 22 carbons
<b>Minimum Value:</b>	carbon chain length of 10 carbons
<b>Application:</b>	<p>This SAR may be applied to monoalkyl (trimethyl) quaternary ammonium surfactants which are dispersible in water. This SAR may be used to estimate toxicity for:</p> <ol style="list-style-type: none"><li>1. monoalkyl cationic surfactants</li><li>2. monoalkyl phosphonium surfactants</li><li>3. monoalkyl sulfonium surfactants</li></ol>
<b>Limitations:</b>	<p>This SAR may be used for monoalkyl quaternary ammonium surfactants where the anionic salt has less than 8 carbons in the alkyl chain. If the alkyl chain contains 8 or more carbons, the cationic surfactant and the anionic surfactant will form a strong ion pair. This ion pair will be much less soluble in water and consequently will be less toxic to snails.</p>
<b>References:</b>	<p>Nabholz JV. 1987. The SAR for monoalkyl (trimethyl) quaternary ammonium surfactants. Washington, DC: Environmental Effects Branch, Health and Environmental Review Division, Office of Toxic Substances, United States Environmental Protection Agency.</p> <p>Knauf W. 1973. Summary of the toxicity of surfactants to aquatic organisms. Tenside Detergents 5:251-255.</p>

**SURFACTANTS, CATIONIC, QUATERNARY AMMONIUM, MONOALKYL**  
9/1993

LIST OF MONOALKYL-TRIMETHYL-AMMONIUM CHLORIDE SURFACTANTS USED TO DEVELOP THE  
SAR FOR QUATERNARY AMMONIUM SURFACTANTS FOR SNAILS

Number of Carbons	Species Tested	Acute LC50 (mg/L)
10	Water snail	100
12	Water snail	23
14	Water snail	3.5
16	Water snail	0.7
18	Water snail	1.0
21	Water snail	1.9

**SURFACTANTS, CATIONIC, QUATERNARY AMMONIUM, DIALKYL**  
9/1993

<b>SAR</b>	<b>SURFACTANTS, CATIONIC, QUATERNARY AMMONIUM, DI-ALKYL</b>
<b>Organism:</b>	Fish
<b>Duration:</b>	96-h
<b>Endpoint:</b>	LC50 and ChV (Mortality)
<b>Equation:</b>	<p>Calculate the average log <math>K_{ow}</math> for the two alkyl groups and use the average value in the SAR equation:</p> $\text{Log 96-h LC50 (mM/L)} = 0.747 - 0.367 \log K_{ow}$ <p>To determine the chronic toxicity value (ChV) of a di-alkyl quaternary ammonium surfactant to fish, divide the 96-hour LC50 value by 26.</p>
<b>Statistics:</b>	$N = 6; R^2 = 0.9$
<b>Maximum Value:</b>	There are no limits on the log $K_{ow}$ values.
<b>Maximum MW:</b>	There are no limits on the molecular weight of the two alkyl groups of the cationic surfactant.
<b>Application:</b>	<p>This SAR may be applied to cationic dialkyl (dimethyl) quaternary ammonium surfactants which are dispersible in water. This SAR may be used to estimate toxicity for:</p> <ol style="list-style-type: none"><li>1. dialkyl cationic surfactants</li><li>2. dialkyl phosphonium surfactants</li><li>3. dialkyl sulfonium surfactants</li></ol>
<b>Limitations:</b>	None.
<b>References:</b>	<p>FDA. Unpublished data.</p> <p>ITC IR-488.</p> <p>USEPA. ECOTOX database. P85-505 Standard Review.</p>

**SURFACTANTS, CATIONIC, QUATERNARY AMMONIUM, DIALKYL**  
9/1993

**SURFACTANTS, CATIONIC, QUATERNARY AMMONIUM, DIALKYL**  
9/1993

<b>SAR</b>	<b>SURFACTANTS, CATIONIC, QUATERNARY AMMONIUM, DI-ALKYL</b>
<b>Organism:</b>	Daphnid
<b>Duration:</b>	48-h
<b>Endpoint:</b>	LC50 AND ChV (Mortality)
<b>Equation:</b>	<p>Calculate the average <math>\log K_{ow}</math> for the two alkyl groups and use the average value in the SAR equation:</p> $\text{Log 48-h LC50 (mM/L)} = 0.874 - 0.462 \log K_{ow}$ <p>To determine the chronic toxicity value (ChV) of a di-alkyl quaternary ammonium surfactant to daphnids, divide the 48-hour LC50 value by 1.8.</p>
<b>Statistics:</b>	$N = 4; R^2 = 0.94$
<b>Maximum Value:</b>	There are no limits on the $\log K_{ow}$ values of the two alkyl groups of the cationic surfactant.
<b>Maximum MW:</b>	There are no limits on the molecular weight of the two alkyl groups of the cationic surfactant.
<b>Application:</b>	<p>This SAR may be applied to cationic dialkyl (dimethyl) quaternary ammonium surfactants which are dispersible in water. This SAR may be used to estimate toxicity for:</p> <ol style="list-style-type: none"><li>1. dialkyl cationic surfactants</li><li>2. dialkyl phosphonium surfactants</li><li>3. dialkyl sulfonium surfactants</li></ol>
<b>Limitations:</b>	None.
<b>References:</b>	<p>FDA. Unpublished data.</p> <p>ITC. IR-488.</p> <p>EPA. ECOTOX database. P85-505 Standard Review.</p>

**SURFACTANTS, CATIONIC, QUATERNARY AMMONIUM, DIALKYL**  
9/1993

## SURFACTANTS, CATIONIC, QUATERNARY AMMONIUM, DIALKYL

9/1993

<b>SAR</b>	<b>SURFACTANTS, CATIONIC, QUATERNARY AMMONIUM, DI-ALKYL</b>
<b>Organism:</b>	Green Algae
<b>Duration:</b>	96-h
<b>Endpoint:</b>	EC50 and ChV
<b>Equation:</b>	<p>Calculate the average <math>\log K_{ow}</math> for the two alkyl groups and use the average value in the SAR equation:</p> $\text{Log 96-h EC50 (mM/L)} = -0.595 - 0.296 \log K_{ow}$ <p>To determine the chronic toxicity value (ChV) of a di-alkyl quaternary ammonium surfactant to green algae, divide the 96-hour EC50 value by 4.</p>
<b>Statistics:</b>	$N = 3; R^2 = 0.99$
<b>Maximum Value:</b>	There are no limits on the $\log K_{ow}$ values of the two alkyl groups of the cationic surfactant.
<b>Maximum MW:</b>	There are no limits on the molecular weight of the two alkyl groups of the cationic surfactant.
<b>Application:</b>	<p>This SAR may be applied to cationic dialkyl (dimethyl) quaternary ammonium surfactants which are dispersible in water. This SAR may be used to estimate toxicity for:</p> <ol style="list-style-type: none"><li>1. dialkyl cationic surfactants</li><li>2. dialkyl phosphonium surfactants</li><li>3. dialkyl sulfonium surfactants</li></ol>
<b>Limitations:</b>	None
<b>References:</b>	<p>FDA. Unpublished data.</p> <p>ITC. IR-488.</p> <p>EPA. ECOTOX database. P85-505 Standard Review.</p>

**SURFACTANTS, CATIONIC, QUATERNARY AMMONIUM, DIALKYL**  
9/1993

**SAR**

**SURFACTANTS, NONIONIC**

**Organism:**

Fish and Daphnid

**Duration:**

96-h, 48-h

**Endpoint:**

LC50 (Mortality) in mg/L

**Equation:**

Determine the number of carbons in the alkyl chains and the number of ethoxylate groups in the surfactant. Determine the toxicity using the appropriate SAR equation based on the length of the carbon chain:

$C = 8; \text{Log LC50} = 0.952 + 0.130 (\text{number of ethoxylates})$

$C = 9; \text{Log LC50} = 0.796 + 0.120 (\text{number of ethoxylates})$

$C = 10; \text{Log LC50} = 0.642 + 0.112 (\text{number of ethoxylates})$

$C = 11; \text{Log LC50} = 0.261 + 0.103 (\text{number of ethoxylates})$

$C = 12; \text{Log LC50} = -0.204 + 0.0996 (\text{number of ethoxylates})$

$C = 13; \text{Log LC50} = -0.388 + 0.092 (\text{number of ethoxylates})$

$C = 14; \text{Log LC50} = -0.480 + 0.0847 (\text{number of ethoxylates})$

$C = 15; \text{Log LC50} = -0.533 + 0.0776 (\text{number of ethoxylates})$

$C = 16; \text{Log LC50} = -0.775 + 0.072 (\text{number of ethoxylates})$

$C = 17; \text{Log LC50} = -1.054 + 0.0674 (\text{number of ethoxylates})$

$C = 18; \text{Log LC50} = -1.290 + 0.063 (\text{number of ethoxylates})$

**Statistics:**

**Maximum Value:**

Maximum carbon chain length of 18; minimum carbon chain length of 8; the maximum number of ethoxylates is 55.

**Application:**

This SAR may be used to estimate the toxicity for the following classes of nonionic surfactants:

1. Alcohol ethoxylate surfactants
2. Alkyl ethoxylate surfactants
3. Nonionic surfactants

Generally, this SAR is expected to be applicable to other nonionic surfactants, such as alcohol ethoxylate-propoxylate surfactants where number of ethoxylates is greater than the number of propoxylates.

## **SURFACTANTS, NONIONIC**

9/1993

### **Limitations:**

When the number of ethoxylates is less than 5, chemicals may begin to act less like surfactants and more like neutral organic chemicals. Alcohol propoxylates and alcohol butoxylates will not act like surfactants; the propoxylate and butoxylate units are not water soluble enough. Alcohol propoxylates and alcohol butoxylates should be treated like neutral organic chemicals.

### **References:**

Nabholz JV. 1988. The structure-activity relationships between nonionic surfactants. Washington, DC: Environmental Effects Branch, Health and Environmental Review Division, Office of Toxic Substances, United States Environmental Protection Agency.

# SURFACTANTS, NONIONIC

9/1993

## LIST OF NONIONIC SURFACTANTS USED TO DEVELOP FISH 96-h AND DAPHNID 48-H LC50 SARS.

Number of Carbons	Number of Ethoxylates	Species	Time (hours)	LC50 (mg/L)
8	12.0	Golden orfe	96	465.0
10	2.5	Rainbow trout	96	5-7
10	5.0	Rainbow trout	96	8-9
12	6.0	Fish spp.	96	2.8
12	9.0	Fish spp.	96	5.4
12	12.0	Fish spp.	96	4.4
12	12.0	Golden orfe	96	4.4
12	15.0	Fish spp.	96	22.0
12.5	2.0	Rainbow trout	96	1-2
12.5	5.3	Rainbow trout	96	1.0
12.5	6.5	Daphnia	24	1.05
12.5	6.5	Rainbow trout	96	2.36
12.5	6.5	Bluegill	96	0.57
12.5	6.5	Daphnia	24	0.57
12.5	6.5	Daphnia	96	1.14
13	6.3	Fathead minnow	24	1.8
13	6.3	Goldfish	24	1.4
13	6.3	Daphnia	48	2.4
13	7.4	Fathead minnow	24	1.8
13	7.4	Goldfish	24	1.4
13	7.4	Daphnia	24	2.3
13	8.0	Goldfish	48	1.4
13	8.0	Harlequin fish	48	1.2
13	8.0	Golden orfe	96	1.8
13	8.0	Rainbow trout	96	0.8
13	8.0	Golden orfe	96	2.7
13	10.5	Harlequin fish	96	1.6-2.8
13	10.5	Rainbow trout	96	1.8
13	10.5	Rainbow trout	96	0.8
13	10.5	Golden orfe	96	4.1
13	10.5	Golden orfe	96	4.1
13	10.5	Goldfish	48	3.0
13	11.0	Golden orfe	48	2.7
13	11.0	Daphnia	24	5.1
13	11.0	Rainbow trout	48	6.2
13.5	3.0	Bluegill	96	1.5
13.5	3.0	Rainbow trout	96	1.3-1.7
13.5	3.0	Rainbow trout	96	3.9
13.5	7.0	Rainbow trout	96	2.7
13.5	9.0	Bluegill	96	2.1
13.5	9.0	Bluegill	96	11.0
13.5	9.0	Channel catfish	96	1.2
13.5	9.0	Daphnia	24	1.71

**SURFACTANTS, NONIONIC**

9/1993

Continued

Number of Carbons	Number of Ethoxylates	Species	Time (hours)	LC50 (mg/L)
13.5	9.0	Bluegill	96	7.8
17	14.0	Minnow	24	3.4
17	14.0	Rainbow trout	96	0.4
17	14.0	Golden orfe	96	2.3
17	14.0	Golden orge	96	2.5
17	14.0	Harlequin fish	96	0.7

## SURFACTANTS, ETHOMEEN

9/1993

### SAR

### SURFACTANTS, ETHOMEEN

**Organism:**

Fish, Daphnid, and Algae

**Duration:**

96-h, 48-h, and 96-h

**Endpoint:**

LC50, LC50, and EC50 (Mortality) in mg/L

**Equation:**

Determine the number of carbons in the alkyl chains and the number of ethoxylate groups in the surfactant. Determine the toxicity using the appropriate SAR equation based on the length of the carbon chain:

$$C = 8; \text{ Log LC50} = 1.022 + 0.122 (\text{number of ethoxylates})$$

$$C = 9; \text{ Log LC50} = 0.794 + 0.116 (\text{number of ethoxylates})$$

$$C = 10; \text{ Log LC50} = 0.553 + 0.112 (\text{number of ethoxylates})$$

$$C = 11; \text{ Log LC50} = 0.335 + 0.104 (\text{number of ethoxylates})$$

$$C = 12; \text{ Log LC50} = 0.107 + 0.098 (\text{number of ethoxylates})$$

$$C = 13; \text{ Log LC50} = -0.102 + 0.092 (\text{number of ethoxylates})$$

$$C = 14; \text{ Log LC50} = -0.348 + 0.086 (\text{number of ethoxylates})$$

$$C = 15; \text{ Log LC50} = -0.566 + 0.079 (\text{number of ethoxylates})$$

$$C = 16; \text{ Log LC50} = -0.706 + 0.074 (\text{number of ethoxylates})$$

$$C = 17; \text{ Log LC50} = -1.057 + 0.069 (\text{number of ethoxylates})$$

$$C = 18; \text{ Log LC50} = -1.316 + 0.063 (\text{number of ethoxylates})$$

**Maximum Value:**

18 carbons in the alkyl chain; 55 ethoxylates

**Maximum MW:**

**Application:**

This SAR may be used to estimate the toxicity of ethomeen surfactants (i.e., ethoxylated beta-amine surfactants) with a carboxylic acid terminus.

**Limitations:**

None.

**References:**

Nabholz JV. 1986. The structure-activity relationships between nonionic surfactants. Washington, DC: Environmental Effects Branch, Health and Environmental Review Division, Office of Toxic Substances, United States Environmental Protection Agency.

## **SURFACTANTS, ETHOMEEN**

9/1993

<b>SAR</b>	<b>THIAZOLINONES, ISO</b>
<b>Organism:</b>	Fish
<b>Duration:</b>	96-h
<b>Endpoint:</b>	LC50 (Mortality)
<b>Equation:</b>	$\text{Log LC50 (mM/L)} = -2.159 - 0.068 \log K_{ow}$
<b>Statistics:</b>	$N = 2; R^2 = 1.0$
<b>Maximum log <math>K_{ow}</math>:</b>	5.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This equation may be used to estimate the toxicity of isothiazolinones or allyl thioamides.
<b>Limitations:</b>	If the log $K_{ow}$ value is greater than 5.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.
<b>References:</b>	United States Environmental Protection Agency (USEPA). 1993. OPPT PMN ECOTOX. Washington, DC: Office of Pollution Prevention and Toxics, USEPA.

**LIST OF ISOTHIAZOLINONES USED TO DEVELOP THE FISH 96-h LC50 SAR**

CHEMICAL	96-H LC50 (mg/L)	Log $K_{ow}$	Ref.
Chemical identity CBI	0.90	0.6	EPA

EPA = USEPA (1993), chemical identity is Confidential Business Information under TSCA.

**THIAZOLINONES, ISO**  
9/1993

<b>SAR</b>	<b>THIAZOLINONES, ISO</b>
<b>Organism:</b>	Daphnid
<b>Duration:</b>	48-h
<b>Endpoint:</b>	LC50 (Mortality)
<b>Equation:</b>	$\text{Log LC50 (mM/L)} = -2.0 - 0.159 \log K_{ow}$
<b>Statistics:</b>	$N = 2; R^2 = 1.0$
<b>Maximum log <math>K_{ow}</math>:</b>	5.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This equation may be used to estimate the toxicity of isothiazolinones or allyl thioamides.
<b>Limitations:</b>	If the log $K_{ow}$ value is greater than 5.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.
<b>References:</b>	United States Environmental Protection Agency (USEPA). 1993. OPPT PMN ECOTOX. Washington, DC: Office of Pollution Prevention and Toxics, USEPA.

**LIST OF ISOTHIAZOLINONES USED TO DEVELOP THE FISH 96-h LC50 SAR**

CHEMICAL	96-H LC50 (mg/L)	Log $K_{ow}$	Ref.
Chemical identity CBI	1.2	0.6	EPA

EPA = USEPA (1993); chemical identity is Confidential Business Information under TSCA.

THIAZOLINONES, ISO  
9/1993

**SAR**                                      **THIAZOLINONES, ISO**

**Organism:**                                      Green Algae  
**Duration:**                                      96-h  
**Endpoint:**                                      EC50

**Equation:**                                       $\text{Log LC50 (mM/L)} = -2.555 - 0.241 \log K_{ow}$

**Statistics:**                                       $N = 2; R^2 = 1.0$

**Maximum log  $K_{ow}$ :**                                      6.4  
**Maximum MW:**                                      1000.0

**Application:**                                      This equation may be used to estimate the toxicity of isothiazolinones or allyl thioamides.

**Limitations:**                                      If the log  $K_{ow}$  value is greater than 6.4, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation.

**References:**                                      United States Environmental Protection Agency (USEPA). 1993. OPPT PMN ECOTOX. Washington, DC: Office of Pollution Prevention and Toxics, USEPA.

**LIST OF ISOTHIAZOLINONES USED TO DEVELOP THE FISH 96-h LC50 SAR**

CHEMICAL	96-H LC50 (mg/L)	Log $K_{ow}$	Ref.
Chemical identity CBI	0.290	0.6	EPA

EPA = USEPA (1993); chemical identity is Confidential Business Information under TSCA.

**THIAZOLINONES, ISO**  
9/1993

<b>SAR</b>	<b>THIAZOLINONES, ISO</b>
<b>Organism:</b>	Green Algae
<b>Duration:</b>	
<b>Endpoint:</b>	Chronic Value
<b>Equation:</b>	$\text{Log LC50 (mM/L)} = -2.938 - 0.270 \log K_{ow}$
<b>Statistics:</b>	$N = 2; R^2 = 1.0$
<b>Maximum log <math>K_{ow}</math>:</b>	8.0
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This equation may be used to estimate the toxicity of isothiazolinones or allyl thioamides.
<b>Limitations:</b>	If the log $K_{ow}$ value is greater than 8.0, or if the compound is solid and the LC50 exceeds the water solubility, no effects expected at saturation
<b>References:</b>	United States Environmental Protection Agency (USEPA). 1993. OPPT PMN ECOTOX. Washington, DC: Office of Pollution Prevention and Toxics, USEPA.

**LIST OF ISOTHIAZOLINONES USED TO DEVELOP THE FISH 96-h LC50 SAR**

CHEMICAL	96-H LC50 (mg/L)	Log $K_{ow}$	Ref.
Chemical identity CBI	0.130	0.6	EPA

EPA = USEPA (1993); chemical identity is Confidential Business Information under TSCA.

THIAZOLINONES, ISO  
9/1993

**SAR**

**THIOLS AND MERCAPTANS**

**Organism:**

Fish

**Duration:**

96-h

**Endpoint:**

LC50 (Mortality)

**Equation:**

$$\text{Log LC50 (mM/L)} = -1.022 - 0.447 \log K_{ow}$$

**Statistics:**

$$N = 4; R^2 = 0.85$$

**Maximum log  $K_{ow}$ :**

6.5

**Maximum MW:**

1000.0

**Application:**

This equation may be used to estimate the toxicity for thiols and mercaptans. Thiols with a carboxylic acid substitution will be about 10 times less toxic than the toxicity value predicted by using this SAR with a log  $K_{ow}$  and molecular weight for the free acid. Therefore, for thiols with a carboxylic acid substitution, predict the toxicity values for the free acid and multiply by 10.

**Limitations:**

For thiols with log  $K_{ow}$  values greater than 4.5, the toxicity prediction may only apply to rainbow trout and other cold water fish species. While a 96-h LC50 value was measured for t-dodecane thiol (log  $K_{ow}$  = 6.2), nonylthiol (log  $K_{ow}$  = 4.9) showed no toxicity at saturation with fathead minnows. The recommended species for testing thiols with log  $K_{ow}$  values greater than 4.5 is rainbow trout using flow-through methods, measured concentrations, and treatment concentrations which do not exceed the aqueous solubility limit of the thiol being tested.

If the log  $K_{ow}$  value is greater than 6.5, or if the compound is solid and the LC50 exceeds the water solubility, use SAR with longer exposure.

**References:**

Bender ME. 1969. The toxicity of the hydrolysis and breakdown products of malathion to the fathead minnow *Pimephales promelas*, Rafinesque. Water Research 3:571-582.

U.S. Environmental Protection Agency. 1991. Toxicity of data gap compounds to fathead minnow (*Pimephales promelas*) and daphnids (*Daphnia magna*). Duluth, MN: Environmental Research Laboratory, Office of Research and Development, USEPA.

U.S. Environmental Protection Agency. 1992. TSCA Sec. 8(e) submission number 994. Washington, DC: Office of Pollution Prevention and Toxics, USEPA.

Verschueren K. 1983. Handbook of environmental data on organic chemicals. 2nd ed. New York, NY: Van Nostrand Reinhold Co

**THIOLS AND MERCAPTANS**  
9/1993

**SAR**

**THIOLS AND MERCAPTANS**

**Organism:**

Daphnid

**Duration:**

48-h

**Endpoint:**

LC50 (Mortality)

**Equation:**

$\text{Log LC50 (mM/L)} = -3.2 - 0.097 \log K_{ow}$

**Statistics:**

$N = 3; R^2 = 0.46$

**Maximum log  $K_{ow}$ :**

5.0

**Maximum MW:**

1000.0

**Application:**

This equation may be used to estimate the toxicity for thiols and mercaptans.

**Limitations:**

If the log  $K_{ow}$  is greater than 5.0, or if the compound is solid and the LC50 exceeds the water solubility, use SAR with longer exposure.

**References:**

U.S. Environmental Protection Agency. 1991. Toxicity of data gap compounds to fathead minnows (*Pimephales promelas*) and daphnids (*Daphnia magna*). Duluth, MN: Environmental Research Laboratory, Office of Research and Development, USEPA.

U.S. Environmental Protection Agency. 1992. TSCA Sec. 8(e) submission number 994. Washington, DC: Office of Pollution Prevention and Toxics, USEPA.



SAR

TRIAZINES, SUBSTITUTED

For fish and daphnid use SAR for **NEUTRAL ORGANICS**;

This category includes substituted triazines which can be aromatic, partially aromatic (or partially saturated), and unsaturated. The nitrogens in the triazine ring may be symmetrical or asymmetrical. Substitutions on the carbons may include but not be limited to: aliphatic alcohols; ketones; benzene and substituted benzenes; aliphatic hydrocarbons, alkyenes and alkynes; free amines and substituted amines; cyclic aliphatic hydrocarbons; halogens; amides; cyanides; ethers; methoxy groups; sulfides; azido groups; and carboxylic acid esters. Substitutions on the nitrogens may include but not be limited to: free amines and substituted amines;  $-N=CH$ ; aliphatic hydrocarbons, alkyenes and alkynes; and benzene and substituted benzenes. Hazard Concerns: many members of this category are commercial herbicides which are used to control both aquatic plants and terrestrial plants. Their mode of toxic action is generally considered to be inhibition of photosynthesis. Many members of this class are toxic to algae at  $< 1$  mg/L and toxic to terrestrial vascular plants at  $< 1$  mg/kg. Members of this group can also be highly toxic to fish and aquatic invertebrates. Toxicity is expected to be related to the octanol/water partition coefficient with respect to fish and aquatic invertebrates, but toxicity to plants may not be related to  $K_{ow}$  when  $\log K_{ow} < 5$ . When the  $\log K_{ow}$  is  $< 5$ , algae and terrestrial plants are expected to be the most sensitive species. As  $\log K_{ow}$  increases, species differences are expected to diminish. At this time there is no formalized SAR for this category for any species. Toxicity predictions will be made using either the closest analog or averaging data for the two closest analogs which bracket the chemical under question.

TRIAZINES, SUBSTITUTED  
9/1993

22-6  
X-5

**SAR**

**UREAS, SUBSTITUTED**

<b>Organism:</b>	Algae
<b>Duration:</b>	4-h
<b>Endpoint:</b>	EC50 (Inhibition of Photosynthesis)
<b>Equation:</b>	$\text{Log EC50 (mM/L)} = -1.29 \log K_{ow} + 0.133$
<b>Statistics:</b>	$N = 12; R^2 = 0.944$
<b>Maximum log <math>K_{ow}</math>:</b>	3.9
<b>Maximum MW:</b>	1000.0
<b>Application:</b>	This SAR may be used to estimate the toxicity for substituted ureas.
<b>Limitations:</b>	If the log $K_{ow}$ value is greater than 3.9 and less than 7.9, use SAR with longer exposure. If the log $K_{ow}$ value is greater than 8.0, no effects expected at saturation.
<b>References:</b>	<p>Wesséls JSC and Van Der Veen R. 1956. The action of some derivatives of phenylurethan and of 3-phenyl-1,1-dimethylurea on the Hill reaction. <i>Biochem. Biophys. Acta</i> 19.</p> <p>Hansch C. 1969. Theoretical considerations of the structure-activity relationship in photosynthesis inhibitors. In: <i>Progress in Photosynthesis Research</i>, Vol. III. Metzner H, ed. pp. 1685-1692.</p>

# UREAS, SUBSTITUTED

9/1993

## LIST OF SUBSTITUTED UREAS USED TO DEVELOP THE ALGAE 4-h EC50 SAR.

CHEMICAL	4-h EC50 (mg/L)	Log K <sub>ow</sub>	Ref.
Ethyl-N-phenylcarbamate (phenylurethan)	5x10 <sup>-4</sup>	*	
Ethyl-N-(3-chlorophenyl)-carbamate	10 <sup>-4</sup>	*	
Ethyl-N-(4-chlorophenyl)-carbamate	10 <sup>-4</sup>	*	
Ethyl-N-(4-nitrophenyl)-carbamate	2x10 <sup>-4</sup>	*	
Allyl-N-phenylcarbamate	5x10 <sup>-4</sup>	*	
Allyl-N-(4-chlorophenyl)-carbamate	8x10 <sup>-5</sup>	*	
Ethyl-N-(3,4-dichlorophenyl)-carbamate	2x10 <sup>-5</sup>	*	
Ethyl-N-(2,5-dichlorophenyl)-carbamate	3x10 <sup>-4</sup>	*	
Benzyl-N-phenylcarbamate	2x10 <sup>-4</sup>	*	
Ethyl-N-(4-hydroxyphenyl)-carbamate	3x10 <sup>-3</sup>	*	
Ethyl-N-(3-hydroxyphenyl)-carbamate	10 <sup>-3</sup>	*	
3-Phenyl-1,1-dimethylurea	4x10 <sup>-5</sup>	*	
3-(4-Chlorophenyl)-1,1-dimethylurea (CMU)	4x10 <sup>-6</sup>	*	
3-(3-Chlorophenyl)-1,1-dimethylurea	2x10 <sup>-6</sup>	*	
3-(3,4-Dichlorophenyl)-1,1-dimethylurea	2x10 <sup>-7</sup>	*	
3-(3,4,5-Trichlorophenyl)-1,1-dimethylurea	2x10 <sup>-7</sup>	*	
3-(4-Nitrophenyl)-1,1-dimethylurea	8x10 <sup>-6</sup>	*	
3-(3-Nitrophenyl)-1,1-dimethylurea	1.3x10 <sup>-5</sup>	*	
3-(4-Trifluoromethylphenyl)-1,1-dimethylurea	4x10 <sup>-6</sup>	*	
3-(3-Trifluoromethylphenyl)-1,1-dimethylurea	6x10 <sup>-4</sup>	*	
4-(3,3-Dimethylureido)-S-trichloromethyl-phenyl-thiosulfonate	4x10 <sup>-7</sup>	*	
3-(4-Methylphenyl)-1,1-dimethylurea	3x10 <sup>-5</sup>	*	
3-(4-Methoxyphenyl)-1,1-dimethylurea	3x10 <sup>-5</sup>	*	
3-(4-Dimethylaminophenyl)-1,1-dimethylurea	2x10 <sup>-4</sup>	*	
3-(4-Acetylaminophenyl)-1,1-dimethylurea	2x10 <sup>-3</sup>	*	

\* = Not available at this time.

## INORGANICS



**Organism:** Aquatic life (freshwater)  
**Duration:** 96-hour  
**Endpoint:** No Observable Effect Concentration (NOEC)  
**Equation:**  $\text{NOEC (mg/L)} = (0.087 \cdot \text{MW})/26.981$   
**Application:** This equation may be used to estimate the acute toxicity of organic and inorganic compounds containing aluminum.  
**Limitations:** This equation is based on the pH dependent Ambient Water Quality Criteria for aluminum. The criteria for pH values between 6.5 and 9.0 were used. If the pH of the solution is less than 6.5.  
**References:** United States Environmental Protection Agency (USEPA). 1980. Ambient Water Quality Criteria for Aluminum. Washington, DC: Office of Water, Criteria and Standards Division.

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**Organism:** Aquatic Life (freshwater)  
**Duration:** 1-hour  
**Endpoint:** No Observable Effect Concentration (NOEC)  
**Equation:**  $\text{NOEC (mg/L)} = (0.750 \cdot \text{MW})/26\ 981$   
**Application:** This equation may be used to estimate the acute toxicity of compounds containing aluminum.  
**Limitations:** This equation is based on the pH dependent Ambient Water Quality Criteria for aluminum. The criteria for pH values between 6.5 and 9.0 were used.  
**References:** United States Environmental Protection Agency (USEPA). 1980. Ambient Water Quality Criteria for Aluminum. Washington, DC: Office of Water, Criteria and Standards Division.

**ALUMINUM**  
9/1993

**Organism:** Aquatic life (freshwater)  
**Duration:**  
**Endpoint:** Acute Value  
**Equation:**  $\text{Acute Value (mg/L)} = (0.088 \cdot \text{MW})/121.75$   
**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing antimony.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1992. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division.

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**Organism:** Aquatic life (freshwater)  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (0.030 \cdot \text{MW})/121.75$   
**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing antimony.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1992. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division.

**ANTIMONY**  
9/1993

**Organism:** Aquatic life (marine)  
**Duration:**  
**Endpoint:** Acute Value  
**Equation:**  $\text{Acute Value (mg/L)} = (1.5 \cdot \text{MW})/121.75$   
**Application:** This equation may be used to estimate the toxicity of both inorganic and organic compounds containing antimony.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1992. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division.

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**Organism:** Aquatic life (marine)  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (0.500 \cdot \text{MW})/121.75$   
**Application:** This equation may be used to estimate the toxicity of organic and inorganic compounds containing antimony  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1992. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division.

**ANTIMONY**  
9/1993

**Organism:** Aquatic life (freshwater)  
**Duration:**  
**Endpoint:** Acute Value  
**Equation:**  $\text{Acute Value (mg/L)} = (0.360 \cdot \text{MW})/74.92$   
**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing arsenic(III).  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

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**Organism:** Aquatic life (freshwater)  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (0.190 \cdot \text{MW})/74.92$   
**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing arsenic(III).  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

ARSENIC(III)  
9/1993

**Organism:** Aquatic life (marine)  
**Duration:**  
**Endpoint:** Acute Value  
**Equation:**  $\text{Acute Value (mg/L)} = (0.069 \cdot \text{MW})/74.92$   
**Application:** This equation may be used to estimate the toxicity of both inorganic and organic compounds containing arsenic(III).  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001

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**Organism:** Aquatic life (marine)  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (0.036 \cdot \text{MW})/74.92$   
**Application:** This equation may be used to estimate the toxicity of organic and inorganic compounds containing arsenic(III)  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

**BORON**  
9/1993

<b>Organism:</b>	Aquatic life (freshwater)
<b>Duration:</b>	
<b>Endpoint:</b>	Acute Value
<b>Equation:</b>	$\text{Acute Value (mg/L)} = (0.0039 \cdot \text{MW})/112.41$
<b>Application:</b>	This equation may be used to estimate the toxicity of both organic and inorganic compounds containing cadmium.
<b>Limitations:</b>	<p>This equation is based on the hardness dependent Water Quality Criteria for cadmium. The criterion for a hardness of 100 mg/L as <math>\text{CaCO}_3</math> was used. For a solution with a hardness of 50 mg/L as <math>\text{CaCO}_3</math>, use the following equation:</p> $\text{Acute Value (mg/L)} = (0.0018 \cdot \text{MW})/112.41$ <p>For a solution with a hardness of 200 mg/L as <math>\text{CaCO}_3</math>, use the following equation:</p> $\text{Acute Value (mg/L)} = (0.0086 \cdot \text{MW})/112.41$
<b>References:</b>	United States Environmental Protection Agency (USEPA). 1986 Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

**CADMIUM**  
9/1993

<b>Organism:</b>	Aquatic life (freshwater)
<b>Duration:</b>	
<b>Endpoint:</b>	Chronic Value (ChV)
<b>Equation:</b>	$\text{ChV (mg/L)} = (0.0011 \cdot \text{MW})/112.41$
<b>Application:</b>	This equation may be used to estimate the toxicity of both organic and inorganic compounds containing cadmium.
<b>Limitations:</b>	<p>This equation is based on the hardness dependent Water Quality Criteria for cadmium. The criterion for a hardness of 100 mg/L as <math>\text{CaCO}_3</math> was used. For solutions with a hardness of 50 mg/L as <math>\text{CaCO}_3</math>, use the following equation:</p> $\text{ChV (mg/L)} = (0.00066 \cdot \text{MW})/112.41$ <p>For solutions with a hardness of 200 mg/L as <math>\text{CaCO}_3</math>, use the following equation:</p> $\text{ChV (mg/L)} = (0.002 \cdot \text{MW})/112.41$
<b>References:</b>	United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

**CADMIUM**  
9/1993

**Organism:** Aquatic life (marine)  
**Duration:**  
**Endpoint:** Acute Value  
**Equation:**  $\text{Acute Value (mg/L)} = (0.043 \cdot \text{MW})/112.41$   
**Application:** This equation may be used to estimate the toxicity of both inorganic and organic compounds containing cadmium.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

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**Organism:** Aquatic life (marine)  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (0.0093 \cdot \text{MW})/112.41$   
**Application:** This equation may be used to estimate the toxicity of organic and inorganic compounds containing cadmium.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

**CADMIUM**  
9/1993

**Organism:** Daphnid  
**Duration:** 48-hour  
**Endpoint:** LC50

**Equation:**  $LC50 \text{ (mg/L)} = (7.4 \cdot MW)/132.9$

**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing cesium.

**Limitations:** None

**References:** United States Environmental Protection Agency (USEPA). 1991. Hazard Profiles for Selected Heavy Metals. Washington, DC: Office of Pollution Prevention and Toxics, Health and Environmental Review Division, Environmental Effects Branch.

**CESIUM**  
9/1993

**Organism:** Aquatic life (freshwater)  
**Duration:**  
**Endpoint:** Acute Value  
**Equation:**  $\text{Acute Value (mg/L)} = (0.019 \cdot \text{MW})/35.45$   
**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing chlorine.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

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**Organism:** Aquatic life (freshwater)  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (0.011 \cdot \text{MW})/35.45$   
**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing chlorine.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

**CHLORINE**  
9/1993

**Organism:** Aquatic life (marine)  
**Duration:**  
**Endpoint:** Acute Value  
**Equation:**  $\text{Acute Value (mg/L)} = (0.013 \cdot \text{MW})/35.45$   
**Application:** This equation may be used to estimate the toxicity of both inorganic and organic compounds containing chlorine.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

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**Organism:** Aquatic life (marine)  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (0.0075 \cdot \text{MW})/35.45$   
**Application:** This equation may be used to estimate the toxicity of organic and inorganic compounds containing chlorine.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

**CHLORINE**  
9/1993

**Organism:** Fish (freshwater)  
**Duration:** 96-hour  
**Endpoint:** LC50

**Equation:**  $LC50 \text{ (mg/L)} = (48.0 \cdot MW)/58.933$

**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing cobalt.

**Limitations:** None

**References:** United States Environmental Protection Agency (USEPA). 1991. Hazard Profiles for Selected Heavy Metals. Washington, DC: Office of Pollution Prevention and Toxics, Health and Environmental Review Division, Environmental Effects Branch.

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**Organism:** Daphnid  
**Duration:** 48-hour  
**Endpoint:** LC50

**Equation:**  $LC50 \text{ (mg/L)} = (1.30 \cdot MW)/58.933$

**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing cobalt.

**Limitations:** None

**References:** United States Environmental Protection Agency (USEPA). 1991. Hazard Profiles for Selected Heavy Metals. Washington, DC: Office of Pollution Prevention and Toxics, Health and Environmental Review Division, Environmental Effects Branch.

**COBALT**  
9/1993

**Organism:** Fish (freshwater)  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (0.0342 \cdot \text{MW})/58.933$   
**Application:** This equation may be used to estimate the toxicity of both inorganic and organic compounds containing cobalt.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1991. Hazard Profiles for Selected Heavy Metals. Washington, DC: Office of Pollution Prevention and Toxics, Health and Environmental Review Division, Environmental Effects Branch.

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**Organism:** Daphnid  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (0.012 \cdot \text{MW})/58.933$   
**Application:** This equation may be used to estimate the toxicity of organic and inorganic compounds containing cobalt.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1991. Hazard Profiles for Selected Heavy Metals. Washington, DC: Office of Pollution Prevention and Toxics, Health and Environmental Review Division, Environmental Effects Branch.

**COBALT**  
9/1993

**Organism:** Fish (marine)  
**Duration:** 96-hour  
**Endpoint:** LC50

**Equation:**  $LC50 \text{ (mg/L)} = (567.0 \cdot MW)/58.933$

**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing cobalt.

**Limitations:** None

**References:** United States Environmental Protection Agency (USEPA). 1991. Hazard Profiles for Selected Heavy Metals. Washington, DC: Office of Pollution Prevention and Toxics, Health and Environmental Review Division, Environmental Effects Branch.

**COBALT**  
9/1993

<b>Organism:</b>	Aquatic life (freshwater)
<b>Duration:</b>	
<b>Endpoint:</b>	Acute Value
<b>Equation:</b>	$\text{Acute Value (mg/L)} = (0.018 \cdot \text{MW}) / 63.546$
<b>Application:</b>	This equation may be used to estimate the toxicity of both organic and inorganic compounds containing copper.
<b>Limitations:</b>	<p>This equation is based on the hardness dependent Water Quality Criteria for copper. The criterion for a hardness of 100 mg/L as <math>\text{CaCO}_3</math> were used. For a solution with a hardness of 50 mg/L as <math>\text{CaCO}_3</math>, use the following equation:</p> $\text{Acute Value (mg/L)} = (0.0092 \cdot \text{MW}) / 63.546$ <p>For a solution with a hardness of 200 mg/L as <math>\text{CaCO}_3</math>, use the following equation:</p> $\text{Acute Value (mg/L)} = (0.034 \cdot \text{MW}) / 63.546$
<b>References:</b>	United States Environmental Protection Agency (USEPA). 1986 Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

**COPPER**  
9/1993

**Organism:** Aquatic life (freshwater)  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (0.012 \cdot \text{MW})/63.546$   
**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing copper.  
**Limitations:** This equation is based on the hardness dependent Water Quality Criteria for copper. The criterion for a hardness of 100 mg/L as  $\text{CaCO}_3$  was used. For solutions with a hardness of 50 mg/L as  $\text{CaCO}_3$ , use the following equation:  
$$\text{ChV (mg/L)} = (0.0065 \cdot \text{MW})/63.546$$
  
For solutions with a hardness of 200 mg/L as  $\text{CaCO}_3$ , use the following equation:  
$$\text{ChV (mg/L)} = (0.0021 \cdot \text{MW})/63.546$$
  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

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**Organism:** Aquatic life (marine)  
**Duration:**  
**Endpoint:** Acute Value  
**Equation:**  $\text{Acute Value (mg/L)} = (0.0029 \cdot \text{MW})/63.546$   
**Application:** This equation may be used to estimate the toxicity of both inorganic and organic compounds containing copper.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

COPPER  
9/1993

<b>Organism:</b>	Aquatic life (freshwater)
<b>Duration:</b>	
<b>Endpoint:</b>	Acute Value
<b>Equation:</b>	$\text{Acute Value (mg/L)} = (1.700 \cdot \text{MW})/51.996$
<b>Application:</b>	This equation may be used to estimate the toxicity of both organic and inorganic compounds containing chromium(III).
<b>Limitations:</b>	<p>This equation is based on the hardness dependent Water Quality Criteria for chromium(III). The criterion for a hardness of 100 mg/L as <math>\text{CaCO}_3</math> was used. For solutions with a hardness of 50 mg/L as <math>\text{CaCO}_3</math>, use the following equation:</p> $\text{Acute Value (mg/L)} = (0.980 \cdot \text{MW})/51.996$ <p>For solutions with a hardness of 200 mg/L as <math>\text{CaCO}_3</math>, use the following equation:</p> $\text{Acute Value (mg/L)} = (3.100 \cdot \text{MW})/51.996$
<b>References:</b>	United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

**CHROMIUM(III)**  
9/1993

**CHROMIUM(III)**  
9/1993

**Organism:** Aquatic life (freshwater)

**Duration:**

**Endpoint:** Chronic Value (ChV)

**Equation:**  $\text{ChV (mg/L)} = (0.210 \cdot \text{MW})/51.996$

**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing chromium(III).

**Limitations:** This equation is based on the hardness dependent Water Quality Criteria for chromium(III). The criterion for a hardness of 100 mg/L as  $\text{CaCO}_3$  was used. For a solution with a hardness of 50 mg/L as  $\text{CaCO}_3$ , use the following equation:

$$\text{ChV (mg/L)} = (0.120 \cdot \text{MW})/51.996$$

For a solution with a hardness of 200 mg/L as  $\text{CaCO}_3$ , use the following equation:

$$\text{ChV (mg/L)} = (0.120 \cdot \text{MW})/51.996$$

**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

**CHROMIUM(III)**  
9/1993

**CHROMIUM(III)**  
9/1993

**Organism:** Eastern Oyster embryos (marine)  
**Duration:** Acute  
**Endpoint:** EC50

**Equation:**  $EC50 \text{ (mg/L)} = (10.3 \cdot MW)/51.996$

**Application:** This equation may be used to estimate the toxicity of both inorganic and organic compounds containing chromium(III).

**Limitations:** None

**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

**CHROMIUM(III)**  
9/1993

**CHROMIUM(VI)**  
9/1993

**Organism:** Aquatic life (freshwater)  
**Duration:**  
**Endpoint:** Acute Value  
**Equation:**  $\text{Acute Value (mg/L)} = (0.016 \cdot \text{MW})/51.996$   
**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing chromium(VI).  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division EPA 440/5-86-001.

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**Organism:** Aquatic life (freshwater)  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (0.011 \cdot \text{MW})/51.996$   
**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing chromium(VI).  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

**CHROMIUM(VI)**  
9/1993

**Organism:** Aquatic life (marine)  
**Duration:**  
**Endpoint:** Acute Value  
**Equation:** Acute Value (mg/L) =  $(1.100 \cdot MW)/51.996$   
**Application:** This equation may be used to estimate the toxicity of both inorganic and organic compounds containing chromium(VI).  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001

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**Organism:** Aquatic life (marine)  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:** ChV (mg/L) =  $(0.050 \cdot MW)/51.996$   
**Application:** This equation may be used to estimate the toxicity of organic and inorganic compounds containing chromium(VI).  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

**CHROMIUM(VI)**  
9/1993

## GERMANIUM

9/1993

<b>Organism:</b>	Fish (freshwater)
<b>Duration:</b>	
<b>Endpoint:</b>	Chronic Value (ChV)
<b>Equation:</b>	$\text{ChV (mg/L)} = (0.003 \cdot \text{MW})/72.6$
<b>Application:</b>	This equation may be used to estimate the toxicity of both organic and inorganic compounds containing germanium.
<b>Limitations:</b>	None
<b>References:</b>	United States Environmental Protection Agency (USEPA). 1991. Hazard Profiles for Selected Heavy Metals. Washington, DC: Office of Pollution Prevention and Toxics, Health and Environmental Review Division, Environmental Effects Branch.

**GERMANIUM**  
9/1993

**Organism:** Daphnid  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (0.180 \cdot \text{MW})/196.967$   
**Application:** This equation may be used to estimate the toxicity of organic and inorganic compounds containing gold.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1991. Hazard Profiles for Selected Heavy Metals. Washington, DC: Office of Pollution Prevention and Toxics, Health and Environmental Review Division, Environmental Effects Branch.

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**Organism:** Green Algae  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (0.125 \cdot \text{MW})/196.967$   
**Application:** This equation may be used to estimate the toxicity of both inorganic and organic compounds containing gold.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1991. Hazard Profiles for Selected Heavy Metals. Washington, DC: Office of Pollution Prevention and Toxics, Health and Environmental Review Division, Environmental Effects Branch

**GOLD**  
9/1993

**Organism:** Aquatic life (freshwater)  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (1.0 \cdot \text{MW})/55.847$   
**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing iron.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

IRON  
9/1993

**Organism:** Daphnid  
**Duration:** 48-hour  
**Endpoint:** LC50

**Equation:**  $LC50 \text{ (mg/L)} = (160.0 \cdot MW)/138.906$

**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing lanthanum.

**Limitations:** None

**References:** United States Environmental Protection Agency (USEPA). 1991. Hazard Profiles for Selected Heavy Metals. Washington, DC: Office of Pollution Prevention and Toxics, Health and Environmental Review Division, Environmental Effects Branch.

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**Organism:** Fish (freshwater)  
**Duration:**  
**Endpoint:** Chronic Value (ChV)

**Equation:**  $ChV \text{ (mg/L)} = (0.0008 \cdot MW)/138.906$

**Application:** This equation may be used to estimate the toxicity of both inorganic and organic compounds containing lanthanum.

**Limitations:** None

**References:** United States Environmental Protection Agency (USEPA). 1991. Hazard Profiles for Selected Heavy Metals. Washington, DC: Office of Pollution Prevention and Toxics, Health and Environmental Review Division, Environmental Effects Branch.

**LANTHANUM**  
9/1993

**Organism:** Green Algae  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (6.4 \cdot \text{MW})/138.906$   
**Application:** This equation may be used to estimate the toxicity of organic and inorganic compounds containing lanthanum.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1991. Hazard Profiles for Selected Heavy Metals. Washington, DC: Office of Pollution Prevention and Toxics, Health and Environmental Review Division, Environmental Effects Branch.

LANTHANUM  
9/1993

**Organism:** Aquatic life (freshwater)  
**Duration:**  
**Endpoint:** Acute Value  
**Equation:**  $\text{Acute Value (mg/L)} = (0.083 \cdot \text{MW})/207.2$   
**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing lead.  
**Limitations:** This equation is based on the hardness dependent Water Quality Criteria for lead. The criterion for a hardness of 100 mg/L as  $\text{CaCO}_3$  was used  
**References:** United States Environmental Protection Agency (USEPA). 1992. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division.

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**Organism:** Aquatic life (freshwater)  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (0.0032 \cdot \text{MW})/207.2$   
**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing lead.  
**Limitations:** This equation is based on the hardness dependent Water Quality Criteria for lead. The criterion for a hardness of 100 mg/L as  $\text{CaCO}_3$  was used.  
**References:** United States Environmental Protection Agency (USEPA). 1992. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division.

**Organism:** Aquatic life (marine)  
**Duration:**  
**Endpoint:** Acute Value  
**Equation:**  $\text{Acute Value (mg/L)} = (0.220 \cdot \text{MW})/207.2$   
**Application:** This equation may be used to estimate the toxicity of both inorganic and organic compounds containing lead.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1992. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division.

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**Organism:** Aquatic life (marine)  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (0.0085 \cdot \text{MW})/207.2$   
**Application:** This equation may be used to estimate the toxicity of organic and inorganic compounds containing lead.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1992. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division.

**LEAD**  
9/1993

**Organism:** Aquatic life (freshwater)  
**Duration:**  
**Endpoint:** Acute Value  
**Equation:**  $\text{Acute Value (mg/L)} = (0.0024 \cdot \text{MW})/200.59$   
**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing mercury.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

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**Organism:** Aquatic life (freshwater)  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (0.00012 \cdot \text{MW})/200.59$   
**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing mercury.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

**MERCURY**  
9/1993

**Organism:** Aquatic life (marine)  
**Duration:**  
**Endpoint:** Acute Value  
**Equation:**  $\text{Acute Value (mg/L)} = (0.0021 \cdot \text{MW})/200.59$   
**Application:** This equation may be used to estimate the toxicity of both inorganic and organic compounds containing mercury.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

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**Organism:** Aquatic life (marine)  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (0.000025 \cdot \text{MW})/200.59$   
**Application:** This equation may be used to estimate the toxicity of organic and inorganic compounds containing mercury.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

**MERCURY**  
9/1993

**Organism:** Fish (freshwater)  
**Duration:** 96-hour  
**Endpoint:** LC50

**Equation:**  $LC50 \text{ (mg/L)} = (553.0 \cdot MW)/95.94$

**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing molybdenum.

**Limitations:** None

**References:** United States Environmental Protection Agency (USEPA). 1991. Hazard Profiles for Selected Heavy Metals. Washington, DC: Office of Pollution Prevention and Toxics, Health and Environmental Review Division, Environmental Effects Branch.

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**Organism:** Fish (freshwater)  
**Duration:**  
**Endpoint:** Chronic Value (ChV)

**Equation:**  $ChV \text{ (mg/L)} = (0.0223 \cdot MW)/95.94$

**Application:** This equation may be used to estimate the toxicity of both inorganic and organic compounds containing molybdenum.

**Limitations:** None

**References:** United States Environmental Protection Agency (USEPA). 1991. Hazard Profiles for Selected Heavy Metals. Washington, DC: Office of Pollution Prevention and Toxics, Health and Environmental Review Division, Environmental Effects Branch.

**MOLYBDENUM**  
9/1993

<b>Organism:</b>	Aquatic life (freshwater)
<b>Duration:</b>	
<b>Endpoint:</b>	Acute Value
<b>Equation:</b>	$\text{Acute Value (mg/L)} = (1.400 \cdot \text{MW})/58.70$
<b>Application:</b>	This equation may be used to estimate the toxicity of both organic and inorganic compounds containing nickel.
<b>Limitations:</b>	<p>This equation is based on the hardness dependent Water Quality Criteria for nickel. The criterion for a hardness of 100 mg/L as <math>\text{CaCO}_3</math> was used. For a solution with a hardness of 50 mg/L as <math>\text{CaCO}_3</math>, use the following equation:</p> $\text{Acute Value (mg/L)} = (0.790 \cdot \text{MW})/58.70$ <p>For a solution with a hardness of 200 mg/L as <math>\text{CaCO}_3</math>, use the following equation:</p> $\text{Acute Value (mg/L)} = (2.500 \cdot \text{MW})/58.70$
<b>References:</b>	United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

NICKEL  
9/1993

<b>Organism:</b>	Aquatic life (freshwater)
<b>Duration:</b>	
<b>Endpoint:</b>	Chronic Value (ChV)
<b>Equation:</b>	$\text{ChV (mg/L)} = (0.160 \cdot \text{MW})/58.70$
<b>Application:</b>	This equation may be used to estimate the toxicity of both organic and inorganic compounds containing nickel.
<b>Limitations:</b>	<p>This equation is based on the hardness dependent Water Quality Criteria for nickel. The criterion for a hardness of 100 mg/L as <math>\text{CaCO}_3</math> was used. For solutions with a hardness of 50 mg/L as <math>\text{CaCO}_3</math>, use the following equation:</p> $\text{ChV (mg/L)} = (0.088 \cdot \text{MW})/58.70$ <p>For solutions with a hardness of 200 mg/L as <math>\text{CaCO}_3</math>, use the following equation:</p> $\text{ChV (mg/L)} = (0.280 \cdot \text{MW})/58.70$
<b>References:</b>	United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

NICKEL  
9/1993

**Organism:** Aquatic life (marine)  
**Duration:**  
**Endpoint:** Acute Value  
**Equation:**  $\text{Acute Value (mg/L)} = (0.075 \cdot \text{MW})/58.70$   
**Application:** This equation may be used to estimate the toxicity of both inorganic and organic compounds containing nickel.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

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**Organism:** Aquatic life (marine)  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (0.0083 \cdot \text{MW})/58.70$   
**Application:** This equation may be used to estimate the toxicity of organic and inorganic compounds containing nickel.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

NICKEL  
9/1993

<b>Organism:</b>	Aquatic life (marine)
<b>Duration:</b>	
<b>Endpoint:</b>	Chronic Value (ChV)
<b>Equation:</b>	$\text{ChV (mg/L)} = (0.0001 \cdot \text{MW})/30.974$
<b>Application:</b>	This equation may be used to estimate the toxicity of organic and inorganic compounds containing phosphorus.
<b>Limitations:</b>	This equation is based on the Water Quality Criteria for yellow (elemental) phosphorus.
<b>References:</b>	United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

**PHOSPHORUS**  
9/1993

<b>Organism:</b>	Daphnid
<b>Duration:</b>	
<b>Endpoint:</b>	Chronic Value (ChV)
<b>Equation:</b>	$\text{ChV (mg/L)} = (0.082 \cdot \text{MW}) / 195.09$
<b>Application:</b>	This equation may be used to estimate the toxicity of both organic and inorganic compounds containing platinum.
<b>Limitations:</b>	None
<b>References:</b>	United States Environmental Protection Agency (USEPA). 1991. Hazard Profiles for Selected Heavy Metals. Washington, DC: Office of Pollution Prevention and Toxics, Health and Environmental Review Division, Environmental Effects Branch.

PLATINUM  
9/1993

**Organism:** Aquatic life (freshwater)  
**Duration:**  
**Endpoint:** Acute Value  
**Equation:**  $\text{Acute Value (mg/L)} = (0.020 \cdot \text{MW})/78.96$   
**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing selenium.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1992. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division.

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**Organism:** Aquatic life (freshwater)  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (0.005 \cdot \text{MW})/78.96$   
**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing selenium.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1992. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division.

**SELENIUM**  
9/1993

**Organism:** Aquatic life (marine)  
**Duration:**  
**Endpoint:** Acute Value  
**Equation:**  $\text{Acute Value (mg/L)} = (0.300 \cdot \text{MW})/78.96$   
**Application:** This equation may be used to estimate the toxicity of both inorganic and organic compounds containing selenium.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1992. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division.

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**Organism:** Aquatic life (marine)  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (0.071 \cdot \text{MW})/78.96$   
**Application:** This equation may be used to estimate the toxicity of organic and inorganic compounds containing selenium.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1992. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division.

**SELENIUM**  
9/1993

**Organism:** Aquatic life (freshwater)  
**Duration:**  
**Endpoint:** Acute Value  
**Equation:**  $\text{Acute Value (mg/L)} = (0.0041 \cdot \text{MW})/107.868$   
**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing silver.  
**Limitations:** This equation is based on the hardness dependent Water Quality Criteria for silver. The criterion for a hardness of 100 mg/L as  $\text{CaCO}_3$  was used.  
**References:** United States Environmental Protection Agency (USEPA). 1992. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division.

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**Organism:** Aquatic life (freshwater)  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (0.00012 \cdot \text{MW})/107.868$   
**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing silver.  
**Limitations:** This equation is based on the hardness dependent Water Quality Criteria for silver. The criterion for a hardness of 100 mg/L as  $\text{CaCO}_3$  was used.  
**References:** United States Environmental Protection Agency (USEPA). 1992. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division.

**SILVER**  
9/1993

**Organism:** Aquatic life (marine)  
**Duration:**  
**Endpoint:** Acute Value  
**Equation:**  $\text{Acute Value (mg/L)} = (0.0023 \cdot \text{MW})/107.868$   
**Application:** This equation may be used to estimate the toxicity of both inorganic and organic compounds containing silver.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1992. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division.

**SILVER**  
9/1993

**Organism:** Aquatic life (freshwater)  
**Duration:** Acute  
**Endpoint:** Lowest Observable Effect Concentration (LOEC)  
**Equation:**  $LOEC \text{ (mg/L)} = (1.4 \cdot MW)/204.37$   
**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing thallium.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

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**Organism:** Aquatic life (freshwater)  
**Duration:** Chronic  
**Endpoint:** Lowest Observable Effect Concentration (LOEC)  
**Equation:**  $LOEC \text{ (mg/L)} = (0.040 \cdot MW)/201.37$   
**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing thallium.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

**THALLIUM**  
9/1993

**Organism:** Aquatic life (marine)  
**Duration:** Acute  
**Endpoint:** Lowest Observed Effect Concentration (LOEC)  
**Equation:**  $LOEC \text{ (mg/L)} = (2.130 \cdot MW)/204.37$   
**Application:** This equation may be used to estimate the toxicity of both inorganic and organic compounds containing thallium.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

THALLIUM  
9/1993

**Organism:** Fish (freshwater)  
**Duration:** 96-hour  
**Endpoint:** LC50

**Equation:**  $LC50 \text{ (mg/L)} = (31.0 \cdot MW)/47.90$

**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing titanium.

**Limitations:** None

**References:** United States Environmental Protection Agency (USEPA). 1991. Hazard Profiles for Selected Heavy Metals. Washington, DC: Office of Pollution Prevention and Toxics, Health and Environmental Review Division, Environmental Effects Branch.

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**Organism:** Daphnid  
**Duration:** 48-hour  
**Endpoint:** EC50

**Equation:**  $EC50 \text{ (mg/L)} = (4.6 \cdot MW)/47.90$

**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing titanium.

**Limitations:** None

**References:** United States Environmental Protection Agency (USEPA). 1991. Hazard Profiles for Selected Heavy Metals. Washington, DC: Office of Pollution Prevention and Toxics, Health and Environmental Review Division, Environmental Effects Branch.

**TITANIUM**  
9/1993

**Organism:** Daphnid  
**Duration:** 48-hour  
**Endpoint:** EC50  
**Equation:**  $EC50 \text{ (mg/L)} = (350.0 \cdot MW)/183.85$   
**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing tungsten.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1991. Hazard Profiles for Selected Heavy Metals. Washington, DC: Office of Pollution Prevention and Toxics, Health and Environmental Review Division, Environmental Effects Branch.

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**Organism:** Fish (freshwater)  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $ChV \text{ (mg/L)} = (15.61 \cdot MW)/183.85$   
**Application:** This equation may be used to estimate the toxicity of both inorganic and organic compounds containing tungsten.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1991. Hazard Profiles for Selected Heavy Metals. Washington, DC: Office of Pollution Prevention and Toxics, Health and Environmental Review Division, Environmental Effects Branch.

**TUNGSTEN**  
9/1993

<b>Organism:</b>	Fish	
<b>Duration:</b>	96-hour	
<b>Endpoint:</b>	LC50	
<b>Equation:</b>	vanadium salts (n=4)	$LC50 \text{ (mg/L)} = (3.9 \cdot MW)/50.942$
	vanadium oxides (n=13)	$LC50 \text{ (mg/L)} = (3.3 \cdot MW)/50.942$
	vanadium complexed with organic acids (n=1)	$LC50 \text{ (mg/L)} = (26.0 \cdot MW)/50.942$
	vanadium sulfate (n=4)	$LC50 \text{ (mg/L)} = (3.9 \cdot MW)/50.942$
	sodium vanadate ( $VO_3$ ) (n=4)	$LC50 \text{ (mg/L)} = (2.5 \cdot MW)/50.942$
	vanadium pentoxide (n=7)	$LC50 \text{ (mg/L)} = (6.1 \cdot MW)/50.942$
	ammonium vanadate ( $VO_3$ ) (n=2)	$LC50 \text{ (mg/L)} = (2.4 \cdot MW)/50.942$
<b>Application:</b>	The appropriate equation may be used to estimate the toxicity of both organic and inorganic compounds containing vanadium.	
<b>Limitations:</b>	This category is not applicable to vanadium-complexed dyes. The toxicity of vanadium salts and weak organic acid complexes is expected to be related to their water solubility. Vanadium is more toxic in soft water than hard water but the relationship is not well defined. These equations are based on toxicity data measured in moderately hard water (150.0 mg/L as $CaCO_3$ ). Strong ion pairs with molecular weights greater than 1000 are not expected to be absorbed by aquatic organisms even if they are water soluble. The boundaries for organovanadium compounds are undefined, but the molecular weight boundary is expected to be less than 1000.	
<b>References:</b>	Nabholz JV. 1993. Vanadium compounds (Unpublished document). Washington, DC: Environmental Effects Branch, Health and Environmental Review Division, Office of Pollution Prevention and Toxics, United States Environmental Protection Agency.	

**VANADIUM**  
9/1993

**Organism:** Daphnid  
**Duration:** 48-hour  
**Endpoint:** LC50

**Equation:** sodium vanadate ( $\text{VO}_3$ ) (~~LC50~~) (mg/L) =  $(4.1 \cdot \text{MW})/50.942$

**Application:** This equation may be used to estimate the toxicity of both organic and inorganic compounds containing vanadium.

**Limitations:** This category is not applicable to vanadium-complexed dyes. The toxicity of vanadium salts and weak organic acid complexes is expected to be related to their water solubility. Vanadium is more toxic in soft water than hard water but the relationship is not well defined. These equations are based on toxicity data measured in moderately hard water (150.0 mg/L as  $\text{CaCO}_3$ ). Strong ion pairs with molecular weights greater than 1000 are not expected to be absorbed by aquatic organisms even if they are water soluble. The boundaries for organovanadium compounds are undefined, but the molecular weight boundary is expected to be less than 1000.

**References:** Nabholz JV. 1993. Vanadium compounds (Unpublished document). Washington, DC: Environmental Effects Branch, Health and Environmental Review Division, Office of Pollution Prevention and Toxics, United States Environmental Protection Agency.

**VANADIUM**  
9/1993

<b>Organism:</b>	Fish
<b>Duration:</b>	
<b>Endpoint:</b>	Chronic Value (ChV)
<b>Equation:</b>	vanadium pentoxide ( $n=3$ ) $ChV \text{ (mg/L)} = (0.670 \cdot MW)/50.942$
<b>Application:</b>	This equation may be used to estimate the toxicity of both organic and inorganic compounds containing vanadium.
<b>Limitations:</b>	This category is not applicable to vanadium-complexed dyes. The toxicity of vanadium salts and weak organic acid complexes is expected to be related to their water solubility. Vanadium is more toxic in soft water than hard water but the relationship is not well defined. These equations are based on toxicity data measured in moderately hard water (150.0 mg/L as $CaCO_3$ ). Strong ion pairs with molecular weights greater than 1000 are not expected to be absorbed by aquatic organisms even if they are water soluble. The boundaries for organovanadium compounds are undefined, but the molecular weight boundary is expected to be less than 1000.
<b>References:</b>	Nabholz JV. 1993. Vanadium compounds (Unpublished document). Washington, DC: Environmental Effects Branch, Health and Environmental Review Division, Office of Pollution Prevention and Toxics, United States Environmental Protection Agency.

**VANADIUM**  
9/1993

<b>Organism:</b>	Green Algae
<b>Duration:</b>	
<b>Endpoint:</b>	No Observable Effect Concentration (NOEC) (increased growth)
<b>Equation:</b>	$\text{ChV (mg/L)} = (0.100 \cdot \text{MW})/50.942$
<b>Application:</b>	This equation may be used to estimate the toxicity of both organic and inorganic compounds containing vanadium. This equation is based on toxicity data for vanadium sulfate and sodium vanadate.
<b>Limitations:</b>	This category is not applicable to vanadium-complexed dyes. The toxicity of vanadium salts and weak organic acid complexes is expected to be related to their water solubility. Vanadium is more toxic in soft water than hard water but the relationship is not well defined. These equations are based on toxicity data measured in moderately hard water (150.0 mg/L as $\text{CaCO}_3$ ). Strong ion pairs with molecular weights greater than 1000 are not expected to be absorbed by aquatic organisms even if they are water soluble. The boundaries for organovanadium compounds are undefined, but the molecular weight boundary is expected to be less than 1000.
<b>References:</b>	Nabholz JV. 1993. Vanadium compounds (Unpublished document). Washington, DC: Environmental Effects Branch, Health and Environmental Review Division, Office of Pollution Prevention and Toxics, United States Environmental Protection Agency.

**VANADIUM**  
9/1993

<b>Organism:</b>	Aquatic life (freshwater)
<b>Duration:</b>	
<b>Endpoint:</b>	Acute Value
<b>Equation:</b>	$\text{Acute Value (mg/L)} = (0.120 \cdot \text{MW})/65.38$
<b>Application:</b>	This equation may be used to estimate the toxicity of both organic and inorganic compounds containing zinc.
<b>Limitations:</b>	<p>This equation is based on the hardness dependent Water Quality Criteria for zinc. The criterion for a hardness of 100 mg/L as <math>\text{CaCO}_3</math> was used. For a solution with a hardness of 50 mg/L as <math>\text{CaCO}_3</math>, use the following equation:</p> $\text{Acute Value (mg/L)} = (0.065 \cdot \text{MW})/65.38$ <p>For a solution with a hardness of 200 mg/L as <math>\text{CaCO}_3</math>, use the following equation:</p> $\text{Acute Value (mg/L)} = (0.210 \cdot \text{MW})/65.38$
<b>References:</b>	United States Environmental Protection Agency (USEPA), 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

ZINC  
9/1993

<b>Organism:</b>	Aquatic life (freshwater)
<b>Duration:</b>	
<b>Endpoint:</b>	Chronic Value (ChV)
<b>Equation:</b>	$\text{ChV (mg/L)} = (0.110 \cdot \text{MW})/65.38$
<b>Application:</b>	This equation may be used to estimate the toxicity of both organic and inorganic compounds containing zinc.
<b>Limitations:</b>	<p>This equation is based on the hardness dependent Water Quality Criteria for zinc. The criterion for a hardness of 100 mg/L as <math>\text{CaCO}_3</math> was used. For solutions with a hardness of 50 mg/L as <math>\text{CaCO}_3</math>, use the following equation:</p> $\text{ChV (mg/L)} = (0.059 \cdot \text{MW})/65.38$ <p>For solutions with a hardness of 200 mg/L as <math>\text{CaCO}_3</math>, use the following equation:</p> $\text{ChV (mg/L)} = (0.190 \cdot \text{MW})/65.38$
<b>References:</b>	United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

ZINC  
9/1993

**Organism:** Aquatic life (marine)  
**Duration:**  
**Endpoint:** Acute Value  
**Equation:**  $\text{Acute Value (mg/L)} = (0.095 \cdot \text{MW})/65.38$   
**Application:** This equation may be used to estimate the toxicity of both inorganic and organic compounds containing zinc.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

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**Organism:** Aquatic life (marine)  
**Duration:**  
**Endpoint:** Chronic Value (ChV)  
**Equation:**  $\text{ChV (mg/L)} = (0.086 \cdot \text{MW})/65.38$   
**Application:** This equation may be used to estimate the toxicity of organic and inorganic compounds containing zinc.  
**Limitations:** None  
**References:** United States Environmental Protection Agency (USEPA). 1986. Quality Criteria for Water. Washington, DC: Office of Water, Criteria and Standards Division. EPA 440/5-86-001.

**ZINC**  
9/1993

<b>Organism:</b>	Fish
<b>Duration:</b>	96-hour
<b>Endpoint:</b>	LC50
<b>Equation:</b>	$LC50 \text{ (mg/L)} = (58.0 \cdot MW)/91.22$
<b>Application:</b>	This equation may be used to estimate the toxicity of both organic and inorganic compounds containing zirconium, including inorganic salts of zirconium, complexes between zirconium and organic acids, and organozirconium compounds, i.e., zirconium covalently-bonded with carbon.
<b>Limitations:</b>	<p>This equation is not applicable to dyes complexed with zirconium. The equation is based on available toxicity data for solution of moderate hardness (i.e., 150 mg/L as <math>CaCO_3</math>). Zirconium is more toxic in soft water than in hard water. Acute toxicity to fish has been shown to increase 13 times as hardness decreases from 400.0 to 20 mg/L.</p> <p>Compounds with molecular weights greater than 1000 are not expected to be absorbed by aquatic organisms even if they are water soluble.</p>
<b>References:</b>	Nabholz JV. 1993. Zirconium compounds (Unpublished internal document). Washington, DC: Environmental Effects Branch, Health and Environmental Review Division, Office of Pollution Prevention and Toxics, United States Environmental Protection Agency.

**ZIRCONIUM**  
9/1993

<b>Organism:</b>	Green Algae
<b>Duration:</b>	96-hour
<b>Endpoint:</b>	EC50
<b>Equation:</b>	$EC50 \text{ (mg/L)} = (2.6 \cdot MW)/91.22$
<b>Application:</b>	This equation is not applicable to dyes complexed with zirconium. The equation is based on available toxicity data for solution of moderate hardness (i.e., 150 mg/L as $CaCO_3$ ). Zirconium is more toxic in soft water than in hard water.
<b>Limitations:</b>	<p>This equation is not applicable to dyes complexed with zirconium. The equation is based on available toxicity data for solution of moderate hardness (i.e., 150 mg/L as <math>CaCO_3</math>). Zirconium is more toxic in soft water than in hard water. Acute toxicity to fish has been shown to increase 13 times as hardness decreases from 400.0 to 20 mg/L.</p> <p>Compounds with molecular weights greater than 1000 are not expected to be absorbed by aquatic organisms even if they are water soluble.</p>
<b>References:</b>	Nabholz JV. 1993. Zirconium compounds (Unpublished internal document). Washington, DC: Environmental Effects Branch, Health and Environmental Review Division, Office of Pollution Prevention and Toxics, United States Environmental Protection Agency.

**ZIRCONIUM**  
9/1993

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