# A Guide to The Globally Harmonized System of Classification and Labeling of Chemicals (GHS)



# Acronyms/Abbreviations

The following list presents some acronyms and abbreviations used in this document. The **6.0** - **Glossary** contains a more complete list.

ANSI: American National Standards Institute **APEC:** Asia-Pacific Economic Cooperation **ASTM:** American Society of Testing and Materials CA: Competent Authority CAS: Chemical Abstract Service **CBI:** Confidential Business Information **CFR:** Code of Federal Regulations CG/HCCS: Coordinating Group for the Harmonization of Chemical Classification Systems **CPSC:** Consumer Product Safety Commission **DOT:** Department of Transportation EINECS: European Inventory of Existing Commercial Chemical Substances **EPA:** Environmental Protection Agency EU: European Union FIFRA: Federal Insecticide, Fungicide and Rodenticide Act GHS: Globally Harmonized System of Classification and Labelling of Chemicals HCS: Hazard Communication Standard IARC: International Agency for the Research on Cancer **IFCS:** International Forum on Chemical Safety ILO: International Labor Organization IOMC: Inter-organization Program on the Sound Management of Chemicals ISO: International Standards Organization **IUPAC:** International Union of Pure and Applied Chemistry  $LD_{50}$ : Lethal dose 50 mg/kg: Milligram per kilogram MSDS: Material Safety Data Sheet NAFTA: North American Free Trade Agreement OSHA: Occupational Safety and Health Administration OECD: The Organization for Economic Cooperation and Development QSARs: Quantitative Structure-Activity Relationships

SDS: Safety Data Sheet SME: Small and medium sized enterprises TFHCL: Task Force on the Harmonization of Classification and Labeling TSCA: Toxic Substances Control Act UN: United Nations UNCED: United Nations Conference on Environment and Development UNCETDG: United Nations Committee of Experts on the Transport of Dangerous Goods UNCETDG/GHS: United Nations Committee of Experts on the Transport of Dangerous Goods and on the Globally Harmonized System of Classification and Labelling of Chemicals UNITAR: United Nations Institute for Training and Research WG: work group WHMIS: Workplace Hazardous Materials Information System WSSD: World Summit on Sustainable Development

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# 1.0 BACKGROUND

The purpose of this document is to describe the United Nations Globally Harmonized System of Classification and Labeling of Chemicals (GHS), why it was developed, and how it relates to the sound management of chemicals. The full official text of the system is available on the web at: <a href="http://www.unece.org/trans/danger/publi/ghs/ghs\_rev00/00files\_e.html">http://www.unece.org/trans/danger/publi/ghs/ghs\_rev00/00files\_e.html</a>

#### 1.1 What is the GHS?

The GHS is an acronym for *The Globally Harmonized System of Classification and Labeling of Chemicals*. The GHS is a system for standardizing and harmonizing the classification and labeling of chemicals. It is a logical and comprehensive approach to:

- Defining health, physical and environmental hazards of chemicals;
- Creating classification processes that use available data on chemicals for comparison with the defined hazard criteria; and
- Communicating hazard information, as well as protective measures, on labels and Safety Data Sheets (SDS).

Many countries already have regulatory systems in place for these types of requirements. These systems may be similar in content and approach, but their differences are significant enough to require multiple classifications, labels and safety data sheets for the same product when marketed in different countries, or even in the same country when parts of the life cycle are covered by different regulatory authorities. This leads to inconsistent protection for those potentially exposed to the chemicals, as well as creating extensive regulatory burdens on companies producing chemicals For example, in the United.States (U.S.) there are requirements for classification and labeling of chemicals for the Consumer Product Safety Commission, the Department of Transportation, the Environmental Protection Agency, and the Occupational Safety and Health Administration.

The GHS itself is not a regulation or a standard. The GHS Document (referred to as "The Purple Book", shown in Figure 1.1) establishes agreed hazard classification and communication provisions with explanatory information

Figure 1.1 GHS Document ("Purple Book")



on how to apply the system. The elements in the GHS supply a mechanism to meet the basic requirement of any hazard communication system, which is to decide if the chemical product produced and/or supplied is hazardous and to prepare a label and/or Safety Data Sheet as appropriate. Regulatory authorities in countries adopting the GHS will thus take the agreed criteria and provisions, and implement them through their own regulatory process and procedures rather than simply incorporating the text of the GHS into their national requirements.

The GHS Document thus provides countries with the regulatory building blocks to develop or modify existing national programs that address classification of hazards and transmittal of information about those hazards and associated protective measures. This helps to ensure the safe use of chemicals as they move through the product life cycle from "cradle to grave."

#### 1.2 Why was the GHS developed?

The production and use of chemicals is fundamental to all economies. The global chemical business is more than a \$1.7 trillion per year enterprise. In the U.S., chemicals are more than a \$450 billion business and exports are greater than \$80 billion per year.

Chemicals directly or indirectly affect our lives and are essential to our food, our health, and our lifestyle. The widespread use of chemicals has resulted in the development of sector-specific regulations (transport, production, workplace, agriculture, trade, and consumer products). Having readily available information on the hazardous properties of chemicals, and recommended control measures, allows the production, transport, use and disposal of chemicals to be managed safely. Thus, human health and the environment are protected.

The sound management of chemicals should include systems through which chemical hazards are identified and communicated to all who are potentially exposed. These groups include workers, consumers, emergency responders and the public. It is important to know what chemicals are present and/or used, their hazards to human health and the environment, and the means to control them. A number of classification and labeling systems, each addressing specific use patterns and groups of chemicals, exist at the national, regional and international levels. The existing hazard classification and labeling systems address potential exposure to chemicals in all the types of use settings listed above.

While the existing laws and regulations are similar, they are different enough to require multiple labels for the same product both within the U.S. and in international trade and to require multiple safety data sheets for the same product in international trade. Several U.S. regulatory agencies and various countries have different requirements for hazard definitions as well as for information to be included on labels or material safety data sheets.

				Α	cute oral toxi	icity LD <sub>50</sub>	(mg/kg)		
Organization/Country Regulation or Standard				Hazard				Low	
	0			< 50 -			< 500		< 5000
ANSI/US/Z129.1		< 50 Highly Taxir		> 50 < 500 Toxic			> 500 < 2,000 Harmful		
OSHAUS/HCS		< 50 Highly Taxia			> 50 < 500 Toxic				
EPA/US/FIFRA	Ta	0≤50 xicity Categor	уI	> 50 < 500 Toxicity Category II			Tax	- 500 ≤ 5,000 ininy Canegory III	> 5,000 Taxinity Category IV
CPS C/US/FHSA		< 50 Highly Taxia		> 50 < 5,000 Taxic					
GHS	≤5	> 5	i≤ <i>5</i> 0	> ;	50 ≤300	> <b>300</b> ≤ 2,0	000	> 2000 ≤ 5000	
DOI/US	< 5 Packaing Chroup I	Pa	5< 30 daing oup II	>	50 < 200 (solid) – Pack 50 < 500 (liquid) – Chrou	ang p⊞		- <b>I</b>	
NEPAUS	≤5 Hanard Category4	> 1	5≤30 Cadegory3	> 50 ≤ 500 Hazard Category 2			> 500 ≤ 2,000 Hazard Category 1	н	> 2,000 àzard Category 0
NPCAUS/HMIS	≤1 Toxicity Rating4	> 1 Toxicity	l≤90 yRating3	> 50 ≤ 500 Toxic ity Rating 2			Т	> 500 ≤ 5,000 orainity Rating 1	> 5,000 Toxicity Rating 0
EU	< 25 Very Toxic		> 25 < 200 Toxic			> 200 < 2,0 Hammful	00		
WHMIS/Carada	Since 100  Since 100  Statistical State 100  State 100  State 100		wisian L	> 50 ≤ 500 Tacia WHMIS Class D. Division 1, Subdivision B					
Australia/NOHS C	< 25 Very Toxic		> 25 < 200 Toxic			> 200 < 2,0 Hammful			
Mexico	< 1 Extremely Tox	ř.	>20<50 HighlyToxic	>50 < 500 Moderately	Taxic		>500 < 5000 Mildly Taxic	•	
Mahysia	<25 Very Toxic				200 to 500 Harmful				
Japan	< 30 Poisanous					300 to 3000 Powerful			
Karea	< 25 Very Toxic		>50 < Tax		>200<2000 Hammful				

Figure 1.2

The numerical values on the hazard index scale in the table are not to scale.

For example, a product may be considered flammable or toxic by one agency or country, but not by another.

We can see by comparing a few hazards how complex it is to comply with all domestic and global regulations. Acute oral toxicity  $(LD_{50})$  is a good example (Figure 1.2). Although most existing systems cover acute toxicity, we can see in the figure that what is considered hazardous varies considerably. These differences allow the same product to be hazardous in one country/system and not in another. At the very least, the same product has different labels and SDSs.

Figure 1.3



# FLAMMABILITY

The numerical values on the hazard index scale in the table are not to scale.

Flammable liquid is another hazard that is covered by most existing systems. As shown in Figure 1.3, the coverage varies between existing systems within the U.S. and globally. This means that the same product can be non-hazardous or hazardous with different labels/SDSs. In Section 4,

Figures 4.1 through 4.7 show the diverse domestic and international labels for a fictitious product (ToxiFlam) which has both oral toxicity and flammability hazards.

These differences in hazards and SDS/labels impact both protection and trade. In the area of protection, users may see different label warnings or safety data sheet information for the same chemical. In the area of trade, the need to comply with multiple regulations regarding hazard classification and labeling is costly and time-consuming. Some multinational companies have estimated that there are over 100 diverse hazard communication regulations for their products globally. For small and medium size enterprises (SMEs) regulatory compliance is complex and costly, and it can act as a barrier to international trade in chemicals.

# **1.3 What was the International Mandate?**

The single most important force that drove the creation of the GHS was the international mandate (Figure 1.4) adopted in the 1992 United Nations Conference on Environment and Development (UNCED), often called the "Earth Summit". The harmonization of classification and labeling of chemicals was one of six program areas that were endorsed by the United Nations General Assembly to Figure 1.4 International Mandate from UNCED Agenda 21, Chapter 19

"A globally harmonized hazard classification and compatible labelling system, including material safety data sheets and easily understandable symbols, should be available, if feasible, by the year 2000."

strengthen international efforts concerning the environmentally sound management of chemicals. It was recognized that an internationally harmonized approach to classification and labeling would provide the foundation for all countries to develop comprehensive national programs to ensure the safe use of chemicals.

# 1.4 How was the GHS developed?

In conjunction with its Convention and Recommendation on Safety in the Use of Chemicals at Work, the International Labor Organization (ILO) studied the tasks required to achieve harmonization. The ILO concluded that there were four major existing systems that needed to be harmonized to achieve a global approach.

No international organization covers all aspects of chemical classification and labeling. A broad scope and extensive expertise and resources were required to develop a system. In order to proceed, several decisions were needed:

(a) what systems would be considered "major" and thus the basis for harmonization, and (b) how could the work be divided to get the best expertise for different aspects. Four existing systems



(Figure #1.5) were deemed to be major and the primary basis for the GHS. While not considered major, requirements of other systems were examined as appropriate, and taken into account as proposals were developed.

A Coordinating Group for the Harmonization of Chemical Classification Systems (CG/HCCS) was created under the Inter-organization Program for the Sound Management of Chemicals (IOMC) and they were charged with coordinating and managing development of the system.

The GC/HCCS worked on a consensus basis and included representatives from major stakeholders, including national governments, industry and workers. They created a set of guiding principles (Figure 1.6). The scope and guiding principles created a common framework for the organizations that were charged with developing the different elements of the system.



- All types of chemicals will be covered;
- All systems will have to be changed;
- Involvement of all stakeholders should be ensured;
- Comprehensibility must be addressed.

In order to get the best expertise and

resources, the work was divided among three technical focal points. Figure 1.7 shows how the work was assigned to the three technical focal points and the overall responsibilities of the Coordinating Group itself. The UN Committee of Experts on Transport of Dangerous Goods was selected as the lead for work on physical hazards, in cooperation with the ILO. Based on their work in the testing guidelines and other chemical issues, the Organization for Economic Cooperation and Development (OECD) was selected for health/environmental hazards and mixtures. ILO has a long history in MSDS/labels, and was selected to be the lead in hazard communication. The OECD and ILO groups also included representatives from governments, industry and workers.





#### 1.5 How will the GHS be maintained and updated?

In October 1999, the United Nations Economic and Social Council decided (resolution 1999/65) to enlarge the mandate of the Committee of Experts on the Transport of Dangerous Goods by reconfiguring it into a Committee of Experts on the Transport of Dangerous Goods and on the Globally Harmonized System of Classification and Labeling of Chemicals (UNCETDG/GHS). At the same time, a new Sub-Committee of Experts on the Globally Harmonized System of Classification and Labeling of chemicals (UNCETDG/GHS).

When the IOMC completed developing the GHS, the system was presented to the UN GHS Sub-Committee, which formally adopted the system at its first session in December 2002. It was subsequently endorsed by the UNCETDG/GHS. The UN Economic and Social Council endorsed the GHS in July 2003.

The Sub-Committee of Experts on the Globally Harmonized System of Classification will:

- Act as custodian of the system, managing and giving direction to the harmonization process,
- Keep the system up-to-date, as necessary, considering the need to introduce changes or updates to ensure its continued relevance,
- Promote understanding and use of the system and encourage feedback,
- Make the system available for worldwide use,
- Make guidance available on the application of the system, and on the interpretation and use of technical criteria to support consistency of application,
- Prepare work programs and submit recommendations to the UNCETDG/GHS.

#### 1.6 When will the GHS be implemented?

There is no international implementation schedule for the GHS. It is likely that different national systems/sectors will require different timeframes for GHS implementation. Existing systems will need to consider phase-in strategies for transition from their current requirements to the new GHS requirements.

Several international bodies have proposed implementation goals. The World Summit on Sustainable Development (WSSD) and the Intergovernmental Forum for Chemical Safety (IFCS) have encouraged countries to implement the new GHS as soon as possible with a view to having the system fully operational by 2008. The Ministers of the Asia-Pacific Economic Cooperation (APEC) have also said that as many APEC economies as possible should implement, on a voluntary basis, the GHS by 2006. Under the North American Free Trade Agreement (NAFTA), the Tri-national Occupational Safety and Health Group and the NAFTA Pesticides Technical Working Group are discussing the GHS.

Some of the major existing systems have begun discussions about GHS implementation and situational analyses comparing existing requirements to GHS requirements. Some countries are considering harmonization to the greatest extent possible between their national sectors.

#### **1.7** What are the benefits?

The basic goal of hazard communication is to ensure that employers, employees and the public are provided with adequate, practical, reliable and comprehensible information on the hazards of chemicals, so that they can take effective preventive and protective measure for their health and safety. Thus, implementation of effective hazard communication provides benefits for **governments**, **companies**, **workers**, and **members of the public**.

The GHS has maximum value if it is accepted in all major regulatory systems for chemical hazard communication. The diversity of hazard definitions is shown in Figures 1.2 and 1.3. The array of domestic and global labels for one product is shown in Figures 4.1 to 4.7. In the USA implementation of the GHS would harmonize hazard definitions and label information among U.S. regulatory agencies (CPSC, DOT, EPA, OSHA, etc.). If the GHS is implemented globally, consistent information will be communicated on labels and SDSs.

It is anticipated that application of the GHS will:

- Enhance the protection of human health and the environment by providing an internationally comprehensible system,
- Provide a recognized framework to develop regulations for those countries without existing systems,
- Facilitate international trade in chemicals whose hazards have been identified on an international basis,
- Reduce the need for testing and evaluation against multiple classification systems.

The tangible benefits to **governments** are:

- Fewer chemical accidents and incidents,
- Lower health care costs,
- Improved protection of workers and the public from chemical hazards,
- Avoiding duplication of effort in creating national systems,
- Reduction in the costs of enforcement,
- Improved reputation on chemical issues, both domestically and internationally.

Benefits to **companies** include:

- A safer work environment and improved relations with employees,
- An increase in efficiency and reduced costs from compliance with hazard communication regulations,
- Application of expert systems resulting in maximizing expert resources and minimizing labor and costs,
- Facilitation of electronic transmission systems with international scope,
- Expanded use of training programs on health and safety,
- Reduced costs due to fewer accidents and illnesses,
- Improved corporate image and credibility.

#### Benefits to workers and members of the public include:

- Improved safety for workers and others through consistent and simplified communications on chemical hazards and practices to follow for safe handling and use,
- Greater awareness of hazards, resulting in safer use of chemicals in the workplace and in the home.

# 2.0 HOW IS THE GHS TO BE APPLIED?

The GHS Classification and Communication elements are the foundation of programs to ensure the safe use of chemicals, as shown in Figure 2.1. The first two steps in any program to ensure the safe use of chemicals are to identify intrinsic hazard(s) (i.e., classification) and then to communicate that information. The design of the GHS communication elements reflect the different needs of various target audiences, such as workers and consumers. To proceed further up the pyramid, some existing national programs also include risk management systems as part of an overall program on the sound management of chemicals. The general goal of these systems is to minimize exposure, resulting in reduced risk. The systems vary in focus and include activities such as establishing exposure limits, recommending exposure monitoring methods and creating engineering controls. However, the target audiences of such systems are generally limited to workplace settings. With or without formal risk management systems, the GHS is designed to promote the safe use of chemicals.



#### 2.1 Are all chemicals covered by the GHS?

The GHS covers all hazardous chemicals. There are no complete exemptions from the scope of the GHS for a particular type of chemical or product. The term "chemical" is used broadly to include substances, products, mixtures, preparations, or any other terms that may be used by existing systems. The goal of the GHS is to identify the intrinsic hazards of chemical substances

and mixtures and to convey hazard information about these hazards. The GHS is not intended to harmonize risk assessment procedures or risk management decisions, as described above.

"Articles" as defined in the OSHA Hazard Communication Standard (HCS) (29 CFR 1910.1200), or by similar definitions, are outside the scope of the GHS. Chemical inventory (e.g., TSCA, EINECS, etc.) and chemical control requirements in various countries are not harmonized by the GHS.

Classification in the GHS is criteria-based, not limiting coverage to a list that can become outdated. It is not anticipated that the GHS will develop or maintain an international classification authority or international classification list. Several countries currently maintain regulatory lists. GHS classification criteria can be used to reclassify chemicals on lists, if desired. Existing lists, such as those provide by organizations that evaluate cancer hazards, could be used in conjunction with the GHS to promote harmonization.

#### 2.2 Will all hazardous chemicals require a GHS label and Safety Data Sheet?

The need for GHS labels and/or Safety Data Sheets is expected to vary by product category or stage in the chemical's lifecycle from research/production to end use. The sequence of lifecycle events is shown in Figure 2.2. For example, pharmaceuticals, food additives, cosmetics and

pesticide residues in food will not be covered by the GHS at the point of consumption, but will be covered where workers may be exposed (workplaces), and in transport. Also, the medical use of human or veterinary pharmaceuticals is generally addressed in package inserts and is not part of existing hazard communication systems. Similarly, foods are generally not labeled under existing hazard communication systems. The exact requirements for labels and Safety Data Sheets will continue to be defined in national regulations. However, national requirements are expected to be consistent with the detailed discussion of scope provided in Chapter 1.1 of the GHS document.



#### 2.3 How will the GHS impact existing regulations?

The GHS is a voluntary international system that imposes no binding treaty obligations on countries. To the extent that countries adopt the GHS into their systems, the regulatory changes

would be binding for covered industries. For countries with existing systems, it is expected that the GHS components will be applied within the framework/infrastructure of existing hazard communication regulatory schemes. For example, exceptions and exemptions found in existing regulations would not be expected to change (e.g., transportation of limited quantities).

*However*, the specific hazard criteria, classification processes, label elements and SDS requirements within an existing regulation will need to be modified to be consistent with the harmonized elements of the GHS. It is anticipated that <u>ALL</u> existing hazard communication systems will need to be changed in order to apply the GHS. For example, in the U.S. EPA and OSHA would be expected to require hazard pictograms/symbols on labels. Canada and the EU would be expected to adopt the GHS pictograms/symbols instead of those currently in use. The transport sector is expected to adopt the changed criteria ( $LD_{50}/LC_{50}$ ) for the GHS Acute Toxicity Categories 1 - 3. OSHA HCS, WHMIS and the EU would all need to change their acute toxicity criteria.

Test data already generated for the classification of chemicals under existing systems should be accepted when classifying these chemicals under the GHS, thereby avoiding duplicative testing and the unnecessary use of test animals.

# 2.4 What is meant by GHS Building Blocks?

The GHS classification and communication requirements can be thought of as a collection of building blocks. In regulatory schemes, coverage and communication of hazards vary by the needs of target audiences/sectors. Accordingly, the GHS was designed to contain the hazard endpoints and communication tools necessary for application to known regulatory schemes. The GHS is structured so that the appropriate elements for classification and communication, which address the target audiences, can be selected.

The full range of harmonized elements is available to everyone, and should be used if a country or organization chooses to cover a certain effect when it adopts the GHS. The full range of these elements does not have to be adopted. Countries can determine which of the building blocks will be applied in different parts of their systems (consumer, workplace, transport, pesticides, etc.). For example, some options for implementing the GHS include:

- Not using a GHS class (e.g., cancer, hazardous to the aquatic environment, etc.);
- Not using a GHS category (normally at the beginning or end of a class, e.g., Acute Toxicity Cat. 5);
- Combining categories (e.g., Acute Toxicity Cat.# 1 and Cat.# 2; Skin Corrosion Cat.1A, 1B and 1C).

# 2.5 How should the GHS Building Blocks by applied?

Appropriate implementation of the GHS means that the hazards covered by a Competent Authority (CA) are covered consistently with the GHS criteria and requirements. The EPA, Health Canada and OSHA are examples of Competent Authorities. Competent Authorities will decide how to apply the various elements of the GHS based on the CA needs and the needs of target audiences. When a regulatory scheme covers something that is in the GHS, and implements the GHS, that coverage should be consistent. Once an endpoint and subclasses are selected, as needed, the GHS classification criteria, assigned label elements and SDS provisions should be followed as specified in the GHS. If a regulatory system covers carcinogenicity, for example, it should follow the harmonized classification scheme, the harmonized label elements and, where appropriate, the SDS. Figure 2.3 shows some of the hazard endpoint/subcategory and hazard communication building block choices for the transport, workplace, consumer and pesticide sectors.



To gain a better understanding of the building block approach, it is helpful to look at the specific sectors/target audiences. The needs and regulations of the various sectors vary depending on the type of chemical and use pattern. Different target audiences or sectors receive and use hazard information in different ways. The primary sectors/target audiences are transport, workplace, consumers and agriculture (pesticides). These sectors are described in more detail below.

#### 2.5.1 Transport

For transport, it is expected that application of the GHS will be similar to application of current transport requirements.

- GHS physical, acute and environmental hazard criteria are expected to be adopted in the transport sector.
- Containers of dangerous goods will have pictograms that address acute toxicity, physical hazards, and environmental hazards.

• GHS hazard communication elements such as signal words, hazard statements and SDS are not expected to be adopted in the transport sector.

# 2.5.2 Workplace

In the workplace, it is expected that most of the GHS elements will be adopted, including;

- GHS physical and health hazard criteria, as appropriate;
- Labels that have the harmonized core information under the GHS (signal words, hazard statements and symbols, etc.);
- Safety Data Sheets;
- Employee training to help ensure effective communication is also anticipated;

All workplace systems may not have the jurisdiction to adopt environmental hazards.

# 2.5.3 Consumer

For the consumer sector, it is expected that labels will be the primary focus of GHS application.

- The appropriate GHS hazard criteria are expected to be adopted;
- These labels will include the core elements of the GHS (signal words, hazard statements and symbols, etc.), subject to some sector-specific considerations in certain systems (e.g., risk-based labeling).

# 2.5.4 Pesticides

For pesticides, it is expected that the GHS will be adopted.

- The appropriate GHS hazard criteria are expected to be adopted;
- Pesticide labels will include the core elements of the GHS (signal words, hazard statements and symbols, etc.), subject to some sector-specific considerations in certain systems.

# 2.6 How will the GHS impact countries without existing regulations?

Developing and maintaining a classification and labeling system is not a simple task. The GHS can be used as a tool for developing national regulations. It is expected that countries that do not have systems will adopt GHS as their basic scheme. The GHS provides the building blocks from which countries can construct chemical safety programs. Although the GHS will facilitate the process, many challenges exist in creating new regulations. For example:

- What is the appropriate legal framework for adopting/implementing the GHS?
- What government agencies should be involved? Are there ministries/agencies ready to implement and maintain the GHS?
- How will stakeholder cooperation and support for implementing the GHS be managed?

Work has begun in international organizations (e.g, UNITAR and ILO) under the guidance of the UN GHS Sub-Committee, to develop technical assistance for developing countries to write new regulations using the GHS elements. Guidance has been developed on how to implement a national GHS action plan. Additionally, pilot implementations have begun in a few countries. The opportunities and challenges learned from the pilot programs will be documented and are expected to facilitate future implementations.

# 3.0 WHAT IS CLASSIFICATION?

Classification is the starting point for hazard communication. It involves the identification of the hazard(s) of a chemical or mixture by assigning a category of hazard/danger using defined criteria. The GHS is designed to be consistent and transparent. It draws a clear distinction between classes and categories in order to allow for **"self classification"**. For many hazards a decision tree approach (e.g., eye irritation) is provided in the GHS Document. For several hazards the GHS criteria are semi-quantitative or qualitative. Expert judgment may be required to interpret these data.

Figure 3.1

#### Figure 3.1 shows the harmonized definition for hazard classification, which can be applied to all hazard categories in the system.

The data used for classification may be obtained from tests, literature, and practical experience. The GHS health and environmental hazard criteria/definitions are test method neutral. Accordingly, tests that determine hazardous properties conducted according to internationally recognized

# Hazard Classification

The term "hazard classification" is used to indicate that only the intrinsic hazardous properties of substances and mixtures are considered and involves the following 3 steps:

- a) Identification of relevant data regarding the hazards of a substance or mixture;
- b) Subsequent review of those data to ascertain the hazards associated with the substance or mixture; and
- c) A decision on whether the substance or mixture will be classified as a hazardous substance or mixture and the degree of hazard, where appropriate, by comparison of the data with agreed hazard classification criteria.

scientific principles can be used for purposes of hazard classification.

The GHS endpoints that cover physical, health and environmental hazards are listed in Figures 3.2 and 3.3, respectively. As mentioned earlier, the GHS hazard definitions are criteria-based. The following information provides an overview of the GHS definitions and classification criteria. It is recommended that the person responsible for GHS implementation consult the GHS Document or "Purple Book" for more complete information.

#### 3.1 What are the GHS Physical Hazards?

The GHS physical hazards criteria, developed by the ILO and UNCETDG, were largely based on the existing criteria used by the UN Model Regulation on the Transport of Dangerous Goods. Therefore, many of the criteria are already being used on a worldwide basis. However, some additions and changes were necessary since the scope of the GHS includes all target audiences. The physical hazards classification process provides specific references to approved test methods and criteria for classification. The GHS physical hazard criteria apply to mixtures. It is assumed that mixtures will be tested for physical hazards.

In general, the GHS criteria for physical hazards are quantitative or semi-quantitative with multiple hazard levels within an endpoint. This is different from several of the existing systems that currently have qualitative criteria for various physical hazards (e.g., organic peroxide criteria

under WHMIS and OSHA HCS). This could make classification under the GHS more consistent.

In developing GHS criteria for physical hazards it was necessary to define physical states. In the GHS,

- a **gas** is a substance or mixture which at 50°C has a vapor pressure greater than 300 kPa; or is completely gaseous at 20°C and a standard pressure of 101.3 kPa.
- a **liquid** is a substance or mixture that is not a gas and which has a melting point or initial melting point of 20°C or less at standard pressure of 101.3 kPa.
- a **solid** is a substance or mixture that does not meet the definitions of a liquid or a gas.

The GHS physical hazards are briefly described below. For many of the physical hazards the GHS Document contains Guidance Sections with practical information to assist in applying the criteria.

#### Figure 3.2

#### Physical Hazards

- Explosives
- Flammable Gases
- Flammable Aerosols
- Oxidizing Gases
- Gases Under Pressure
- Flammable Liquids
- Flammable Solids
- Self-Reactive Substances
- Pyrophoric Liquids
- Pyrophoric Solids
- Self-Heating Substances
- Substances which, in contact with water, emit flammable gases
- Oxidizing Liquids
- Oxidizing Solids
- Organic Peroxides
- Corrosive to Metals

#### 3.1.1 Explosives

An explosive substance (or mixture) is a solid or liquid which is in itself capable by chemical reaction of producing gas at such a temperature and pressure and at such a speed as to cause damage to the surroundings. Pyrotechnic substances are included even when they do not evolve gases. A pyrotechnic substance (or mixture) is designed to produce an effect by heat, light, sound, gas or smoke or a combination of these as the result of non-detonative, self-sustaining, exothermic chemical reactions.

Classification as an explosive and allocation to a division is a three-step process:

- Ascertain if the material has explosive effects (Test Series 1);
- Acceptance procedure (Test Series 2 to 4);
- Assignment to one of six hazard divisions (Test Series 5 to 7).

Explosive properties are associated with certain chemical groups that can react to give very rapid increases in temperature or pressure.

The GHS provides a screening procedure that is aimed at identifying the presence of such reactive groups and the potential for rapid energy release. If the screening procedure identifies the substance or mixture to be a potential explosive, the acceptance procedure has to be

#### Table 3.1 Explosives

Division	Characteristics				
1.1	Mass explosion hazard				
1.2	Projection hazard				
1.3	Fire hazard or minor projection hazard				
1.4	No significant hazard				
1.5	Very insensitive substances with mass explosion hazard				
1.6	Extremely insensitive articles with no mass explosion hazard				

performed.

Substances, mixtures and articles are assigned to one of six divisions, 1.1 to 1.6, depending on the type of hazard they present. See, *UN Manual of Tests and Criteria* Part I Test Series 2 to 7. Currently, only the transport sector uses six categories for explosives.

# 3.1.2 Flammable Gases

Flammable gas means a gas having a flammable range in air at 20°C and a standard pressure of 101.3 kPa. Substances and mixtures of this hazard class are assigned to one of two hazard categories on the basis of the outcome of the test or calculation method (ISO 10156:1996).

# 3.1.3 Flammable Aerosols

Aerosols are any gas compressed, liquefied or dissolved under pressure within a non-refillable container made of metal, glass or plastic, with or without a liquid, paste or powder. The container is fitted with a release device allowing the contents to be ejected as solid or liquid particles in suspension in a gas, as a foam, paste or powder or in a liquid or gaseous state.

Aerosols should be considered for classification as either a Category 1 or Category 2 Flammable Aerosol if they contain any component classified as flammable according to the GHS criteria for flammable liquids, flammable gases, or flammable solids. Classification is based on:

- Concentration of flammable components;
- Chemical heat of combustion (mainly for transport/storage);
- Results from the foam test (foam aerosols) (mainly for worker/consumer);
- Ignition distance test (spray aerosols) (mainly for worker/consumer);
- Enclosed space test (spray aerosols) (mainly for worker/consumer).

Aerosols are considered:

- Nonflammable, if the concentration of the flammable components  $\leq$  1% and the heat of combustion is < 20 kJ/g.
- Extremely flammable, if the concentration of the flammable components >85% and the heat of combustion is  $\geq$  30 kJ/g to avoid excessive testing.

See the UN Manual of Tests and Criteria for the test method.

# 3.1.4 Oxidizing Gases

Oxidizing gas means any gas which may, generally by providing oxygen, cause or contribute to the combustion of other material more than air does. Substances and mixtures of this hazard class are assigned to a single hazard category on the basis that, generally by providing oxygen, they cause or contribute to the combustion of other material more than air does. The test method is ISO 10156:1996. Currently, several workplace hazard communication systems cover oxidizers (solids, liquids, gases) as a class of chemicals.

#### 3.1.5 Gases under Pressure

Gases under pressure are gases that are contained in a receptacle at a pressure not less than 280 Pa at 20°C or as a refrigerated liquid. This endpoint covers four types of gases or gaseous mixtures to address the effects of sudden release of pressure or freezing which may lead to serious damage to people, property, or the environment independent of other hazards the gases may pose.

For this group of gases, the following information is required:

- vapor pressure at 50°C:
- physical state at 20°C at standard ambient pressure;
- critical temperature.

Criteria that use the physical state or compressed gases will be a different classification basis for some workplace systems.

#### Table 3.2 Gases under Pressure

Group	Criteria		
Compressed gas	Entirely gaseous at - 50°C		
Liquefied gas	Partially liquid at temperatures > - 50°C		
Refrigerated liquefied gas	Partially liquid because of its low temperature		
Dissolved gas	Dissolved in a liquid phase solvent		

Data can be found in the literature,

and calculated or determined by testing. Most pure gases are already classified in the UN Model Regulations. Gases are classified, according to their physical state when packaged, into one of four groups as shown in Table 3.2.

#### **3.1.6 Flammable Liquids**

Flammable liquid means a liquid having a flash point of not more than 93°C. Substances and mixtures of this hazard class are assigned to one of four hazard

categories on the basis of the

#### Table 3.3 Flammable Liquids

Category	Criteria			
1	Flash point < 23°C (73°F) and initial boiling point $\leq$ 35°C (95°F)			
2	Flash point $<23\ ^\circ C\ (73\ ^\circ F)$ and initial boiling point $>35\ ^\circ C\ (95\ ^\circ F)$			
3	Flash point $\geq$ 23 °C (73°F) and $\leq$ 60 °C (140°F)			
4	Flash point $>$ 60 °C (140°F) and $\leq$ 93 °C (200°F)			

flash point and boiling point (See Table 3.3). Flash Point is determined by closed cup methods as provided in the GHS document, Chapter 2.5, paragraph 11.

#### 3.1.7 Flammable Solids

Flammable solids are solids that are readily combustible, or may cause or contribute to fire through friction. Readily combustible solids are powdered, granular, or pasty substances which are dangerous if they can be easily ignited by brief contact with an ignition source, such as a burning match, and if the flame spreads rapidly.

Table 3.4 Flammable Solids

Category	Criteria			
1	Metal Powders: burning time $\leq$ 5 minutes			
	Others: wetted zone does not stop fire & burning time < 45 seconds or burning > 2.2 mm/second			
2	Metal Powders: burning time > 5 and ≤ 10 minutes Others: wetted zone stop fire for at least 4 minutes & burning time < 45 seconds or burning rate > 2.2 mm/second			

Substances and mixtures of this hazard class are assigned to one of two hazard categories (Table 3.4) on the basis of the outcome of the UN Test N.1 (*UN Manual of Tests and Criteria*). The tests include burning time, burning rate and behavior of fire in a wetted zone of the test sample.

#### 3.1.8 Self-Reactive Substances

Self-reactive substances are thermally unstable liquids or solids liable to undergo a strongly exothermic thermal decomposition even without participation of oxygen (air). This definition excludes materials classified under the GHS as explosive, organic peroxides or as oxidizing. These materials may have similar properties, but such hazards are addressed in their specific endpoints. There are exceptions to the self-reactive classification for material: (i) with heat of decomposition <300 J/g or (ii) with self-accelerating decomposition temperature (SADT) > 75°C for a 50 kg package.

Substances and mixtures of this hazard class are assigned to one of the seven 'Types', A to G, on the basis of the outcome of the UN Test Series A to H (*UN Manual of Tests and Criteria*). Currently, only the transport sector uses seven categories for self-reactive substances (Table 3.5).

Туре	Criteria						
А	Can detonate or deflagrate rapidly, as packaged.						
В	Possess explosive properties and which, as packaged, neither detonates nor deflagrates rapidly, but is liable to undergo a thermal explosion in that package.						
С	Possess explosive properties when the substance or mixture as packaged cannot detonate or deflagrate rapidly or undergo a thermal explosion.						
D	• Detonates partially, does not deflagrate rapidly and shows no violent effect when heated under confinement; or						
	<ul> <li>Does not detonate at all, deflagrates slowly and shows no violent effect when heated under confinement; or</li> </ul>						
	Does not detonate or deflagrate at all and shows a medium effect when heated under confinement.						
E	Neither detonates nor deflagrates at all and shows low or no effect when heated under confinement.						
F	Neither detonates in the cavitated bubble state nor deflagrates at all and shows only a low or no effect when heated under confinement as well as low or no explosive power.						
G	Neither detonates in the cavitated state nor deflagrates at all and shows no effect when heated under confinement nor any explosive power, provided that it is thermally stable (self-accelerating decomposition temperature is 60°C to 75°C for a 50 kg package), and, for liquid mixtures, a diluent having a boiling point not less than 150°C is used for desensitization.						

# **Pyrophorics**

# 3.1.9 Pyrophoric Liquids

A pyrophoric liquid is a liquid which, even in small quantities, is liable to ignite within five minutes after coming into contact with air. Substances and mixtures of this hazard class are assigned to a single hazard category on the basis of the outcome of the UN Test N.3 (*UN Manual of Tests and Criteria*).

#### 3.1.10 Pyrophoric Solids

A pyrophoric solid is a solid which, even in small quantities, is liable to ignite within five minutes after coming into contact with air. Substances and mixtures of this hazard class are assigned to a single hazard category on the basis of the outcome of the UN Test N.2 (*UN Manual of Tests and Criteria*).

#### 3.1.11 Self-Heating Substances

A self-heating substance is a solid or liquid, other than a pyrophoric substance, which, by reaction with air and without energy supply, is liable to self-heat. This endpoint differs from a pyrophoric substance in that it will ignite only when in large amounts (kilograms) and after long periods of time (hours or days). Substances and mixtures of this hazard class are assigned to one of two hazard categories on the basis of the outcome of the UN Test N.4 (*UN Manual of Tests and Criteria*).

#### 3.1.12 Substances which on Contact with Water Emit Flammable Gases

Substances that, in contact with water, emit flammable gases are solids or liquids which, by

interaction with water, are liable to become spontaneously flammable or to give off flammable gases in dangerous quantities. Substances and mixtures of this hazard class are assigned to one of three hazard categories on the basis of test results (UN Test N.5 UN Manual of Tests and Criteria) which measure gas evolution and speed of evolution.

Table 3.6 Substances which on Contact with Water Emit Flammable Gases					
Category Criteria					
1	≥10 L/kg/1 minute				
2	≥20 L/kg/1 hour + < 10 L/kg/1 min				
3	$\geq$ 1 L/kg/1 hour + < 20 L/kg/1 hour				
Not classified < 1 L/kg/1 hour					

# 3.1.13 Oxidizing Liquids

An oxidizing liquid is a liquid which, while in itself not necessarily combustible, may, generally by yielding oxygen, cause or contribute to the combustion of other material. Substances and mixtures of this hazard class are assigned to one of three hazard categories on the basis of test results (UN Test O.2 *UN Manual of Tests and Criteria*) which measure ignition or pressure rise time compared to defined mixtures.

#### 3.1.14 Oxidizing Solids

An oxidizing solid is a solid which, while in itself not necessarily combustible, may, generally by yielding oxygen, cause or contribute to the combustion of other material. Substances and mixtures of this hazard class are assigned to one of three hazard categories on the basis of test results (UN Test O.1 *UN Manual of Tests and Criteria*) which measure mean burning time and are compared to defined mixtures. Currently, several workplace hazard communication systems cover oxidizers (solids, liquids, gases) as a class of chemicals.

#### **3.1.15 Organic Peroxides**

An organic peroxide is an organic liquid or solid which contains the bivalent -0-0- structure and may be considered a derivative of hydrogen peroxide, where one or both of the hydrogen atoms have been replaced by organic radicals. The term also includes organic peroxide formulations (mixtures). Such substances and mixtures may:

- be liable to explosive decomposition;
- burn rapidly;
- be sensitive to impact or friction;
- react dangerously with other substances.

Substances and mixtures of this hazard class are assigned to one of seven 'Types', A to G, on the basis of the outcome of the UN Test Series A to H (*UN Manual of Tests and Criteria*). Currently, only the transport sector uses seven categories for organic peroxides.

Туре	Criteria
Α	Can detonate or deflagrate rapidly, as packaged.
В	Possess explosive properties and which, as packaged, neither detonates nor deflagrates rapidly, but is liable to undergo a
	thermal explosion in that package.
С	Posses explosive properties when the substance or mixture as packaged cannot detonate or deflagrate rapidly or undergo a
	thermal explosion.
D	• Detonates partially, does not deflagrate rapidly and shows no violent effect when heated under confinement; or
	• Does not detonate at all, deflagrates slowly and shows no violent effect when heated under confinement; or
	<ul> <li>Does not detonate or deflagrate at all and shows a medium effect when heated under confinement.</li> </ul>
E	Neither detonates nor deflagrates at all and shows low or no effect when heated under confinement.
F	Neither detonates in the cavitated bubble state nor deflagrates at all and shows only a low or no effect when heated under
	confinement as well as low or no explosive power.
G	Neither detonates in the cavitated state nor deflagrates at all and shows no effect when heated under confinement nor any explosive power, provided that it is thermally stable (self-accelerating decomposition temperature is 60°C to 75°C for a 50 kg package), and, for liquid mixtures, a diluent having a boiling point not less than 150°C is used for desensitization.

#### Table 3.7 Organic Peroxides

#### 3.1.16 Substances Corrosive to Metal

A substance or a mixture that by chemical action will materially damage, or even destroy, metals is termed 'corrosive to metal'. These substances or mixtures are classified in a single hazard category on the basis of tests (Steel: ISO 9328 (II): 1991 - Steel type P235; Aluminum: ASTM G31-72 (1990) – non-clad types 7075-T6 or AZ5GU-T66). The GHS criteria are a corrosion rate on steel or aluminum surfaces exceeding 6.25 mm per year at a test temperature of 55°C.

The concern in this case is the protection of metal equipment or installations in case of leakage (e.g., plane, ship, tank), not material compatibility between the container/tank and the product. This hazard is not currently covered in all systems.

#### **3.2** What are the GHS Health and Environmental Hazards?

The GHS health and environmental hazard criteria represent a harmonized approach for existing classification systems (see Figure 3.3). The work at the OECD to develop the GHS criteria included:

- A thorough analysis of existing classification systems, including the scientific basis for a system and its criteria, its rationale and an explanation of the mode of use;
- A proposal for harmonized criteria for each category. For some categories the harmonized approach was easy to develop because the existing systems had similar approaches. In cases where the approach was different, a compromise consensus proposal was developed.
- Health and environmental criteria were established for substances and mixtures.

#### Figure 3.3



#### The GHS Health and Environmental Endpoints

The following paragraphs briefly describe the GHS health and environmental endpoints. The criteria for classifying substances are presented first. Then the GHS approach to classifying mixtures is briefly discussed. It is recommended that the person responsible for GHS implementation consult the GHS Document or "Purple Book" for more complete information.

#### 3.2.1 Acute Toxicity

Five GHS categories have been included in the GHS Acute Toxicity scheme from which the appropriate elements relevant to transport, consumer, worker and environment protection can be selected. Substances are assigned to one of the five toxicity categories on the basis of  $LD_{50}$  (oral, dermal) or  $LC_{50}$  (inhalation). The  $LC_{50}$  values are based on 4-hour tests in animals. The GHS provides guidance on

converting 1-hour

inhalation test results to a 4hour equivalent. The five categories are shown in the Table 3.8 Acute Toxicity.

Category 1, the most severe toxicity category, has cut-off values currently used primarily by the transport sector for classification for packing groups. Some Competent Authorities may consider combining Acute Categories 1 and 2. Category 5 is for chemicals which are of relatively low

Acute toxicity	Cat. 1	Cat. 2	Cat. 3	Cat. 4	Category 5		
Oral (mg/kg)	≤ 5	> 5 ≤ 50	> 50 ≤ 300	> 300 ≤ 2000	Criteria: • Anticipated oral LD50 between 2000		
Dermal (mg/kg)	≤ 50	> 50 ≤ 200	> 200 ≤ 1000	> 1000 ≤ 2000	<ul> <li>and 5000 mg/kg;</li> <li>Indication of significant effect in humans;*</li> <li>Any mortality at class 4;*</li> <li>Significant clinical signs at class 4;*</li> <li>Indications from other studies.*</li> <li>*If assignment to a more hazardous class is not warranted.</li> </ul>		
Gases (ppm)	≤ 100	> 100 ≤ 500	> 500 ≤ 2500	> 2500 ≤ 5000			
Vapors (mg/l)	≤ 0.5	> 0.5 ≤ 2.0	> 2.0 ≤ 10	> 10 ≤ 20		signs at class 4;*	signs at class 4;*
Dust & mists (mg/l)	≤ 0.05	> 0.05 ≤ 0.5	> 0.5 ≤ 1.0	> 1.0 ≤ 5			

Table 3.8 Acute Toxicity

acute toxicity but which, under certain circumstances, may pose a hazard to vulnerable populations. Criteria other than LD50/LC50 data are provided to identify substances in Category 5 unless a more hazardous class is warranted.

#### 3.2.2 Skin Corrosion

Skin corrosion means the production of irreversible damage to the skin following the application of a test substance for up to 4 hours. Substances and mixtures in this hazard class are assigned to a single harmonized corrosion category. For Competent Authorities, such as transport packing groups, needing more than one designation for corrosivity, up to three subcategories are provided within the corrosive category. See the Skin Corrosion/Irritation Table 3.9.

Several factors should be considered in determining the corrosion potential before testing is initiated:

- Human experience showing irreversible damage to the skin;
- Structure/activity or structure property relationship to a substance or mixture already classified as corrosive;
- pH extremes of  $\leq 2$  and  $\geq 11.5$  including acid/alkali reserve capacity.

Skin Corrosion			Skin Irritation	Mild Skin Irritation
Category 1			Category 2	Category 3
Destruction of dermal tissue: visible necrosis in at least one			Reversible adverse effects	Reversible adverse
	animal			effects in dermal tissue
Subcategory 1A Exposure < 3 min. Observation < 1 hr,	e < 3 min. Exposure < 1 hr. Exposure < 4 hrs.		Draize score: $\geq 2.3 < 4.0$ or persistent inflammation	Draize score: $\geq 1.5 < 2.3$

#### 3.2.3 Skin Irritation

Skin irritation means the production of reversible damage to the skin following the application of a test substance for up to 4 hours. Substances and mixtures in this hazard class are assigned to a single irritant category. For those authorities, such as pesticide regulators, wanting more than one designation for skin irritation, an additional mild irritant category is provided. See the Skin Corrosion/Irritation Table 3.9.

Several factors should be considered in determining the irritation potential before testing is initiated:

- Human experience or data showing reversible damage to the skin following exposure of up to 4 hours;
- Structure/activity or structure property relationship to a substance or mixture already classified as an irritant.

#### 3.2.4 Eye Effects

Several factors should be considered in determining the *serious eye damage* or *eye irritation* potential before testing is initiated:

- Accumulated human and animal experience;
- Structure/activity or structure property relationship to a substance or mixture already classified;
- pH extremes like ≤ 2 and ≥ 11.5 that may produce serious eye damage.

Serious eye damage means the production of tissue damage in the eye, or serious physical decay of vision, following application of a test substance to the front surface of the eye, which is not fully reversible within 21 days of application. Substances and mixtures in this hazard class are assigned to a single harmonized category.

#### Table 3.10 Eye Effects

Category 1 Serious eye damage	Category 2 Eye Irritation	
Irreversible damage 21 days after exposure	Reversible adverse effects on cornea, iris, conjunctiva	
Draize score: Corneal opacity $\ge 3$ Iritis $> 1.5$	Draize score: Corneal opacity ≥ 1 Iritis ≥ 1 Redness ≥ 2 Chemosis ≥ 2	
	Irritant Subcategory 2A Reversible in 21 days	Mild Irritant Subcategory 2B Reversible in 7 days

**Eye irritation** means changes in the eye following the application of a test substance to the front surface of the eye, which are fully reversible within 21 days of application. Substances and mixtures in this hazard class are assigned to a single harmonized hazard category. For authorities, such as pesticide regulators, wanting more than one designation for eye irritation, one of two subcategories can be selected, depending on whether the effects are reversible in 21 or 7 days.

#### 3.2.5 Sensitization

**Respiratory sensitizer** means a substance that induces hypersensitivity of the airways following inhalation of the substance. Substances and mixtures in this hazard class are assigned to one hazard category.

**Skin sensitizer** means a substance that will induce an allergic response following skin contact. The definition for "skin sensitizer" is equivalent to "contact sensitizer". Substances and mixtures in this hazard class are assigned to one hazard category. Consideration should be given to classifying substances which cause immunological contact urticaria (an allergic disorder) as contact sensitizers.

# 3.2.6 Germ Cell Mutagenicity

Mutagen means an agent giving rise to an increased occurrence of mutations in populations of cells and/or organisms. Substances and mixtures in this hazard class are assigned to one of two hazard categories. Category 1 has two subcategories. See the Germ Cell Mutagenicity (Table 3.11) below.

Category 1 Known / Presumed		Category 2 Suspected / Possible
Known to produce h	eritable mutations in human germ cells	May induce heritable mutations in human     germ cells
Known to produce heritable mutations in human germ cells         Subcategory 1A         Positive evidence from         epidemiological studies         Positive results in:         • In vivo heritable germ cell tests in         mammals         • Human germ cell tests         • In vivo somatic mutagenicity tests,         combined with some evidence of germ cell		<ul> <li>Positive evidence from tests in mammals and somatic cell tests</li> <li>In vivo somatic genotoxicity supported by in vitro mutagenicity</li> </ul>

# 3.2.7 Carcinogenicity

Carcinogen means a chemical substance or a mixture of chemical substances which induce cancer or increase its incidence. Substances and mixtures in this hazard class are assigned to one of two hazard categories. Category 1 has two subcategories. The Carcinogenicity Guidance Section in the GHS Document includes comments about IARC.

Table 3.12 Carcinogenicity

Cate	Category 2	
Known or Pres	Suspected Carcinogen	
Subcategory 1A Known Human Carcinogen Based on human evidence	Subcategory 1B Presumed Human Carcinogen Based on demonstrated animal carcinogenicity	Limited evidence of human or animal carcinogenicity

#### 3.2.8 Reproductive Toxicity

Reproductive toxicity includeds adverse effects on sexual function and fertility in adult males and females, as well as developmental toxicity in offspring. Substances and mixtures with

reproductive and/or developmental effects are assigned to one of two hazard categories, 'known or presumed' and 'suspected'. Category 1 has two subcategories for reproductive and developmental effects. Materials which cause concern for the health of breastfed children have a separate category, Effects on or Via Lactation.

Table 3.13 Reproductive Toxicity

	-	•	
Category 1		Category 2 Suspected	Additional Category
Known or presumed to cause effects on human reproduction or on development		Human or animal evidence	Effects on or via lactation
Category 1A Known Based on human evidence	Known Presumed Based on human Based on		

#### 3.2.9 Target Organ Systemic Toxicity (TOST): Single Exposure & Repeated Exposure

The GHS distinguishes between single and repeat exposure for Target Organ Effects. Some existing systems distinguish between single and repeat exposure for these effects and some do not. All significant health effects, not otherwise specifically included in the GHS, that can impair function, both reversible and irreversible, immediate and/or delayed are included in the non-lethal target organ/systemic toxicity class (TOST). Narcotic effects and respiratory tract irritation are considered to be target organ systemic effects following a single exposure.

Substances and mixtures of the single exposure target organ toxicity hazard class are assigned to one of three hazard categories in Table 3.14.

	0 1	
Category 1	Category 2	Category 3
<ul> <li>Significant toxicity in humans</li> <li>Reliable, good quality human case studies or epidemiological studies</li> <li>Presumed significant toxicity in humans</li> <li>Animal studies with significant and/or severe toxic effects relevant to humans at generally low exposure (guidance)</li> </ul>	<ul> <li>Presumed to be harmful to human health</li> <li>Animal studies with significant toxic effects relevant to humans at generally moderate exposure (guidance)</li> <li>Human evidence in exceptional cases</li> </ul>	Transient target organ effects <ul> <li>Narcotic effects</li> <li>Respiratory tract irritation</li> </ul>

Table 3.14 TOST: Single Exposure

Substances and mixtures of the repeated exposure target organ toxicity hazard class are assigned to one of two hazard categories in Table 3.15. Table 3.15 TOST: Repeated Exposure

In order to help reach a decision about whether a substance should be classified or not, and to what degree it would be classified (Category 1 vs. Category 2), dose/concentration 'guidance values' are provided in the

Category 1 Category 2 Significant toxicity in humans Presumed to be harmful to human health Reliable, good quality human case studies Animal studies with significant toxic or epidemiological studies effects relevant to humans at generally moderate exposure Presumed significant toxicity in humans (guidance) Animal studies with significant and/or severe Human evidence in exceptional toxic effects relevant to humans at generally cases low exposure (guidance)

GHS. The guidance values and ranges for single and repeated doses are intended only for guidance purposes. This means that they are to be used as part of the weight of evidence approach, and to assist with decisions about classification. They are not intended as strict demarcation values. The guidance value for repeated dose effects refer to effects seen in a standard 90-day toxicity study conducted in rats. They can be used as a basis to extrapolate equivalent guidance values for toxicity studies of greater or lesser duration.

#### 3.2.10 Aspiration Hazard

Aspiration toxicity includes severe acute effects such as chemical pneumonia, varying degrees of pulmonary injury or death following aspiration. Aspiration is the entry of a liquid or solid directly through the oral or nasal cavity, or indirectly from vomiting, into the trachea and lower

respiratory system. Some hydrocarbons (petroleum distillates) and certain chlorinated hydrocarbons have been shown to pose an aspiration hazard in humans. Primary alcohols, and ketones

Table 3.16 Aspiration Toxicity

Category 1: Known (regarded) human	Category 2: Presumed human
- human evidence	- Based on animal studies
<ul> <li>hydrocarbons with kinematic viscosity ≤ 20.5 mm<sup>2</sup>/s at 40° C.</li> </ul>	<ul> <li>surface tension, water solubility, boiling point</li> </ul>
	<ul> <li>kinematic viscosity ≤ 14 mm2/s at 40° C</li> <li>&amp; not Category 1</li> </ul>

have been shown to pose an aspiration hazard only in animal studies.

Substances and mixtures of this hazard class are assigned to one of two hazard categories this hazard class on the basis of viscosity.

#### 3.3 Environmental Hazards

#### 3.3.1 Hazardous to the Aquatic Environment

The harmonized criteria are considered suitable for packaged goods in both supply and use in multi-modal transport schemes. Elements of it may be used for bulk land transport and bulk marine transport under MARPOL (International Convention for the Prevention of Pollution from Ships) insofar as this uses aquatic toxicity. Two Guidance Documents (Annexes 8 and 9 of the GHS Document) cover issues such as data interpretation and the application of the criteria to special substances. Considering the complexity of this endpoint and the breadth of the application, the Guidance Annexes are important in the application of the harmonized criteria.

#### 3.3.1.1 Acute Aquatic Toxicity

Acute aquatic toxicity means the intrinsic property of a material to cause injury to an aquatic organism in a short-term exposure. Substances and mixtures of this hazard class are assigned to one of three toxicity categories on the basis of acute toxicity data:  $LC_{50}$  (fish) or  $EC_{50}$  (crustacea) or  $ErC_{50}$  (for algae or other aquatic plants). In some regulatory systems these acute toxicity categories may be subdivided or extended for certain sectors.

#### 3.3.1.2 Chronic Aquatic Toxicity

Chronic aquatic toxicity means the potential or actual properties of a material to cause adverse effects to aquatic organisms during exposures that are determined in relation to the lifecycle of the organism. Substances and mixtures in this hazard class are assigned to one of four toxicity categories on the basis of acute data *and* environmental fate data:  $LC_{50}$  (fish) or  $EC_{50}$  (crustacea) or  $ErC_{50}$  (for algae or other aquatic plants) *and* degradation/bioaccumulation.

While experimentally derived test data are preferred, where no experimental data are available, validated Quantitative Structure Activity Relationships (QSARs) for aquatic toxicity and log KOW may be used in the classification process. The log  $K_{OW}$  is a surrogate for a measured Bioconcentration Factor (BCF), where such a measured BCF value would always take precedence.

Chronic Category IV is considered a "safety net" classification for use when the available data do not allow classification under the formal criteria, but there are some grounds for concern.

Acute Cat. I		Acute Cat. II		Acute Cat. III	
Acute toxicity ≤1.00 mg/l		Acute toxicity > 1.00 but $\leq$ 10.0 mg/l		Acute toxicity > 10.0 but < 100 mg/l	
$\begin{array}{l} \mbox{Chronic Cat. I} \\ \mbox{Acute toxicity} \\ \leq 1.00 \mbox{ mg/l and lack of rapid} \\ \mbox{degradability and log } K_{ow} \geq 4 \\ \mbox{unless BCF} < 500 \end{array}$	$\label{eq:chronic Cat. II} Acute toxicity \\ > 1.00 \mbox{ but } \le 10.0 \mbox{ mg/l and law} \\ rapid \mbox{ degradability and log } K_{ow} \\ \mbox{ unless BCF} < 500 \mbox{ and unless} \\ \mbox{ chronic toxicity} > 1 \mbox{ mg/l} \end{cases}$	$\geq$ 4 rapid degradability	xicity mg/I and lack of and log $K_{ow} \ge 4$ 00 and unless	$\label{eq:chronic Cat. IV} \begin{array}{c} \mbox{Acute toxicity} \\ > 100 \mbox{ mg/l and lack of rapid} \\ \mbox{degradability and log } K_{ow} \geq 4 \\ \mbox{unless BCF} < 500 \mbox{ and unless} \\ \mbox{chronic toxicity} > 1 \mbox{ mg/l} \end{array}$	

Table 3.17 Acute & Chronic Aquatic Toxicity

#### 3.4 What is the GHS approach to classifying mixtures?

For consistency and understanding the provisions for classifying mixtures, the GHS defines certain terms. These working definitions are for the purpose of evaluating or determining the hazards of a product for classification and labeling.

*Substance*: Chemical elements and their compounds in the natural state or obtained by any production process, including any *additive* necessary to preserve the stability of the product and any *impurities* deriving from the process used, but excluding any solvent which may be separated without affecting the stability of the substance or changing its composition.

*Mixture:* Mixtures or solutions composed of two or more substances in which they do not react.

*Alloy:* An alloy is a metallic material, homogeneous on a macroscopic scale, consisting of two or more elements so combined that they cannot be readily separated by mechanical means. Alloys are considered to be mixtures for the purpose of classification under the GHS.

Where impurities, additives or individual constituents of a substance or mixture have been identified and are themselves classified, they should be taken into account during classification if they exceed the cutoff value/concentration limit for a given hazard class.

As mentioned previously, the GHS physical

hazard criteria apply to mixtures. It is assumed that mixtures will be tested for physical hazards. Each health and environmental endpoint chapter in the GHS contains specific criteria for classifying mixtures as well as substances. The GHS Document or "Purple Book" should be consulted for complete information on classifying mixtures.

The process established for classifying a mixture allows the use of (a) available data for the mixture itself and/or (b) similar mixtures and/or



(c) data for ingredients of the mixture. The GHS approach to the classification of mixtures for health and environmental hazards is tiered, and is dependent upon the amount of information available for the mixture itself and for its components. The process for the classification of mixtures is based on the following steps:

(1) Where test data are available for the mixture itself, the classification of the mixture will be based on that data (See exception for carcinogens, mutagens & reproductive toxins in the GHS Document); (2) Where test data are not available for the mixture itself, then the appropriate bridging principles (as described below) in the specific chapter should be used; (3) If (i) test data are not available for the mixture itself, and (ii) the bridging principles cannot be applied, then use the calculation or cutoff values described in the specific endpoint to classify the mixture.

# 3.5 What are bridging principles?

Bridging principles are an important concept in the GHS for classifying untested mixtures. When a mixture has not been tested, but there are sufficient data on the components and/or similar tested mixtures, these data can be used in accordance with the following bridging principles:

- **Dilution**: If a mixture is diluted with a diluent that has an equivalent or lower toxicity, then the hazards of the new mixture are assumed to be equivalent to the original.
- **Batching**: If a batch of a complex substance is produced under a controlled process, then the hazards of the new batch are assumed to be equivalent to the previous batches.
- **Concentration of Highly Toxic Mixtures**: If a mixture is severely hazardous, then a concentrated mixture is also assumed to be severely hazardous
- **Interpolation within One Toxic Category: Mixtures** having component concentrations within a range where the hazards are known are assumed to have those known hazards.
- **Substantially Similar Mixtures**: Slight changes in the concentrations of components are not expected to change the hazards of a mixture and substitutions involving toxicologically similar components are not expected to change the hazards of a mixture
- Aerosols: An aerosol form of a mixture is assumed to have the same hazards as the tested, non-aerosolized form of the mixture unless the propellant affects the hazards upon spraying.

All bridging principles do not apply to every health and environmental endpoint. Consult each endpoint to determine which bridging principles apply.

When the bridging principles do not apply or ca not be used, the health and environmental hazards of mixtures are estimated based on component information. In the GHS, the methodology used to estimate these hazards varies by endpoint. The GHS Document or "Purple Book" should be consulted for more complete information on classifying mixtures. Figure 3.5 summarizes the GHS mixtures approach for the various health and environmental endpoints.

# 3.6 What testing is required?

The GHS itself does not include requirements for testing substances or mixtures. Therefore, there is no requirement under the GHS to generate test data for any hazard class. Some parts of regulatory systems may require data to be generated (e.g., for pesticides), but these requirements are not related specifically to the GHS. The GHS criteria for determining health and environmental hazards are test method neutral, allowing different approaches as long as they are scientifically sound and validated according to international procedures and criteria already

referred to in existing systems. Test data already generated for the classification of chemicals under existing systems should be accepted when classifying these chemicals under the GHS, thereby avoiding duplicative testing and the unnecessary use of test animals. The GHS physical hazard criteria are linked to specific test methods. It is assumed that mixtures will be tested for physical hazards.

# Figure 3.5 GHS Mixtures

Hazard Endpoint	<b>Classification Approach</b>	Bridging Principles	Comments
Acute toxicity	Acute Toxicity Estimate (ATE): 2 formulas	All	Conversion values, relevant components usually at $\geq 1\%$
Serious Eye Damage & Eye Irritation	Mostly additivity approach, sometimes cutoffs	All	Relevant components usually at $\geq 1\%$ , exceptions for certain chemical classes
Skin corrosion & Skin Irritation	Mostly additivity approach, sometimes cutoffs	All	Relevant components usually at $\geq 1\%$ , exceptions for certain chemical classes
Skin Sensitization	Cutoffs with CA options	Dilution, Batching, Substantially similar mixtures, Aerosols	
Respiratory Sensitization	Cutoffs with CA options	Dilution, Batching, Substantially similar mixtures, Aerosols	
Germ Cell Mutagenicity	Cutoffs	Dilution, Batching, Substantially similar mixtures	Mixture test data only case-by case
Carcinogenicity	Cutoffs with CA options	Dilution, Batching, Substantially similar mixtures	Mixture test data only case-by-case
Reproductive Toxicity	Cutoffs with CA options	Dilution, Batching, Substantially similar mixtures	Mixture test data only case-by-case
Target Organ Systemic Toxicity	Cutoffs with CA options	All	
Aspiration Toxicity	Cutoffs	Dilution, Batching, Concentration of highly toxic mixtures, Interpolation within one toxicity category, Substantially similar mixtures	

Hazard Endpoint	<b>Classification Approach</b>	Bridging Principles	Comments
	Additivity Formula (Acute		Relevant components usually at $\geq$ 1%,
Aquatic	only); Summation Method	Concentration of highly toxic	Mixture test data only case-by-case for
Environment	(Acute or Chronic);	mixtures, Interpolation within	chronic
	Combination of Additivity	one toxicity category,	
	Formula & Summation	Substantially similar mixtures	
	Method		
#### 4. HAZARD COMMUNICATION

Section 3, explained that classification is the starting point for the GHS. Once a chemical has been classified, the hazard(s) must be communicated to target audiences. As in existing systems, labels and Safety Data Sheets are the main tools for chemical hazard communication. They identify the hazardous properties of chemicals that may pose a health, physical or environmental hazard during normal handling or use. The goal of the GHS is to identify the intrinsic hazards found in chemical substances and mixtures, and to convey information about these hazards.

The international mandate for the GHS included the development of a harmonized hazard communication system, including labeling, Safety Data Sheets and easily understandable symbols, based on the classification criteria developed for the GHS.

#### 4.1 What factors influenced development of the GHS communication tools?

Early in the process of developing the GHS communication tools, several significant issues were recognized. One of the most important was comprehensibility of the information provided. After all, the aim of the system is to present hazard information in a manner that the intended audience can easily understand and that will thus minimize the possibility of adverse effects resulting from exposure. The GHS identifies some guiding principles to assist in this process:

- Information should be conveyed in more than one way, e.g., text and symbols;
- The comprehensibility of the components of the system should take account of existing studies and literature as well as any evidence gained from testing;
- The phrases used to indicate degree (severity) of hazard should be consistent across the health, physical and environmental hazards.

Comprehensibility is challenging for a single culture and language. Global harmonization has numerous complexities. Some factors that affected the work include:

- Different philosophies in existing systems on how and what should be communicated;
- Language differences around the world;
- Ability to translate phrases meaningfully;
- Ability to understand and appropriately respond to symbols/pictograms.

These factors were considered in developing the GHS communication tools. The GHS Purple Book includes a comprehensibility-testing instrument in Annex 6.

#### 4.2 Labels

#### 4.2.1 What does a label look like?

Existing systems have labels that look different for the same product. We know that this leads to worker confusion, consumer uncertainty and the need for additional resources to maintain different systems. In the U.S. as well as in other countries, chemical products are regulated by sector/target audience. Different agencies regulate the workplace, consumers, agricultural chemicals and transport. Labels for these sectors/target audiences vary both in the U.S. and globally.

In order to understand the value of the GHS and its benefits to all stakeholders, it is instructive to look at the different labels for one fictional product. In the U.S. the product, ToxiFlam, which has a flash point of 120°F and has an oral LD50 of 275 mg/kg, has different labels for different sectors/target audiences. Label examples as seen in the U.S.A. are shown first, followed by international examples.

#### 4.2.2 USA Examples:

#### Workplace and Workers

In the U.S., regulatory requirements for workplace labels are 'performance oriented'. This results at a minimum in a straightforward label that has a product identity, hazard statement and supplier identification (Figure 4.1). Some products can also have additional labeling requirements depending on their end use.

However, many companies follow the voluntary ANSI Z129.1 Precautionary Labeling Standard for workplace labeling and often use it also for Figure 4.1

ToxiFlam TOXIC COMBUSTIBLE LIQUID AND VAPOR

My Company, My Street, MyTown NJ 00000 Tel: 444 999 9999

labeling consumer products. The American National Standards Institute (ANSI) standard includes several label elements that are core to the GHS as well as other helpful elements to assist users in safe handling (Figure 4.2).

#### Figure 4.2

#### ToxiFlam (Contains XYZ)

## WARNING ! HARMFUL IF SWALLOWED, FLAMMABLE LIQUID AND VAPOR

Do not taste or swallow. Do not take internally. Wash thoroughly after handling. Keep away from heat, sparks and flame. Keep container closed. Use only with adequate ventilation.

**FIRST AID:** If swallowed, do NOT induce vomiting unless directed to do so by medical personnel. Never give anything by mouth to an unconscious person.

**In Case of Fire**, use water fog, dry chemical, CO<sub>2</sub>, or alcohol foam. Water may be ineffective.

Flash Point =  $120^{\circ}$ F. Residue vapor may explode or ignite on ignition; do not cut, drill, grind, or weld on or near this container.

See Material Safety Data Sheet for further details regarding safe use of this product.

My Company, My Street, MyTown NJ 00000, Tel: 444 999 9999

Figure 4.3

In several countries consumer products are regulated separately from workplace chemicals. In the U.S. the CPSC regulates consumer products. Consumer products have required label elements, but only the signal words are specified. The ANSI labeling standard is often used in developing consumer labels.

#### **ToxiFlam** (Contains XYZ) WARNING! TOXIC, COMBUSTIBLE LIQUID AND VAPOR

Do not taste or swallow. Do not take internally. Wash thoroughly after handling. Keep away from heat, sparks and flame. Keep container closed. Use only with adequate ventilation.

#### **FIRST AID**

If swallowed, do NOT induce vomiting unless directed to do so by medical personnel. Never give anything by mouth to an unconscious person.

Keep out of the reach of children.

My Company, My Street, MyTown NJ 00000, Tel: 444 999 9999

#### Transport and Emergency Responders

For hazardous products being transported, outer containers have required label elements, product identifier and hazard symbols. Transportation requirements are in addition to workplace or end use label requirements.





#### Agricultural Chemicals and Pesticides

In many systems, agricultural chemicals often have special label requirements. In the U.S. the EPA is the agency covering these chemicals. A pesticide product with the same hazards as ToxiFlam would have a label developed using FIFRA requirements. FIFRA has requirements for product identity, chemical identity, signal word, hazard statements, and precautionary measures including first aid.

Fiaure 4.5

	ToxiFlam Active/Inerts: Contains XYZ %		
	KEEP OUT OF THE REACH OF CHILDREN		
WARNING: M or using tobacc	PRECAUTIONARY STATEMENTS - HAZARDS TO HUMANS AND DOMESTIC ANIMALS: WARNING: May be fatal if swallowed. Wash thoroughly with soap and water after handling and before eating, drinking or using tobacco . PHYSICAL AND CHEMICAL HAZARDS: Combustible. Do not use or store near heat or open flame.		
If swallowed	<ul> <li>-Call a poison control center or doctor immediately for treatment advice.</li> <li>-Have person sip a glass of water if able to swallow.</li> <li>-Do not induce vomiting unless told to do so by a poison control center or doctor.</li> <li>-Do not give anything by mouth to an unconscious person.</li> </ul>		
EPA Est. No.	My Company, My Street, MyTown AZ 00000, Tel: 444 999 9999 5840-AZ-1 EPA Reg. No. 3120-280		

#### 4.2.3 International Examples

All the previous examples are specific to the U.S. Many companies do business globally. So in addition to the U.S. regulations, these companies would need to comply with the corresponding regulations in the countries to which they export products. Canada and the EU are two existing systems that were considered in the development of the GHS. To illustrate the differences in labeling, it is interesting to examine an EU and Canadian label for ToxiFlam.

#### European Union Label

Labels in the EU have chemical identity, symbols, and R/S (Risk and Safety) phrases which are hazard statements, precautionary measures and first aid.

Figure 4.6



#### Canadian Workplace Hazardous Materials Identification System (WHMIS) Label

The WHMIS label requires product identifier, hazard symbol, hazard statement, precautionary measures, first aid, MSDS statement and supplier identification. In addition to these common label elements, WHMIS requires a hatched border.



#### 4.3 What are the GHS label elements?

Some GHS label elements have been standardized (identical with no variation) and are directly related to the endpoints and hazard level. Other label elements are harmonized with common definitions and/or principles. See Figure 4.8 for an illustration of the GHS label elements.

The standardized label elements included in the GHS are:

- **Symbols (hazard pictograms)**: Convey health, physical and environmental hazard information, assigned to a GHS hazard class and category.
- **Signal Words:** "Danger" or "Warning" are used to emphasize hazards and indicate the relative level of severity of the hazard, assigned to a GHS hazard class and category.
- **Hazard Statements:** Standard phrases assigned to a hazard class and category that describe the nature of the hazard.

The symbols, signal words, and hazard statements have all been standardized and assigned to specific hazard categories and classes, as appropriate. This approach makes it easier for countries to implement the system and should make it easier for companies to comply with regulations based on the GHS. The prescribed symbols, signal words, and hazard statements can be readily selected from Annex 1 of the GHS Purple Book. These standardized elements are not subject to variation, and should appear on the GHS label as indicated in the GHS for each hazard category/class in the system. The use of symbols, signal words or hazard statements other than those that have been assigned to each of the GHS hazards would be contrary to harmonization.



The Section numbers refer to the sections in the GHS Document or "Purple Book".

#### 4.3.1 Symbols/Pictograms

The GHS symbols have been incorporated into pictograms for use on the GHS label. Pictograms include the harmonized hazard symbols plus other graphic elements, such as borders, background patterns or colors which are intended to convey specific information. For transport, pictograms (Table 4.10) will have the background, symbol and colors currently used in the UN Recommendations on the Transport of Dangerous Goods, Model Regulations. For other sectors, pictograms (Table 4.9) will have a black symbol on a white background with a red diamond frame. A black frame may be used for shipments within one country. Where a transport pictogram appears, the GHS pictogram for the same hazard should not appear.

#### 4.3.2 Signal Words

The signal word indicates the relative degree of severity a hazard. The signal words used in the GHS are

"**Danger**" for the more severe hazards, and "**Warning**" for the less severe hazards.

Signal words are standardized and assigned to the hazard categories within endpoints. Some lower level hazard categories do not use signal words. Only one signal word corresponding to the class of the most severe hazard should be used on a label.

#### 4.3.3 Hazard Statements

Hazard statements are standardized and assigned phrases that describe the hazard(s) as determined by hazard classification. An appropriate statement for each GHS hazard should be included on the label for products possessing more than one hazard.

The assigned label elements are provided in each hazard chapter of the Purple Book as well as in Annexes 1 & 2. Figure 4-11 illustrates the assignment of standardized GHS label elements for the acute oral toxicity categories.

Figure 4.9



Figure 4.10

Transport "Pictograms"				
Flammable Liquid	Flammable solid	Pyrophorics		
Flammable Gas Flammable Aerosol	Self-Reactive Substances	(Spontaneously Combustible) Self-Heating Substances		
Substances, which in contact with water, emit flammable gases (Dangerous When Wet)	Oxidizing Gases Oxidizing Liquids Oxidizing Solids	Explosive Divisions 1.1, 1.2, 1.3		
1.4	1.5	1.6		
Explosive Division 1.4	Explosive Division 1.5	Explosive Division 1.6		
2				
Compressed Gases	Acute Toxicity (Poison): Oral, Dermal, Inhalation	Corrosive		
MARINE POLLUTANT	5.2			
Marine Pollutant	Organic Peroxides			

Figure 4	4.1	1
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ACUTE ORAL TOXICITY – Annex 1					
	Category 1	Category 2	Category 3	Category 4	Category 5
LD <sub>50</sub>	≤5 mg/kg	> 5 < 50 mg/kg	≥ 50 < 300 mg/kg	≥ 300 < 2000 mg/kg	≥ 2000 < 5000 mg/kg
Pictogram					No symbol
Signal word	Danger	Danger	Danger	Warning	Warning
Hazard statement	Fatal if swallowed	Fatal if swallowed	Toxic if swallowed	Harmful if swallowed	May be harmful if swallowed

Other GHS label elements include:

- **Precautionary Statements and Pictograms:** Measures to minimize or prevent adverse effects.
- **Product Identifier (ingredient disclosure):** Name or number used for a hazardous product on a label or in the SDS.
- **Supplier identification:** The name, address and telephone number should be provided on the label.
- Supplemental information: non-harmonized information.

#### 4.3.4 Precautionary Statements and Pictograms

Precautionary information supplements the hazard information by briefly providing measures to be taken to minimize or prevent adverse effects from physical, health or environmental hazards. First aid is included in precautionary information. The GHS label should include appropriate precautionary information. Annex 3 of the GHS Purple Book includes precautionary statements and pictograms that can be used on labels.

Annex 3 includes four types of precautionary statements covering: prevention, response in cases of accidental spillage or exposure, storage, and disposal. The precautionary statements have been linked to each GHS hazard statement and type of hazard. The goal is to promote consistent use of precautionary statements. Annex 3 is guidance and is expected to be further refined and developed over time.

#### 4.3.5 Product Identifier (Ingredient Disclosure)

A product identifier should be used on a GHS label and it should match the product identifier used on the SDS. Where a substance or mixture is covered by the UN Model Regulations on the Transport of Dangerous Goods, the UN proper shipping name should also be used on the package.

The GHS label for a substance should include the chemical identity of the substance (name as determined by IUPAC, ISO, CAS or technical name). For mixtures/alloys, the label should include the chemical identities of all ingredients that contribute to acute toxicity, skin corrosion or serious eye damage, germ cell mutagenicity, carcinogenicity, reproductive toxicity, skin or respiratory sensitization, or Target Organ Systemic Toxicity (TOST), when these hazards appear on the label. Where a product is supplied exclusively for workplace use, the Competent Authority may give suppliers discretion to include chemical identities on the SDS, in lieu of including them on labels. The Competent Authority rules for confidential business information (CBI) take priority over the rules for product identification.

#### 4.3.6 Supplier Identification

The name, address and telephone number of the manufacturer or supplier of the product should be provided on the label.

#### 4.3.7 Supplemental Information

Supplemental label information is non-harmonized information on the container of a hazardous product that is not required or specified under the GHS. In some cases this information may be required by a Competent Authority or it may be additional information provided at the discretion of the manufacturer/distributor. The GHS provides guidance to ensure that supplemental information does not lead to wide variation in information or undermine the GHS information. Supplemental information may be used to provide further detail that does not contradict or cast doubt on the validity of the standardized hazard information. It also may be used to provide information about hazards not yet incorporated into the GHS. The labeler should have the option of providing supplementary information related to the hazard, such as physical state or route of exposure, with the hazard statement.

#### 4.4 How are multiple hazards handled on labels?

Where a substance or mixture presents more than one GHS hazard, there is a GHS precedence scheme for pictograms and signal words. For substances and mixtures covered by the UN Recommendations on the Transport of Dangerous Goods, Model Regulations, the precedence of symbols for physical hazards should follow the rules of the UN Model Regulations. For health hazards the following principles of precedence apply for symbols:

(a) if the skull and crossbones applies, the exclamation mark should not appear;

(b) if the corrosive symbol applies, the exclamation mark should not appear where it is used for skin or eye irritation;

(c) if the health hazard symbol appears for respiratory sensitization, the exclamation mark should not appear where it is used for skin sensitization or for skin or eye irritation.

If the signal word 'Danger' applies, the signal word 'Warning' should not appear. All assigned hazard statements should appear on the label. The Competent Authority may choose to specify the order in which they appear.

#### 4.5 Is there a specific GHS label format / layout?

The GHS hazard pictograms, signal word and hazard statements should be located together on the label. The actual label format or layout is not specified in the GHS. National authorities may choose to specify where information should appear on the label or allow supplier discretion.

Figure 4.12 shows an example of a GHS label for the fictional product 'ToxiFlam'. The core GHS label elements are expected to replace the need for the array of different labels shown earlier for ToxiFlam. (Figure 4.8 also illustrates the GHS label elements.)

Figure 4.12 Example GHS Inner Container Label (e.g., bottle inside a shipping box)



There has been discussion about the size of GHS pictograms and that a GHS pictogram might be confused with a transport pictogram or "diamond". Transport pictograms (Table 4.10) are different in appearance than the GHS pictograms (Table 4.9). Annex 7 of the Purple Book explains how the GHS pictograms are expected to be proportional to the size of the label text. So that generally the GHS pictograms would be smaller than the transport pictograms.

Several arrangements for GHS labels are also provided in Annex 7 of the Purple Book. Figure 4.13 shows an arrangement for a combination packaging with an outer shipping box and inner bottles. The shipping box has a transportation

## Figure 4.13 Combination Packaging (Outer box with inner bottles)



pictogram. The inner bottles have a GHS label with a GHS pictogram.

For a container such as a 55 gallon drum, the transport required markings and pictograms may be combined with the GHS label elements or presented separately. In Figure 4.14 a

label arrangement for a single packaging such as a 55 gallon drum is shown. Pictograms and markings required by the transport regulations as well as GHS label and non-duplicative GHS pictogram are shown on the drum.

Figure 4.14 Combination Packaging (Outer box with inner bottles)



A label merging the transportation requirements and the GHS requirements into one label for the fictional product "ToxiFlam" is shown in Figure 4.15. This combined type label could also be used on a 55 gallon drum.

Figure 4.15 Example GHS Outer Container Label (55 gallon/200 liter drum)



#### 4.6 What about risk?

Competent Authorities may vary the application of the components of the GHS by the type of product (industrial, pesticide, consumer, etc.) or the stage in the lifecycle (workplace, farm, retail store, etc.). Once a chemical is classified, the likelihood of adverse effects may be considered in deciding what informational or other steps should be taken for a given product or use setting. Annex 5 of the GHS Purple Book includes a discussion of an example of how risk-based labeling could be considered for chronic health effects of consumer products in the consumer use setting.

#### 4.7 Are workplace containers covered in the GHS ?

Products falling within the scope of the GHS will carry the GHS label at the point where they are supplied to the workplace, and that label should be maintained on the supplied container in the workplace. The GHS label or label elements can also be used for workplace containers (e.g., storage tanks). However, the Competent Authority can allow employers to use alternative means of giving workers the same information in a different written or displayed format when such a format is more appropriate to the workplace and communicates the information as effectively as the GHS label. For example, label information could be displayed in the work area, rather than on the individual containers. Some examples of workplace situations where chemicals may be transferred from supplier containers include: containers for laboratory testing, storage vessels, piping or process reaction systems or temporary containers where the chemical will be used by one worker within a short timeframe.

#### 4.8 What is the GHS Safety Data Sheet (SDS)?

The (Material) Safety Data Sheet (SDS) provides comprehensive information for use in workplace chemical management. Employers and workers use the SDS as sources of information about hazards and to obtain advice on safety precautions. The SDS is product related and, usually, is not able to provide information that is specific for any given workplace where the product may be used. However, the SDS information enables the employer to develop an active program of worker protection measures, including training, which is specific to the individual workplace and to consider any measures that may be necessary to protect the environment. Information in a SDS also provides a source of information for other target audiences such as those involved with the transport of dangerous goods, emergency responders, poison centers, those involved with the professional use of pesticides and consumers.

The SDS should contain 16 headings (Figure 4.14). The GHS MSDS headings, sequence and content are similar to the ISO, EU and ANSI MSDS/SDS requirements, except that the order of sections 2 and 3 have been reversed. The SDS should provide a clear description of the data used to identify the hazards. Figure 4.14 and the GHS Purple Book provide the minimum information that is required in each section of the SDS. Examples of draft GHS SDSs are provided in Appendix B of this guidance document.

The revised Purple Book contains guidance on developing a GHS SDS (Annex 4). Other resources for SDSs include:

- ILO Standard under the Recommendation 177 on Safety in the Use of Chemicals at Work,
- International Standard 11014-1 (1994) of the International Standard Organization (ISO) and ISO Safety Data Sheet for Chemical Products 11014-1: 2003 DRAFT,
- American National Standards Institute (ANSI) Standard Z400.1,
- European Union SDS Directive 91/155/-EEC.

1	<b>T1 4.6</b> 4 0.41 1.4	
1.	Identification of the substance	• GHS product identifier.
	or mixture and of the supplier	• Other means of identification.
		• Recommended use of the chemical and
		restrictions on use.
		• Supplier's details (including name, address, phone
		number, etc.).
		Emergency phone number.
2.	Hazards identification	• GHS classification of the substance/mixture and
		any national or regional information.
		• GHS label elements, including precautionary
		statements. (Hazard symbols may be provided as a
		graphical reproduction of the symbols in black and
		white or the name of the symbol, e.g., flame, skull
		and crossbones.)
		• Other hazards which do not result in classification
		(e.g., dust explosion hazard) or are not covered by
		the GHS.
3.	Composition/information on	Substance
	ingredients	• Chemical identity.
		• Common name, synonyms, etc.
		• CAS number, EC number, etc.
		• Impurities and stabilizing additives which are
		themselves classified and which contribute to the
		classification of the substance.
		Mixture
		• The chemical identity and concentration or
		concentration ranges of all ingredients which are
		hazardous within the meaning of the GHS and are
		present above their cutoff levels.
		<b>NOTE</b> : For information on ingredients, the competent
		authority rules for CBI take priority over the rules for
		product identification.
4.	First aid measures	• Description of necessary measures, subdivided
		according to the different routes of exposure, i.e.,
		inhalation, skin and eye contact, and ingestion.
		• Most important symptoms/effects, acute and
		delayed.
		• Indication of immediate medical attention and
		special treatment needed, if necessary.

#### Figure 4.14

#### Minimum information for an SDS

5	Finafighting magging	
5.	Firefighting measures	<ul> <li>Suitable (and unsuitable) extinguishing media.</li> <li>Specific hazards arising from the chemical (e.g., nature of any hazardous combustion products).</li> <li>Special protective equipment and precautions for</li> </ul>
		firefighters.
6.	Accidental release measures	• Personal precautions, protective equipment and emergency procedures.
		<ul> <li>Environmental precautions.</li> </ul>
		• Methods and materials for containment and cleaning up.
7.	Handling and storage	Precautions for safe handling.
		<ul> <li>Conditions for safe storage, including any incompatibilities.</li> </ul>
8.	Exposure controls/personal	Control parameters, e.g., occupational exposure
	protection.	limit values or biological limit values.
		• Appropriate engineering controls.
		• Individual protection measures, such as personal
-		protective equipment.
9.	Physical and chemical	• Appearance (physical state, color, etc.).
	properties	• Odor.
		• Odor threshold.
		• pH.
		• melting point/freezing point.
		• initial boiling point and boiling range.
		• flash point.
		• evaporation rate.
		• flammability (solid, gas).
		• upper/lower flammability or explosive limits.
		<ul> <li>vapor pressure.</li> <li>vapor density.</li> </ul>
		<ul><li>vapor density.</li><li>relative density.</li></ul>
		<ul> <li>solubility(ies).</li> </ul>
		<ul> <li>partition coefficient: n-octanol/water.</li> </ul>
		<ul> <li>autoignition temperature.</li> </ul>
		<ul> <li>decomposition temperature.</li> </ul>
10.	Stability and reactivity	<ul> <li>Chemical stability.</li> </ul>
	· · ·	<ul> <li>Possibility of hazardous reactions.</li> </ul>
		• Conditions to avoid (e.g., static discharge, shock
		or vibration).
		• Incompatible materials.
		Hazardous decomposition products.

11.	Toxicological information	<ul> <li>Concise but complete and comprehensible description of the various toxicological (health) effects and the available data used to identify those effects, including:</li> <li>information on the likely routes of exposure (inhalation, ingestion, skin and eye contact);</li> <li>Symptoms related to the physical, chemical and toxicological characteristics;</li> </ul>
		<ul> <li>Delayed and immediate effects and also chronic effects from short- and long-term exposure;</li> <li>Numerical measures of toxicity (such as acute toxicity estimates).</li> </ul>
12.	Ecological information	<ul> <li>Ecotoxicity (aquatic and terrestrial, where available).</li> <li>Persistence and degradability.</li> <li>Bioaccumulative potential.</li> <li>Mobility in soil.</li> <li>Other adverse effects.</li> </ul>
13.	Disposal considerations	• Description of waste residues and information on their safe handling and methods of disposal, including the disposal of any contaminated packaging.
14.	Transport information	<ul> <li>UN Number.</li> <li>UN Proper shipping name.</li> <li>Transport Hazard class(es).</li> <li>Packing group, if applicable.</li> <li>Marine pollutant (Yes/No).</li> <li>Special precautions which a user needs to be aware of or needs to comply with in connection with transport or conveyance either within or outside their premises.</li> </ul>
15.	Regulatory information	• Safety, health and environmental regulations specific for the product in question.
16.	Other information including information on preparation and revision of the SDS	

#### 4.9 What is the difference between the GHS SDS and existing MSDSs/SDSs?

SDSs are in use globally. So it is useful to have an understanding of the similarities and differences in the existing MSDS/SDS content and format and the GHS SDS content and format. A table comparing MSDS/SDS content/format is provided in Appendix A of this guidance document.

#### 4.10 When should SDSs and labels be updated?

All hazard communication systems should specify a means of responding in an appropriate and timely manner to new information and updating labels and SDS information accordingly. Updating should be carried out promptly on receipt of the information that necessitates the revision. The Competent Authority may choose to specify a time limit within which the information should be revised.

Suppliers should respond to "new and significant" information they receive about a chemical hazard by updating the label and safety data sheet for that chemical. New and significant information is any information that changes the GHS classification and leads to a change in the label information or information that may affect the SDS.

#### 4.11 How does the GHS address Confidential Business Information (CBI)?

Confidential business information (CBI) will not be harmonized under the GHS. National authorities should establish appropriate mechanisms for CBI protection. The GHS established CBI principles which include:

- CBI provisions should not compromise the health and safety of users;
- CBI claims should be limited to the names of chemicals and their concentrations in mixtures;
- Mechanisms should be established for disclosure in emergency and non-emergency situations.

#### 4.12 Does the GHS address training?

The GHS states in Chapter 1.4, Section1.4.9, the importance of training all target audiences to recognize and interpret label and/or SDS information, and to take appropriate action in response to chemical hazards. Training requirements should be appropriate for and commensurate with the nature of the work or exposure. Key target audiences include workers, emergency responders and also those responsible for developing labels and SDSs. To varying degrees, the training needs of additional target audiences have to be addressed. These should include training for persons involved in transport and strategies required for educating consumers in interpreting label information on products that they use.

#### 5. **REFERENCES**

References for Section 1.

- ANSI Z129.1: American National Standard for Hazardous Industrial Chemicals-Precautionary Labeling.
- Australia: Australia Worksafe, National Occupational Health and Safety Commission, Approved Criteria for Classifying Hazardous Substances (1994).
- CPSC FHSA: U.S. CPSC, 16 CFR 1500, FHSA regulations.
- DOT: U.S. DOT, 49 CFR Part 173, Subpart D.
- EPA FIFRA: U.S. EPA, 40 CFR Part 156, FIFRA regulations.
- EU: Council Directive 92/32/European Economic Community, amending for the 7th time, Directive 67/548/European Economic Community, approximation of the laws, regulations and administrative provisions on the classification, packaging and labeling of dangerous preparations.
- GHS: Globally Harmonized System of Classification and Labelling of Chemicals, United Nations, 1<sup>st</sup> Revised Edition 2005.
- IATA: International Air Transport Association's Dangerous Goods Regulations.
- ICAO: International Civil Aviation Organization's Technical Instructions for the Safe Transport Of Dangerous Goods By Air.
- IMO: International Maritime Organization's International Maritime Dangerous Goods (IMDG) Code.
- Japan: Japanese Official Notice of Ministry of Labor No. 60 "Guidelines for Labeling of the

Danger and Hazards of Chemical Substances".

- Korea: Korean Ministry of Labor Notice 1997-27 "Preparation of MSDS and Labelling Regulation".
- Malaysia: Malaysian Occupational Safety and Health Act (1994), Act 514 and Regulations (1994).
- Mexico: Dario Oficial (March 30, 1996) NORMA Oficial Mexicana NOM-114-STPS-1994.
- NFPA: National Fire Protection Association, 704 Standard, System for the Identification of Fire Hazards of Materials, 2001.
- NPCA HMIS: National Paint and Coatings Association, Hazardous Materials Identification System, 2001.
- OSHA HCS: U.S. DOL, OSHA, 29 CFR 1910.1200.
- WHMIS: Controlled Products Regulation, Hazardous Products Act, Canada Gazette, Part II, Vol. 122, No. 2, 1987.

#### References for Section 2.0:

GHS Chapter 1.1	Purpose, Scope and Application of the GHS.
GHS Chapter 1.3	Classification of Hazardous Substances and Mixtures.

#### References for Section 3.0:

GHS Chapter 1.3. Classification of Hazardous Substances and Mixtures.

- GHS Part 2. Physical Hazards.
- GHS Part 3. Health Hazards.
- GHS Part 4. Environmental Hazards

GHS Annex 8. An Example of Classification in the GHS.

GHS Annex 9. Guidance on Hazards to the Aquatic Environment.

GHS Annex 10. Guidance on Transformation/Dissolution of Metals and Metal Compounds in Aqueous Media

#### References for Section 4:

GHS Chapter 1.4. Hazard Communication: Labelling.

- GHS Chapter 1.5. Hazard Communication: Safety Data Sheets.
- GHS Annex 1 Allocation of Label Elements.
- GHS Annex 2 Classification and Labelling Summary Tables.
- GHS Annex 3 Precautionary Statements and Precautionary Pictograms.
- GHS Annex 4 Guidance on the preparation of Safety Data Sheets
- GHS Annex 5 Consumer Product Labelling Based on the Likelihood of Injur.
- GHS Annex 6 Comprehensibility Testing Methodology.
- GHS Annex 7 Examples of Arrangements of GHS Label Elements.

#### References for Government and Private Standards:

#### Canada

Hazardous Products Act: Controlled Products Regulations; Consumer Chemical and Container Regulations, 2001 Pest Control Products Act; Transportation of Dangerous Goods Act.

Health Canada GHS Website: www.healthcanada.ca/ghs;

#### **European Union (EU)**

Directive 67/548/EEC (consolidated, 7th revision).

- Directive 2001/59/EC adapting to technical progress for the 28th time Council Directive 67/548/EEC.
- Manual of decisions, implementation for the sixth and seventh amendments to Directive 67/548/EEC on dangerous substances.
- Directive 1999/45/EC of the European Parliament and of the Council of 31 May 1999 related to the classification, packaging and labelling of dangerous preparations.
- Commission Directive 91/155/EEC defining and laying down the detailed arrangements for the system of specific information relating to dangerous preparations (SDS.)
- Directive 2001/58/EC (amending Directive 91/155/EEC) defining and laying down the detailed arrangements for the system of specific information relating to dangerous preparations (SDS).

EU GHS web site: http://europa.eu.int/comm/enterprise/reach/ghs\_en.htm

#### Standards

- American National Standard for Hazardous Industrial Chemicals Precautionary Labeling (ANSI Z-129.1-2000).
- American National Standard for Hazardous Industrial Chemicals MSDS Preparation (ANSI Z400.1-2004).
- ISO 11014-1:2003 DRAFT Safety Data Sheet for Chemical Products.

#### UN GHS

- Globally Harmonized System of Classification and Labelling of Chemicals (GHS) ("The Purple Book"), United Nations, 2005 First Revised Edition, available at <u>www.unece.org/trans/danger/publi/ghs/ghs\_rev01/01files\_e.html</u> or from United Nations Publications (publications@un.org)
- UN GHS website: www.unece.org/trans/danger/publi/ghs/ghs\_welcome\_e.htm

#### **UN TRANSPORT**

UN Recommendations on the Transport of Dangerous Goods, Model Regulations (14th Revised Edition 2005).

UN Recommendations on the Transport of Dangerous Goods, Manual of Tests and Criteria, 4<sup>th</sup> Revised Edition

#### USA

OSHA Hazard Communication Standard 29 CFR 1910.1200.

CPSC Consumer Product Safety Act (15 U.S.C. 2051 *et seq.*) and Federal Hazardous Substances Act (15 U.S.C. 1261 *et seq.*).
(FIFRA) Federal Insecticide, Fungicide, and Rodenticide Act (7 U.S.C. 136 *et seq.*).
US EPA Label Review Manual (3rd Edition, August 2003) EPA 735-B-03-001.
Federal Hazardous Materials Transportation Law (49 U.S.C. 5101 *et seq.*).
USA websites:
www.osha.gov/SLTC/hazardcommunications/global.html
www.epa.gov/oppfead1/international/globalharmon.htm
http://hazmat.dot.gov/regs/intl/globharm.htm

#### **GHS Focal Point websites:**

ILO - www.ilo.org/public/english/protection/safework/ghs/index.htm

OECD - www.oecd.org/department/0,2688,en\_2649\_34371\_1\_1\_1\_1\_1,00.html

#### 6.0 GLOSSARY

*Aerosols* means any non-refillable receptacles made of metal, glass or plastics and containing a gas compressed, liquefied or dissolved under pressure, with or without a liquid, paste or powder, and fitted with a release device allowing the contents to be ejected as solid or liquid particles in suspension in a gas, as a foam, paste or powder or in a liquid state or in a gaseous state. Aerosol includes aerosol dispensers.

*Alloy* means a metallic material, homogeneous the naked eye, consisting of two or more elements so combined that they cannot be readily separated by mechanical means. Alloys are considered to be mixtures for the purpose of classification under the GHS.

*Aspiration* means the entry of a liquid or solid chemical product into the trachea and lower respiratory system directly through the oral or nasal cavity, or indirectly from vomiting;

ASTM means the "American Society of Testing and Materials".

BCF means "bioconcentration factor".

BOD/COD means "biochemical oxygen demand/chemical oxygen demand".

CA means "competent authority".

*Carcinogen* means a chemical substance or a mixture of chemical substances which induce cancer or increase its incidence.

CAS means "Chemical Abstract Service".

CBI means "confidential business information".

*Chemical identity* means a name that will uniquely identify a chemical. This can be a name that is in accordance with the nomenclature systems of the International Union of Pure and Applied Chemistry (IUPAC) or the Chemical Abstracts Service (CAS), or a technical name.

*Competent authority* means any national body(ies) or authority(ies) designated or otherwise recognized as such in connection with the Globally Harmonized System of Classification and Labelling of Chemicals (GHS).

*Compressed gas* means a gas which when packaged under pressure is entirely gaseous at -50°C; including all gases with a critical temperature  $\leq$  -50°C.

*Contact sensitizer* means a substance that will induce an allergic response following skin contact. The definition for "contact sensitizer" is equivalent to "skin sensitizer".

*Corrosive to metal* means a substance or a mixture which by chemical action will materially damage, or even destroy, metals.

*Criteria* means the technical definition for the physical, health and environmental hazards;

*Critical temperature* means the temperature above which a pure gas cannot be liquefied, regardless of the degree of compression.

Dermal Corrosion: see skin corrosion; Dermal irritation: see skin irritation.

*Dissolved gas* means a gas which when packaged under pressure is dissolved in a liquid phase solvent.

 $EC_{50}$  means the effective concentration of a substance that causes 50% of the maximum response.

*EC Number or (ECN*) is a reference number used by the European Communities to identify dangerous substances, in particular those registered under EINECS.

ECOSOC means the "Economic and Social Council of the United Nations".

EINECS means "European Inventory of Existing Commercial Chemical Substances".

*End Point* means physical, health and environmental hazards;

 $ErC_{50}$  means EC<sub>50</sub> in terms of reduction of growth rate.

EU means "European Union".

*Explosive article* means an article containing one or more explosive substances.

*Explosive substance* means a solid or liquid substance (or mixture of substances) which is in itself capable by chemical reaction of producing gas at such a temperature and pressure and at such a speed as to cause damage to the surroundings. Pyrotechnic substances are included even when they do not emit gases.

*Eye irritation* means the production of changes in the eye following the application of test substance to the front surface of the eye, which are fully reversible within 21 days of application.

*Flammable gas* means a gas having a flammable range with air at 20°C and a standard pressure of 101.3 kPa.

*Flammable liquid* means a liquid having a flash point of not more than 93°C.

*Flammable solid* means a solid which is readily combustible, or may cause or contribute to fire through friction.

*Flash point* means the lowest temperature (corrected to a standard pressure of 101.3 kPa) at which the application of an ignition source causes the vapors of a liquid to ignite under specified test conditions.

*Gas* means a substance which (i) at 50 °C has a vapor pressure greater than 300 kPa; or (ii) is completely gaseous at 20 °C at a standard pressure of 101.3 kPa.

*GESAMP* means "the Joint Group of Experts on the Scientific Aspects of Marine Environmental Protection of IMO/FAO/UNESCO/WMO/WHO/IAEA/UN/UNEP."

GHS means "the Globally Harmonized System of Classification and # Labeling of Chemicals".

*Hazard category* means the division of criteria within each hazard class, e.g., oral acute toxicity includes five hazard categories and flammable liquids includes four hazard categories. These categories compare hazard severity within a hazard class and should not be taken as a comparison of hazard categories more generally.

*Hazard class* means the nature of the physical, health or environmental hazard, e.g., flammable solid carcinogen, oral acute toxicity.

*Hazard statement* means a statement assigned to a hazard class and category that describes the nature of the hazards of a hazardous product, including, where appropriate, the degree of hazard;

IARC means the "International Agency for the Research on Cancer".

ILO means the "International Labor Organization".

IMO means the "International Maritime Organization".

*Initial boiling point* means the temperature of a liquid at which its vapor pressure is equal to the standard pressure (101.3 kPa), i.e., the first gas bubble appears.

IOMC means the "Inter-organization Program on the Sound Management of Chemicals".

IPCS means the "International Program on Chemical Safety".

ISO means International Standards Organization.

IUPAC means the "International Union of Pure and Applied Chemistry".

*Label* means an appropriate group of written, printed or graphic information elements concerning a hazardous product, selected as relevant to the target sector(s), that is affixed to, printed on, or attached to the immediate container of a hazardous product, or to the outside packaging of a hazardous product.

*Label element* means one type of information that has been harmonized for use in a label, e.g., pictogram, signal word.

 $LC_{50}$  (50% lethal concentration) means the concentration of a chemical in air or of a chemical in water which causes the death of 50% (one-half) of a group of test animals.

 $LD_{50}$  means the amount of a chemical, given all at once, which causes the death of 50% (one half) of a group of test animals.

 $L(E)C_{50}$  means LC<sub>50</sub> or EC<sub>50</sub>.

*Liquefied gas* means a gas which when packaged under pressure, is partially liquid at temperatures above  $-50^{\circ}$ C. A distinction is made between.

- (i) High pressure liquefied gas: a gas with a critical temperature between -50°C and +65°C; and
- (ii) Low pressure liquefied gas: a gas with a critical temperature above  $+65^{\circ}$ C.

*Liquid* means a substance or mixture which at 50°C has a vapor pressure of not more than 300 kPa (3 bar), which is not completely gaseous at 20 °C and at a standard pressure of 101.3 kPa, and which has a melting point or initial melting point of 20°C or less at a standard pressure of 101.3 kPa. A viscous substance or mixture for which a specific melting point cannot be determined shall be subjected to the ASTM D 4359-90 test; or to the test for determining fluidity (penetrometer test) prescribed in section 2.3.4 of Annex A of the European Agreement concerning the International Carriage of Dangerous Goods by Road (ADR).

MARPOL means the "International Convention for the Prevention of Pollution from Ships".

*Mixture* means a mixture or a solution composed of two or more substances in which they do not react.

*MSDS* means "Material Safety Data Sheet" and in this document is used interchangeably with Safety Data Sheet (SDS).

*Mutagen* means an agent giving rise to an increased occurrence of mutations in populations of cells and /or organisms.

*Mutation* means a permanent change in the amount or structure of the genetic material in a cell;

NGO means "non-governmental organization".

NOEC means the "no observed effect concentration".

OECD means "The Organization for Economic Cooperation and Development".

*Organic peroxide* means a liquid or solid organic substance which contains the bivalent -0-0structure and may be considered a derivative of hydrogen peroxide, where one or both of the hydrogen atoms have been replaced by organic radicals. The term also includes organic peroxide formulation (mixtures).

*Oxidizing gas* means any gas which may, generally by providing oxygen, cause or contribute to the combustion of other material more than air does.

*Oxidizing liquid* means a liquid which, while in itself not necessarily combustible, may, generally by yielding oxygen, cause, or contribute to, the combustion of other material.

*Oxidizing solid* means a solid which, while in itself not necessarily combustible, may, generally by yielding oxygen, cause, or contribute to, the combustion of other material.

QSAR means "quantitative structure-activity relationships".

*Pictogram* means a graphical composition that may include a symbol plus other graphic elements, such as a border, background pattern or color that is intended to convey specific information.

*Precautionary statement* means a phrase (and/or pictogram) that describes recommended measures that should be taken to minimize or prevent adverse effects resulting from exposure to a hazardous product, or improper storage or handling of a hazardous product.

*Product identifier* means the name or number used for a hazardous product on a label or in the SDS. It provides a unique means by which the product user can identify the substance or mixture within the particular use setting (e.g. transport, consumer or workplace).

*Pyrophoric liquid* means a liquid which, even in small quantities, is liable to ignite within five minutes after coming into contact with air.

*Pyrophoric solid* means a solid which, even in small quantities, is liable to ignite within five minutes after coming into contact with air.

Pyrotechnic article means an article containing one or more pyrotechnic substances;

*Pyrotechnic substance* means a substance or mixture of substances designed to produce an effect by heat, light, sound, gas or smoke or a combination of these as the result of non-detonative, self-sustaining exothermic (heat-related) chemical reactions.

*Readily combustible solid* means powdered, granular, or pasty substance or mixture which is dangerous if it can be easily ignited by brief contact with an ignition source, such as a burning match, and if the flame spreads rapidly.

*Recommendations on the Transport of Dangerous Goods, Manual of Tests and Criteria* means the latest revised edition of the United Nations publication bearing this title, and any published amendment thereto.

**Recommendations on the Transport of Dangerous Goods, Model Regulations** means the latest revised edition of the United Nations publication bearing this title, and any published amendment thereto.

*Refrigerated liquefied gas* means a gas which when packaged is made partially liquid because of its low temperature.

*Respiratory sensitizer* means a substance that induces hypersensitivity of the airways following inhalation of the substance.

*RID* means The Regulations concerning the International Carriage of Dangerous Goods by Rail [Annex 1 to Appendix B (Uniform Rules concerning the Contract for International Carriage of Goods by Rail) (CIM) of COTIF (Convention concerning international carriage by rail)], as amended.

SAR means "Structure Activity Relationship".

*SDS* means "Safety Data Sheet" and in this document is used interchangeably with Material Safety Data Sheet (MSDS).

*Self-Accelerating Decomposition Temperature (SADT)* means the lowest temperature at which self-accelerating decomposition may occur with substance as packaged.

*Self-heating substance* means a solid or liquid substance, other than a pyrophoric substance, which, by reaction with air and without energy supply, is liable to self-heat; this substance differs from a pyrophoric substance in that it will ignite only when in large amounts (kilograms) and after long periods of time (hours or days).

*Self-reactive substance* means a thermally unstable liquid or solid substance liable to undergo a strongly exothermic decomposition even without participation of oxygen (air). This definition excludes substances or mixtures classified under the GHS as explosive, organic peroxides or as oxidizing.

*Serious eye damage* means the production of tissue damage in the eye, or serious physical decay of vision, following application of a test substance to the front surface of the eye, which is not fully reversible within 21 days of application.

*Signal word* means a word used to indicate the relative level of severity of hazard and alert the reader to a potential hazard on the label. The GHS uses 'Danger' and 'Warning' as signal words.

*Skin corrosion* means the production of irreversible damage to the skin following the application of a test substance for up to 4 hours.

*Skin irritation* means the production of reversible damage to the skin following the application of a test substance for up to 4 hours.

*Skin sensitizer* means a substance that will induce an allergic response following skin contact. The definition for "skin sensitizer" is equivalent to "contact sensitizer".

*Solid* means a substance or mixture which does not meet the definitions of a liquid or gas.

SPR means "Structure Property Relationship".

*Substance* means chemical elements and their compounds in the natural state or obtained by any production process, including any additive necessary to preserve the stability of the product and any impurities deriving from the process used, but excluding any solvent which may be separated without affecting the stability of the substance or changing its composition.

*Substance which, in contact with water, emits flammable gases* means a solid or liquid substance or mixture which, by interaction with water, is liable to become spontaneously flammable or to give off flammable gases in dangerous quantities.

*Supplemental label element* means any additional non-harmonized type of information supplied on the container of a hazardous product that is not required or specified under the GHS. In some cases this information may be required by other competent authorities or it may be additional information provided at the discretion of the manufacturer/distributor.

Symbol means a graphical element intended to succinctly convey information.

*Technical name* means a name that is generally used in commerce, regulations and codes to identify a substance or mixture, other than the IUPAC or CAS name, and that is recognized by the scientific community. Examples of technical names include those used for complex mixtures (e.g., petroleum fractions or natural products), pesticides (e.g., ISO or ANSI systems), dyestuffs (Color Index system) and minerals.

UNCED means the "United Nations Conference on Environment and Development".

*UNCETDG/GHS* means the "United Nations Committee of Experts on the Transport of Dangerous Goods and on the Globally Harmonized System of Classification and Labelling of Chemicals".

UNITAR means the "United Nations Institute for Training and Research";

*UNSCEGHS* means the "United Nations Sub-Committee of Experts on the Globally Harmonized System of Classification and Labelling of Chemicals".

*UNSCETDG* means the "United Nations Sub-Committee of Experts on the Transport of Dangerous Goods".

# 7.0 APPENDICES

# A. Comparison of MSDS/SDS Elements

### **B. GHS MSDS Examples**



# The following tables provide a comparison of MSDS elements for the following:

- ♦ Globally Harmonized System<sup>1</sup>
- ISO Safety Data Sheet for Chemical Products 11014-1: 2003 DRAFT<sup>2</sup>
- ♦ ANSI MSDS Preparation Z400.1- 2004<sup>3</sup>
- OSHA Hazard Communication Standard 29#CFR#1910.1200<sup>4</sup>

	MSDS Comparison			
MSDS Sections	GHS SDS <sup>1</sup>	ISO MSDS <sup>2</sup>	ANSI MSDS <sup>3</sup>	OSHA MSDS <sup>4</sup>
1. Product and company identification	<ul> <li>GHS product identifier.</li> <li>Other means of identification.</li> <li>Recommended use of the chemical and restrictions on use.</li> <li>Supplier's details (including name, address, phone number etc).</li> <li>Emergency phone number.</li> </ul>	<ul> <li>GHS product identifier.</li> <li>Other means of identification.</li> <li>Recommended use of the chemical and restrictions on use.</li> <li>Supplier's details (including name, address, phone number etc).</li> <li>Emergency phone number.</li> </ul>	<ul> <li>Product identity same as on label</li> <li>Product name, product code</li> <li>name, address and telephone number of supplier</li> <li>emergency telephone number</li> </ul>	<ul> <li>Product identity same as on label.</li> <li>Name address and telephone number of the manufacturer, distributor, employer or other responsible party.</li> </ul>
2. Hazards identification	<ul> <li>GHS classification of the substance/mixture and any regional information.</li> <li>GHS label elements, including precautionary statements. (Hazard symbols may be provided as a graphical reproduction of the symbols in black and white or the name of the symbol, e.g., flame, skull and crossbones.)</li> <li>Other hazards which do not result in classification (e.g., dust explosion hazard) or are not covered by the GHS.</li> </ul>	<ul> <li>GHS classification of the substance/mixture and any regional information.</li> <li>GHS label elements, including precautionary statements. (Hazard symbols may be provided as a graphical reproduction of the symbols in black and white or the name of the symbol, e.g., flame, skull and crossbones.)</li> <li>Other hazards which do not result in classification (e.g., dust explosion hazard) or are not covered by the GHS.</li> </ul>	<ul> <li>Emergency Overview         <ul> <li>(description of product and most significant immediate physical, health and environmental concerns)</li> <li>OSHA Regulatory Status</li> <li>Potential health effects                 (information on adverse human health effects and symptoms, relevant route(s) and length of exposure, type and severity of effects, target organs, medical symptoms that are aggravated by exposure)</li> <li>if listed as a carcinogen by OSHA, IARC, NTP</li> <li>environmental effects</li> </ul> </li> </ul>	<ul> <li>health hazards including acute and chronic effects, listing target organs or systems</li> <li>signs &amp; symptoms of exposure</li> <li>conditions generally recognized as aggravated by exposure</li> <li>primary routes of exposure</li> <li>if listed as a carcinogen by OSHA, IARC, NTP</li> <li>physical hazards, including the potential for fire, explosion, and reactivity</li> </ul>

MSDS Comparison				
MSDS Sections	GHS SDS <sup>1</sup>	ISO MSDS <sup>2</sup>	ANSI MSDS <sup>3</sup>	OSHA MSDS <sup>4</sup>
3. Composition/in- formation on ingredients	<ul> <li>Substance</li> <li>Chemical identity</li> <li>Common name, synonyms, etc.</li> <li>CAS number, EC number, etc.</li> <li>Impurities and stabilizing additives which are themselves classified and which contribute to the classification of the substance.</li> <li>Mixture</li> <li>The chemical identity and concentration or concentration ranges of all ingredients which are hazardous within the meaning of the GHS and are present above their cut-off levels.</li> <li>Cut-off level for reproductive toxicity, carcinogenicity and category 1 mutagenicity is ≥ 0.1%</li> <li>Cut-off level for all other hazard classes is ≥ 1%</li> <li>Note: For information on ingredients, the competent authority rules for CBI take priority over the rules for product identification</li> </ul>	<ul> <li>Substance</li> <li>Chemical identity</li> <li>Common name, synonyms etc.</li> <li>CAS number, EC number, etc.</li> <li>Impurities and stabilizing additives which are themselves classified and which contribute to the classification of the substance.</li> <li>Mixture</li> <li>The chemical identity and concentration or concentration ranges of all ingredients which are hazardous within the meaning of the GHS and are present above their cut-off levels.</li> <li>Cut-off level for reproductive toxicity, carcinogenicity and category 1 mutagenicity is ≥ 0.1%</li> <li>Cut-off level for all other hazard classes is ≥ 1%</li> </ul>	<ul> <li>common chemical name(s)</li> <li>generic name(s)</li> <li>synonyms</li> <li>CAS number(s)</li> <li>components or impurities contributing to the hazard (name, concentration)</li> </ul>	<ul> <li>Chemical and common name of ingredients contributing to known hazards</li> <li>For untested mixtures, the chemical &amp; common name of ingredients at 1% or more that present a health hazard and those that present a physical hazard in the mixture</li> <li>Ingredients at 0.1% or greater, if carcinogens</li> </ul>

	MSDS Comparison				
MSDS Sections	GHS SDS <sup>1</sup>	ISO MSDS <sup>2</sup>	ANSI MSDS <sup>3</sup>	OSHA MSDS <sup>4</sup>	
4. First-aid measures	<ul> <li>Description of necessary measures, subdivided according to the different routes of exposure, i.e., inhalation, skin and eye contact and ingestion.</li> <li>Most important symptoms/effects, acute and delayed.</li> <li>Indication of immediate medical attention and special treatment needed, if necessary.</li> </ul>	<ul> <li>Description of necessary measures, subdivided according to the different routes of exposure, i.e., inhalation, skin and eye contact and ingestion.</li> <li>Most important symptoms/effects, acute and delayed.</li> <li>Indication of immediate medical attention and special treatment needed, if necessary.</li> </ul>	<ul> <li>first aid procedures by route of exposure, i.e., inhalation, skin contact, eye contact, ingestion</li> <li>important symptoms and effects useful for diagnostic treatment</li> <li>antidotes</li> <li>notes to a physician</li> </ul>	<ul> <li>emergency &amp; first aid procedures</li> </ul>	
5. Firefighting measures	<ul> <li>Suitable (and unsuitable) extinguishing media.</li> <li>Specific hazards arising from the chemical (e.g., nature of any hazardous combustion products).</li> <li>Special protective equipment and precautions for fire-fighters.</li> </ul>	<ul> <li>Suitable (and unsuitable) extinguishing media.</li> <li>Specific hazards arising from the chemical (e.g., nature of any hazardous combustion products).</li> <li>Special protective equipment and precautions for firefighters.</li> </ul>	<ul> <li>Qualitative flammable and reactivity properties</li> <li>suitable extinguishing media</li> <li>unsuitable extinguishing media</li> <li>Guidance to firefighters</li> <li>Specific hazards arising from the chemical</li> <li>Protective equipment and precautions for firefighters</li> </ul>	<ul> <li>generally applicable control measures</li> <li>flammable property information such as flashpoint</li> <li>physical hazards including the potential for fire, explosion, and reactivity</li> </ul>	
6. Accidental release measures	<ul> <li>Personal precautions, protective equipment and emergency procedures.</li> <li>Environmental precautions.</li> <li>Methods and materials for containment and cleaning up.</li> </ul>	<ul> <li>Personal precautions, protective equipment and emergency procedures.</li> <li>Environmental precautions.</li> <li>Methods and materials for containment and cleaning up.</li> </ul>	<ul> <li>Clean-up technique</li> <li>Personal Precautions</li> <li>Environmental Precautions</li> <li>containment technique</li> <li>regulatory information</li> </ul>	<ul> <li>procedures for clean up of spills and leaks</li> </ul>	

	MSDS Comparison				
<b>MSDS Sections</b>	GHS SDS <sup>1</sup>	ISO MSDS <sup>2</sup>	ANSI MSDS <sup>3</sup>	OSHA MSDS <sup>4</sup>	
7. Handling and storage	<ul> <li>Precautions for safe handling.</li> <li>Conditions for safe storage, including any incompatibilities.</li> </ul>	<ul> <li>Precautions for safe handling.</li> <li>Conditions for safe storage, including any incompatibilities.</li> </ul>	<ul> <li><i>handling</i></li> <li>measures to prevent exposure and release, prevent fire or explosion and ensure precautions for safe handling</li> <li><i>storage</i></li> <li>storage conditions and technical measures for safe storage</li> <li>incompatibilities</li> <li>suitable/non suitable packaging material</li> </ul>	<ul> <li>Precautions for safe handling &amp; use, including appropriate hygenic practices.</li> </ul>	
8. Exposure controls/ personal protection	<ul> <li>Control parameters (e.g., occupational exposure limit values or biological limit values).</li> <li>Appropriate engineering controls.</li> <li>Individual protection measures, such as personal protective equipment.</li> </ul>	<ul> <li>Control parameters (e.g., occupational exposure limit values or biological limit values).</li> <li>Appropriate engineering controls.</li> <li>Individual protection measures, such as personal protective equipment.</li> </ul>	<ul> <li>exposure guidelines (limit values)</li> <li>engineering controls to minimize hazards</li> <li>personal protective equipment (respiratory, hand, eye, skin and body protection)</li> <li>General Hygiene Considerations</li> </ul>	<ul> <li>General applicable control measures</li> <li>appropriate engineering controls and work practices</li> <li>protective measures during maintenance &amp; repair</li> <li>personal protective equipment</li> <li>permissible exposure levels, threshold limit values, listed by OSHA, ACGIH, or established company limits.</li> </ul>	

MSDS Comparison					
MSDS Sections	GHS SDS <sup>1</sup>	ISO MSDS <sup>2</sup>	ANSI MSDS <sup>3</sup>	OSHA MSDS <sup>4</sup>	
9. Physical and chemical properties	<ul> <li>Appearance (physical state, colour, etc.)</li> <li>Odour</li> <li>Odour threshold</li> <li>pH</li> <li>melting point/freezing point</li> <li>initial boiling point and boiling range</li> <li>flash point: <ul> <li>evaporation rate</li> <li>flammability (solid, gas)</li> <li>upper/lower flammability or explosive limits</li> <li>vapour pressure</li> <li>vapour density</li> <li>relative density:</li> <li>solubility(ies)</li> <li>partition coefficient: <i>n</i>-octanol/water</li> <li>auto-ignition temperature</li> <li>decomposition temperature</li> </ul> </li> </ul>	<ul> <li>Appearance (physical state, colour, etc.)</li> <li>Odour</li> <li>Odour threshold</li> <li>pH</li> <li>melting point/freezing point</li> <li>initial boiling point and boiling range</li> <li>flash point:</li> <li>evaporation rate</li> <li>flammability (solid, gas)</li> <li>upper/lower flammability or explosive limits</li> <li>vapour pressure</li> <li>vapour density</li> <li>relative density:</li> <li>solubility(ies)</li> <li>partition coefficient: <i>n</i>-octanol/water:</li> <li>auto-ignition temperature</li> <li>decomposition temperature</li> </ul>	<ul> <li>appearance (color, physical form, shape)</li> <li>odor/odor threshold</li> <li>physical state</li> <li>pH</li> <li>melting/freezing point(specify which)</li> <li>initial boiling point and boiling range</li> <li>flash point</li> <li>evaporation rate</li> <li>flammability (solid, gas)</li> <li>upper/lower flammability or explosive limits</li> <li>vapor pressure</li> <li>vapor density</li> <li>specific gravity or relative density</li> <li>solubility(ies) (specify solvent, e.g., water)</li> <li>partition coefficient: n- octanol/water</li> <li>auto-ignition temperature</li> <li>other relevant data</li> </ul>	<ul> <li>characteristics of hazardous chemicals such as vapor pressure &amp; density.</li> <li>physical hazards including the potential for fire, explosion, and reactivity.</li> </ul>	
10. Stability and reactivity	<ul> <li>Chemical stability.</li> <li>Possibility of hazardous reactions.</li> <li>Conditions to avoid (e.g., static discharge, shock or vibration).</li> <li>Incompatible materials,</li> <li>Hazardous decomposition products.</li> </ul>	<ul> <li>Chemical stability.</li> <li>Possibility of hazardous reactions.</li> <li>Conditions to avoid (e.g., static discharge, shock or vibration).</li> <li>Incompatible materials.</li> <li>Hazardous decomposition products.</li> </ul>	<ul> <li>Physical hazards</li> <li>chemical stability</li> <li>conditions to avoid</li> <li>Incompatible Materials</li> <li>hazardous decomposition products</li> <li>Possibility of Hazardous Reactions</li> </ul>	<ul> <li>organic peroxides, pyrophoric, unstable # (reactive), or water- reactive hazards</li> <li>physical hazards, including reactivity and hazardous polymerization</li> </ul>	
	MSDS Comparison				
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MSDS Sections	GHS SDS <sup>1</sup>	ISO MSDS <sup>2</sup>	ANSI MSDS <sup>3</sup>	OSHA MSDS <sup>4</sup>	
11. Toxicological information	<ul> <li>Concise but complete and comprehensible description of the various toxicological (health) effects and the available data used to identify those effects, including:</li> <li>Information on the likely routes of exposure (inhalation, ingestion, skin and eye contact);</li> <li>Symptoms related to the physical, chemical and toxicological characteristics;</li> <li>Delayed and immediate effects and also chronic effects from short- and long-term exposure;.</li> <li>Numerical measures of toxicity (such as acute toxicity estimates).</li> </ul>	<ul> <li>Concise but complete and comprehensible description of the various toxicological (health) effects and the available data used to identify those effects, including:</li> <li>Information on the likely routes of exposure (inhalation, ingestion, skin and eye contact);</li> <li>Symptoms related to the physical, chemical and toxicological characteristics;</li> <li>Delayed and immediate effects and also chronic effects from short- and long-term exposure;</li> <li>Numerical measures of toxicity (such as acute toxicity estimates).</li> </ul>	<ul> <li>Toxicological information: human, animal, and in vitro data, SAR</li> <li>acute dose effects: single/ short-term exposures. (e.g., LD50, LC50).</li> <li>Repeated dose effects: (e.g., NOAEL)</li> <li>Irritation/Corrosivity</li> <li>Sensitization (skin and respiratory)</li> <li>Carcinogenicity</li> <li>Neurological effects</li> <li>Genetic effects (e.g., mutagenicity)</li> <li>Reproductive effects</li> <li>Developmental effects</li> <li>Target organ effects</li> </ul>	<ul> <li>See also Section 2 [health hazards Including acute and chronic effects, listing target organs or systems</li> <li>signs &amp; symptoms of exposure</li> <li>primary routes of exposure</li> <li>if listed as a carcinogen by OSHA, IARC, NTP]</li> </ul>	
12. Ecological information	<ul> <li>Ecotoxicity (aquatic and terrestrial, where available).</li> <li>Persistence and degradability</li> <li>Bioaccumulative potential</li> <li>Mobility in soil</li> <li>Other adverse effects</li> </ul>	<ul> <li>Ecotoxicity (aquatic and terrestrial, where available).</li> <li>Persistence and degradability</li> <li>Bioaccumulative potential</li> <li>Mobility in soil</li> <li>Other adverse effects</li> </ul>	<ul> <li>ecotoxicity acute and longterm (fish, invertebrates)</li> <li>persistence / degradability</li> <li>bioaccumulation / bioconcentration</li> <li>mobilty: air, soil, water</li> <li>Other adverse effects</li> </ul>	- No present requirements.	

	MSDS Comparison				
MSDS Sections	GHS SDS <sup>1</sup>	ISO MSDS <sup>2</sup>	ANSI MSDS <sup>3</sup>	OSHA MSDS <sup>4</sup>	
13. Disposal considerations	- Description of waste residues and information on their safe handling and methods of disposal, including any contaminated packaging.	- Description of waste residues and information on their safe handling and methods of disposal, including any contaminated packaging.	<ul> <li>safe and environmentally preferred waste management of the material and/or its container</li> <li>classification under applicable law</li> </ul>	<ul> <li>No present requirements,</li> <li>See section 7,</li> </ul>	
14. Transport information	<ul> <li>UN number.</li> <li>UN Proper shipping name.</li> <li>Transport Hazard class(es).</li> <li>Packing group, if applicable.</li> <li>Marine pollutant (Y/N).</li> <li>Special precautions which a user needs to be aware of or needs to comply with in connection with transport or conveyance either within or outside their premises.</li> </ul>	<ul> <li>UN number.</li> <li>UN Proper shipping name.</li> <li>Transport Hazard class(es).</li> <li>Packing group, if applicable.</li> <li>Marine pollutant (Y/N).</li> <li>Special precautions which a user needs to be aware of or needs to comply with in connection with transport or conveyance either within or outside their premises.</li> </ul>	<ul> <li>proper shipping name</li> <li>hazard class(es)</li> <li>identification number</li> <li>packing group</li> <li>hazardous substances</li> <li>marine pollutants (Y/N)</li> <li>IMDG classification</li> <li>TDG classification</li> <li>ICAO/IATA classification</li> <li>RID/ADR classification</li> </ul>	- No present requirements,	
15. Regulatory information	<ul> <li>Safety, health and environmental regulations specific for the product in question.</li> </ul>	- Safety, health and environmental regulations specific for the product in question.	<ul> <li>U.S .federal regulations</li> <li>international regulations</li> <li>U.S. state regulations</li> </ul>	- No present requirements.	
16. Other information	- Other information including information on preparation and revision of the SDS.	- Other information including information on preparation and revision of the SDS.	<ul> <li>label text</li> <li>hazard rating and rating system</li> <li>information on preparation and revision of safety data sheet</li> <li>Key/legend</li> </ul>	- Date of preparation of MSDS or date of last change	

1. Globally Harmonized System of Classification and Labelling of Chemicals (GHS), United Nations, 2005.

2. ISO 11014-1:2003 DRAFT Safety Data Sheet for Chemical Products.

3. American National Standard for Hazardous Industrial Chemicals-MSDS Preparation (ANSI Z-400.1-2004).

4. U.S. DOL, OSHA, 29 CFR 1910.1200, HAZCOM.

**Appendix B** 

# MSDS Examples (Fictional Products)

- **B-1: Bondit**
- **B-2:** Chemical Stuff

# **Appendix B-1**

# Bondit

(GHS MSDS Example)

## Appendix B-1 MSDS for **BONDIT**

# 1. Identification

Name of the product: Bondit

Recommended use: General adhesive.

**Producer:** GHS Ltd., UK – London, SE, Southwarkbridge 1

**Telephone no.** +44 171717 555.555 5, **Emergency no.** +44 171717 333 333 3

# 2. Hazard(s) identification

<b>Classification:</b>	Flammable liquid, Category 2	
	Eye irritation, Category 2A	
	Hazardous to the aquatic environment, Acute Category 3	

#### Labelling:

Symbol: Signal word:	Flame, Exclamation mark Danger
Hazard statement:	Highly flammable liquid and vapour. Causes severe eye irritation. Harmful to aquatic life.
Precautionary statements:	Keep container tightly closed. Keep away from heat/sparks/open flame. – No smoking. Wear protective gloves and eye/face protection. Ground/Bond container and receiving equipment. Use explosion-proof electrical/ventilating/lighting/ equipment. Take precautionary measures against static discharge. Use only non-sparking tools. Store in cool/well-ventilated place. Avoid release to the environment.

# 3. Composition / Information on ingredients

Chemical identity: Component A 70-80% Common name: Solvent A Numbers of identity: CAS-Nr.:111111-11-1 Impurities: None

Chemical identity: Component C 20-25% Common name: Not applicable Numbers of identity: CAS-Nr.: 44444-44-4 Impurities: none

# 4. First-aid measures

#### Inhalation:

Remove person to fresh air. If respiratory irritation, dizziness, nausea, or unconsciousness occurs, seek immediate medical assistance. If breathing has stopped, give artificial respiration.

#### Skin contact:

Wash the contaminated area with soap and water. Remove contaminated clothing and wash before reuse. If irritation develops, get medical attention.

#### Eye contact:

Hold eyelids apart and flush eyes with plenty of water for at least 15 minutes. Get medical attention.

#### **Ingestion:**

If swallowed, do NOT induce vomiting. Seek immediate medical attention.

# 5. Firefighting measures

**Suitable extinguishing media:** Foam, extinguishing powder, carbon dioxide, water fog. In case of fire, cool endangered containers with water fog.

Unsuitable extinguishing media: High pressure water jet.

Specific hazards in case of fire: None are known.

**Special protective equipment and precaution for fire fighters:** For fires in enclosed areas, wear self-contained breathing apparatus. Do not inhale combustion gases.

### 6. Accidental release measures

#### **Personal precautions:**

Depending on extent of release, consider the need for fire fighters/emergency responders with adequate personal protective equipment for cleaning up.

Do not eat, drink or smoke while cleaning up. Use a self-contained respirator, a mask with filter (type A class 3) or a filtering mask (e.g., EN 405). Wear protective clothing, safety glasses and impervious gloves (e.g., neoprene gloves). Ensure adequate ventilation. Avoid all sources of ignition, hot surfaces and open flames (see also Section 7).

#### **Environmental precautions:**

Prevent spills from entering storm sewers or drains and contact with soil.

#### Methods and materials for containment and cleaning up:

Eliminate all ignition sources. Runoff may create fire or explosion hazard in sewer system. Absorb on fire retardant, liquid-absorbing material (treated sawdust, diatomaceous earth, sand). Shovel up and dispose of at an appropriate waste disposal facility in accordance with current applicable laws and regulations, and product characteristics at time of disposal (see also Section 13).

# 7. Handling and storage

#### **Precautions for safe handling:**

Avoid contact with eyes. Avoid prolonged repeated skin contact and breathing mists/vapours.

Use in well-ventilated area away from all ignition sources. Switch off all electrical devices such as parabolic heaters, hotplates, storage heaters etc. in good time for them to have cooled down before commencing work. Do not smoke; do not weld. Do not empty waste into sanitary drains. Take measures to prevent the build up of electrostatic charge.

#### Conditions for safe storage, including incompatibilities:

Storage containers must be grounded and bonded. Store away from all ignition sources in a cool area equipped with an automatic sprinkling system. Ensure adequate ventilation. Store at temperatures between +5 and  $+50^{\circ}$ C. Store only in the original container.

# 8. Exposure controls / personal protection

#### Information on the system design:

Draw off vapours directly at the point of generation and exhaust from the work area. In the case of regular work, provide bench-mounted extraction equipment.

#### **Exposure Limits:**

		Т	WA	ST	EL
Component Name (CAS-No.)	Reference	ppm	mg/m3	ppm	mg/m3
	UK OEL	500	1200		
Component C (4444-44-4)	German MAK	200	950		

#### Ventilation:

Use in well-ventilated area with local exhaust.

#### **Respiratory protection:**

Approved respiratory equipment must be used when airborne concentrations are unknown or exceed the exposure limits. When processing large amounts, use a light duty construction compressed air line breathing apparatus (e.g., in accordance with EN1835), a mask with filter (type A class 3, colour brown) or a filtering half mask (e.g., in accordance with EN 405) when there is inadequate ventilation.

#### **Eye protection:**

Safety glasses with side shields or chemical goggles must be worn.

#### Skin protection:

If prolonged or repeated skin contact is likely, neoprene gloves should be worn. Good personal hygiene practices should always be followed.

# 9. Physical and chemical properties

**Physical state:** Liquid **Colour:** Colourless, transparent **Odour:** Solvent. ester-like **Odour threshold:** Not available pH-value: Not applicable Melting point: Not available **Freezing Point:** Not available 56°C **Initial boiling point:** Flash point: - 22°C DIN 51755 **Evaporation rate:** Not available Flammability (solid, gas): Not applicable

Explosion limits: Vapour pressure:		
Vapour density:	Not available	
<b>Relative density:</b>	0.89 g/cm3 at 20°C	
Solubility:	Partially soluble in water at 20°C	
Partition coefficient:	Log Kow = 3.3	
Auto-ignition temperature:	Not available	
<b>Decomposition temperature:</b>	Not available	

# 10. Stability and reactivity

Chemical stability: No decomposition, if used according to specifications.
Possibility of hazardous reactions: None are known.
Conditions to avoid: Heat, sparks, flame and build up of static electricity.
Materials to avoid: Halogens, strong acids, alkalies and ozidizers.
Hazardous decomposition products: None are known.

# **11.** Toxicological information

Acute Toxicity:			
Test	Results	Basis	
Oral Toxicity (Rats)	Not Classified	Based on Ingredients	
Dermal Toxicity (Rats)	Not Classified	Product Test Data	
Inhalation Toxicity, Vapor (Rats)	Not Classified	Based on Testing of Similar Materials	
Eye Irritation (Rabbits)	Eye Irritant Category 2A	Based on Testing of Similar Materials	
Dermal Irritation (Rabbits)	Not Classified	Product Test Data	

#### Summary Comments: May cause severe eye irritation like ocular lesions, which are reversible.

#### Subchronic/Chronic Toxicity:

Test	Results	Comments
	Not Classified: Negative response in Bueller, guinea pig test. 0% animals considered positive.	Product Test Data

**Summary Comments**: Component A may have a drying effect on the skin; frequent or prolonged contact may cause flaking or cracking of the skin.

# **12.** Ecological information

**Persistence and degradability:** The total of the organic components contained in the product is not classified as "readily biodegradable" (OECD-301 A-F). However, this product is expected to be inherently biodegradable.

#### **Bio-accumulative potential;**

There is no evidence to suggest bioaccumulation will occur.

**Mobility:** Accidental spillage may lead to penetration in the soil and groundwater. However, there is no evidence that this would cause adverse ecological effects.

#### **Aquatic Toxicity:**

Test	Results Comments	
Acute Toxicity	Acute Category 3: 96 hr. $LC_{50} = 65 \text{ mg/L}$	Product Test Data

# 13. Disposal considerations

#### Waste Disposal:

Product is suitable for burning in an enclosed, controlled burner for fuel value or disposal by supervised incineration. Such burning may be limited by local regulation. The product is suitable for processing at an appropriate government waste disposal facility. Use of these methods is subject to user compliance with applicable laws and regulations and consideration of product characteristics at time of disposal.

Recommended European waste code (EWC): 080406

# **14.** Transport information

UN-number: 1993 UN proper shipping name: Flammable Liquid, N.O.S. (Contains Component C) Transport hazard class: 3 Packing group: II Marine Pollutant: No

# **15. Regulatory information**

**Inventory Status:** All components are on TSCA, EINECS/ELINCS, AICS, and DSL.

#### German:

Regulations governing combustible liquids (German-VbF) class: AI

German water endangering class (WGK) = 1, slightly water-endangering product (manufacturer classification.)

#### **Australian Regulations:**

AS 1940 Class: PGII Poisons Schedule: S5

#### **U.S. Regulations:**

#### U.S. Superfund Amendments and Reauthorization Act (SARA) Title III:

#### SARA (311/312) HAZARD CATEGORIES: FIRE, ACUTE

SARA 313: This product contains the following SARA 313 Toxic Release Chemicals.

Chemical Name	CAS Number	<b>Concentration</b>
Component A	111111-11-1	70-80%
Component C	4444-44-4	20-25%

The following product components are cited on the lists below:

Chemical Name	CAS Number	List Citations
Component A	111111-11-1	NJ RTK, TSCA 12(b)
Component C	4444-44-4	Prop. 65, NJ RTK

# **16. Other information**

#### Abbreviations and acronyms:

UK OES = United Kingdom Occupational Exposure Standards German MAK = Germany Maximum Allowable Concentration

#### MSDS Prepration date: July 1, 2005

The information contained herein is accurate to the best of our knowledge. My Company makes no warranty of any kind, express or implied, concerning the safe use of this material in your process or in combination with other substances.

**Appendix B-2** 

# **Chemical Stuff**

(GHS MSDS Example)

# Appendix B-2 MSDS for Chemical Stuff

#### GHS SAFETY DATA SHEET

### 1. Identification

Chemical Stuff
Methyltoxy Solution
000-00-0
Organic Synthesis
My Company
My Street, Mytown, TX 00000

General Information: 7	13-000-0000
------------------------	-------------

Transportation Emergency Number: CHEMTREC: 800-424-9300

#### 2. Hazards Identification

#### **GHS Classification:**

Health	Environmental	Physical
Acute Toxicity – Category 2 (inhalation), Category 3 (oral/dermal) Eye Corrosion - Category 1 Skin Corrosion – Category 1 Skin Sensitization – Category 1 Mutagenicity – Category 2 Carcinogenicity – Category 1B Reproductive/Developmental – Category 2 Target Organ Toxicity (Repeated) – Category 2	Aquatic Toxicity – Acute 2	Flammable Liquid – Category 2

#### GHS Label:

Symbols: flame, skull and crossbones, corrosion, health hazard		
Hazard Statements	Precautionary Statements	
DANGER!	Do not eat, drink or use tobacco when using this	
Highly Flammable Liquid and Vapor.	product.	
Fatal if inhaled.	Do not breathe mist/vapors.	
Causes severe skin burns and eye damage.	Keep container tightly closed.	
May cause allergic skin reaction.	Keep away from heat/sparks/open flame. – No	
Toxic if swallowed and in contact with skin	smoking.	
May cause cancer.	Wear respiratory protection, protective gloves and	
Suspected of damaging the unborn child.	eye/face protection.	
Suspected of causing genetic defects.	Use only in a well-ventilated area.	
May cause damage to cardiovascular, respiratory,	Take precautionary measures against static	
nervous, and gastrointestinal systems and liver and	discharge.	
blood through prolonged or repeated exposure.	Use only non-sparking tools.	
Toxic to aquatic life.	Store container tightly closed in cool/well-ventilated	
	place.	
	Wash thoroughly after handling.	

#### 3. Composition / Information on Ingredients

Component	CAS Number	Weight %
Methyltoxy	000-00-0	80
(See Section 8 for Exposure Limits)		

#### 4. First Aid Measures

**Eye:** Eye irritation. Flush immediately with large amounts of water for at least 15 minutes. Eyelids should be held away from the eyeball to ensure thorough rinsing. Get immediate medical attention.

**Skin:** Itching or burning of the skin. Immediately flush the skin with plenty of water while removing contaminated clothing and shoes. Get immediate medical attention. Wash contaminated clothing before reuse.

**Inhalation:** Nasal irritation, headache, dizziness, nausea, vomiting, heart palpitations, breathing difficulty, cyanosis, tremors, weakness, red flushing of face, irritability. Remove exposed person from source of exposure to fresh air. If not breathing, clear airway and start cardiopulmonary resuscitation (CPR). Avoid mouth-to-mouth resuscitation.

**Ingestion:** Get immediate medical attention. Do not induce vomiting unless directed by medical personnel.

#### 5. Fire Fighting Measures

**Suitable Extinguishing Media:** Use dry chemical, foam, or carbon dioxide to extinguish fire. Water may be ineffective but should be used to cool fire-exposed containers, structures and to protect personnel. Use water to dilute spills and to flush them away from sources of ignition.

**Fire Fighting Procedures**: Do not flush down sewers or other drainage systems. Exposed firefighters must wear NIOSH-approved positive pressure self-contained breathing apparatus with full-face mask and full protective clothing.

**Unusual Fire and Explosion Hazards:** Dangerous when exposed to heat or flame. Will form flammable or explosive mixtures with air at room temperature. Vapor or gas may spread to distant ignition sources and flash back. Vapors or gas may accumulate in low areas. Runoff to sewer may cause fire or explosion hazard. Containers may explode in heat of fire. Vapors may concentrate in confined areas. Liquid will float and may reignite on the surface of water.

**Combustion Products:** Irritating or toxic substances may be emitted upon thermal decomposition. Thermal decomposition products may include oxides of carbon and nitrogen.

#### 6: Accidental Release Measures

Keep unnecessary people away; isolate hazard area and deny entry. Stay upwind; keep out of low areas. (Also see Section 8).

Vapor protective clothing should be worn for spills and leaks. Shut off ignition sources; no flares, smoking or flames in hazard area. Small spills: Take up with sand or other noncombustible absorbent material and place into containers for later disposal. Large spills: Dike far ahead of liquid spill for later disposal.

Do not flush to sewer or waterways. Prevent release to the environment if possible. Refer to Section 15 for spill/release reporting information.

#### 7. Handling and Storage

#### Handling

Do not get in eyes, on skin or on clothing. Do not breathe vapors or mists. Keep container closed. Use only with adequate ventilation. Use good personal hygiene practices. Wash hands before eating, drinking, smoking. Remove contaminated clothing and clean before re-use. Destroy contaminated belts and shoes and other items that cannot be decontaminated.

Keep away from heat and flame. Keep operating temperatures below ignition temperatures at all times. Use non-sparking tools.

#### Storage

Store in tightly closed containers in cool, dry, well-ventilated area away from heat, sources of ignition and incompatibles. Ground lines and equipment used during transfer to reduce the possibility of static spark-initiated fire or explosion. Store at ambient or lower temperature. Store out of direct sunlight. Keep containers tightly closed and upright when not in use. Protect against physical damage.

Empty containers may contain toxic, flammable and explosive residue or vapors. Do not cut, grind, drill, or weld on or near containers unless precautions are taken against these hazards.

#### 8. Exposure Controls / Personal Protection

#### **Exposure Limits**

Component	OS	НА	
-	TWA	STEL	
Methyltoxy	3 ppm	C 15 ppm	
	(skin)	(15 min.)	

**Engineering Controls:** Local exhaust ventilation may be necessary to control air contaminants to their exposure limits. The use of local ventilation is recommended to control emissions near the source. Provide mechanical ventilation for confined spaces. Use explosion-proof ventilation equipment.

#### **Personal Protective Equipment (PPE)**

**Eye Protection:** Wear chemical safety goggles and face shield. Have eye-wash stations available where eye contact can occur.

**Skin Protection:** Avoid skin contact. Wear gloves impervious to conditions of use. Additional protection may be necessary to prevent skin contact including use of apron, face shield, boots or full body protection. A safety shower should be located in the work area. Recommended protective materials include: Butyl rubber and for limited contact Teflon. **Respiratory Protection:** If exposure limits are exceeded, NIOSH approved respiratory protection should be worn. A NIOSH approved respirator for organic vapors is generally acceptable for concentrations up to 10 times the PEL. For higher concentrations, unknown concentrations and for oxygen deficient atmospheres, use a NIOSH approved air-supplied respirator. Engineering controls are the preferred means for controlling chemical exposures. Respiratory protection may be needed for non-routine or emergency situations. Respiratory protection must be provided in accordance with OSHA 29 CFR 1910.134.

#### 9. Physical and Chemical Properties

Lower Flammability Limit: >3.00%

Specific Gravity: 0.82g/ml @ 20°C

% Volatile: 100

Viscosity: 0.3 cP @ 25°C

Acetate =1)

**Upper Flammability Limit:** <15.00%

Evaporation Rate (Water=1): 5(Butyl

**Octanol/Water Partition Coefficient:** 

Flashpoint: 2°C (35°F) Autoignition Temperature: 480°C (896°F) Boiling Point: 77°C (170.6°F) @ 760 mm Hg Melting Point: -82°C Vapor Pressure: 100.0 mm Hg @ 23°C

Vapor Density(Air=1): 1.7; air = 1 % Solubility in Water: 10 @ 20°C

Pour Point: NAlog K<sub>ow</sub>: 0.5Pour Point: NApH: 7, 8% aqueous solutionMolecular Formula: MixtureMolecular Weight: MixtureOdor/Appearance: Clear, colorless liquid with mild, pungent odor.

# 10. Stability and Reactivity

**Stability/Incompatibility:** Incompatible with ammonia, amines, bromine, strong bases and strong acids.

Hazardous Reactions/Decomposition Products: Thermal decomposition products may include oxides of carbon and nitrogen.

#### **11. Toxicological Information**

**Signs and Symptoms of Overexposure:** Eye and nasal irritation, headache, dizziness, nausea, vomiting, heart palpitations, difficulty breathing, cyanosis, tremors, weakness, itching or burning of the skin.

#### Acute Effects:

**Eye Contact:** may cause severe conjunctival irritation and corneal damage.

**Skin Contact:** may cause reddening, blistering or burns with permanent damage. Harmful if absorbed through the skin. May cause allergic skin reaction.

Inhalation: may cause severe irritation with possible lung damage (pulmonary edema).

**Ingestion:** may cause severe gastrointestinal burns.

**Target Organ Effects:** May cause gastrointestinal (oral), respiratory tract, nervous system and blood effects based on experimental animal data. May cause cardiovascular system and liver effects.

**Chronic Effects:** based on experimental animal data, may cause changes to genetic material; adverse effects on the developing fetus or on reproduction at doses that were toxic to the mother. Methyltoxy is classified by IARC as group 2B and by NTP as reasonably anticipated to be a human carcinogen. OSHA regulates Methyltoxy as a potential carcinogen.

**Medical Conditions Aggravated by Exposure:** preexisting diseases of the respiratory tract, nervous system, cardiovascular system, liver or gastrointestinal tract.

#### **Acute Toxicity Values**

Oral  $LD_{50}$  (Rat) = 100 mg/kg Dermal  $LD_{50}$  (Rabbit) = 225-300 mg/kg Inhalation  $LC_{50}$  (Rat) = 200 ppm/4 hr., 1100 ppm vapor/1 hr

#### **12. Ecological Information**

 $LC_{50}$  (Fathead Minnows) = 9 mg/L/96 hr. EC<sub>50</sub> (Daphnia) = 8.6 mg/L/48 hr.

Bioaccumulation is not expected to be significant. This product is readily biodegradable.

#### **13. Disposal Considerations**

As sold, this product, when discarded or disposed of, is a hazardous waste according to Federal regulations (40 CFR 261). It is listed as Hazardous Waste Number Z000, listed due to its toxicity. The transportation, storage, treatment and disposal of this waste material must be conducted in compliance with 40 CFR 262, 263, 264, 268 and 270. Disposal can occur only in properly permitted facilities. Refer to state and local requirements for any additional requirements, as these may be different from Federal laws and regulations. Chemical additions, processing or otherwise altering this material may make waste management information presented in the MSDS incomplete, inaccurate or otherwise inappropriate.

#### **14. Transport Information**

U.S. Department of Transportation (DOT) Proper Shipping Name: Methyltoxy Hazard Class: 3, 6.1 UN/NA Number: UN0000 Packing Group: PG 2 Labels Required: Flammable Liquid and Toxic

International Maritime Organization (IMDG) Proper Shipping Name: Methyltoxy Hazard Class: 3 Subsidiary 6.1 UN/NA Number: UN0000

#### Packing Group: PG 2 Labels Required: Flammable Liquid and Toxic

#### **15. Regulatory Information**

#### U.S. Federal Regulations

#### Comprehensive Environmental Response and Liability Act of 1980 (CERCLA):

The reportable quantity (RQ) for this material is 1000 pounds. If appropriate, immediately report to the National Response Center (800/424-8802) as required by U.S. Federal Law. Also contact appropriate state and local regulatory agencies.

**Toxic Substances Control Act (TSCA):** All components of this product are included on the TSCA inventory.

**Clean Water Act (CWA):** Methyltoxy is a hazardous substance under the Clean Water Act. Consult Federal, State and local regulations for specific requirements.

**Clean Air Act (CAA):** Methyltoxy is a hazardous substance under the Clean Air Act. Consult Federal, State and local regulations for specific requirements.

Superfund Amendments and Reauthorization Act (SARA) Title III Information:

SARA Section 311/312 (40 CFR 370) Hazard Categories:			
Immediate Hazard: X	Delayed Hazard: X	Fire Hazard: X	
Pressure Hazard:	Reactivity Hazard:		

This product contains the following toxic chemical(s) subject to reporting<br/>requirements of SARA Section 313 (40 CFR 372)<br/>Component:Maximum %CAS Number:Maximum %

		Muximum /
Methyltoxy	000-00-0	80

#### **State Regulations**

**California:** This product contains the following chemicals(s) known to the State of California to cause cancer, birth defects or reproductive harm:

Component:	CAS NUMBER:	<u>%</u>
Methyltoxy	000-00-0	80

#### International Regulations

**Canadian Environmental Protection Act:** All of the components of this product are included on the Canadian Domestic Substances list (DSL).

#### Canadian Workplace Hazardous Materials Information System (WHMIS):

Class B-2 Flammable Liquid

Class D-1-B Toxic

Class D-2-A Carcinogen Class D-2-B Chronic Toxin

Class E Corrosive

This product has been classified in accordance with the hazard criteria of the Controlled Products Regulations and the MSDS contains all the information required by the Controlled Products Regulations.

**European Inventory of Existing Chemicals (EINECS):** All of the components of this product are included on EINECS.

**EU Classification**: F Highly Flammable; T Toxic; N Dangerous to the Environment **EU Risk (R) and Safety (S) Phrases**:

R11: Highly flammable.
R23/24/25: Toxic by inhalation, in contact with skin and if swallowed.
R37/38: Irritating to respiratory system and skin.
R41: Risk of serious damage to eyes.
R43: May cause sensitization by skin contact.
R45: May cause cancer.
R51/53: Toxic to aquatic organisms, may cause long-term adverse effects in the aquatic environment.
S53: Avoid exposure - obtain special instructions before use.
S16: Keep away from sources of ignition - No Smoking.
S45: In case of accident or if you feel unwell, seek medical advice immediately (show the label where possible).
S9: Keep container in a well-ventilated place.
S36/37: Wear suitable protective clothing and gloves.
S57: Use appropriate container to avoid environmental contamination.

#### 16. Other Information

National Fire Protection Association (NFPA) Ratings: This information is intended solely for the use of individuals trained in the NFPA system.

Health: 3 Flammability: 3 Reactivity: 0

Revision Indicator: New MSDS

**Disclaimer:** The information contained herein is accurate to the best of our knowledge. My Company makes no warranty of any kind, express or implied, concerning the safe use of this material in your process or in combination with other substances.