SAFETY DATA SHEETS

This SDS packet was issued with item: 078240220

The safety data sheets (SDS) in this packet apply to one or more components included in the items listed below. Items listed below may require one or more SDS. Please refer to invoice for specific item number(s).

078073963



Merck Animal Health One Merck Dr. Whitehouse Station, NJ 08889

MATERIAL SAFETY DATA SHEET

Merck Animal Health urges each user or recipient of this MSDS to read the entire data sheet to become aware of the hazards associated with this material.

SECTION 1. IDENTIFICATION OF SUBSTANCE AND CONTACT INFORMATION

MSDS NAME:	Beuthanasia-D Solution
SYNONYM(S):	Beuthanasia-D Special Beuthanasia-D Injection
MSDS NUMBER:	SP000354
EMERGENCY NUMBER(S):	(908) 423-6000 (24/7/36) English Only
	Transportation Emergencies - CANUTEC: (613) 996-6666 (Canada)
INFORMATION:	Animal Health Technical Services: (888) 306-0069 (Canada)
MERCK MSDS HELPLINE:	(800) 770-8878 (US and Canada) (908) 473-3371 (Worldwide) Monday to Friday, 9am to 5pm (US Eastern Time)

SECTION 2. HAZARDS IDENTIFICATION

EMERGENCY OVERVIEW

Solution Clear, Pink Odor unknown Toxic if swallowed. May be toxic by inhalation. May cause allergic reactions in susceptible individuals. Prolonged exposure may cause serious health effects. Causes effects to: central nervous system respiratory system brain cardiovascular system Causes birth defects. May cause effects to: gastrointestinal tract blood immune system liver kidney Harmful to fish and aquatic organisms.

POTENTIAL HEALTH EFFECTS:

SECTION 2. HAZARDS IDENTIFICATION

The following summary is based upon available information about the individual ingredients of the mixture, or of the expected properties of the mixture. Only information about the ingredients that are expected to contribute significantly to the potential health hazard profile of the formulation(s) are presented.

This product is intended to cause euthanasia in dogs upon administration intravenously. Euthanasia is due to cerebral death in conjunction with respiratory arrest and circulatory collapse. Central nervous system depression and hypotension may also occur.

Pentobarbital sodium is a short-acting barbiturate used as a sedative, preanesthetic, and sleeping aid. Barbiturates may be habit-forming, and tolerance, psychological, or physical dependence may occur especially following prolonged use of high concentrations. Barbiturates are central nervous system and respiratory depressants. Effects that may be seen following acute exposure include slurred speech, confusion, poor judgement, irritability, insomnia, or incoordination. Effects that may be seen following exposure to high concentrations include severe confusion, decrease or loss of reflexes, severe drowsiness, fever, hypothermia, shortness of breath or troubled breathing, slow heartbeat, severe weakness, respiratory depression, pneumonia, congestive heart failure, renal failure, coma, respiratory arrest, or death.

Barbiturates readily cross the placenta following oral administration. Barbiturates have been associated with an increased risk of congenital heart disease, facial abnormalities, and other birth defects; however, no effects have been observed in women exposed to pentobarbital. In addition, newborns that were chronically exposed to bariturates in utero may exhibit withdrawl symptoms such as hyperactivity and tremors.

Phenytoin, often administered as phenytoin sodium, is an anticonvulsant and antiarrhythmic agent. Phenytoin is a central nervous system depressant. Acute effects from exposure may include nausea, vomiting, gastrointestinal pain, loss of appetite, dizziness, staggering, blurred vision, nystagmus (involuntary movement of the eye), drowsiness, pupil dilation, hyperactive tendon reflexes, tremor, increased or decreased activity, hallucinations, confusion, respiratory depression, breathing difficulties, or coma. Hypersensitivity reactions, sometimes fatal, have been reported after chronic therapy. General symptoms of potential reactions include fever, general discomfort, rash, facial swelling, skin redness, lymph node effects, hepatitis, anemia, pharyngitis, diarrhea, anorexia, kidney inflammation, and acute inflammation of the lungs. Phenytoin may also invoke autoimmune dysfunction, swelling of the gums, psychological disorders, or effects on the liver or blood.

Phenytoin freely pases through the placenta. Human teratogenicity (birth defects) has been reported in women who received phenytoin treatment, and phenytoin has been linked to Fetal Hydantoin Syndrome (FHS). Phenytoin is a teratogen in animals.

Propylene glycol is considered to be relatively non-toxic. It is a mild irritant to the eyes and has been reported to irritate the skin. It may cause skin sensitization resulting in allergic contact dermatitis in susceptible individuals. Inhalation exposure to saturated and supersaturated atmospheres of propylene glycol for prolonged periods of time produced no adverse effects. Propylene glycol may cause nervous system depression, acidosis, stupor, and seizures after chronic ingestion.

LISTED CARCINOGENS

INGREDIENT	CAS NUMBER	OSHA	IARC	NTP	ACGIH
Phenytoin Sodium	630-93-3		2B		
Ethyl Alcohol	64-17-5			K	A3

Phenytoin: IARC has classified phenytoin as a Group 2B (possibly carcinogenic to humans).

Ethanol (ethyl alcohol): IARC (International Agency for Research on Cancer) has classified Alcoholic Beverages as Group 1 (indicating in their evaluation that the agent is carcinogenic to humans). However, occupational handling or manufacturer's specified use of this product is not expected to result in relevant exposures.

SECTION 3. COMPOSITION AND INFORMATION ON INGREDIENTS

PRODUCT USE:

Veterinary product

Mixture.

CHEMICAL FORMULA:

The formulation for this product is proprietary information. Only hazardous ingredients in concentrations of 1% or greater and/or carcinogenic ingredients in concentrations of 0.1% or greater are listed in the Chemical Composition table. Active ingredients in any concentration are listed. For additional information about carcinogenic ingredients see Section 2.

CHEMICAL COMPOSITION

INGREDIENT	CAS NUMBER	PERCENT
Pentobarbital Sodium	57-33-0	39
Phenytoin Sodium	630-93-3	5
Propylene Glycol	57-55-6	10-20
Ethyl Alcohol	64-17-5	<10
Benzyl Alcohol	100-51-6	<10

Latest Revision Date: 07-Oct-2011

This MSDS is written to provide health and safety information for individuals who will be handling the final product formulation during research, manufacturing, and distribution. For health and safety information for individual ingredients used during manufacturing, refer to the appropriate MSDS for each ingredient. Refer to the package insert or product label for handling guidance for the consumer.

SECTION 4. FIRST AID MEASURES

INHALATION:	Remove to fresh air. If any trouble breathing, get immediate medical attention. Administer artificial respiration if breathing has ceased. If irritation or symptoms occur or persist, consult a physician.
SKIN CONTACT:	In case of skin contact, while wearing protective gloves, carefully remove any contaminated clothing, including shoes, and wash skin thoroughly with soap and water. If irritation or symptoms occur or persist, consult a physician.
EYE CONTACT:	In case of eye contact, immediately rinse eyes thoroughly with plenty of water. If wearing contact lenses, remove only after initial rinse, and continue rinsing eyes for at least 15 minutes. If irritation occurs or persists, consult a physician.
INGESTION:	DO NOT induce vomiting. Do not attempt to give anything by mouth to a seizing, drowsy or unconscious person. If alert, rinse mouth, drink a glass of water and IMMEDIATELY consult a physician.

SECTION 5. FIRE FIGHTING MEASURES

FLAMMABILITY DATA:

Flash Point:

Not determined (liquids) or not applicable (solids).

SPECIAL FIRE FIGHTING PROCEDURES:

Wear full protective clothing and self-contained breathing apparatus (SCBA).

SUITABLE EXTINGUISHING MEDIA:

Carbon dioxide (CO2), extinguishing powder or water spray.

See Section 9 for Physical and Chemical Properties.

SECTION 6. ACCIDENTAL RELEASE MEASURES

PERSONAL PRECAUTIONS:

Wear appropriate personal protective equipment as specified in Section 8. Keep personnel away from the clean-up area.

SPILL RESPONSE / CLEANUP:

All spills should be handled according to site requirements and based on precautions cited in the MSDS. In the case of liquids, use proper absorbent materials. For laboratories and small-scale operations, incidental spills within a hood or enclosure should be cleaned by using a HEPA filtered vacuum or wet cleaning methods as appropriate. For large dry or liquid spills or those spills outside enclosure or hood, appropriate emergency response personnel should be notified. In manufacturing and large-scale operations, HEPA vacuuming prior to wet mopping or cleaning is required.

See Sections 9 and 10 for additional physical, chemical, and hazard information.

SECTION 7. HANDLING AND STORAGE

HANDLING:

Keep containers adequately sealed during material transfer, transport, or when not in use. Wash face, hands, and any exposed skin after handling. Do not eat, drink, or smoke when using this substance or mixture.

Appropriate handling of this material is dependent on many factors, including physical form, duration and frequency of process or task, and effectiveness of engineering controls. Site-specific risk assessments should be conducted to determine the feasibility and the appropriateness of all exposure control measures. See Section 8 (Exposure Controls) for additional guidance.

STORAGE:

Store in a cool, dry, well ventilated area.

See Section 8 for exposure controls and additional safe handling information.

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SECTION 8. EXPOSURE CONTROLS AND PERSONAL PROTECTION

The following guidance applies to the handling of the active ingredient(s) in this formulation.

OCCUPATIONAL EXPOSURE BAND (OEB):

Pentobarbital: OEB 4: 1-10 mcg/m³. Materials in an OEB 4 category are considered high health hazards. The OEB is range of airborne concentrations expressed as an 8-hour Time Weighted Average (8-hr. TWA) and is intended to be used with Industrial Hygiene Risk Assessment to assist with industrial hygiene sampling and selection of proper controls for worker protection. Consult your site safety and industrial hygiene staff for guidance on handling and control strategies.

Phenytoin Sodium: OEB 3: 10-100 mcg/m³. Materials in an OEB 3 category are considered moderate health hazards. The OEB is a range of airborne concentrations expressed as an 8-hour Time Weighted Average (8-hr. TWA) and is intended to be used with Industrial Hygiene Risk Assessment to assist with industrial hygiene sampling and selection of proper controls for worker protection. Consult your site safety and industrial hygiene staff for guidance on handling and control strategies.

ESTIMATED TARGET AIRBORNE CONCENTRATION (ETAC):

A range of airborne concentrations expressed as an 8-hour Time Weighted Average (8-hr. TWA). The ETAC is intended to be used with Industrial Hygiene Risk Assessment to assist with industrial hygiene sampling and selection of proper controls for worker protection.

ETAC: 10-50 mcg/m³

A range of airborne concentrations expressed as an 8-hour Time Weighted Average (8-hr. TWA). The ETAC is intended to be used with Industrial Hygiene Risk Assessment to assist with industrial hygiene sampling and selection of proper controls for worker protection.

ETAC: 1-10 mcg/m³

HHC/OEG NOTATION(S):

Phenytoin Sodium: This material has a notation of "A" for its ability to cause immediate allergic hypersensitivity reactions or anaphylaxis.

EXPOSURE CONTROLS

The health hazard risks of handling this material are dependent on many factors, including physical form, duration and frequency of process or task, and effectiveness of engineering controls. Site-specific risk assessments should be conducted to determine the feasibility and the appropriateness of all exposure control measures. Exposure controls for normal operating or routine procedures follow a tiered strategy. Engineering controls are the preferred means of long-term or permanent exposure control. If engineering controls are not feasible, appropriate use of personal protective equipment (PPE) may be considered as alternative control measures. Exposure controls for non-routine operations must be evaluated and addressed as part of the site-specific risk assessment.

RECOMMENDED PERSONAL PROTECTIVE EQUIPMENT (PPE):

Respiratory Protection:	Respiratory protective equipment (RPE) may be required for certain laboratory and large-scale manufacturing tasks if potential airborne breathing zone concentrations of substances exceed the relevant exposure limit(s). Workplace risk assessment should be completed before specifying and implementing RPE usage. Potential exposure points and pathways, task duration and frequency, potential employee contact with the substance, and the ability of the substance to be rendered airborne during specific tasks should be evaluated. Initial and ongoing strategies of quantitative exposure measurement should be obtained as required by the workplace risk assessment. All RPE must conform to local and regional specifications for efficacy and performance. Consult your site or corporate health and safety professional for additional guidance.
Skin Protection:	Gloves that provide an appropriate barrier to the skin are recommended if there is potential for contact with this material. Consult your site safety staff for guidance.
Eye Protection:	Safety glasses with side shields. Use of goggles or full face protection may be required based on hazard, potential for contact, or level of exposure. Consult your site safety staff for guidance.
Body Protection:	In small-scale or laboratory operations, lab coats or equivalent protection is required. Disposable Tyvek or other dust impermeable suit should be considered based on procedure or level of exposure. Use of additional PPE such as shoe coverings, gauntlets, hood, or head covering may be necessary. Consult your site safety staff for guidance.
	In large-scale or manufacturing operations, disposable Tyvek or other dust impermeable suit is recommended and based on level of exposure. Use of additional PPE such as shoe coverings, gauntlets, hood, or head covering may be necessary. Consult your site safety staff for guidance.

EXPOSURE LIMIT VALUES

INGREDIENT	CAS NUMBER	ACGIH TLV (TWA)	OSHA PEL (TWA)
Ethyl Alcohol	64-17-5		1000 ppm
			1900 mg/m ³

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INGREDIENT	CAS NUMBER	ACGIH TLV (STEL / SKIN)	ACGIH TLV (CEIL)	OSHA PEL (STEL / SKIN)	OSHA PEL (CEIL)
Ethyl Alcohol	64-17-5	1000 ppm			

SECTION 9. PHYSICAL AND CHEMICAL PROPERTIES

FORM: COLOR: ODOR: SOLUBILITY:

Water:

Solution Clear, Pink Odor unknown

Not determined

See Section 5 for flammability/explosivity information.

SECTION 10. STABILITY AND REACTIVITY

STABILITY/ REACTIVITY:

Stable under normal conditions.

INCOMPATIBLE MATERIALS / CONDITIONS TO AVOID:

Open flames and high temperatures.

HAZARDOUS DECOMPOSITION PRODUCTS / REACTIONS:

Carbon oxides (COx).

SECTION 11. TOXICOLOGICAL INFORMATION

The toxicological properties of the mixture(s) have not been fully characterized in humans or animals. The information presented below pertains to the following individual ingredients in this formulation, unless indicated otherwise.

ACUTE TOXICITY DATA

INHALATION:

Propylene glycol caused no adverse effects in monkeys or rats following exposure to saturated atmospheres for prolonged periods of time.

SKIN:

Propylene glycol: Dermal LD50: 20.8 g/kg (rabbit)

Propylene glycol was irritating in a human patch test. Propylene glycol was not irritating to the skin of rabbits, guinea pigs and swine.

EYE:

Propylene glycol was slightly irritating to the eyes of rabbits.

ORAL:

Pentobarbital Sodium: Oral LD50: 118 mg/kg (rat); 65 mg/kg (dog)

Phenytoin Sodium: Oral LD50: 1530 mg/kg (rat); 165-490 mg/kg (mouse) Toxic doses of phenytoin sodium in animals produce mydriasis, nystagmus, salivation, incoordination, and ataxia. Muscular spasticity, rigidity, tremors, convulsive movements, and opisthotonos has preceded death from respiratory failure.

Propylene glycol: Oral LD50: 21 to 33.7 g/kg (rat), 10 to 20 g/kg (dog)

Propylene glycol caused dyspnea, cramps, loss of equilibrium, depression, analgesia, and death after prolonged moribund state in mice at doses ranging from 23.9 to 31.8 g/kg. In rabbits, 1 to 1.5 g/kg propylene glycol reduced intraocular pressure by raising the osmotic pressure of blood.

DERMAL AND RESPIRATORY SENSITIZATION:

Propylene glycol did not cause sensitization in a human patch test.

ADDITIONAL INFORMATION:

This product is intended for euthansia in dogs upon intravenous administration. Cerebral death in conjunction with respiratory arrest and circulatory collapse is expected.

REPEAT DOSE TOXICITY DATA

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SUBCHRONIC / CHRONIC TOXICITY:

Phenytoin effected the peripheral nervous system when given to female rats orally at doses of 300 mg/jg/day for 180 days. Increased thickness of craniofacial bones measured by increases of histomorphometric (osteoblast number, bone mineral apposition rate) and biochemical (skeletal alkaline phosphatase activity, osteocalcin concentrations) parameters of bone formation were observed in rats given phenytoin at doses of 5 mg/kg/day for 36 days by intraperitoneal injection.

Propylene glycol caused no adverse effects in monkeys or rats exposed to saturated vapor concentrations for 12 to 18 months. Rats exposed to 25 or 50% (7.7 and 13.2 g/kg/day) propylene glycol in water died within 69 days in a 140 day study. In a separate study, a diet of 30% propylene glycol was not well tolerated in young rats, and dams could not bring their young to weaning; diets containing 40, 50, or 60% propylene glycol were lethal after a few days.

REPRODUCTIVE / DEVELOPMENTAL TOXICITY:

Pentobarbital (base) induced a number of anomalies in mice; however, it was not shown to be teratogenic in rats, rabbits, or guinea pigs.

Phenytoin is a teratogen and fetotoxin in rats. It is a teratogen in mice and rabbits, and fetotoxic in monkeys at doses that were also maternally toxic. Phenytoin is not teratogenic in dogs or cats.

Rabbits were administered phenytoin sodium by oral gavage at doses of 150 mg/kg on gestation day 14-16 or 300 mg/kg on gestation days 15-16. Fetuses were examined shortly after the last dose on Day 16. The following effects were observed in the fetuses: digital areas of the limb plates showed edema and dilated blood vessels, vascular disruption occurred with hemorrhages, mesenchymal necrosis, amputation of digits, superficial hemorrhage in the fontal and nasal region, and intracranial and superficial hemorrhage in the central nevous system. Rats were administered phenytoin sodium through intraperitoneal injections at doses of 10, 50, or 100 mg/kg on Day 17 of gestation. There were no adverse effects on pregnancy or neonatal survival in the 10 and 50 mg/kg group. In the 100 mg/kg group, total fetal loss was observed in 50% of the dams, and in the remaining dams, delivery was delayed. In monkeys, oral administration of 60 to 600 mg/kg of phenytoin during gestational days 21 to 50 resulted in dose dependent maternal toxicity, and an increase in embryonic loss. In mice, phenytoin induced cleft palated when administered subcutaneously at doses up to 50 mg/kg from days 9 to 15 of gestation.

Propylene glycol caused decreased food consumption, retarded growth, smaller litters, changes in breeding patterns, and inhibited weaning in rats that were fed 30% propylene glycol through six generations; however, this may have been due to nutritional insufficiency. Propylene glycol was not teratogenic in rabbits, monkeys or chickens.

MUTAGENICITY / GENOTOXICITY:

Pentobarbital (base) was positive in the mouse micronucleus assay, mouse cell DNA inhibition test, hamster cytogenetic assay, and in the hamster dominant lethal test.

Studies with phenytoin showed no induction of micronuclei, chromosomal aberrations, or aneuploidy in human lymphocytes in vivo. There was an increase of polyploidy in one study, and sister chromatid exchange in three of seven studies. Neither chromosomal aberrations nor aneuploidy were induced in human bone marrow. Phenytoin induced mutations in Salmonella typhimurium in the presence of a metabolic activation system in one study, but was negative in Drosophila or mammalian cells in vitro assays in the absence of a metabolic system. Aneuploidy was induced in one study in primary mouse embryonic fibroblasts in vitro. Cell transformation was induced in Syrian hamster embryo. Phenytoin inhibited gap-junctional intercellular communication.

Propylene glycol was negative in a bacterial mutagenicity study (Ames).

CARCINOGENICITY:

This material or product has not been evaluated for carcinogenicity.

IARC has classified phenytoin as a Group 2B (agent is a possible human carcinogen) based on sufficient evidence in animals.

Phenytoin sodium was tested in three strains of mice at oral doses of 60 mg/kg/day for 168 days. There was an increase of thymic lymphomas in two strains of mice, and in the other strain, there was an increase of generatlized lymphomas. In another study, mice administered intraperitoneal injections of 0.6 mg/mouse over 66 days showed an increase in tumors: thymic, mesenteric, and leukemia.

Propylene glycol was not carcinogenic when applied to the skin, or when given orally in mice and rats.

SECTION 12. ECOLOGICAL INFORMATION

There are no data for the final product or its formulation(s). The information presented below pertains to the following ingredient(s).

ECOTOXICITY DATA

INGREDIENT ECOTOXICITY

Pentobarbital Sodium: 96-hr LC50 (fathead minnow): 49.5 mg/L

Propylene glycol: 96-hr LC50 (sheepshead minnow): 23,800 mg/L Propylene glycol: 48-hr EC50 (daphnid): >43,500 mg/L Propylene glycol: 72-hr EC50 (green algae): >19,000 mg/L

ENVIRONMENTAL DATA

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SECTION 13. DISPOSAL CONSIDERATIONS

MATERIAL WASTE:

Disposal must be in accordance with applicable federal, state/provincial, and/or local regulations. Incineration is the preferred method of disposal, when appropriate. Operations that involve the crushing or shredding of waste materials or returned goods must be handled to meet the recommended exposure limit(s).

PACKAGING AND CONTAINERS:

Disposal must be in accordance with applicable federal, state/provincial, and/or local regulations.

SECTION 14. TRANSPORT INFORMATION

This material is not subject to the transportation regulations of DOT, IATA, IMO, and the ADR.

SECTION 15. REGULATORY INFORMATION

WHMIS CLASSIFICATIONS:

This product has been classified in accordance with the hazard criteria on the Controlled Products Regulations and the MSDS contains all the information required by the Controlled Products Regulations. The final packaged product is not subject to WHMIS classification. The following classification applies to the bulk formulation handled in the workplace.

Controlled Product Class:	
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D1B: Toxic D2A: Very Toxic

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TSCA LISTING

INGREDIENT	TSCA
Phenytoin Sodium	Х
Propylene Glycol	Х
Ethyl Alcohol	Х
Benzyl Alcohol	Х

SECTION 16. OTHER INFORMATION

Although reasonable care has been taken in the preparation of this document, we extend no warranties and make no representations as to the accuracy or completeness of the information contained therein, and assume no responsibility regarding the suitability of this information for the user's intended purposes or for the consequence of its use. Each individual should make a determination as to the suitability of the information for their particular purpose(s).

DEPARTMENT ISSUING MSDS:	Global Safety & the Environr Merck & Co., Inc. One Merck Drive Whitehouse Station, NJ 088	
MERCK MSDS HELPLINE:	(800) 770-8878 (US and Car (908) 473-3371 (Worldwide) Monday to Friday, 9am to 5p	,
MSDS CREATION DATE: SUPERSEDES DATE:	03-Mar-1992 03-Sep-2009	
SECTIONS CHANGED (CAN SUBFORMAT): SIGNIFICANT CHANGES (CAN SUBFORMAT):	8 ETAC, OEB	
MSDS NAME: Beuthanasia-D Solution		MSDS NUMBER: SP000354
Latest Revision Date: 07-Oct-2011	Page 7 of 8	



Versior 5.1	n Revision Date: 04/12/2018		OS Number: 1678-00009	Date of last issue: 02/27/2018 Date of first issue: 05/12/2016			
SECTION 1. IDENTIFICATION							
Pr	oduct name	:	: Pentobarbital Sodium / Phenytoin Formulation				
Ma	anufacturer or supplie	er's deta	ails				
Co	ompany name of suppli	er :	Merck & Co., Inc				
Ac	ldress	:	2000 Galloping H Kenilworth - New	ill Road Jersey - U.S.A. 07033			
Te	elephone	:	908-740-4000				
Τe	elefax	:	908-735-1496				
Er	nergency telephone	:	: 1-908-423-6000				
E-	mail address	:	EHSDATASTEWARD@merck.com				
Re	ecommended use of t	ne chen	nical and restriction	ons on use			
Re	ecommended use	:	Veterinary produc	ct			

SECTION 2. HAZARDS IDENTIFICATION

GHS classification in accordance with 29 CFR 1910.1200

Flammable liquids	:	Category 3
Acute toxicity (Oral)	:	Category 3
Carcinogenicity	:	Category 2
Reproductive toxicity	:	Category 2
Specific target organ systemic toxicity - single exposure	:	Category 1 (Central nervous system)
Specific target organ systemic toxicity - repeated exposure	:	Category 1 (Central nervous system)
GHS label elements		
Hazard pictograms	:	
Signal Word	:	Danger
Hazard Statements	:	H226 Flammable liquid and vapor. H301 Toxic if swallowed. H351 Suspected of causing cancer.



ersion 1	Revision Date: 04/12/2018	SDS Number: 671678-00009	Date of last issue: 02/27/2018 Date of first issue: 05/12/2016
		H370 Causes o H372 Causes o	ed of damaging fertility or the unborn child. damage to organs (Central nervous system). damage to organs (Central nervous system) ged or repeated exposure.
Precautionary Statements		P202 Do not ha and understood P210 Keep awa No smoking. P233 Keep con P241 Use explo ment. P242 Use only P243 Take pres P260 Do not br P264 Wash ski P270 Do not ea	ay from heat/sparks/open flames/hot surfaces. Itainer tightly closed. Dision-proof electrical/ ventilating/ lighting/ equip non-sparking tools. cautionary measures against static discharge. reathe mist or vapors. In thoroughly after handling. at, drink or smoke when using this product. Itective gloves/ protective clothing/ eye protectio
		POISON CENT P303 + P361 + all contaminate	P330 IF SWALLOWED: Immediately call a ER/doctor. Rinse mouth. P353 IF ON SKIN (or hair): Take off immediate d clothing. Rinse skin with water/shower. exposed: Call a POISON CENTER or doctor/
		Storage: P403 + P235 S P405 Store loc	tore in a well-ventilated place. Keep cool. ked up.
		Disposal:	of contents/ container to an approved waste dis

Vapors may form explosive mixture with air.

SECTION 3. COMPOSITION/INFORMATION ON INGREDIENTS

Substance / Mixture : Mixture

Hazardous ingredients

Chemical name	CAS-No.	Concentration (% w/w)
Pentobarbital sodium	57-33-0	>= 30 - < 50
Propylene glycol	57-55-6	>= 10 - < 20
Ethanol	64-17-5	>= 10 - < 20
Phenytoin sodium	630-93-3	>= 5 - < 10
Benzyl alcohol	100-51-6	>= 1 - < 5



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SECTION	4. FIRST AID MEASUR	ES				
Gener	al advice	:		cident or if you feel unwell, seek medical ely., When symptoms persist or in all cases of cal advice.		
lf inha	led	:	If inhaled, remove Get medical atter			
In cas	e of skin contact	:	In case of contact, immediately flush skin with plenty of water. Remove contaminated clothing and shoes. Get medical attention. Wash clothing before reuse. Thoroughly clean shoes before reuse.			
In cas	e of eye contact	:		vater as a precaution. ation if irritation develops and persists.		
lf swa	llowed	:	If swallowed, DO NOT induce vomiting. Call a physician or poison control center immediately. Rinse mouth thoroughly with water. Never give anything by mouth to an unconscious person.			
	mportant symptoms ffects, both acute and ed	:	Causes damage	ising cancer. naging fertility or the unborn child.		
Protec	Protection of first-aiders		and use the recor	ers should pay attention to self-protection, mmended personal protective equipment al for exposure exists.		
Notes	Notes to physician		Treat symptomati	ically and supportively.		
ECTION	5. FIRE-FIGHTING ME	ASL	IRES			
Suitab	le extinguishing media	:	Water spray Alcohol-resistant Carbon dioxide (0 Dry chemical			
Unsuit media	table extinguishing	:	High volume wate	er jet		
Specif fightin	fic hazards during fire g	:	 Do not use a solid water stream as it may scatter and fire. Flash back possible over considerable distance. Vapors may form explosive mixtures with air. Exposure to combustion products may be a hazard to 			
Hazar ucts	dous combustion prod-	:	Carbon oxides Nitrogen oxides (Metal oxides	NOx)		



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Specific extinguishing meth- ods		:	Use extinguishing measures that are appropriate to local cumstances and the surrounding environment. Use water spray to cool unopened containers. Remove undamaged containers from fire area if it is safe so. Evacuate area.		
	ial protective equipment e-fighters	:		e, wear self-contained breathing apparatus. tective equipment.	
SECTION	6. ACCIDENTAL RELE	ASI	E MEASURES		
tive e	onal precautions, protec- quipment and emer- y procedures	:		tective equipment. ling advice and personal protective	
Envir	onmental precautions	:	Prevent further le Prevent spreadin oil barriers). Retain and dispos	e environment must be avoided. akage or spillage if safe to do so. g over a wide area (e.g., by containment or se of contaminated wash water. should be advised if significant spillages ned.	
	ods and materials for inment and cleaning up	:	Soak up with iner Suppress (knock jet. For large spills, p containment to ke can be pumped, s container. Clean up remaini absorbent. Local or national disposal of this m employed in the o determine which	Is should be used. t absorbent material. down) gases/vapors/mists with a water spray rovide diking or other appropriate eep material from spreading. If diked material store recovered material in appropriate ng materials from spill with suitable regulations may apply to releases and laterial, as well as those materials and items cleanup of releases. You will need to regulations are applicable. 15 of this SDS provide information regarding	

SECTION 7. HANDLING AND STORAGE

Technical measures	:	See Engineering measures under EXPOSURE CONTROLS/PERSONAL PROTECTION section.
Local/Total ventilation	:	Use with local exhaust ventilation. Use only in an area equipped with explosion-proof exhaust ventilation if advised by assessment of the local exposure potential



Version 5.1	Revision Date: 04/12/2018	SDS Number: 671678-00009	Date of last issue: 02/27/2018 Date of first issue: 05/12/2016
Advice on safe handling		Do not swallo Avoid contact Avoid prolong Handle in acc practice, base assessment Non-sparking Keep contain Keep away fr Take precaut	
Conditions for safe storage		Store locked Keep tightly o Keep in a coo Store in acco	
Ma	erials to avoid	Strong oxidizi Organic pero Flammable so Pyrophoric lic Pyrophoric so Self-heating s	xides blids Juids blids substances and mixtures and mixtures which in contact with water emit

SECTION 8. EXPOSURE CONTROLS/PERSONAL PROTECTION

Components	CAS-No.	Value type (Form of	Control parame- ters / Permissible	Basis
		exposure)	concentration	
Pentobarbital sodium	57-33-0	TWA	40µg/m3 (OEB3)	Internal
		Wipe limit	400µg/100cm2	Internal
Propylene glycol	57-55-6	TWA	10 mg/m ³	US WEEL
Ethanol	64-17-5	TWA	1,000 ppm 1,900 mg/m ³	NIOSH REL
		STEL	1,000 ppm	ACGIH
		TWA	1,000 ppm 1,900 mg/m ³	OSHA Z-1
Phenytoin sodium	630-93-3	TWA	50 µg/m3 (OEB3)	Internal
		Wipe limit	500 µg/100 cm2	Internal
Benzyl alcohol	100-51-6	TWA	10 ppm	US WEEL

Ingredients with workplace control parameters

Engineering measures

: Use appropriate engineering controls and manufacturing



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		less quick c All engineer design and protect prod Containmen are required the compou containmen	s to control airborne concentrations (e.g., drip- onnections). ing controls should be implemented by facility operated in accordance with GMP principles to lucts, workers, and the environment. It technologies suitable for controlling compounds to control at source and to prevent migration of nd to uncontrolled areas (e.g., open-face t devices). en handling.
Perso	onal protective equip	ment	
	iratory protection	: General and maintain va concentratio unknown, a Follow OSH use NIOSH/ by air purify hazardous o supplied res release, exp	d local exhaust ventilation is recommended to por exposures below recommended limits. Where ons are above recommended limits or are opropriate respiratory protection should be worn. A respirator regulations (29 CFR 1910.134) and MSHA approved respirators. Protection provided ing respirators against exposure to any chemical is limited. Use a positive pressure air spirator if there is any potential for uncontrolled posure levels are unknown, or any other where air purifying respirators may not provide otection.
Hand	protection		
Ma	aterial	: Chemical-re	esistant gloves
Re	emarks		ouble gloving. Take note that the product is which may impact the selection of hand
Еуе р	protection	If the work e mists or aer Wear a face	glasses with side shields or goggles. environment or activity involves dusty conditions, osols, wear the appropriate goggles. eshield or other full face protection if there is a direct contact to the face with dusts, mists, or
Skin a	and body protection	Additional b task being p disposable s	m or laboratory coat. ody garments should be used based upon the performed (e.g., sleevelets, apron, gauntlets, suits) to avoid exposed skin surfaces. riate degowning techniques to remove potentially ed clothing.
Hygie	ne measures	located clos When using Wash conta The effective	eye flushing systems and safety showers are e to the working place. do not eat, drink or smoke. minated clothing before re-use. e operation of a facility should include review of controls, proper personal protective equipment,



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				wning and decontamination procedures, monitoring, medical surveillance and the tive controls.
SECTION	9. PHYSICAL AND CH	EMIC		S
Appe	arance	:	liquid	
Color		:	pink	
Odor		:	No information a	vailable.
Odor	Threshold	:	No data available	9
рН		:	No data available	9
Meltir	ng point/freezing point	:	No data available	9
Initial range	boiling point and boiling	:	No data available	9
Flash	point	:	111 - 140 °F / 44	- 60 °C
Evap	oration rate	:	No data available	9
Flam	mability (solid, gas)	:	Not applicable	
Flam	mability (liquids)	:	Not applicable	
	r explosion limit / Upper nability limit	:	No data available	9
	r explosion limit / Lower nability limit	:	No data available	9
Vapo	r pressure	:	No data available	9
Relat	ive vapor density	:	No data available	9
Relat	ive density	:	No data available	2
Dens	ity	:	No data available	9
	ility(ies) ater solubility	:	No data available	9
	ion coefficient: n- ol/water	:	No data available	9
Autoi	gnition temperature	:	No data available	9
Deco	mposition temperature	:	No data available	9



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Viscosity Viscosity, kinematic		:	No data availa	ble
Explos	sive properties	:	Not explosive	
Oxidizing properties		:	The substance	e or mixture is not classified as oxidizing.
Molecular weight		:	No data availa	ble
Particle size		:	No data availa	ble

SECTION 10. STABILITY AND REACTIVITY

Reactivity	:	Not classified as a reactivity hazard.
Chemical stability	:	Stable under normal conditions.
Possibility of hazardous reac- tions	:	Flammable liquid and vapor. Vapors may form explosive mixture with air. Can react with strong oxidizing agents.
Conditions to avoid	:	Heat, flames and sparks.
Incompatible materials	:	Oxidizing agents
Hazardous decomposition products	:	No hazardous decomposition products are known.

SECTION 11. TOXICOLOGICAL INFORMATION

Information on likely routes of exposure

Inhalation Skin contact Ingestion Eve contact	0.	
Eye contact Acute toxicity Toxic if swallowed.		
Product:		
Acute oral toxicity	:	Acute toxicity estimate: 298.5 mg/kg Method: Calculation method
Acute inhalation toxicity	:	Acute toxicity estimate: > 200 mg/l Exposure time: 4 h Test atmosphere: dust/mist Method: Calculation method
Components:		
Pentobarbital sodium: Acute oral toxicity	:	LD50 (Rat): 118 mg/kg



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		L	.D50 (Mouse):	239 mg/kg	
		L	.D50 (Rabbit):	175 mg/kg	
		L	.D50 (Dog): 68	5 mg/kg	
	lene glycol: oral toxicity	: L	.D50 (Rat): > {	5,000 mg/kg	
Acute	inhalation toxicity	E	.C50 (Rabbit): Exposure time Test atmosphe	4 h	
Acute	dermal toxicity	A	· · /	> 2,000 mg/kg he substance or mixture has no acute dermal	
Ethan	ol:				
Acute	oral toxicity		.D50 (Rat): > /lethod: OECD	5,000 mg/kg 9 Test Guideline 401	
Acute	cute inhalation toxicity		: LC50 (Rat): 124.7 mg/l Exposure time: 4 h Test atmosphere: vapor		
Pheny	toin sodium:				
Acute	oral toxicity	: L	D50 (Mouse):	150 - 490 mg/kg	
Benzy	l alcohol:				
Acute	oral toxicity	: L	.D50 (Rat): 1,6	S20 mg/kg	
Acute	inhalation toxicity	E	C50 (Rat): > 4 Exposure time Test atmosphe Method: OECE	4 h	
	orrosion/irritation				
	assified based on ava onents:	ilable in	formation.		
Specie Metho Result	d	: 0	Rabbit DECD Test Gu No skin irritatio		
Ethan Specie Metho Result	es d	: 0	Rabbit DECD Test Gu lo skin irritatio		



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Benz	yl alcohol:		
Speci	ies	: Rabbit	
Metho		: OECD Test Gu	iideline 404
Resu	lt	: No skin irritatio	n
Serio	ous eye damage/eye	irritation	
	lassified based on ava		
<u>Com</u>	ponents:		
	ylene glycol:		
Speci Resu		: Rabbit	-
Metho		: No eye irritation : OECD Test Gu	
Metho	Ju	. OECD Test Gt	
Ethar	nol:		
Speci		: Rabbit	
Resu			s, reversing within 21 days
Metho	Da	: OECD Test Gu	lideline 405
Benz	yl alcohol:		
Speci		: Rabbit	
Resu			s, reversing within 21 days
N/Ioth/			
Metho			ideline 405
Resp	iratory or skin sensi		
Resp Skin	iratory or skin sensi sensitization	itization	
Resp Skin	iratory or skin sensi	itization	
Resp Skin Not c	iratory or skin sensi sensitization	itization ailable information.	
Resp Skin Not cl Resp Not cl	iratory or skin sensi sensitization lassified based on ava iratory sensitization lassified based on ava	itization ailable information.	
Resp Skin Not cl Resp Not cl	iratory or skin sensi sensitization lassified based on ava iratory sensitization	itization ailable information.	
Resp Skin Not c Resp Not c <u>Com</u>	iratory or skin sensi sensitization lassified based on ava iratory sensitization lassified based on ava ponents: ylene glycol:	itization ailable information. ailable information.	
Resp Skin Not c Resp Not c Com Prop	iratory or skin sensi sensitization lassified based on ava iratory sensitization lassified based on ava ponents: ylene glycol: Type	itization ailable information. ailable information. : Maximization T	
Resp Skin Not c Resp Not c Com Prop Test Route	iratory or skin sensi sensitization lassified based on ava iratory sensitization lassified based on ava ponents: ylene glycol: Type es of exposure	itization ailable information. ailable information. : Maximization T : Skin contact	
Resp Skin Not c Resp Not c <u>Com</u> Test	iratory or skin sensi sensitization lassified based on ava iratory sensitization lassified based on ava ponents: ylene glycol: Type es of exposure ies	itization ailable information. ailable information. : Maximization T	
Resp Skin Not c Resp Not c Com Prop Test Route Speci Resu	iratory or skin sensi sensitization lassified based on ava iratory sensitization lassified based on ava ponents: ylene glycol: Type es of exposure ies lt	itization ailable information. ailable information. : Maximization T : Skin contact : Guinea pig	
Resp Skin Not cl Resp Not cl <u>Com</u> Prop Test Route Speci Resu	iratory or skin sensi sensitization lassified based on ava iratory sensitization lassified based on ava ponents: ylene glycol: Type es of exposure ies lt	itization ailable information. ailable information. : Maximization T : Skin contact : Guinea pig : negative	- est
Resp Skin Not cl Resp Not cl Com Prop Test Route Speci Resu Ethar	iratory or skin sensi sensitization lassified based on ava iratory sensitization lassified based on ava ponents: ylene glycol: Type es of exposure les lt nol: Type	itization ailable information. ailable information. : Maximization T : Skin contact : Guinea pig : negative : Local lymph no	
Resp Skin Not c Resp Not c Com Prop Test Route Speci Resu Ethar Test Route	iratory or skin sensi sensitization lassified based on avai iratory sensitization lassified based on avai ponents: ylene glycol: Type es of exposure ies lt nol: Type es of exposure	itization ailable information. ailable information. : Maximization T : Skin contact : Guinea pig : negative : Local lymph no : Skin contact	- est
Resp Skin Not c Resp Not c Com Prop Test Route Speci Resu Ethar Route Speci	iratory or skin sensi sensitization lassified based on ava iratory sensitization lassified based on ava ponents: ylene glycol: Type es of exposure ies lt nol: Type es of exposure ies	itization ailable information. ailable information. : Maximization T : Skin contact : Guinea pig : negative : Local lymph no : Skin contact : Skin contact : Mouse	- est
Resp Skin Not c Resp Not c Com Prop Test Route Speci Resu Ethar Test Route	iratory or skin sensi sensitization lassified based on ava iratory sensitization lassified based on ava ponents: ylene glycol: Type es of exposure les lt nol: Type es of exposure ies	itization ailable information. ailable information. : Maximization T : Skin contact : Guinea pig : negative : Local lymph no : Skin contact	- est
Resp Skin Not c Resp Not c Com Prop Test Route Speci Resu Ethar Test Route Speci Resu	iratory or skin sensi sensitization lassified based on avai iratory sensitization lassified based on avai ponents: ylene glycol: Type es of exposure ies lt nol: Type es of exposure ies lt yl alcohol:	itization ailable information. ailable information. : Maximization T : Skin contact : Guinea pig : negative : Local lymph no : Skin contact : Mouse : negative	⁻ est ode assay (LLNA)
Resp Skin Not c Resp Not c Com Prop Test Route Speci Resu Ethar Test Route Speci Resu Ethar Test	iratory or skin sensi sensitization lassified based on avai iratory sensitization lassified based on avai ponents: ylene glycol: Type es of exposure les lt nol: Type es of exposure les lt yl alcohol: Type	itization ailable information. ailable information. : Maximization T : Skin contact : Guinea pig : negative : Local lymph no : Skin contact : Mouse : negative : negative	⁻ est ode assay (LLNA)
Resp Skin Not c Resp Not c Com Prop Test Route Speci Resu Ethar Test Route Speci Resu Ethar Test Route	iratory or skin sensi sensitization lassified based on avai iratory sensitization lassified based on avai ponents: ylene glycol: Type es of exposure les lt nol: Type es of exposure les lt yl alcohol: Type es of exposure	itization ailable information. ailable information. : Maximization T : Skin contact : Guinea pig : negative : Local lymph no : Skin contact : Mouse : negative : negative	⁻ est ode assay (LLNA)
Resp Skin Not c Resp Not c Com Prop Test Route Speci Resu Ethar Test Route Speci Resu Ethar Test	iratory or skin sensi sensitization lassified based on ava iratory sensitization lassified based on ava ponents: ylene glycol: Type es of exposure ies lt nol: Type es of exposure ies lt yl alcohol: Type es of exposure ies	itization ailable information. ailable information. : Maximization T : Skin contact : Guinea pig : negative : Local lymph no : Skin contact : Mouse : negative : negative	⁻ est



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Resul	lt	: ne	gative			
Not cl	a cell mutagenicity lassified based on ava conents:	ilable info	rmation.			
	ylene glycol: toxicity in vitro		st Type: Bact sult: negative	erial reverse mutation assay (AMES)		
Geno	toxicity in vivo	cyt Sp Ap	 Test Type: Mammalian erythrocyte micronucleus test (in viv cytogenetic assay) Species: Mouse Application Route: Intraperitoneal injection Result: negative 			
Ethar	nol:					
	toxicity in vitro		st Type: In vit sult: negative	ro mammalian cell gene mutation test		
			st Type: Bact sult: negative	erial reverse mutation assay (AMES)		
Geno	toxicity in vivo	Sp Ap	st Type: Rode ecies: Mouse plication Rou sult: equivoca	te: Ingestion		
Phen	ytoin sodium:					
	toxicity in vitro		st Type: Bact sult: negative	erial reverse mutation assay (AMES)		
		Re	sult: negative			
		Re	marks: Based	d on data from similar materials		
		ma	st Type: In vit Ilian cells sult: positive	ro sister chromatid exchange assay in mam-		
		Re	marks: Based	d on data from similar materials		
Geno	toxicity in vivo	cyt Sp Ap Re	ogenetic assa ecies: Mouse plication Rou sult: negative	te: Ingestion		
		cyt Sp	ogenetic test ecies: Rat	igenicity (in vivo mammalian bone-marrow , chromosomal analysis) te: Intraperitoneal injection		



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		Result: negat Remarks: Ba	ive sed on data from similar materials
		change Species: Mou	ammalian bone marrow sister chromatid ex- ise oute: Intraperitoneal injection
		Result: positiv	
	n cell mutagenicity - ssment	: Weight of evi cell mutagen.	dence does not support classification as a gerr
Benz	yl alcohol:		
	toxicity in vitro	: Test Type: Ba Result: negat	acterial reverse mutation assay (AMES) ive
Geno	toxicity in vivo	cytogenetic a Species: Mou	use oute: Intraperitoneal injection
Carci	inogenicity		
	inogenicity ected of causing cance	ır.	
Susp	• •	ır.	
Suspo <u>Com</u>	ected of causing cance	ır.	
Suspe <u>Com</u> Prop Speci	ected of causing cance ponents: ylene glycol: ies	: Rat	
Suspo <u>Com</u> Prop Speci Applio	ected of causing cance ponents: ylene glycol: ies cation Route	: Rat : Ingestion	
Suspo <u>Com</u> Prop Speci Applio	ected of causing cance ponents: ylene glycol: ies cation Route sure time	: Rat	
Suspe Comj Prop Speci Applic Expos Resu	ected of causing cance ponents: ylene glycol: ies cation Route sure time	: Rat : Ingestion : 2 Years	
Suspection Comp Propy Speci Applic Expose Resu Phen Speci	ected of causing cance <u>ponents:</u> ylene glycol: ies cation Route sure time lt ytoin sodium: ies	: Rat : Ingestion : 2 Years : negative : Rat	
Suspect Com Prop Speci Applic Expos Resu Phen Speci Applic	ected of causing cance ponents: ylene glycol: ies cation Route sure time lt hytoin sodium: ies cation Route	: Rat : Ingestion : 2 Years : negative : Rat : Ingestion	
Suspect Com Prop Speci Applic Expos Resu Phen Speci Applic Expos	ected of causing cance ponents: ylene glycol: ies cation Route sure time It ytoin sodium: ies cation Route sure time	: Rat : Ingestion : 2 Years : negative : Rat : Ingestion : 2 Years	
Suspect Com Propy Speci Applic Expos Resu Phen Speci Applic Expos Resu	ected of causing cance ponents: ylene glycol: ies cation Route sure time It ytoin sodium: ies cation Route sure time	: Rat : Ingestion : 2 Years : negative : Rat : Ingestion	
Suspection Comp Special Applical Expose Result Applical Expose Result Target Special	ected of causing cance ponents: ylene glycol: ies cation Route sure time It ytoin sodium: ies cation Route sure time lt et Organs ies	: Rat : Ingestion : 2 Years : negative : Rat : Ingestion : 2 Years : positive : Liver : Mouse	
Suspection Comp Special Applie Expose Resul Phen Special Applie Expose Resul Targe Special Applie	ected of causing cance ponents: ylene glycol: ies cation Route sure time It ytoin sodium: ies cation Route sure time It et Organs ies cation Route	: Rat : Ingestion : 2 Years : negative : Rat : Ingestion : 2 Years : positive : Liver : Mouse : Ingestion	
Suspect Comp Propy Speci Applic Expose Resu Phen Speci Applic Expose Resu Targe Speci Applic	ected of causing cance ponents: ylene glycol: ies cation Route sure time It ytoin sodium: ies cation Route sure time It et Organs ies cation Route sure time lt	: Rat : Ingestion : 2 Years : negative : Rat : Ingestion : 2 Years : positive : Liver : Mouse : Ingestion : 2 Years	
Suspection Comp Propy Specia Applica Expose Resul Phen Specia Applica Expose Resul Targe Specia Applica Expose Resul Targe Specia Applica Expose Resul Targe Specia Applica Expose Resul Targe Resul Specia Applica Expose Resul Targe Resul Specia Applica Expose Resul Targe	ected of causing cance ponents: ylene glycol: ies cation Route sure time It ytoin sodium: ies cation Route sure time It et Organs ies cation Route sure time lt	: Rat : Ingestion : 2 Years : negative : Rat : Ingestion : 2 Years : positive : Liver : Mouse : Ingestion	
Suspection Comp Prop Specia Applica Expose Resul Target Specia Applica Expose Resul Target Resul Target	ected of causing cance ponents: ylene glycol: ies cation Route sure time It ytoin sodium: ies cation Route sure time It et Organs ies cation Route sure time It et Organs nogenicity - Assess-	: Rat : Ingestion : 2 Years : negative : Rat : Ingestion : 2 Years : positive : Liver : Mouse : Ingestion : 2 Years : positive : Liver	nce of carcinogenicity in animal studies
Suspection Comp Propy Specia Applica Expose Resul Target Specia Applica Expose Resul Target Carcia ment	ected of causing cance ponents: ylene glycol: ies cation Route sure time It ytoin sodium: ies cation Route sure time It et Organs ies cation Route sure time It et Organs nogenicity - Assess-	: Rat : Ingestion : 2 Years : negative : Rat : Ingestion : 2 Years : positive : Liver : Mouse : Ingestion : 2 Years : positive : Liver	nce of carcinogenicity in animal studies



Application Route Exposure time Method Result		SDS Number: Date of last issue: 02/27/2018 671678-00009 Date of first issue: 05/12/2016 : Ingestion : 103 weeks : OECD Test Guideline 451 : negative				
OSHA		nt of this product present at levels greater than or equal to 0.1% is st of regulated carcinogens.				
NTP	Reasonably Phenytoin so	anticipated to be a human carcinogen dium 630-93-3				
-	ductive toxicity cted of damaging fertil	ity or the unborn child.				
<u>Comp</u>	onents:					
Pento	barbital sodium:					
Repro sessm	ductive toxicity - As- ient	: Some evidence of adverse effects on development, based or animal experiments.				
Propy	lene glycol:					
Effects	s on fertility	: Test Type: Three-generation reproduction toxicity study Species: Mouse Application Route: Ingestion Result: negative				
Effects	s on fetal development	: Test Type: Embryo-fetal development Species: Mouse Application Route: Ingestion Result: negative				
Ethan	ol:					
Effects	s on fertility	: Test Type: Two-generation reproduction toxicity study Species: Mouse Application Route: Ingestion Result: negative				
Pheny	/toin sodium:					
-	s on fertility	: Species: Rat Application Route: Ingestion Fertility: LOAEL: 10 mg/kg body weight Result: positive				
Effects	s on fetal development	 Test Type: Embryo-fetal development Species: Rabbit Application Route: Ingestion Developmental Toxicity: LOAEL: 150 mg/kg body weight Result: positive 				



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		Test Type: Embryo-fetal development Species: Monkey Application Route: Ingestion Result: positive	
Repro sessn	oductive toxicity - As- nent	: Some evidence of adverse effects on sexual function and fertility, and/or on development, based on animal experim	
Benz	yl alcohol:		
	s on fertility	 Test Type: Fertility/early embryonic development Species: Rat Application Route: Ingestion Result: negative Remarks: Based on data from similar materials 	
Effect	s on fetal development	: Test Type: Embryo-fetal development Species: Mouse Application Route: Ingestion Result: negative	
STOT	-single exposure		
	es damage to organs (C	entral nervous system).	
<u>Com</u>	oonents:		
Pento	barbital sodium:		
Targe	es of exposure et Organs ssment	 Ingestion Central nervous system Causes damage to organs. 	
STOT	-repeated exposure		
		entral nervous system) through prolonged or repeated exposu	re.
<u>Com</u>	oonents:		
Phen	ytoin sodium:		
	es of exposure et Organs	 Ingestion Central nervous system 	

Repeated dose toxicity

Components:

Assessment

Propylene glycol:

Species	: Rat, male
NOAEL	: 1,700 mg/kg
Application Route	: Ingestion
Exposure time	: 2 y

:

Ethanol:

Species

centrations of 10 mg/kg bw or less.

Shown to produce significant health effects in animals at con-



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NOAEL LOAEL Application Route Exposure time		 1,280 mg/kg 3,156 mg/kg Ingestion 90 Days 				
Pheny	toin sodium:					
Exposi	L ation Route ure time Organs	:	Mouse 30 mg/kg Ingestion 13 Weeks Liver Based on data fro	om similar materials		
Benzy	l alcohol:					
NOAEI Applica Exposi	Species NOAEL Application Route Exposure time Method		Rat 1.072 mg/l inhalation (dust/mist/fume) 28 Days OECD Test Guideline 412			
-	tion toxicity ssified based on availa	ble	information.			
Experi	ence with human exp	osu	Ire			
Comp	onents:					
Pentol Ingesti	parbital sodium: on	:		nouth, mood swings, Dizziness, Headache, nervous system effects, Sweating		
Pheny	toin sodium:					
Ingesti	Ingestion		nervous system e	ea, constipation, confusion, Vomiting, centra effects, Dizziness, insomnia, Blood disorders remors, anorexia		
ECTION 1	2. ECOLOGICAL INFO	DRN	IATION			
Ecoto	kicity					
Comp	onents:					
	parbital sodium: y to fish	:	LC50 (Pimephale Exposure time: 9	es promelas (fathead minnow)): 49.5 mg/l 6 h		
	ana duaali					
Propy Toxicit	y to fish	:	LC50 (Oncorhyno Exposure time: 9	chus mykiss (rainbow trout)): 40,613 mg/l 6 h		



ersion I	Revision Date: 04/12/2018		S Number: 1678-00009	Date of last issue: 02/27/2018 Date of first issue: 05/12/2016
aquatic	invertebrates		Exposure time: 48	3 h
Toxicity	∕ to algae	:	ErC50 (Skeletone Exposure time: 72 Method: OECD T	
	y to daphnia and other invertebrates (Chron- ity)	:	NOEC (Ceriodapl Exposure time: 7	nnia dubia (water flea)): 13,020 mg/l d
Toxicity	y to microorganisms	:	NOEC (Pseudomonas putida): > 20,000 mg/l Exposure time: 18 h	
Ethanc	bl:			
Toxicity	/ to fish	:	LC50 (Pimephale Exposure time: 96	s promelas (fathead minnow)): > 1,000 mg 5 h
	y to daphnia and other invertebrates	:	EC50 (Ceriodaph Exposure time: 48	nia (water flea)): > 1,000 mg/l 3 h
Toxicity	Toxicity to algae		ErC50 (Chlorella Exposure time: 72	vulgaris (Fresh water algae)): 275 mg/l 2 h
			EC10 (Chlorella v Exposure time: 72	rulgaris (Fresh water algae)): 11.5 mg/l 2 h
	/ to daphnia and other invertebrates (Chron- ity)	:	NOEC (Daphnia r Exposure time: 9	nagna (Water flea)): 9.6 mg/l d
Toxicity	y to microorganisms	:	EC50 (Pseudomo Exposure time: 16	onas putida): 6,500 mg/l 5 h
Pheny	toin sodium:			
Ecotox	cicology Assessment			
Acute a	aquatic toxicity	:	Toxic effects can	not be excluded
Chronic	c aquatic toxicity	:	Toxic effects can	not be excluded
Benzyl	alcohol:			
Toxicity	y to fish	:	LC50 (Pimephales promelas (fathead minnow)): 460 mg/l Exposure time: 96 h	
	y to daphnia and other invertebrates	:	EC50 (Daphnia magna (Water flea)): 230 mg/l Exposure time: 48 h Method: OECD Test Guideline 202	
Toxicity	∕ to algae	:	EC50 (Pseudokiro mg/l Exposure time: 72 Method: OECD T	



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			mg/l Exposure time: 7	irchneriella subcapitata (green algae)): 310 72 h Fest Guideline 201		
	ity to daphnia and other tic invertebrates (Chron- icity)	:	: NOEC (Daphnia magna (Water flea)): 51 mg/l Exposure time: 21 d Method: OECD Test Guideline 211			
Persi	stence and degradabil	ity				
<u>Com</u>	ponents:					
	ylene glycol: egradability	:	Result: Readily b Biodegradation: Exposure time: 2 Method: OECD 1	98.3 %		
Ethar Biode	n ol: egradability	:	Result: Readily b Biodegradation: Exposure time: 2	84 %		
	Benzyl alcohol: Biodegradability		Result: Readily b Biodegradation: Exposure time: 1	92 - 96 %		
Bioad	ccumulative potential					
<u>Com</u>	ponents:					
Partiti	ylene glycol: ion coefficient: n- ol/water	:	log Pow: -1.07			
	nol: ion coefficient: n- ol/water	:	log Pow: -0.35			
Partiti	yl alcohol: ion coefficient: n- ol/water	:	log Pow: 1.05			
No da Othe i	lity in soil ata available r adverse effects ata available					



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SECTION	13. DISPOSAL CONS	IDEF	RATIONS	
•	sal methods from residues	:	Dispose of in acc	ordance with local regulations.
Contaminated packaging		:	handling site for r Empty containers Do not pressurize expose such con sources of ignitio death.	e should be taken to an approved waste ecycling or disposal. Fretain residue and can be dangerous. e, cut, weld, braze, solder, drill, grind, or tainers to heat, flame, sparks, or other n. They may explode and cause injury and/or pecified: Dispose of as unused product.

SECTION 14. TRANSPORT INFORMATION

International Regulations

UNRTDG UN number Proper shipping name	:	UN 1993 FLAMMABLE LIQUID, N.O.S.
Class Packing group Labels	:	(Ethanol, Pentobarbital sodium) 3 III 3
IATA-DGR UN/ID No. Proper shipping name	:	UN 1993 Flammable liquid, n.o.s. (Ethanol, Pentobarbital sodium)
Class Packing group Labels Packing instruction (cargo aircraft) Packing instruction (passen- ger aircraft)		3 III Flammable Liquids 366 355
IMDG-Code UN number Proper shipping name Class	-	UN 1993 FLAMMABLE LIQUID, N.O.S. (Ethanol, Pentobarbital sodium) 3
Packing group Labels EmS Code Marine pollutant	:	5 III 3 F-E, <u>S-E</u> no
Transport in bulk according	* **	Annox II of MARROL 73/78 and the IRC

Transport in bulk according to Annex II of MARPOL 73/78 and the IBC Code

Not applicable for product as supplied.

Domestic regulation

49 CFR UN/ID/NA number : NA 1993



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Proper shipping name			: Combustible liquid, n.o.s. (Ethanol, Pentobarbital sodium)		
Class	;	: CBL			
Packing group		: 111			
Labels		: None			
ERG Code		: 128			
Marine pollutant		: no			
Rema	arks	liters. Not reg to 119 gallon unless other	es only to containers over 119 gallons or 450 gulated if shipped in packages less than or equal s (450 liters). If transporting by vessel or aircraft, means of transportation is impracticable, then the t be shipped as a flammable liquid.		

Special precautions for user

The transport classification(s) provided herein are for informational purposes only, and solely based upon the properties of the unpackaged material as it is described within this Safety Data Sheet. Transportation classifications may vary by mode of transportation, package sizes, and variations in regional or country regulations.

SECTION 15. REGULATORY INFORMATION

EPCRA - Emergency Planning and Community Right-to-Know

CERCLA Reportable Quantity

This material does not contain any components with a CERCLA RQ.

SARA 304 Extremely Hazardous Substances Reportable Quantity

This material does not contain any components with a section 304 EHS RQ.

SARA 302 Extremely Hazardous Substances Threshold Planning Quantity

This material does not contain any components with a section 302 EHS TPQ.

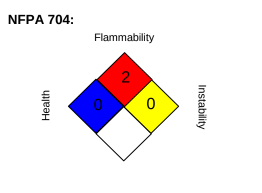
SARA 31	1/312 Hazards :	Flammable (gases, aeroso Acute toxicity (any route o Carcinogenicity Reproductive toxicity Specific target organ toxic	f exposure)	
SARA 31	l 3 :	The following components established by SARA Title		
		Pentobarbital sodium	57-33-0	>= 30 - < 50 %
US State	Regulations			
Pennsylv	vania Right To Know			
	Pentobarbital sodium Water Propylene glycol Ethanol Phenytoin sodium Benzyl alcohol			57-33-0 7732-18-5 57-55-6 64-17-5 630-93-3 100-51-6



Version 5.1	Revision Date: 04/12/2018		S Number: 678-00009	Date of last issue: 0 Date of first issue: 0	
W/ kno to t	California Prop. 65 WARNING: This product can expose you to chemicals including Phenytoin sodium, which is/are known to the State of California to cause cancer, and Pentobarbital sodium, which is/are known to the State of California to cause birth defects or other reproductive harm. For more information go to www.P65Warnings.ca.gov.				
California List of Hazardous Substances64-17-5Ethanol630-93-3					
California Permissible Exposure Limits for Chemical Contaminants Ethanol 64-17-5					64-17-5
The ingredients of this product are reported in the following inventories: AICS : not determined					
DS	SL	: 1	not determined		
IEC	CSC	: 1	not determined		

SECTION 16. OTHER INFORMATION

Further information



Special hazard.

Full text of other abbreviations

HMIS® IV:

HEALTH	*	4
FLAMMABILITY		2
PHYSICAL HAZARD		0

HMIS® ratings are based on a 0-4 rating scale, with 0 representing minimal hazards or risks, and 4 representing significant hazards or risks. The "*" represents a chronic hazard, while the "/" represents the absence of a chronic hazard.

ACGIH	:	USA. ACGIH Threshold Limit Values (TLV)
NIOSH REL	:	USA. NIOSH Recommended Exposure Limits
OSHA Z-1	:	USA. Occupational Exposure Limits (OSHA) - Table Z-1 Lim-
		its for Air Contaminants
US WEEL	:	USA. Workplace Environmental Exposure Levels (WEEL)
ACGIH / STEL	:	Short-term exposure limit
NIOSH REL / TWA	:	Time-weighted average concentration for up to a 10-hour
		workday during a 40-hour workweek
OSHA Z-1 / TWA	:	8-hour time weighted average
US WEEL / TWA	:	8-hr TWA



Version	Revision Date:	SDS Number:	Date of last issue: 02/27/2018
5.1	04/12/2018	671678-00009	Date of first issue: 05/12/2016

AICS - Australian Inventory of Chemical Substances; ASTM - American Society for the Testing of Materials; bw - Body weight; CERCLA - Comprehensive Environmental Response, Compensation, and Liability Act; CMR - Carcinogen, Mutagen or Reproductive Toxicant; DIN - Standard of the German Institute for Standardisation; DOT - Department of Transportation; DSL - Domestic Substances List (Canada); ECx - Concentration associated with x% response; EHS - Extremely Hazardous Substance; ELx - Loading rate associated with x% response; EmS - Emergency Schedule; ENCS - Existing and New Chemical Substances (Japan); ErCx - Concentration associated with x% growth rate response; ERG - Emergency Response Guide; GHS - Globally Harmonized System; GLP - Good Laboratory Practice; HMIS - Hazardous Materials Identification System; IARC -International Agency for Research on Cancer; IATA - International Air Transport Association; IBC - International Code for the Construction and Equipment of Ships carrying Dangerous Chemicals in Bulk; IC50 - Half maximal inhibitory concentration; ICAO - International Civil Aviation Organization; IECSC - Inventory of Existing Chemical Substances in China; IMDG - International Maritime Dangerous Goods; IMO - International Maritime Organization; ISHL - Industrial Safety and Health Law (Japan); ISO - International Organisation for Standardization; KECI - Korea Existing Chemicals Inventory; LC50 - Lethal Concentration to 50 % of a test population; LD50 - Lethal Dose to 50% of a test population (Median Lethal Dose); MARPOL - International Convention for the Prevention of Pollution from Ships; MSHA - Mine Safety and Health Administration; n.o.s. - Not Otherwise Specified; NFPA - National Fire Protection Association; NO(A)EC - No Observed (Adverse) Effect Concentration; NO(A)EL - No Observed (Adverse) Effect Level; NOELR - No Observable Effect Loading Rate; NTP - National Toxicology Program; NZIoC - New Zealand Inventory of Chemicals; OECD - Organization for Economic Co-operation and Development; OPPTS - Office of Chemical Safety and Pollution Prevention; PBT - Persistent, Bioaccumulative and Toxic substance; PICCS - Philippines Inventory of Chemicals and Chemical Substances; (Q)SAR - (Quantitative) Structure Activity Relationship; RCRA - Resource Conservation and Recovery Act; REACH - Regulation (EC) No 1907/2006 of the European Parliament and of the Council concerning the Registration, Evaluation, Authorisation and Restriction of Chemicals; RQ - Reportable Quantity; SADT - Self-Accelerating Decomposition Temperature; SARA - Superfund Amendments and Reauthorization Act; SDS - Safety Data Sheet; TCSI - Taiwan Chemical Substance Inventory; TSCA - Toxic Substances Control Act (United States); UN - United Nations; UNRTDG -United Nations Recommendations on the Transport of Dangerous Goods; vPvB - Very Persistent and Very Bioaccumulative

Sources of key data used to	:	Internal technical data, data from raw material SDSs, OECD
compile the Material Safety Data Sheet		eChem Portal search results and European Chemicals Agen- cy, http://echa.europa.eu/
Data Sheet		cy, mp.//echa.eu/opa.eu/

Revision Date : 04/12/2018

The information provided in this Safety Data Sheet is correct to the best of our knowledge, information and belief at the date of its publication. The information is designed only as a guidance for safe handling, use, processing, storage, transportation, disposal and release and shall not be considered a warranty or quality specification of any type. The information provided relates only to the specific material identified at the top of this SDS and may not be valid when the SDS material is used in combination with any other materials or in any process, unless specified in the text. Material users should review the information and recommendations in the specific context of their intended manner of handling, use, processing and storage, including an assessment of the appropriateness of the SDS material in the user's end product, if applicable.

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