SAFETY DATA SHEETS

This SDS packet was issued with item: 078778447

The safety data sheets (SDS) in this packet apply to the individual products listed below. Please refer to invoice for specific item number(s).

078778439 078778454 078882556 078941619



Merck & Co., Inc. 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:	Florfenicol (300 mg/mL)-Flunixin (16.5 mg/mL)-2-Pyrrolidone Injectable Solution		
SYNONYM(S):	RESFLOR (2-Pyrrolidone) Injectable Solution RESFLOR Injectable Solution - Reformulation RESFLOR (2-Pyrrolidone) Cattle Injectable RESFLOR Gold		
MSDS NUMBER:	SP001649		
EMERGENCY NUMBER(S):	(908) 423-6000 (24/7/365) English Only Emergencies - CHEMTREC: (800) 424-9300 (Inside Continental USA) (703) 527-3887 (Outside Continental USA)		
MERCK MSDS HELPLINE:	(800) 770-8878 (US and Canada) (908) 473-3371 (Worldwide) Monday to Friday, 9am to 5pm (US Eastern Time)		

The brand-names or trademarks indicated by CAPITAL LETTERS in this [M]SDS are the property of, licensed to, promoted or distributed by Merck & Co., Inc., its subsidiaries or related companies.

EMERGENCY OVERVIEW

Solution Yellow-brown Odor unknown Toxic by inhalation. May be harmful if swallowed. Irritating to eyes. May cause allergic reactions in susceptible individuals. May cause effects to: gastrointestinal tract respiratory system blood liver kidnev male reproductive system fetus May cause impaired fertility. May cause developmental effects. Toxic to aquatic organisms. May cause long-term adverse effects in the aquatic environment.

POTENTIAL HEALTH EFFECTS:

The toxicological properties of the mixture(s) have not been fully characterized in humans or animals. However, there are data to describe the toxicological properties of the individual ingredients. The following summary is based upon available information about the individual ingredients of the mixture(s), or of the expected properties of the mixture(s).

Florfenicol is a broad spectrum antibiotic used in veterinary products. Florfenicol may cause allergic reactions in susceptible individuals. Based on animal studies, florfenicol may cause slight eye irritation, constipation, changes in blood cell counts, changes in stool, or liver effects. It may also cause developmental effects or effects to male reproductive organs.

Flunixin meglumine is a potent non-narcotic, non-steroidal agent with pain killing, anti-inflammatory, and fever-reducing activity. Based on animal studies, flunixin meglumine may cause severe eye irritation or irreversible ocular effects. It may also cause irritation of the skin, mucous membranes, respiratory tract, and gastrointestinal tract. Repeated dermal contact to high concentrations may cause severe skin irritation. Prolonged inhalation may produce serious lung effects. Repeated ingestion or inhalation of high doses may cause internal bleeding, predominantly of the gastrointestinal tract.

Glyceryl triacetate may cause slight to moderate eye irritation based on animal studies.

2-Pyrrolidone may cause fetal effects based on animal studies.

Malic acid is a relatively strong acid. It may cause severe eye irritation, and skin and mucous membrane irritation.

LISTED CARCINOGENS

No carcinogens or potential carcinogens listed by OSHA, IARC, NTP or ACGIH are present in concentrations >0.1% in this mixture.

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

MSDS NAME: Florfenicol (300 mg/mL)-Flunixin (16.5 mg/mL)-2-Pyrrolidone Injectable Solution Latest Revision Date: 20-Jan-2012

MSDS NUMBER: SP001649

Page 2 of 9

INGREDIENT	CAS NUMBER	PERCENT
Florfenicol	73231-34-2	25
Glyceryl Triacetate	102-76-1	40-50
2-Pyrrolidone	616-45-5	20-30
Malic Acid	6915-15-7	<10
Flunixin Meglumine	42461-84-7	2.2

ADDITIONAL INFORMATION:	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. Administer artificial respiration if breathing has ceased. IMMEDIATELY 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:	Rinse mouth and drink a glass of water. Do not induce vomiting unless under the direction of a qualified medical professional or Poison Control Center. If symptoms persist, consult a physician.
NOTE TO PHYSICIAN:	This product contains florfenicol, a broad spectrum antibiotic which may cause allergic reactions in susceptible individuals. Flunixin meglumine is a potent Non-Steroidal Anti-inflammatory Drug (NSAID), and overexposure may cause gastrointestinal irritation and bleeding, kidney and central nervous system effects.

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.

ENVIRONMENTAL PRECAUTIONS:

This product is toxic to aquatic organisms. Do not allow product to reach ground water, water course, sewage or drainage systems.

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.

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):

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.

OCCUPATIONAL EXPOSURE GUIDELINE (OEG):

An Occupational Exposure Guideline (OEG) of 80 mcg/m³ (8-hr TWA) has been established for Florfenicol. Consult your site safety and industrial hygiene professional(s) for additional guidance.

An Occupational Exposure Guideline (OEG) of 18 mcg/m³ (8-hr TWA) has been established for flunixin. Consult your site safety and industrial hygiene professional(s) for additional guidance.

OEB/OEL NOTATION(S):

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

Flunixin Meglumine: This material has a notation of "C" for corrosivity.

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.
NAME: Florfenicol (300 mg/mL)-Flun	xin MSDS NUMBER: SP001649

MSDS NAME: Florfenicol (300 mg/mL)-Flunixin (16.5 mg/mL)-2-Pyrrolidone Injectable Solution Latest Revision Date: 20-Jan-2012

EXPOSURE LIMIT VALUES

See Occupational Exposure Guideline (OEG) listed above.

SECTION 9. PHYSICAL AND CHEMICAL PROPERTIES

FORM: COLOR: ODOR: SPECIFIC GRAVITY: SOLUBILITY: Water: Solution Yellow-brown Odor unknown 1.22

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: None known.

HAZARDOUS DECOMPOSITION PRODUCTS / REACTIONS:

No dangerous decomposition is expected if used according to manufacturer's specifications.

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

Florfenicol: 96-hr LC50 (bluegill): >830 mg/L Florfenicol: 96-hr LC50 (trout): >780 mg/L Florfenicol: 48-hr EC50 (daphnid): >330 mg/L Florfenicol: Algae maximum cell density: MIC = 1.5 mg/L Florfenicol: Algae maximum growth rate: MIC >2.9 mg/L

Flunixin meglumine: 96-hr LC50 (trout): 9.2 mg/L Flunixin meglumine: 96-hr LC50 (bluegill): 46 mg/L Flunixin meglumine: 48-hr EC50 (Daphnia): 25 mg/L Flunixin meglumine: 72 hr IC50 (Algae): 36-120 mg/L

ENVIRONMENTAL DATA

OTHER INGREDIENT ENVIRONMENTAL DATA:

Florfenicol: log Pow (log octanol/water partition coefficient): 2.36 Florfenicol: Solubility 1.32 mg/ml at pH 7 Florfenicol: Biodegrability: Not readily biodegradable but there is evidence of inherent biodegradability.

Flunixin Meglumine: log Pow (log octanol/water partition coefficient): 1.34

Glyceryl Triacetate: log Pow (log octanol/water partition coefficient): 0.25 Glyceryl triacetate is not expected to bioaccumulate or bioconcentrate in aquatic organisms. The estimated BCF is 1.3. It is expected to hydrolyse in soil and water to rapidly biodegradable products.

Malic Acid: log Pow (log octanol/water partition coefficient): -1.26

ENVIRONMENTAL FATE AND EFFECTS:

Photolytic half-life of Florfenicol in synthetic humic water (SHW) or pure water (PW) was 196 days in SHW and 171 days in PW.

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, and not to the mixture(s).

MSDS NAME: Florfenicol (300 mg/mL)-Flunixin (16.5 mg/mL)-2-Pyrrolidone Injectable Solution Latest Revision Date: 20-Jan-2012

MSDS NUMBER: SP001649

Page 5 of 9

ACUTE TOXICITY DATA

INHALATION:

Florfenicol: No mortality occurred in rats exposed to florfenicol for 4 hours at 0.28 mg/L (the maximum concentration tested). Clinical effects included dry rales, anogenital staining, secretory discharge, soft stool, and decreased body weights. These effects were seen immediately or up to one-week post exposure. Some effects did not resolve by study termination.

Flunixin Meglumine: Inhalation LC50 (4hr): <0.52 mg/L (rat)

Mortality occurred in all rats (10/10) between days 3 and 6 following a single 4-hour exposure to an average analytical concentration of 0.52 mg/L (maximum attainable exposure). Signs exhibited following exposure included lacrimation, nasal discharge, dried red material around facial area, and yellow anogenital staining. Significant weight loss was noted following exposure in all animals.

SKIN:

Florfenicol was not irritating to rabbit skin.

Flunixin meglumine: Slightly irritating

Flunixin meglumine produced mild, transient dermal irritation in rabbits. Dose-related skin irritation effects were observed in rabbits during a 21-day repeat skin application study (see below under Subchronic to Chronic Toxicity).

Glyceryl triacetate was not irritating when absorbed through the skin of guinea pigs.

2-Pyrrolidone was not irritating to the skin of rabbits.

Malic acid was moderately irritating to the skin of rabbits, and strongly irritating to the skin of guinea pigs.

EYE:

Florfenicol was slightly irritating to the eyes of rabbits.

Flunixin Meglumine: Severely irritating

All six animals exhibited severe conjunctival irritation including redness, swelling, discharge, and necrosis, as well as corneal opacity, ulceration and iridial damage. Severe ocular irritation was irreversible in most animals.

Rabbits treated with 0.1 ml of undiluted glyceryl triacetate exhibited slight to moderate irritation, but produced no effect when immediately rinsed for six minutes.

2-Pyrrolidone was not irritating to the eyes of rabbits.

Malic acid was severely irritating to the eyes of rabbits.

ORAL:

Florfenicol: Oral LD50: >2000 mg/kg (rat, mouse).

Dogs (one animal/sex) were administered successive oral doses of florfenicol that ranged from 160 to 1280 mg/kg. No clinical effects occurred at doses as high as 640 mg/kg. At 640 mg/kg, the only female died from inhalation of vomitus. Vomiting or soft stool occurred at 640 to 1280 mg/kg.

Flunixin Meglumine: Oral LD50: 53 to 157 mg/kg (rat), 176 to 249 mg/kg (male mouse, female estimated)

Flunixin (free acid): Oral LD50: 468.3 mg/kg (guinea pig)

Common effects observed in acute oral studies across species include gastrointestinal effects (perforation/ulceration and hemorrhage), hypoactivity, pallor, spleen enlargement, congestion of kidneys, lungs, or gastrointestinal tract, and respiratory distress. Necropsy of animals that died from flunixin meglumine revealed abnormalities of the brain, epididymides, abdominal cavity, thymus, liver, mesenteric lymph nodes, esophagus, mesentery, pancreas, and lungs. No signs of toxicity were observed following acute oral administration of 100 & 200 mg/kg to rhesus monkeys. However, 1 of 3 monkeys died following administration of 300 mg/kg. That monkey showed lethargy, prostration, and salivation prior to death, and signs of hyperemic mucosa in gastrointestinal tract and lungs at necropsy. Flunixin administered orally to mice at a dose of 300 mg/kg (100x the projected clinical dose) caused slight tremors and ataxia which resolved within 24 hours. Effects from acute oral and IV treatment of horses with 1.1 mg/kg flunixin were limited to sporadic incidence of fecal occult blood.

Glyceryl triacetate: Oral LD50: 3000-12800 mg/kg (rat); 1100-6100 mg/kg (mouse). Symptoms observed during the determination of the oral LD50 in rats and mice were weakness and ataxia.

2-Pyrrolidone: Oral LD50: 328-6500 mg/kg (rat); 6500 mg/kg (guinea pig)

Malic Acid: Oral LD50: > 3200 mg/kg (rat); 1600-3200 mg/kg (mouse)

Signs of acute poisoning in rats and mice are weakness, retraction of the abdomen, respiratory distress, and cyanosis.

DERMAL AND RESPIRATORY SENSITIZATION:

Flunixin Meglumine was found not to be sensitizing in guinea pigs when tested by intradermal induction at 1% and topically at 100%.

Glyceryl triacetate was not a skin sensitizer in guinea pigs.

ADDITIONAL INFORMATION:

Florfenicol: Intaperitoneal LD50: 1913-2253 mg/kg (rat)

Clinical signs of toxicity noted in rats treated with 1000 to 2000 mg/kg florfenicol included hypoactivity, wet or stained urogenital hair, chromorhinorrhea, or discolored stool. Abnormal pathological findings in rats included white, granular foci on surface of the liver or intestines, or pale or friable kidneys (high dose group).

REPEAT DOSE TOXICITY DATA

SUBCHRONIC / CHRONIC TOXICITY:

Florfenicol was administered orally to dogs, rats, and mice at dosages as high as 100 to 400 mg/kg/day for up to 13 weeks. Effects including decreased body weight, changes in liver weight or liver enzyme levels, changes in testicular weight, testicular atrophy, decreased white blood cell counts, and decreased hemoglobin levels were observed at high dosages. Cellular changes in the liver or lymph nodes of rats and mice, and histopathologic changes in the brain and spinal cord of dogs were also noted at these high dosages. Although some effects were reversible after a 4-week withdrawal from treatment, testicular effects in rats persisted. Intramuscular injections of 45 mg/kg of florfenicol in swine produced diarrhea, injection site lesions, decreased body weight, decreased food and water consumption, changes in serum electrolytes and proteins, decreased red blood cell and white blood cell counts, decreased spleen weight, and decreased kidney weight.

In 52-week oral toxicity studies in dogs and rats, high dosages of florfenicol (12 and 48 mg/kg/day, respectively) increased liver weight and produced cellular changes in the gall bladder of dogs. In rats, florfenicol at the high dosage reduced body weight gain, reduced testicular weight, induced changes in hematologic and clinical chemistry parameters, and increased the incidence of testicular tubular atrophy. In two-year chronic studies in mice and rats, florfenicol caused similar effects as those observed in other long-term studies including reduced body weight gain, reduced red blood cell count, reduced hemoglobin levels, and testicular effects such as small testes, tubular atrophy and aspermatogenesis in both the high dosage rats (48 mg/kg/day) and mice (200 mg/kg/day).

Repeat oral dosing studies have been performed with flunixin across multiple species. The most common adverse effect seen in these studies is gastrointestinal irritation/ulceration and bleeding as indicated by blood in the stools. Other common adverse effects observed across species from oral, IV or IM routes of exposure include nephrotoxicity, emesis, anorexia, and bleeding. Blood cell count changes, blood coagulation effects, and immune organ effects were observed secondary to gastrointestinal erosion and bleeding. Liver, nervous system and behavioral effects were also noted in mice. In addition to ulceration and bleeding, significant mortality was observed in rats at 8 and 16 mg/kg dosed for six weeks. [6-week oral toxicity NOAEL: 2 mg/kg (rats); 90-day oral toxicity NOAEL: 5 mg/kg (monkeys), 3.0 mg/kg (rats); one year oral toxicity NOEL: 1 mg flunixin/kg (rats)]

In several 21-day repeat skin application studies in rabbits using up to 80 mg/kg flunixin meglumine or the free acid in spray or cream formulations, no conclusive treatment-related toxicity could be established. The incidence and severity of dermal irritation increased in a dose-related manner with severe irritation seen at 80 mg/kg/day.

Rats exposed to inhalation doses of glyceryl triacetate at 250 ppm for 13 weeks, and saturated vapors for 5 days, produced no symptoms or histopathological effects.

REPRODUCTIVE / DEVELOPMENTAL TOXICITY:

In a two-generation reproductive study, oral administration as high as 12 mg/kg/day of florfenicol reduced epididymal weights, decreased pup survival, and reduced lactation index in rats [NOAEL: 3 mg/kg/day].

There was no evidence of teratogenicity in rats administered florfenicol at dosages of 4, 12 or 40 mg/kg/day. Slight maternal toxicity, evidenced by decreased food and water consumption, was observed above 4 mg/kg/day. At 40 mg/kg/day, an increased incidence of delayed ossification and decreased fetal weight occurred. The NOAEL for maternal and fetal toxicity in rats was determined to be 4 mg florfenicol/kg/day.

Two teratogenicity studies were performed in mice. In the first study, the mice were administered florfenicol at dosages of 40, 120, or 400 mg/kg by gavage on days 6-15 of gestation. Florfenicol produced embryolethality at the 400 mg/kg/day dose level, which was evidenced by the high incidence of intrauterine deaths. Significant decreases in mean fetal body weight, soft tissue defects, and retarded skeletal ossification were also observed at 400 mg/kg/day. Skeletal ossification was less pronounced, in a dose-related fashion, at the lower doses tested (40 and 120 mg/kg/day). A developmental NOAEL could not be determined for these data [NOAEL for maternal: 120 mg/kg]. In the second teratogenicity study, florfenicol was retested at lower administered dosages of 1, 3, or 60 mg/kg/day. Maternal effects were limited to a slight increase in water consumption at the 60 mg/kg/day dose. There was no evidence of any adverse effects on the embryo/fetus at doses as high as 60 mg/kg/day in this study. However, based upon the retarded skeletal ossification effects observed in the first study at 40 mg/kg/day the NOAEL for the two studies combined was determined to be between 3 and 40 mg/kg/day.

Reproductive and teratology studies in rats, mice and rabbits were performed with flunixin. Although significant maternal toxicity, including mortality, was reported, these studies indicate that flunixin does not affect offspring development, male or female fertility, or mating behavior. A slight increase in the length of gestation and difficult labor with an increase in stillbirths were observed. No evidence of any drug-related teratogenic effects were observed. Maternal toxicity observed in these studies was consistent with those findings in acute and repeated dose oral toxicity studies with the addition of pale eyes, ears and extremities. [Reproductive or developmental NOELs ranged from 2-21 mg/kg in studies with multiple species. Maternal toxicity NOELs ranged from 3-9 mg/kg in these studies].

2-Pyrrolidone was embryotoxic but not teratogenic in mice exposed through oral and intraperitoneal routes of exposure. Maternal toxicity and malformations were observed in rats orally administered 1900 mg/kg/day. In a study conducted in rats, the inhalation of 150 ppm of 2-pyrrolidone on days 7 to 20 of gestation was associated with decreased pup weights and delays in developmental milestones.

MUTAGENICITY / GENOTOXICITY:

Florfenicol was negative in a bacterial mutagenicity study (Ames), a mammalian mutagenicity study (mouse lymphoma), a bone marrow micronucleus assay, an in vitro chromosomal aberration assay in CHO cells, a cytogenetics assay in bone marrow, and an unscheduled DNA synthesis assay in rat hepatocytes.

Flunixin meglumine was negative in the Ames and mouse micronucleus assays. It was positive in mouse lymphoma L5178Y cells, both in the absence and presence of S-9 metabolic activation and in the chromosomal abberation assay in CHO cells in vitro both in the absence and presence of S-9 metabolic activation. It has been reported to alter cellular DNA and caused primary DNA damage in E. coli. Flunixin free acid yielded the same results as flunixin meglumine. However, it was inconclusive in the bacterial repair assay in E.coli whereas flunixin meglumine was strongly positive. The meglumine moiety (N-methyl-D glucamine) was negative in all studies performed except the micronucleus study in which it was positive in one study and negative in a second.

2-Pyrrolidone was not mutagenic in a bacterial mutagenicity assay (Ames).

Malic acid was negative in bacterial mutagenicity studies (Ames), either in the absence or presence of metabolic activation. Malic acid was not clastogenic in Chinese hamester fibroblast cells.

CARCINOGENICITY:

Florfenicol was not carcinogenic in a 2-year study in rats administered dosages up to 48 mg/kg/day for 5 days a week or in mice at dosages up to 200 mg/kg/day for 5 days per week.

Flunixin meglumine had no carcinogenic effects or increase in tumor incidence relative to controls in either a 104-week study in rats administered 2, 4 and 8 mg flunixin meglumine/kg/day in the diet, or in mice administered 0.6, 2.0 and 6.0 mg flunixin meglumine/kg/day in the diet for 97 weeks. Significant toxicity observed in rats and mice included decreased body weights, increased mortality (high dose groups) and dose-related increases in gastrointestinal lesions in all treated groups. Compound-related lesions observed at necropsy included dose-related gastrointestinal ulcers, ulcer perforation with secondary peritonitis and adhesion formation, and large or edematous lymph nodes. Dose-related nonproliferative lesions were present in the gastrointestinal tract and mesenteric lymph node. Necrosis and ulceration of the mucosa, transmural necrosis, mucosal and mural inflammation, lymphoid hyperplasia, peritonitis and abscess formation were present. Inflammatory lesions and necrosis secondary to the peritonitis were present in other abdominal organs. Splenomegaly (enlarged spleens) were observed at necropsy in mice and were significant in the high dose group only. [Rat NOEL for tumor formation = 8 mg flunixin meglumine/kg/day and the LOEL = 2 mg flunixin meglumine/kg/day based on GI lesions. Mouse NOEL for tumor formation = 6.0 mg flunixin meglumine/kg/day; Toxicity NOEL = 0.6 mg flunixin meglumine/kg/day].

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.

SPECIAL ENVIRONMENTAL HANDLING PROCEDURES:

This product contains materials that are harmful to the environment. Do not allow undiluted/unneutralized product to reach ground water, water course, sewage or drainage systems.

SECTION 14. TRANSPORT INFORMATION

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

SECTION 15. REGULATORY INFORMATION

TSCA LISTING

INGREDIENT	TSCA
Glyceryl Triacetate	Х
2-Pyrrolidone	Х
Malic Acid	Х

Substances not included in the table above are TSCA exempt or not regulated under TSCA.

U.S. STATE REGULATIONS

INGREDIENT	California Proposition 65	CARTK	NJRTK	CTRTK	MARTK
2-Pyrrolidone					Х

MSDS NAME: Florfenicol (300 mg/mL)-Flunixin (16.5 mg/mL)-2-Pyrrolidone Injectable Solution Latest Revision Date: 20-Jan-2012

INGREDIENT	PARTK	MNRTK	MIRTK	RIRTK
2-Pyrrolidone	Х			

Fields in the above tables that do not contain data indicate that those materials have not been listed by local regulations.

X: Listed on applicable state hazardous substance or right-to-know lists.

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).

The brand-names or trademarks indicated by CAPITAL LETTERS in this [M]SDS are the property of, licensed to, promoted or distributed by Merck & Co., Inc., its subsidiaries or related companies.

DEPARTMENT ISSUING MSDS:	Global Safety & the Environment Merck & Co., Inc. One Merck Drive Whitehouse Station, NJ 08889
MERCK MSDS HELPLINE:	(800) 770-8878 (US and Canada) (908) 473-3371 (Worldwide) Monday to Friday, 9am to 5pm (US Eastern Time)
MSDS CREATION DATE:	13-Apr-2006
SUPERSEDES DATE:	07-May-2010
SECTIONS CHANGED (US SUBFORMAT): SIGNIFICANT CHANGES (US SUBFORMAT):	2, 11 OEB





Versio 6.2	n Revision Date: 06/05/2018		05 Number: 058-00011	Date of last issue: 04/12/2018 Date of first issue: 11/04/2014
SECTI	ON 1. IDENTIFICATION			
Pi	roduct name	:	Florfenicol / Flunix	kin Formulation
	anufacturer or supplier's			
C	ompany name of supplier	:	Merck & Co., Inc	
A	ddress	:	2000 Galloping H Kenilworth - New	ill Road Jersey - U.S.A. 07033
Т	elephone	:	908-740-4000	
Te	elefax	:	908-735-1496	
E	mergency telephone	:	1-908-423-6000	
E	-mail address	:	EHSDATASTEW	ARD@merck.com
R	ecommended use of the c	hen	nical and restriction	ons on use
R	ecommended use	:	Veterinary produc	t

SECTION 2. HAZARDS IDENTIFICATION

GHS classification in accordance with 29 CFR 1910.1200

2

:

Danger

Acute toxicity (Oral)	:	Category 4
Acute toxicity (Inhalation)	:	Category 4
Eye irritation	:	Category 2A
Reproductive toxicity	:	Category 2
Specific target organ systemic toxicity - repeated exposure	:	Category 1 (Liver, Brain, Testes, Spinal cord, Blood, gallbladder, Gastrointestinal tract, Kidney)

GHS label elements

Hazard pictograms

Signal Word



Hazard Statements	:	H302 + H332 Harmful if swallowed or if inhaled. H319 Causes serious eye irritation. H361fd Suspected of damaging fertility. Suspected of damaging the unborn child.
		H372 Causes damage to organs (Liver, Brain, Testes, Spinal cord, Blood, gallbladder, Gastrointestinal tract, Kidney) through prolonged or repeated exposure.

SAFETY DATA SHEET



Florfenicol / Flunixin Formulation

ersion 2	Revision Date: 06/05/2018	SDS Number: 28058-00011	Date of last issue: 04/12/2018 Date of first issue: 11/04/2014
Preca	utionary Statements	P202 Do not h and understoo P260 Do not b P264 Wash sk P270 Do not e P271 Use only	reathe mist or vapors. in thoroughly after handling. at, drink or smoke when using this product. outdoors or in a well-ventilated area. otective gloves/ protective clothing/ eye protection
		CENTER/doct P304 + P340 - and keep com CENTER/doct P305 + P351 - for several mir to do. Continue P308 + P313 I attention.	 P330 IF SWALLOWED: Call a POISON or if you feel unwell. Rinse mouth. P312 IF INHALED: Remove person to fresh air fortable for breathing. Call a POISON or if you feel unwell. P338 IF IN EYES: Rinse cautiously with water nutes. Remove contact lenses, if present and easy e rinsing. F exposed or concerned: Get medical advice/ f eye irritation persists: Get medical advice/ atten-
		Storage: P405 Store loc	sked up.
		Disposal: P501 Dispose posal plant.	of contents/ container to an approved waste dis-
	hazards known.		

Substance / Mixture	:	Mixture	
---------------------	---	---------	--

Hazardous ingredients

Chemical name	CAS-No.	Concentration (% w/w)
Florfenicol	73231-34-2	>= 20 - < 30
2-Pyrrolidone	616-45-5	>= 20 - < 30
Malic Acid	6915-15-7	>= 1 - < 5
1-Deoxy-1-(methylamino)-D-glucitol	42461-84-7	>= 1 - < 3
2-[2-methyl-3-		
(perfluoromethyl)anilino]nicotinate		

SECTION 4. FIRST AID MEASURES

General advice

: In the case of accident or if you feel unwell, seek medical



Vers 6.2	ion	Revision Date: 06/05/2018		0S Number: 058-00011	Date of last issue: 04/12/2018 Date of first issue: 11/04/2014
				advice immediate doubt seek medic	ly., When symptoms persist or in all cases of al advice.
	lf inhale	ed	:	If inhaled, remove If not breathing, g If breathing is diffi Get medical atten	ive artificial respiration. cult, give oxygen.
	In case	of skin contact	:	of water. Remove contamir Get medical atten Wash clothing bet	
	In case	of eye contact	:	for at least 15 min	ove contact lens, if worn.
	lf swalle	owed	:	Get medical atten Rinse mouth thore	
		nportant symptoms ects, both acute and d	:	unborn child.	
	Protect	ion of first-aiders	:	and use the recor	ers should pay attention to self-protection, nmended personal protective equipment I for exposure exists.
	Notes t	o physician	:	Treat symptomati	cally and supportively.
SEC	TION 5	. FIRE-FIGHTING ME	ASL	JRES	
	Suitable	e extinguishing media	:	Water spray Alcohol-resistant f Carbon dioxide (C Dry chemical	
	Unsuita media	ble extinguishing	:	None known.	
	Specific fighting	c hazards during fire	:	Exposure to comb	oustion products may be a hazard to health.
	Hazard ucts	ous combustion prod-	:	Carbon oxides Fluorine compour Nitrogen oxides (I	





Version 6.2	Revision Date: 06/05/2018		0S Number: 058-00011	Date of last issue: 04/12/2018 Date of first issue: 11/04/2014
Spec ods	Specific extinguishing meth- ds		cumstances and Use water spray	g measures that are appropriate to local cir- the surrounding environment. to cool unopened containers. ged containers from fire area if it is safe to do
	ial protective equipment e-fighters	:		e, wear self-contained breathing apparatus. tective equipment.
SECTION	6. ACCIDENTAL RELE	AS	EMEASURES	
tive e	onal precautions, protec- equipment and emer- y procedures	:		tective equipment. ling advice and personal protective imendations.
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.
	Methods and materials for containment and cleaning up		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 Sections 13 and	t absorbent material. 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 naterial, as well as those materials and items cleanup of releases. You will need to regulations are applicable. 15 of this SDS provide information regarding ational requirements.
SECTION	7. HANDLING AND ST	OR	AGE	
Tech	nical measures	:		measures under EXPOSURE RSONAL PROTECTION section.
Loca	I/Total ventilation	:	Use with local ex	haust ventilation.

Do Do Avo Har prao ass	not breathe vapors or spray mist. not swallow. not get in eyes. id prolonged or repeated contact with skin. dle in accordance with good industrial hygiene and safety stice, based on the results of the workplace exposure essment p container tightly closed.
---------------------------------------	--

4 / 24



Version 6.2	Revision Date: 06/05/2018	SDS Number: 28058-00011	Date of last issue: 04/12/2018 Date of first issue: 11/04/2014
		Take care to pre environment.	vent spills, waste and minimize release to the
Condi	tions for safe storage	Store locked up. Keep tightly clos Keep in a cool, w	labeled containers. ed. vell-ventilated place. nce with the particular national regulations.
Materials to avoid		: Do not store with Strong oxidizing Organic peroxide Explosives Gases	•

SECTION 8. EXPOSURE CONTROLS/PERSONAL PROTECTION

• •	•			
Components	CAS-No.	Value type	Control parame-	Basis
		(Form of	ters / Permissible	
		exposure)	concentration	
Florfenicol	73231-34-2	TWA	100 µg/m3 (OEB	Internal
			2)	
1-Deoxy-1-(methylamino)-D- glucitol 2-[2-methyl-3-	42461-84-7	TWA	40 µg/m3 (OEB 3)	Internal
(perfluorome-				
thyl)anilino]nicotinate				
		Wipe limit	400 µg/100 cm²	Internal

Ingredients with workplace control parameters

Engineering measures :	Use appropriate engineering controls and manufacturing technologies to control airborne concentrations (e.g., drip- less quick connections). All engineering controls should be implemented by facility design and operated in accordance with GMP principles to protect products, workers, and the environment. Containment technologies suitable for controlling compounds are required to control at source and to prevent migration of the compound to uncontrolled areas (e.g., open-face containment devices). Minimize open handling.
Personal protective equipment	
Respiratory protection :	General and local exhaust ventilation is recommended to maintain vapor exposures below recommended limits. Where

Spiratory protection C General and local exhaust ventilation is recommended to maintain vapor exposures below recommended limits. Where concentrations are above recommended limits or are unknown, appropriate respiratory protection should be worn. Follow OSHA respirator regulations (29 CFR 1910.134) and use NIOSH/MSHA approved respirators. Protection provided by air purifying respirators against exposure to any hazardous chemical is limited. Use a positive pressure air supplied respirator if there is any potential for uncontrolled release, exposure levels are unknown, or any other



Version 6.2	Revision Date: 06/05/2018	SDS Nu 28058-0		Date of last issue: 04/12/2018 Date of first issue: 11/04/2014
			mstance whe juate protecti	ere air purifying respirators may not provide on.
Hand	protection			
Ma	aterial	: Cher	nical-resistar	nt gloves
Re	emarks	: Cons	sider double (gloving.
Eye protection		lf the miste Wea	e work enviro s or aerosols r a faceshield ntial for direc	ses with side shields or goggles. nment or activity involves dusty conditions, , wear the appropriate goggles. d or other full face protection if there is a t contact to the face with dusts, mists, or
Skin and body protection		Addi task dispo Use	tional body g being perforr osable suits)	aboratory coat. arments should be used based upon the ned (e.g., sleevelets, apron, gauntlets, to avoid exposed skin surfaces. legowning techniques to remove potentially thing.
Hygie	ne measures	locat Whe Was The engin appr indus	ed close to the n using do no h contaminat effective ope neering contro opriate degor	lushing systems and safety showers are ne working place. ot eat, drink or smoke. ed clothing before re-use. ration of a facility should include review of ols, proper personal protective equipment, wning and decontamination procedures, monitoring, medical surveillance and the tive controls.

SECTION 9. PHYSICAL AND CHEMICAL PROPERTIES

Appearance	:	liquid
Color	:	yellow
Odor	:	No information available.
Odor Threshold	:	No data available
рН	:	No data available
Melting point/freezing point	:	No data available
Initial boiling point and boiling range	:	No data available
Flash point	:	No data available
Evaporation rate	:	No data available

SAFETY DATA SHEET



Florfenicol / Flunixin Formulation

Ver 6.2	sion	Revision Date: 06/05/2018		S Number: 958-00011	Date of last issue: 04/12/2018 Date of first issue: 11/04/2014
	Flamm	ability (solid, gas)	:	Not applicable	
	Flamm	ability (liquids)	:	No data available	
		explosion limit / Upper ability limit	:	No data available	
		explosion limit / Lower ability limit	:	No data available	
	Vapor	pressure	:	No data available	
	Relativ	e vapor density	:	No data available	
	Relativ	e density	:	1.22	
	Density	y	:	No data available	
	Solubil Wat	ity(ies) ter solubility	:	No data available	
	Partitio octano	n coefficient: n- I/water	:	Not applicable	
	Autoigr	nition temperature	:	No data available	
	Decom	position temperature	:	No data available	
	Viscosi Visc	ity cosity, kinematic	:	No data available	
	Explos	ive properties	:	Not explosive	
	Oxidizi	ng properties	:	The substance or	mixture is not classified as oxidizing.
	Particle	e size	:	Not applicable	

SECTION 10. STABILITY AND REACTIVITY

Reactivity	:	Not classified as a reactivity hazard.
Chemical stability	:	Stable under normal conditions.
Possibility of hazardous reac- tions	:	Can react with strong oxidizing agents.
Conditions to avoid	:	None known.
Incompatible materials	:	Oxidizing agents
Hazardous decomposition products	:	No hazardous decomposition products are known.





ersion 2	Revision Date: 06/05/2018		9S Number: 058-00011	Date of last issue: 04/12/2018 Date of first issue: 11/04/2014
	11. TOXICOLOGICAL I	NFC	ORMATION	
Inforr Inhala	nation on likely routes	of e	exposure	
	contact			
Inges				
•	ontact			
	e toxicity ful if swallowed or if inha	led.		
Produ	uct:			
Acute	oral toxicity	:	Acute toxicity esti Method: Calculati	mate: 1,320 mg/kg on method
Acute	inhalation toxicity	:	Acute toxicity esti	
			Exposure time: 4	
			Test atmosphere: Method: Calculation	
<u>Com</u>	<u>oonents:</u>			
	enicol:			
Acute	oral toxicity	:	LD50 (Rat): > 2,0	00 mg/kg
			LD50 (Mouse): >	2,000 mg/kg
			LD50 (Dog): > 1,2	280 mg/kg
Acute	inhalation toxicity	:	LC50 (Rat): > 0.2	8 mg/l
			Exposure time: 4	
Acute	e dermal toxicity	:	Remarks: No data	a available
Acute	toxicity (other routes of	:		
admir	nistration)		Application Route	: Intraperitoneal
			LD50 (Mouse): 10	
			Application Route	: Intravenous
2-Pyr	rolidone:			
Acute	oral toxicity	:	LD50 (Rat): > 2,0	
			Method: OECD To	est Guideline 401
Acute	inhalation toxicity	:	LC0 (Rat): 0.061	mg/l
	-		Exposure time: 8	h
			Test atmosphere:	vapor
Acute	e dermal toxicity	:	LD50 (Rabbit): > 2	
			Method: OECD To	
			toxicity	substance or mixture has no acute derma

SAFETY DATA SHEET



rsion	Revision Date: 06/05/2018		0S Number: 058-00011	Date of last issue: 04/12/2018 Date of first issue: 11/04/2014	
Malic	Acid:				
Acute	oral toxicity	:	LD50 (Rat): 3,50	00 mg/kg	
Acute inhalation toxicity :		 LC50 (Rat): > 1.306 mg/l Exposure time: 4 h Test atmosphere: dust/mist Method: OECD Test Guideline 403 Assessment: The substance or mixture has no acute inha tion toxicity Remarks: Based on data from similar materials 			
Acute	dermal toxicity	:	LD50 (Rat): 20,0 Remarks: Based	000 mg/kg I on data from similar materials	
1-Dec	oxy-1-(methylamino)-D-	-glu	citol 2-[2-methy	-3-(perfluoromethyl)anilino]nicotinate:	
	oral toxicity	:	LD50 (Rat): 53 -		
			LD50 (Mouse):	176 - 249 mg/kg	
			LD50 (Guinea p	ig): 488.3 mg/kg	
			LD50 (Monkey):	300 mg/kg	
Acute	inhalation toxicity	:	LC50 (Rat): < 0. Exposure time: 4 Test atmosphere	4 h	
	toxicity (other routes of histration)	:	LD50 (Rat): 59.4 Application Rou	4 - 185.3 mg/kg te: Intraperitoneal	
			LD50 (Mouse): ⁻ Application Rou	164 - 363 mg/kg te: Intraperitoneal	
Skin	corrosion/irritation				
Not cl	assified based on availa	ble	information.		
Com	oonents:				
	enicol:				
	es	:	Rabbit		
Speci Resul		:	No skin irritation		
Resul		:	No skin irritation		
Resul 2-Pyr Speci	t rolidone: es	:	Rabbit		
Resul 2-Pyr Speci Metho	r olidone: es od	•	Rabbit OECD Test Guid	deline 404	
Resul 2-Pyr Speci	r olidone: es od	:	Rabbit	deline 404	
Resul 2-Pyr Speci Metho Resul	r olidone: es od	:	Rabbit OECD Test Guid	deline 404	
Resul 2-Pyr Speci Metho Resul Malic Speci	It rolidone: es od It Acid: es	:	Rabbit OECD Test Guid No skin irritation Rabbit	deline 404	
Resul 2-Pyr Speci Metho Resul	It rolidone: es od It Acid: es od	:	Rabbit OECD Test Guid No skin irritation	deline 404 deline 404	





sion	Revision Date: 06/05/2018	SDS Number: 28058-00011	Date of last issue: 04/12/2018 Date of first issue: 11/04/2014
1-Dec	oxy-1-(methylamino)	-D-glucitol 2-[2-meth	yl-3-(perfluoromethyl)anilino]nicotinat
Speci		: Rabbit	
Resu	lt	: Mild skin irritat	ION
	us eye damage/eye		
	es serious eye irritatio ponents:	JII.	
	enicol:		
Speci		: Rabbit	
Resul		: Mild eye irritat	on
2-Pyr	rolidone:		
Speci		: Rabbit	
Resu	lt	: Irritation to eye	es, reversing within 7 days
Malic	Acid:		
Speci		: Rabbit	
Resu			es, reversing within 21 days
Metho Rema		: OECD Test G	udeline 405 from similar materials
1-Dec Speci Resul	es	: Rabbit	yl-3-(perfluoromethyl)anilino]nicotinat ects on the eye
Resu	it.	. Ineversible en	
Resp	iratory or skin sens	itization	
	sensitization	- 11 - 1- 1- f	
	lassified based on av		
-	iratory sensitization lassified based on ava		
	oonents:		
	enicol:		
Test		: Maximization	Foot
Speci		: Guinea pig	
Resul		: negative	
2-Pyr	rolidone:		
Test ⁻		: Local lymph ne	ode assay (LLNA)
	es of exposure	: Skin contact	
Speci		: Mouse	videline 400
			uideline 429
Metho			-
	lt	: negative	from similar materials

SAFETY DATA SHEET



Florfenicol / Flunixin Formulation

ersion 2	Revision Date: 06/05/2018		t issue: 04/12/2018 t issue: 11/04/2014			
Malic	Acid:					
Test Type Routes of exposure Species Method Result Remarks		 Maximization Test Skin contact Guinea pig OECD Test Guideline 406 negative Based on data from similar materials 				
1-Deo	xy-1-(methylamino	-D-glucitol 2-[2-methyl-3-(perfluoro	methyl)anilino]nicotinate:			
Specie	s of exposure es sment	 Maximization Test Dermal Guinea pig Does not cause skin sensitization negative 	tion.			
	cell mutagenicity assified based on av	ailable information.				
Comp	oonents:					
	enicol:					
Genot	oxicity in vitro	: Test Type: Bacterial reverse n Result: negative	nutation assay (AMES)			
		Test Type: DNA damage and thesis in mammalian cells (in Test system: rat hepatocytes Result: negative				
		Test Type: In vitro mammaliar Test system: mouse lymphom Result: negative				
		Test Type: Chromosome aber Test system: Chinese hamste Result: positive				
Genot	oxicity in vivo	: Test Type: Micronucleus test Species: Mouse Cell type: Bone marrow Application Route: Oral Result: negative				
2-Pyr	rolidone:					
Genot	oxicity in vitro	: Test Type: In vitro mammaliar Method: OECD Test Guideline Result: negative				
		Test Type: Chromosome aber Method: OECD Test Guideline Result: negative				

11 / 24



ersion 2	Revision Date: 06/05/2018	SDS Number: 28058-00011	Date of last issue: 04/12/2018 Date of first issue: 11/04/2014
		Method: OEC Result: negat	CD Test Guideline 471 ive
Genotoxicity in vivo		cytogenetic a Species: Mou Application R	use oute: Intraperitoneal injection CD Test Guideline 474
Malic	Acid:		
Genot	oxicity in vitro	: Test Type: C Result: negat	hromosome aberration test in vitro ive
1-Deo	oxy-1-(methylamino)	-D-glucitol 2-[2-met	hyl-3-(perfluoromethyl)anilino]nicotinate:
	oxicity in vitro	-	acterial reverse mutation assay (AMES)
		Test Type: in Test system: Result: positi	mouse lymphoma cells
			hromosomal aberration Chinese hamster ovary cells ve
		Test Type: in Test system: Result: positi	Escherichia coli
Genot	oxicity in vivo	: Test Type: M Species: Mou Application R Result: negat	oute: Oral
	cell mutagenicity - sment	: Weight of evi cell mutagen.	dence does not support classification as a gern
Carci	nogenicity		
	assified based on ava	ailable information.	
Comp	oonents:		
Florfe	nicol:		
Specie		: Rat	
	ation Route	: oral (gavage) : 2 Years	
Resul	sure time t	: negative	
	t Organs	: Liver, Testes	
Specie	es	: Mouse	
Applic	ation Route	: oral (gavage)	
	sure time	: 2 Years	



ersion 2	Revision Date: 06/05/2018	SDS Number: 28058-00011	Date of last issue: 04/12/2018 Date of first issue: 11/04/2014		
Resul Targe	t t Organs	: negative : Testes, Bloo	d		
1-Dec	oxy-1-(methylami	no)-D-glucitol 2-[2-me	hyl-3-(perfluoromethyl)anilino]nicotinate:		
Speci	es	: Rat			
	ation Route	: oral (feed)			
	sure time	: 104 w			
LOAE		: 2 mg/kg body	y weight		
Resul	=	: negative	n al tra at		
Rema	t Organs ırks	: Gastrointesti : Significant to	xicity observed in testing		
Speci		: Mouse			
	ation Route	: oral (feed)			
	sure time	: 97 w			
NOAE Resul		: 0.6 mg/kg bo	ay weight		
	t Organs	: negative : Gastrointesti	nal tract		
Rema			xicity observed in testing		
IARC			esent at levels greater than or equal to 0.1% is or confirmed human carcinogen by IARC.		
OSH/		No component of this product present at levels greater than or equal to on OSHA's list of regulated carcinogens.			
NTP			esent at levels greater than or equal to 0.1% is ated carcinogen by NTP.		
Suspe	oductive toxicity ected of damaging ponents:	fertility. Suspected of d	amaging the unborn child.		
Florfe	enicol:				
Effect	s on fertility	Species: Rat Application F Fertility: LOA			
Effect	s on fetal develop	ment : Test Type: E Species: Rat	mbryo-fetal development		
		General Toxi Embryo-fetal Result: No te	city Maternal: NOAEL: 4 mg/kg body weight toxicity.: LOAEL: 40 mg/kg body weight eratogenic effects., Fetotoxicity. e effects were seen only at maternally toxic dos-		
		Species: Mo Application F General Toxi	mbryo-fetal development use Route: oral (gavage) city Maternal: NOAEL: 120 mg/kg body weight toxicity.: LOAEL: 40 mg/kg body weight		



Vers 6.2	sion	Revision Date: 06/05/2018		9S Number: 058-00011	Date of last issue: 04/12/2018 Date of first issue: 11/04/2014
				Result: Fetotoxicil	у.
	Reproductive toxicity - As- sessment		:	fertility, based on	adverse effects on sexual function and animal experiments., Some evidence of development, based on animal
	2-Pyrro	olidone:			
	Effects	on fertility	:	Species: Rat Application Route Result: negative	: Ingestion
	Effects on fetal development		:	Test Type: Embry Species: Rat Application Route Result: negative	o-fetal development : Ingestion
	Malic A	Acid:			
		on fertility	:	Test Type: Two-g Species: Rat Application Route Result: negative	eneration reproduction toxicity study : Ingestion
	Effects	on fetal development	:	Test Type: Embry Species: Rat Application Route Result: negative	o-fetal development : Ingestion
	1-Deox	xy-1-(methylamino)-D	-alu	citol 2-[2-methyl-:	B-(perfluoromethyl)anilino]nicotinate:
		on fertility	:	Test Type: Two-g Species: Rat Application Route General Toxicity F Symptoms: No fet	eneration reproduction toxicity study : Oral Parent: LOAEL: 1 - 1.5 mg/kg body weight al abnormalities. on fertility and early embryonic
	Effects	on fetal development	:	Embryo-fetal toxic Result: Embryotox offspring were det	: Oral Maternal: LOAEL: 2 mg/kg body weight ity.: NOAEL: 2 mg/kg body weight kic effects and adverse effects on the ected only at high maternally toxic doses o-fetal development
				General Toxicity M Embryo-fetal toxic Result: Embryotox	Aaternal: LOAEL: 3 mg/kg body weight ity.: NOAEL: 3 mg/kg body weight kic effects and adverse effects on the ected only at high maternally toxic doses





ersion 2	Revision Date: 06/05/2018	SDS Number: 28058-00011	Date of last issue: 04/12/2018 Date of first issue: 11/04/2014	
STO	-single exposure			
	lassified based on av	ailable information.		
Com	oonents:			
		,	yl-3-(perfluoromethyl)anilino]nicotinate:	
Asses	ssment	: May cause res	spiratory irritation.	
STO	-repeated exposure	e		
		s (Liver, Brain, Testes, onged or repeated exp	Spinal cord, Blood, gallbladder, Gastrointestina osure.	
<u>Com</u>	oonents:			
Florfe	enicol:			
	et Organs		estes, Spinal cord, Blood, gallbladder	
Asses	ge to organs through prolonged or repeated			
1-Dec	oxy-1-(methylamino)-D-glucitol 2-[2-meth	yl-3-(perfluoromethyl)anilino]nicotinate:	
	et Organs		al tract, Kidney, Blood	
Asses	Assessment : Causes damage to organs through prolonged or repe exposure.			
Repe	ated dose toxicity			
-	oonents:			
Florfe	enicol:			
Speci	65	: Dog		
	00	. Dog		
NOAE	ΞL	: 3 mg/kg		
Expo	EL sure time	: 3 mg/kg : 13 Weeks	Drain Chinal and	
Expo	ΞL	: 3 mg/kg : 13 Weeks	Brain, Spinal cord	
Expos Targe Speci	EL sure time et Organs es	: 3 mg/kg : 13 Weeks : Liver, Testes, : Mouse	Brain, Spinal cord	
Expos Targe Speci NOA	EL sure time et Organs es EL	: 3 mg/kg : 13 Weeks : Liver, Testes, : Mouse : 200 mg/kg	Brain, Spinal cord	
Expos Targe Speci NOAE Expos	EL sure time et Organs Es EL sure time	: 3 mg/kg : 13 Weeks : Liver, Testes, : Mouse : 200 mg/kg : 13 Weeks	Brain, Spinal cord	
Expos Targe Speci NOAE Expos	EL sure time et Organs es EL	: 3 mg/kg : 13 Weeks : Liver, Testes, : Mouse : 200 mg/kg	Brain, Spinal cord	
Expos Targe Speci NOAE Expos Targe Speci	EL sure time et Organs es EL sure time et Organs	: 3 mg/kg : 13 Weeks : Liver, Testes, : Mouse : 200 mg/kg : 13 Weeks : Liver, Testes : Rat	Brain, Spinal cord	
Expos Targe Speci NOAE Expos Targe Speci NOAE	EL sure time et Organs EL sure time et Organs EL	: 3 mg/kg : 13 Weeks : Liver, Testes, : Mouse : 200 mg/kg : 13 Weeks : Liver, Testes : Rat : 30 mg/kg	Brain, Spinal cord	
Expos Targe Speci NOAE Expos Targe Speci NOAE Expos	EL sure time et Organs EL sure time et Organs EL sure time	: 3 mg/kg : 13 Weeks : Liver, Testes, : Mouse : 200 mg/kg : 13 Weeks : Liver, Testes : Rat : 30 mg/kg : 13 Weeks	Brain, Spinal cord	
Expos Targe Speci NOAE Expos Targe Speci NOAE Expos	EL sure time et Organs EL sure time et Organs EL	: 3 mg/kg : 13 Weeks : Liver, Testes, : Mouse : 200 mg/kg : 13 Weeks : Liver, Testes : Rat : 30 mg/kg	Brain, Spinal cord	
Expos Targe Speci NOAE Expos Targe Speci Speci Speci	EL sure time et Organs EL sure time et Organs EL sure time et Organs	 3 mg/kg 13 Weeks Liver, Testes, Mouse 200 mg/kg 13 Weeks Liver, Testes Rat 30 mg/kg 13 Weeks Liver, Testes Liver, Testes 	Brain, Spinal cord	
Expos Targe Speci NOAE Expos Targe NOAE Expos Targe Speci NOAE	EL sure time et Organs EL sure time et Organs EL sure time et Organs EL	 3 mg/kg 13 Weeks Liver, Testes, Mouse 200 mg/kg 13 Weeks Liver, Testes Rat 30 mg/kg 13 Weeks Liver, Testes Liver, Testes 30 mg/kg 3 mg/kg 3 mg/kg 	Brain, Spinal cord	
Expos Targe Speci NOAE Expos Targe Speci NOAE Speci NOAE LOAE	EL sure time et Organs EL sure time et Organs EL sure time et Organs EL sure time et Organs	 3 mg/kg 13 Weeks Liver, Testes, Mouse 200 mg/kg 13 Weeks Liver, Testes Rat 30 mg/kg 13 Weeks Liver, Testes Liver, Testes Dog 3 mg/kg 12 mg/kg 	Brain, Spinal cord	
Expos Targe Speci NOAE Expos Targe Speci NOAE Expos Speci NOAE LOAE Expos	EL sure time et Organs EL sure time et Organs EL sure time et Organs EL	 3 mg/kg 13 Weeks Liver, Testes, Mouse 200 mg/kg 13 Weeks Liver, Testes Rat 30 mg/kg 13 Weeks Liver, Testes Liver, Testes 30 mg/kg 3 mg/kg 3 mg/kg 		
Expos Targe Speci NOAE Expos Targe Speci NOAE Expos Targe Speci NOAE Expos Targe Speci	EL sure time et Organs EL sure time et Organs EL sure time et Organs EL sure time et Organs EL sure time et Organs	 3 mg/kg 13 Weeks Liver, Testes, Mouse 200 mg/kg 13 Weeks Liver, Testes Rat 30 mg/kg 13 Weeks Liver, Testes Dog 3 mg/kg 12 mg/kg 52 Weeks Liver, gallblade Rat 		
Expos Targe Speci NOAE Expos Targe Speci NOAE Expos Targe Speci NOAE Expos Targe	EL sure time et Organs EL sure time et Organs EL sure time et Organs EL sure time et Organs EL sure time et Organs	 3 mg/kg 13 Weeks Liver, Testes, Mouse 200 mg/kg 13 Weeks Liver, Testes Rat 30 mg/kg 13 Weeks Liver, Testes Dog 3 mg/kg 12 mg/kg 52 Weeks Liver, gallblade Rat 1 mg/kg 		
Expos Targe Speci NOAE Expos Targe Speci NOAE Expos Targe Speci NOAE Expos Targe	EL sure time et Organs EL sure time et Organs EL sure time et Organs EL sure time et Organs EL sure time et Organs	 3 mg/kg 13 Weeks Liver, Testes, Mouse 200 mg/kg 13 Weeks Liver, Testes Rat 30 mg/kg 13 Weeks Liver, Testes Dog 3 mg/kg 12 mg/kg 52 Weeks Liver, gallblade Rat 		

15 / 24



sion	Revision Date: 06/05/2018	SDS Number: 28058-00011	Date of last issue: 04/12/2018 Date of first issue: 11/04/2014				
Targe	t Organs	: Testes					
2-Pyrı	rolidone:						
Specie		: Rat					
NOAE		: 207 mg/kg					
	ation Route	: Ingestion					
	ure time	: 3 Months					
Metho	D	: OECD Test Gu	lideline 408				
1-Deo	xy-1-(methylamino)	-D-glucitol 2-[2-meth	yl-3-(perfluoromethyl)anilino]nicotinate:				
Specie	es	: Rat					
NOAE		: 2 mg/kg					
LOAE		: < 4 mg/kg					
	ation Route	: Oral					
	ure time t Organs	: 6 w : Gastrointestina	al tract				
raige	Ulgans	. Gastionitestine					
Specie		: Rat					
NOAE		: 1 mg/kg					
	ation Route	: Oral					
	ure time t Organs	-	: 1 y : Gastrointestinal tract, Kidney				
raige	Ulgans	. Gastronitestina					
Specie		: Monkey					
NOAE		: 15 mg/kg					
	ation Route	: Oral					
	ure time t Organs	: 90 d : Gastrointestina	l tract Blood				
raige	Ulgans	. Gastonitestine					
Specie		: Rabbit					
LOAE		: 80 mg/kg					
	ation Route	: Dermal					
Sympt	ure time	: 21 d : Severe irritatio	n				
Oymp		. Gevere initatio					
Specie		: Dog					
LOAE	—	: 11 mg/kg					
	ation Route	: Oral					
	ure time t Organs	: 9 d : Gastrointestina	al tract				
Sympt		: Vomiting					
		· · · · · · · · · · · · · · · · · · ·					
-	ation toxicity						
	assified based on av						
-	ience with human e	exposure					
	onents:						
			yl-3-(perfluoromethyl)anilino]nicotinate:				
Inhala	tion	: Symptoms: res	piratory tract irritation				
01.1	ontact	: Symptoms: Sk	in irritation				

16 / 24



ersion .2	Revision Date: 06/05/2018		0S Number: 058-00011	Date of last issue: 04/12/2018 Date of first issue: 11/04/2014	
Eye c	contact	:	Symptoms: Sever	e irritation	
Inges	Ingestion		: Symptoms: Gastrointestinal disturbance, bleeding, hyperter sion, Kidney disorders		
ECTION	12. ECOLOGICAL INFO	ORN	ΙΑΤΙΟΝ		
Ecoto	oxicity				
<u>Com</u>	ponents:				
	enicol: ity to fish	:	LC50 (Lepomis m Exposure time: 96 Method: FDA 4.11		
			LC50 (Oncorhync Exposure time: 96 Method: FDA 4.17		
	ity to daphnia and other tic invertebrates	:	EC50 (Daphnia m Exposure time: 48 Method: OECD Te		
Toxic	ity to algae	:	EC50 (Pseudokiro mg/l Exposure time: 14 Method: FDA 4.07		
			NOEC (Pseudokin mg/l Exposure time: 14 Method: FDA 4.07		
			IC50 (Skeletonem Exposure time: 72 Method: ISO 1025		
			NOEC (Skeletone Exposure time: 72 Method: ISO 1025		
			EC50 (Lemna gib Exposure time: 7 Method: OECD To		
			NOEC (Lemna gil Exposure time: 7 Method: OECD To		
			EC50 (Navicula p Exposure time: 72 Method: OECD Te		



Version 6.2	Revision Date: 06/05/2018		0S Number: 058-00011	Date of last issue: 04/12/2018 Date of first issue: 11/04/2014
			NOEC (Navicula Exposure time: 72 Method: OECD T	
			EC50 (Anabaena Exposure time: 72 Method: OECD T	
			NOEC (Anabaena Exposure time: 72 Method: OECD T	
Toxic icity)	Toxicity to fish (Chronic tox- icity)		NOEC (Pimephale Exposure time: 32 Method: OECD T	
	ity to daphnia and other ic invertebrates (Chron- icity)		NOEC (Daphnia r Exposure time: 2′ Method: OECD T	
2-Pvr	rolidone:			
-	ity to fish	:	LC50 (Danio rerio Exposure time: 96 Method: OECD T	
	ity to daphnia and other ic invertebrates	:	EC50 (Daphnia m Exposure time: 48	nagna (Water flea)): > 500 mg/l 3 h
Toxic	ity to algae	:	EC50 (Desmodes Exposure time: 72	smus subspicatus (green algae)): > 500 mg/l 2 h
			EC10 (Desmodes Exposure time: 72	smus subspicatus (green algae)): 22.2 mg/l 2 h
Toxic	ity to microorganisms	:	EC50: > 1,000 mg Exposure time: 30 Method: OECD Te) min
Malic	Acid:			
Toxic	ity to fish	:	Exposure time: 96 Method: OECD T	
	ity to daphnia and other ic invertebrates	:	EC50 (Daphnia m Exposure time: 48	nagna (Water flea)): 240 mg/l 3 h
Toxic	ity to algae	:	mg/l Exposure time: 72 Method: OECD T	



Versi 6.2	on	Revision Date: 06/05/2018		S Number: 058-00011	Date of last issue: 04/12/2018 Date of first issue: 11/04/2014
				mg/l Exposure time: 72 Method: OECD Te	
-	Toxicity	to microorganisms	:	EC50: > 300 mg/l Exposure time: 3 Method: OECD Te Remarks: Based o	
	1-Deox	v-1-(methvlamino)-D-	-alu	citol 2-[2-methyl-3	-(perfluoromethyl)anilino]nicotinate:
	Toxicity		:		acrochirus (Bluegill sunfish)): 28 mg/l
				LC50 (Oncorhync Exposure time: 96 Method: FDA 4.11	
		to daphnia and other invertebrates	:	EC50 (Daphnia m Exposure time: 48 Method: FDA 4.08	
-	Toxicity	to algae	:	NOEC (Microcysti Exposure time: 13 Method: FDA 4.01	
				NOEC (Selenastru Exposure time: 12	um capricornutum (green algae)): 96 mg/l : d
I	Persist	ence and degradabili	ity		
	Compo	nents:			
	2-Pyrro	lidone:			
I	Biodegr	adability	:	Result: Readily bio Biodegradation: 9 Exposure time: 9	08 %
I	Malic A	cid:			
I	Biodegr	adability	:		3%
	1-Deox	y-1-(methylamino)-D-	-glu	citol 2-[2-methyl-3	-(perfluoromethyl)anilino]nicotinate:
		in water	:	Hydrolysis: 0 %(2	



Version 6.2	Revision Date: 06/05/2018		S Number:)58-00011	Date of last issue: 04/12/2018 Date of first issue: 11/04/2014
Bioa	ccumulative potential			
Com	ponents:			
Florf	enicol:			
	ion coefficient: n- ol/water	:	log Pow: 0.373	
2-Pyr	rrolidone:			
	ion coefficient: n- ol/water	:	log Pow: -0.71	
Malic	Acid:			
	ion coefficient: n- ol/water	:	log Pow: -1.26	
1-Dec	oxy-1-(methylamino)-D)-glu	citol 2-[2-methyl-	3-(perfluoromethyl)anilino]nicotinate:
	ion coefficient: n- ol/water	:	log Pow: 1.34	
Mobi	lity in soil			
<u>Com</u>	ponents:			
Distri	oxy-1-(methylamino)-E bution among environ- al compartments)-glu :	c itol 2-[2-methyl- log Koc: 1.92	3-(perfluoromethyl)anilino]nicotinate:
Othe	r adverse effects			
No da	ata available			
ECTION	13. DISPOSAL CONSI	DER	ATIONS	
Dispo	osal methods			
Wast	e from residues	:	Dispose of in acc	ordance with local regulations.
Conta	aminated packaging	:	handling site for r	should be taken to an approved waste ecycling or disposal. pecified: Dispose of as unused product.
SECTION	14. TRANSPORT INFO	DRM/	ATION	
Interi	national Regulations			
UNR				
	umber er shipping name	:	UN 3082 ENVIRONMENTA N.O.S.	ALLY HAZARDOUS SUBSTANCE, LIQUID,

20 / 24

(Florfenicol)

: 9



Versior 6.2	n	Revision Date: 06/05/2018		9S Number: 058-00011	Date of last issue: 04/12/2018 Date of first issue: 11/04/2014	
	Packing group Labels		:	 9		
IAT A-DGR UN/ID No. Proper shipping name Class Packing group Labels Packing instruction (cargo aircraft) Packing instruction (passen-			 UN 3082 Environmentally hazardous substance, liquid, n.o.s. (Florfenicol) 9 III Miscellaneous 964 964 			
	er airc nviron	raft) mentally hazardous	:	yes		
U	IMDG-Code UN number : UI Proper shipping name : EN N.		UN 3082 ENVIRONMENTALLY HAZARDOUS SUBSTANCE, LIQUID, N.O.S. (Florfenicol)			
Pa La Er	abels mS Co	l group ode pollutant	:	: 9 : III : 9 : F-A, S-F : yes		
Transport in bulk according to Annex II of MARPOL 73/78 an Not applicable for product as supplied.			OL 73/78 and the IBC Code			
		tic regulation	1- 1			
UI Pr CI Pa La EF	roper s lass acking abels RG Co	IA number shipping name group ode pollutant		(Florfenicol) 9 III CLASS 9 171 yes(Florfenicol) Above applies on liters., Shipment b however it may be	azardous substance, liquid, n.o.s. y to containers over 119 gallons or 450 y ground under DOT is non-regulated; shipped per the applicable hazard cilitate multi-modal transport involving ICAO	

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.

SAFETY DATA SHEET



Florfenicol / Flunixin Formulation

Version	Revision Date:	SDS Number:	Date of last issue: 04/12/2018
6.2	06/05/2018	28058-00011	Date of first issue: 11/04/2014

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 311/312 Hazards	: Acute toxicity (any route of exposure) Serious eye damage or eye irritation Reproductive toxicity Specific target organ toxicity (single or repeated exposure)
SARA 313	: This material does not contain any chemical components with known CAS numbers that exceed the threshold (De Minimis) reporting levels established by SARA Title III, Section 313.

US State Regulations

Pennsylvania Right To Know	
Triacetin	102-76-1
2-Pyrrolidone	616-45-5
Florfenicol	73231-34-2

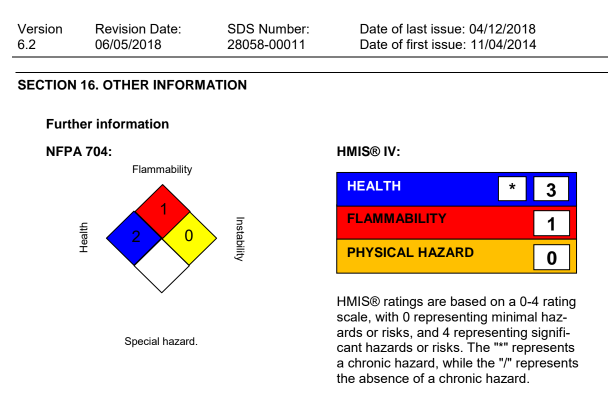
California Prop. 65

This product does not contain any chemicals known to the State of California to cause cancer, birth, or any other reproductive defects.

The ingredients of this product are reported in the following inventories:

AICS	:	not determined
DSL	:	not determined
IECSC	:	not determined





Full text of other abbreviations

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 -





Version 6.2	Revision Date: 06/05/2018		0S Number: 058-00011	Date of last issue: 04/12/2018 Date of first issue: 11/04/2014	
	d Nations Recommenda ery Bioaccumulative	ation	s on the Transport	of Dangerous Goods; vPvB - Very Persistent	
Sources of key data used to : compile the Material Safety Data Sheet		Internal technical data, data from raw material SDSs, OECD eChem Portal search results and European Chemicals Agen- cy, http://echa.europa.eu/			
Revis	ion Date	:	06/05/2018		
inform	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				

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.

US / Z8