

## **SAFETY DATA SHEETS**

**This SDS packet was issued with item:**

078914243

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

078914401

## SAFETY DATA SHEET



GlaxoSmithKline

### 1. IDENTIFICATION OF THE SUBSTANCE/PREPARATION AND OF THE COMPANY/UNDERTAKING

<b>Material</b>	<b>ZANTAC INJECTION</b>	
<b>Synonyms</b>	ZANTAC INJECTION 25MG/ML * ZANTAC 2 INYECTABLE * ZANTAC AMPOLLA * ZANTAC AMPULLE * ZANTAC INFUSIONKONCENTRAT * ZANTAC INJECTIEVLOEISTOF * ZANTAC INJEKCIJAS * ZANTAC INJEKCIO * ZANTAC INJEKSJONSVAESKE * ZANTAC INJEKTIONESTE * ZANTAC INJEKTIJONSVATSKA * ZANTAC INYECTABLE * ZANTAC SOLUCAO INJECTAVEL * ZANTAC SOLUCION INYECTABLE * ZANTAC ZAWIESINA DO INIEKJIE * NDC NO 0173-0362-38 * NDC NO 0173-0363-00 * NDC NO 0173-0363-01 * RANITIDINE HYDROCHLORIDE, FORMULATED PRODUCT	
<b>Company Name</b>	GlaxoSmithKline, Corporate Environment, Health & Safety 980 Great West Road Brentford, Middlesex TW8 9GS UK UK General Information: +44-20-8047-5000 Transport Emergency (EU) +44-1865-407333 Medical Emergency +1-612-221-3999, Ext 221 Information and Advice: US number, available 24 hours Multi-language response  GlaxoSmithKline, Corporate Environment, Health & Safety 2200 Renaissance Blvd, Suite 105 King of Prussia, PA 19406 US US General Information: +1-888-825-5249 Transport Emergency (non EU) +1-703-527-3887 US number, available 24 hours Multi-language response	

### 2. COMPOSITION / INFORMATION ON INGREDIENTS

Ingredients	CAS RN	Percentage
RANITIDINE HYDROCHLORIDE	66357-59-3	3
NON-HAZARDOUS INGREDIENTS	Unassigned	97

### 3. HAZARDS IDENTIFICATION

<b>Fire and Explosion</b>	Expected to be non-combustible.
<b>Health</b>	Handling this product in its final form presents minimal risk from occupational exposure. Caution - Pharmaceutical agent. Health effects information is based on hazards of components. Respiratory allergen. May produce allergic skin reactions.

<b>Environment</b>	No information is available about the potential of this product to produce adverse environmental effects.
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#### 4. FIRST-AID MEASURES

<b>Ingestion</b>	Never attempt to induce vomiting. Do not attempt to give any solid or liquid by mouth if the exposed subject is unconscious or semi-conscious. Wash out the mouth with water. If the exposed subject is fully conscious, give plenty of water to drink. Obtain medical attention.
<b>Inhalation</b>	Physical form suggests that risk of inhalation exposure is negligible.
<b>Skin Contact</b>	Using appropriate personal protective equipment, remove contaminated clothing and flush exposed area with large amounts of water. Obtain medical attention if skin reaction occurs, which may be immediate or delayed.
<b>Eye Contact</b>	Wash immediately with clean and gently flowing water. Continue for at least 15 minutes. Obtain medical attention.

#### NOTES TO HEALTH PROFESSIONALS

<b>Medical Treatment</b>	Medical treatment in cases of overexposure should be treated as an overdose of a histamine antagonist (H2-type). Treat according to locally accepted protocols. For additional guidance, refer to the current prescribing information or to the local poison control information centre.
<b>Medical Conditions Caused or Aggravated by Exposure</b>	Refer to prescribing information for detailed description of medical conditions caused by or aggravated by overexposure to this product.
<b>Antidotes</b>	No specific antidotes are recommended.

#### 5. FIRE-FIGHTING MEASURES

<b>Fire and Explosion Hazards</b>	This product is non-combustible, although the packaging is combustible.
<b>Extinguishing Media</b>	Water is recommended for fires involving packaging.
<b>Special Firefighting Procedures</b>	For single units (packages): No special requirements needed. For larger amounts (multiple packages/pallets) of product: Since toxic, corrosive or flammable vapours might be evolved from fires involving this product and associated packaging, self contained breathing apparatus and full protective equipment are recommended for firefighters. If possible, contain and collect firefighting water for later disposal.
<b>Hazardous Combustion Products</b>	Toxic, corrosive or flammable thermal decomposition products are expected when the product is exposed to fire.

#### 6. ACCIDENTAL RELEASE MEASURES

<b>Personal Precautions</b>	Wear protective clothing and equipment consistent with the degree of hazard.
<b>Environmental Precautions</b>	Prevent entry into waterways, sewers, surface drainage systems and poorly ventilated areas.
<b>Clean-up Methods</b>	Spread an inert absorbent on the spill and place in a suitable, properly labelled container for recovery or disposal.
<b>Decontamination Procedures</b>	No specific decontamination or detoxification procedures have been identified for this product.

#### 7. HANDLING AND STORAGE

##### HANDLING

**General Requirements** No special control measures required for the normal handling of this product. Normal room ventilation is expected to be adequate for routine handling of this product.

**STORAGE** No storage requirements necessary for occupational hazards. Follow product information storage instructions to maintain efficacy.

## 8. EXPOSURE CONTROLS/PERSONAL PROTECTION

<b>INGREDIENT</b>	RANITIDINE HYDROCHLORIDE
<b>GSK Occupational Hazard Category</b>	3
<b>GSK Occupational Exposure Limit</b>	50 mcg/m <sup>3</sup> (15 MIN STEL) SKIN SENSITISER, RESPIRATORY SENSITISER

### ENGINEERING CONTROLS

**Exposure Controls** An Exposure Control Approach (ECA) is established for operations involving this material based upon the OEL/Occupational Hazard Category and the outcome of a site- or operation-specific risk assessment. Refer to the Exposure Control Matrix for more information about how ECA's are assigned and how to interpret them.

### PERSONAL PROTECTIVE EQUIPMENT

**Eye Protection** Wear approved safety glasses with side shields if eye contact is possible.

**Other Equipment or Procedures** An eye wash station should be available. Wash hands and arms thoroughly after handling. Wear appropriate clothing to avoid skin contact.

## 9. PHYSICAL AND CHEMICAL PROPERTIES

### Appearance

**Physical Form** Solution.

**pH of Aqueous Solutions** 6.8 to 7.1

## 10. STABILITY AND REACTIVITY

**Stability** DO NOT FREEZE - dispose of properly if frozen.

**Conditions to Avoid** None for normal handling of this product.

## 11. TOXICOLOGICAL INFORMATION

**Oral Toxicity** Not expected to be toxic following ingestion.

**Skin Effects** Irritation is not expected following direct contact.

**Eye Effects** Irritation is not expected following direct contact with eyes.

**Target Organ Effects** No specific target organ effects have been identified.

**Sensitisation** Assessment based upon effects of structurally similar substances. Allergic skin reactions might occur following dermal exposure. Assessment based upon information from human exposure.

**Genetic Toxicity** Not expected to be genotoxic under occupational exposure conditions.

**Carcinogenicity** No components are listed as carcinogens by GSK, IARC, NTP or US OSHA.

**Reproductive Effects** Not expected to produce adverse effects on fertility or development under occupational exposure conditions.

**Pharmacological Effects**

This product contains active ingredient(s) with the following activity: a histamine antagonist (H2 sub-type).  
It is an agent intended for the treatment of gastric ulcers.  
Adverse effects of overexposure might include: altered heart rate and pulse; symptoms of hypersensitivity (such as skin rash, hives, itching, and difficulty breathing); temporary decrease in white blood cell counts; coughing; headache; increased mucous secretion.

**Other Adverse Effects**

None known for occupational exposure.

## 12. ECOLOGICAL INFORMATION

**\* Summary**

This material contains an active pharmaceutical ingredient that has been tested, and no environmental effects have been identified. No information is available about the potential of this product to produce adverse environmental effects. Local regulations and procedures should be consulted prior to environmental release.

Specific information on the active pharmaceutical ingredient is provided below.

### ECOTOXICITY

#### Aquatic

**\* Activated Sludge Respiration**

This material contains an active pharmaceutical ingredient that is not toxic to activated sludge microorganisms.

IC50: > 1000 mg/l, 3 Hours, Activated sludge

**\* Algal**

This material contains an active pharmaceutical ingredient that is not toxic to algae.

IC50: 167 mg/l, 72 Hours, Selenastrum capricornutum, green algae

NOEL: 56 mg/l, 72 Hours, Selenastrum capricornutum, green algae

**\* Daphnid**

This material contains an active pharmaceutical ingredient that is not toxic to daphids.

EC50: 730 mg/l, 48 Hours, Daphnia magna, Static test

NOEL: 347 mg/l, 48 Hours, Daphnia magna, Static test

**\* Fish**

This material contains an active pharmaceutical ingredient that is not toxic to fish.

Juvenile Oncorhynchus mykiss, rainbow trout

EC50: > 112 mg/l, 14 Days, Flow-through test

Juvenile Oncorhynchus mykiss, rainbow trout

NOEL: 112 mg/l, 14 Days, Flow-through test

### MOBILITY

**\* Solubility**

This material contains an active pharmaceutical ingredient that for environmental fate predictions has solubility in water.

**\* Volatility**

This material contains an active pharmaceutical ingredient that will not readily enter into the air from hard surfaces or from a container of the pure substance.

Henry's Law Constant 2.30E-11 atm m<sup>3</sup>/mol, Estimated at 24 C

**\* Adsorption**

This material contains an active pharmaceutical ingredient that is likely to adsorb to soil or sediment. It may persist in soil or sediment if released directly to the environment.

Soil Sediment Sorption 2.51 to 4.49 at pH 5 to 7  
(log Koc):

- \* Partitioning** This material contains an active pharmaceutical ingredient with octanol/water partition coefficient data that suggests that for environmental fate predictions the active pharmaceutical ingredient will not have the tendency to distribute into fats.

#### PERSISTENCE/DEGRADATION

- \* Hydrolysis** This material contains an active pharmaceutical ingredient that has been shown to be chemically stable in water. Hydrolysis is unlikely to be a significant depletion mechanism.  
Half-Life, Neutral: > 1 Years, Measured
- \* Photolysis** This material contains an active pharmaceutical ingredient that has been shown to be chemically unstable in water when exposed to light. Aqueous photolysis may be a significant depletion mechanism.  
Half-Life, Aqueous: 70 Minutes, Measured, Lake water  
UV/Visible Spectrum: 313 nm at pH 7, Measured
- \* Biodegradation** This material contains an active pharmaceutical ingredient that is not readily biodegradable but is inherently biodegradable (as defined by 1993 OECD Testing Guidelines) and is not expected to persist in the environment.  
Aerobic - Ready  
Percent Degradation: < 1 %, 28 days, Modified Sturm test.  
Aerobic - Inherent  
Percent Degradation: 43 %, 28 days, Modified Zahn-Wellens, primary biodegradation, loss of parent., Activated sludge  
Percent Degradation: 2 %, 28 days, Modified Zahn-Wellens, DOC removal., Activated sludge  
Anaerobic  
Percent Degradation: 12 %, 35 days  
Aerobic - Soil  
Percent Degradation: 3 to 10 %, 67 days
- \* BIOACCUMULATION** This material contains an active pharmaceutical ingredient that will not have a tendency to bioaccumulate in the food chain.  
Bioconcentration Factor: 1

### 13. DISPOSAL CONSIDERATIONS

- Disposal Recommendations** Collect for recycling or recovery if possible. The disposal method for rejected products/returned goods must ensure that they cannot be re-sold or re-used.
- Regulatory Requirements** Observe all local and national regulations when disposing of this product.

### 14. TRANSPORT INFORMATION

The SDS should accompany all shipments for reference in the event of spillage or accidental release. Only authorised persons trained and competent in accordance with appropriate national and international regulatory requirements may prepare dangerous goods for transport.

#### UN Classification and Labelling

- Transport Information** Transportation and shipping of this product is not restricted. It has no known, significant hazards requiring special packaging or labelling for air, maritime, US or European ground transport purposes.

### 15. REGULATORY INFORMATION

The information included below is an overview of the major regulatory requirements. It should not be considered to be an exhaustive summary. Local regulations should be consulted for additional requirements.

### EU Classification and Labelling

Exempt from requirements of EU Dangerous Preparations directive - product regulated as a medicinal product, cosmetic product or medical device.

For waste disposal purpose, this product should be classified in line with the European Waste Catalogue criteria.

### US OSHA Standard (29 CFR Part 1910.1200)

**Classification** This product is classified as hazardous according to the OSHA Hazard Communication Standard.

### Other US Regulations

**TSCA Status** Exempt

## 16. OTHER INFORMATION

**References** GSK Hazard Determination

**Date Approved/Revised** 18-Nov-2004

**SDS Version Number** 8

### SDS Sections Updated

#### Sections

ECOLOGICAL INFORMATION

#### Subsections

Activated Sludge Respiration  
Adsorption  
Algal  
Algal Degradation  
Bioaccumulation  
Biodegradation  
Daphnid  
Distribution  
Earthworm  
Ecotoxicity  
Fish  
Hydrolysis  
Microbial Growth Inhibition  
Microtox  
Mobility  
Other Adverse Effects  
Other Species - Aquatic  
Other Species - Terrestrial  
Partitioning  
Persistence/Degradation  
Photolysis  
Solubility  
Summary  
Volatility

The information and recommendations in this safety data sheet are, to the best of our knowledge, accurate as of the date of issue. Nothing herein shall be deemed to create any warranty, express or implied. It is the responsibility of the user to determine the applicability of this information and the suitability of the material or product for any particular purpose.

## 1. Identification

<b>Product identifier</b>	<b>ZANTAC INJECTION</b>
<b>Other means of identification</b>	Not available.
<b>Synonym(s)</b>	ZANTAC INJECTION 25 MG/ML * ANTAC INJECTION * AZANTAC INJECTION * SOSTRIL INJECTION * ZANTIC INJECTION * ZINETAC INJECTION * RANITIDINE HYDROCHLORIDE, FORMULATED PRODUCT
<b>Recommended use</b>	Medicinal Product
<b>Recommended restrictions</b>	No other uses are advised.
<b>Manufacturer/Importer/Supplier/Distributor information</b>	
<b>Manufacturer</b>	

GlaxoSmithKline US  
 5 Moore Drive  
 Research Triangle Park, NC 27709 USA  
 US General Information (normal business hours): +1-888-825-5249  
 Email Address: msds@gsk.com  
 Website: www.gsk.com  
 EMERGENCY PHONE NUMBERS -  
 TRANSPORT EMERGENCIES::  
 US / International toll call +1 703 527 3887  
 available 24 hrs/7 days; multi-language response

## 2. Hazard(s) identification

<b>Classified hazards</b>	Exempt from requirements - product regulated as a medicinal product, cosmetic product or medical device.
<b>Label elements</b>	Exempt from requirements - product regulated as a medicinal product, cosmetic product or medical device.
<b>Hazard(s) not otherwise classified (HNOC)</b>	Exempt from requirements - product regulated as a medicinal product, cosmetic product or medical device.

## 3. Composition/information on ingredients

<b>Mixtures</b>			
<b>Hazardous components</b>			
Chemical name	Common name and synonyms	CAS number	%
RANITIDINE HYDROCHLORIDE	AH 19065AB N,N-DIMETHYL-5-(2-(1-METHYLAMINO-2-N AMINE HYDROCHLORIDE 54 (GW ACN)	66357-59-3	3
Other components below reportable levels			97.0

\*Designates that a specific chemical identity and/or percentage of composition has been withheld as a trade secret.

## 4. First-aid measures

<b>Inhalation</b>	In case of accident by inhalation: remove casualty to fresh air and keep at rest. If not breathing, give artificial respiration. Oxygen or artificial respiration if needed. Do not use mouth-to-mouth method if victim inhaled the substance. Induce artificial respiration with the aid of a pocket mask equipped with a one-way valve or other proper respiratory medical device. Get medical attention immediately.
<b>Skin contact</b>	Immediately flush with plenty of water for at least 15 minutes while removing contaminated clothing and shoes. Get medical attention immediately. For minor skin contact, avoid spreading material on unaffected skin.
<b>Eye contact</b>	Rinse thoroughly with plenty of water for at least 15 minutes and consult a physician.



<b>Ingestion</b>	Call a physician or poison control center immediately. Only induce vomiting at the instruction of medical personnel. Never give anything by mouth to an unconscious person.
<b>Most important symptoms/effects, acute and delayed</b>	May cause allergic skin reaction. May cause allergic respiratory reaction. Sensitization. Direct contact with eyes may cause temporary irritation. The following adverse effects have been noted with therapeutic use of this material: decrease in heart rate; decrease in blood pressure; temporary decrease in white blood cell counts; coughing; headache; increased mucous secretion.
<b>Indication of immediate medical attention and special treatment needed</b>	Provide general supportive measures and treat symptomatically. Symptoms may be delayed. No specific antidotes are recommended. Treat according to locally accepted protocols. For additional guidance, refer to the local poison control information centre.
<b>General information</b>	Ensure that medical personnel are aware of the material(s) involved, and take precautions to protect themselves. Wash contaminated clothing before reuse. The need for pre-placement and periodic health surveillance must be determined by risk assessment. Following assessment, if the risk of exposure is considered significant then exposed individuals should receive health surveillance focused on detecting respiratory symptoms and including respiratory function testing. In the event of overexposure, individuals should receive post exposure health surveillance focused on detecting respiratory conditions and other allergy symptoms.

## 5. Fire-fighting measures

<b>Suitable extinguishing media</b>	Water fog. Foam. Dry chemical powder. Carbon dioxide (CO <sub>2</sub> ).
<b>Unsuitable extinguishing media</b>	None known.
<b>Specific hazards arising from the chemical</b>	During fire, gases hazardous to health may be formed.
<b>Special protective equipment and precautions for firefighters</b>	Self-contained breathing apparatus and full protective clothing must be worn in case of fire.
<b>Fire-fighting equipment/instructions</b>	Move containers from fire area if you can do so without risk.

## 6. Accidental release measures

<b>Personal precautions, protective equipment and emergency procedures</b>	Keep unnecessary personnel away. Keep people away from and upwind of spill/leak. Keep out of low areas. Wear appropriate protective equipment and clothing during clean-up. Do not touch damaged containers or spilled material unless wearing appropriate protective clothing. Avoid inhalation of vapors or mists. Ensure adequate ventilation. Local authorities should be advised if significant spillages cannot be contained. For personal protection, see section 8 of the MSDS.
<b>Methods and materials for containment and cleaning up</b>	Large Spills: Stop the flow of material, if this is without risk. Dike the spilled material, where this is possible. Cover with plastic sheet to prevent spreading. Absorb in vermiculite, dry sand or earth and place into containers. Prevent entry into waterways, sewer, basements or confined areas. Following product recovery, flush area with water.  Small Spills: Wipe up with absorbent material (e.g. cloth, fleece). Clean surface thoroughly to remove residual contamination.  Never return spills in original containers for re-use. For waste disposal, see section 13 of the MSDS.
<b>Environmental precautions</b>	Avoid discharge into drains, water courses or onto the ground.

## 7. Handling and storage

<b>Precautions for safe handling</b>	Avoid breathing mist or vapor. Avoid contact with skin. Avoid prolonged exposure. Provide adequate ventilation. Wear appropriate personal protective equipment. Observe good industrial hygiene practices.
<b>Conditions for safe storage, including any incompatibilities</b>	Store in original tightly closed container. Store in a cool, dry place out of direct sunlight. Store in a well-ventilated place. Store away from incompatible materials (see Section 10 of the MSDS).

## 8. Exposure controls/personal protection

### Occupational exposure limits

GSK Components	Type	Value	Note
RANITIDINE HYDROCHLORIDE (CAS 66357-59-3)	15 MIN STEL	50 mcg/m <sup>3</sup>	SKIN SENSITISER
		50 mcg/m <sup>3</sup>	RESPIRATORY SENSITISER
	OHC	3	
<b>Biological limit values</b>	No biological exposure limits noted for the ingredient(s).		

<b>Appropriate engineering controls</b>	An Exposure Control Approach (ECA) is established for operations involving this material based upon the OEL/Occupational Hazard Category and the outcome of a site- or operation-specific risk assessment. Good general ventilation (typically 10 air changes per hour) should be used. Ventilation rates should be matched to conditions. If applicable, use process enclosures, local exhaust ventilation, or other engineering controls to maintain airborne levels below recommended exposure limits. If exposure limits have not been established, maintain airborne levels to an acceptable level.
<b>Individual protection measures, such as personal protective equipment</b>	
<b>Eye/face protection</b>	Avoid contact with eyes. Face-shield. Wear a full-face respirator, if needed. Eye wash fountain is recommended.
<b>Hand protection</b>	The choice of an appropriate glove does not only depend on its material but also on other quality features and is different from one producer to the other. Glove selection must take into account any solvents and other hazards present.
<b>Other</b>	Wear suitable protective clothing.
<b>Respiratory protection</b>	Do not breathe dust/fume/gas/mist/vapors/spray. Wear positive pressure self-contained breathing apparatus (SCBA).
<b>Thermal hazards</b>	Wear appropriate thermal protective clothing, when necessary.
<b>General hygiene considerations</b>	For advice on suitable monitoring methods, seek guidance from a qualified environment, health and safety professional. Always observe good personal hygiene measures, such as washing after handling the material and before eating, drinking, and/or smoking. Routinely wash work clothing and protective equipment to remove contaminants. Contaminated work clothing should not be allowed out of the workplace.

## 9. Physical and chemical properties

### Appearance

<b>Physical state</b>	Liquid.
<b>Form</b>	Solution.
<b>Color</b>	Not available.

**Odor** Not available.

**Odor threshold** Not available.

**pH** 6.8 - 7.1

**Melting point/freezing point** Not available.

**Initial boiling point and boiling range** Not available.

**Flash point** Not available.

**Evaporation rate** Not available.

**Flammability (solid, gas)** Not available.

### Upper/lower flammability or explosive limits

**Flammability limit - lower (%)** Not available.

**Flammability limit - upper (%)** Not available.

**Explosive limit - lower (%)** Not available.

**Explosive limit - upper (%)** Not available.

**Vapor pressure** Not available.

**Vapor density** Not available.

**Relative density** Not available.

**Solubility(ies)** Not available.

**Partition coefficient (n-octanol/water)** Not available.

**Auto-ignition temperature** Not available.

**Decomposition temperature** Not available.

**Viscosity** Not available.

## 10. Stability and reactivity

**Reactivity** The product is stable and non-reactive under normal conditions of use, storage and transport.

**Chemical stability** Material is stable under normal conditions.

<b>Possibility of hazardous reactions</b>	No dangerous reaction known under conditions of normal use.
<b>Conditions to avoid</b>	Contact with incompatible materials.
<b>Incompatible materials</b>	Strong oxidizing agents.
<b>Hazardous decomposition products</b>	Irritating and/or toxic fumes and gases may be emitted upon the products decomposition.

## 11. Toxicological information

### Information on likely routes of exposure

<b>Ingestion</b>	May be harmful if swallowed.
<b>Inhalation</b>	May cause allergy or asthma symptoms or breathing difficulties if inhaled.
<b>Skin contact</b>	May cause an allergic skin reaction.
<b>Eye contact</b>	Direct contact with eyes may cause temporary irritation.

**Symptoms related to the physical, chemical and toxicological characteristics** Sensitization. The following adverse effects have been noted with therapeutic use of this material: decrease in heart rate; decrease in blood pressure; temporary decrease in white blood cell counts; coughing; headache; increased mucous secretion.

### Information on toxicological effects

**Acute toxicity** May cause allergic skin reaction.

Components	Species	Test Results
RANITIDINE HYDROCHLORIDE (CAS 66357-59-3)		
<b>Acute</b>		
<i>Oral</i>		
LD50	Rat	> 1000 mg/kg

\* Estimates for product may be based on additional component data not shown.

**Skin corrosion/irritation** Prolonged skin contact may cause temporary irritation.

#### Irritation Corrosion - Skin

RANITIDINE HYDROCHLORIDE	Acute dermal irritation; OECD 404, Primary dermal irritation index = 0 Result: Negative Species: Rabbit
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**Serious eye damage/eye irritation** Avoid contact with eyes.

#### Eye

RANITIDINE HYDROCHLORIDE	Acute ocular irritation; OECD 405, Kay and Calandra score = 3 Result: Minimal Irritant Species: Rabbit IRE Assay Result: Negative; not likely to be a severe irritant Species: Rabbit
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**Respiratory sensitization** May cause allergy or asthma symptoms or breathing difficulties if inhaled.

RANITIDINE HYDROCHLORIDE	Occupational exposure Result: Positive Species: Human
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**Skin sensitization** May cause an allergic skin reaction.

#### Sensitization

RANITIDINE HYDROCHLORIDE	Occupational exposure Result: Positive Species: Human Optimisation Test Result: Weak sensitiser Species: Guinea pig
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**Germ cell mutagenicity** No data available to indicate product or any components present at greater than 0.1% are mutagenic or genotoxic.

RANITIDINE HYDROCHLORIDE	Ames Assay, GLP assay Result: Negative Chromosomal Aberration Assay In Vitro, human lymphocytes, Ranitidine bismuth citrate tested Result: Positive Chromosomal Aberration Assay In Vivo; germ cells, Maximum dose = 1000 mg/kg Result: Negative Species: Mouse
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GreenScreen Assay  
 Result: Negative  
 Micronucleus Test  
 Result: Negative  
 Species: Rat  
 Mouse Lymphoma Cell (L5178Y) Mutation Assay, GLP assay  
 Result: Negative  
 SOS/umu Assay  
 Result: Negative  
 Unscheduled DNA Synthesis in vivo, Maximum dose = 200 mg/kg  
 Result: Negative  
 Species: Rat  
 Organ: Stomach  
 Yeast Mutation Assay  
 Result: Negative

**Carcinogenicity** This product is not considered to be a carcinogen by IARC, ACGIH, NTP, or OSHA.  
 RANITIDINE HYDROCHLORIDE 2 year bioassay, Maximum dose = 2000 mg/kg/day  
 Result: Negative  
 Species: Mouse  
 2 year bioassay, Maximum dose = 2000 mg/kg/day  
 Result: Negative  
 Species: Rat

**Reproductive toxicity** This product is not expected to cause reproductive or developmental effects.  
 RANITIDINE HYDROCHLORIDE Embryo-foetal development - Oral  
 Result: Foetal NOAEL = 100 mg/kg/day (maximum dose);  
 Maternal NOAEL = 25 mg/kg/day (decreased weight gain at 50 and 100 mg/kg/day)  
 Species: Rat  
 Embryo-foetal development - Oral  
 Result: NOAEL = 100 mg/kg/day (maximum dose)  
 Species: Rabbit  
 Fertility  
 Result: NOAEL / fertility = 100 mg/kg/day (male) and 200 mg/kg/day (female) (maximum doses)  
 Species: Rat

**Specific target organ toxicity - single exposure** None known.

**Specific target organ toxicity - repeated exposure** None known.

**Aspiration hazard** Not available.

**Chronic effects** Prolonged inhalation may be harmful.

**Further information** Caution - Pharmaceutical agent.

## 12. Ecological information

**Ecotoxicity** The product is not classified as environmentally hazardous. However, this does not exclude the possibility that large or frequent spills can have a harmful or damaging effect on the environment.

Components		Species	Test Results
RANITIDINE HYDROCHLORIDE (CAS 66357-59-3)			
Aquatic			
Acute			
Activated Sludge Respiration	IC50	Residential sludge	> 1000 mg/l, 3 hours, OECD 209
Algae	EC50	Green algae (Selenastrum capricornutum)	167 mg/l, 72 hours, OECD 201
	NOEC	Green algae (Selenastrum capricornutum)	56 mg/l, 72 hours
Crustacea	EC50	Water flea (Daphnia magna)	730 mg/l, 48 hours, Static test, OECD 202
	NOEC	Water flea (Daphnia magna)	347 mg/l, 48 hours, Static test
Fish	EC50	Rainbow trout (Juvenile Oncorhyncus mykiss)	> 112 mg/l, 14 days, Flow-through test, OECD 203
	NOEC	Rainbow trout (Juvenile Oncorhyncus mykiss)	112 mg/l, 14 days, Flow-through test

Components		Species	Test Results
Chronic Crustacea	LOEC	Water flea (Ceriodaphnia dubia)	100 mg/l, 8 days, Static renewal test, EPA 1002
	NOEC	Water flea (Ceriodaphnia dubia)	32 mg/l, 8 days
	* Estimates for product may be based on additional component data not shown.		
Persistence and degradability			
Photolysis			
Half-life (Photolysis-aqueous)			
	RANITIDINE HYDROCHLORIDE		70 Minutes Measured, Lake water
UV/visible spectrum wavelength			
	RANITIDINE HYDROCHLORIDE		313 nm Measured, pH 7
Hydrolysis			
Half-life (Hydrolysis-neutral)			
	RANITIDINE HYDROCHLORIDE		> 1 Years Measured
Biodegradability			
Percent degradation (Aerobic biodegradation-soil)			
	RANITIDINE HYDROCHLORIDE		3 - 10 %, 67 days
Percent degradation (Anaerobic biodegradation)			
	RANITIDINE HYDROCHLORIDE		12 %, 35 days
Bioaccumulative potential		Not available.	
Partition coefficient n-octanol / water (log Kow)			
	RANITIDINE HYDROCHLORIDE		0.0815
Mobility in soil			
Adsorption			
Soil/sediment sorption - log Koc			
	RANITIDINE HYDROCHLORIDE		2.51 - 4.49, pH 5-7
Mobility in general			
Volatility			
Henry's law			
	RANITIDINE HYDROCHLORIDE		0 atm m^3/mol, 24 C Estimated
Distribution			
Octanol/water distribution coefficient log DOW			
	RANITIDINE HYDROCHLORIDE		-1.09, pH 7 -2.5, pH 5 0.14, pH 9
Other adverse effects		Not available.	

### 13. Disposal considerations

<b>Disposal instructions</b>	Collect and reclaim or dispose in sealed containers at licensed waste disposal site. Dispose of contents/container in accordance with local/regional/national/international regulations.
<b>Local disposal regulations</b>	Dispose in accordance with all applicable regulations.
<b>Hazardous waste code</b>	The waste code should be assigned in discussion between the user, the producer and the waste disposal company.
<b>Waste from residues / unused products</b>	Dispose of in accordance with local regulations. Empty containers or liners may retain some product residues. This material and its container must be disposed of in a safe manner (see: Disposal instructions).
<b>Contaminated packaging</b>	Empty containers should be taken to an approved waste handling site for recycling or disposal. Since emptied containers may retain product residue, follow label warnings even after container is emptied.

### 14. Transport information

<b>DOT</b>	Not regulated as a dangerous good.
<b>IATA</b>	Not regulated as a dangerous good. Read safety instructions, SDS and emergency procedures before handling.
<b>IMDG</b>	Not regulated as a dangerous good.

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

MARPOL Annex II applies to liquids used in a ship's operation that pose a threat to the marine environment. These materials may not be transported in bulk.

## 15. Regulatory information

### US federal regulations

This product is a "Hazardous Chemical" as defined by the OSHA Hazard Communication Standard, 29 CFR 1910.1200.

#### TSCA Section 12(b) Export Notification (40 CFR 707, Subpt. D)

Not regulated.

#### CERCLA Hazardous Substance List (40 CFR 302.4)

Not listed.

#### US. OSHA Specifically Regulated Substances (29 CFR 1910.1001-1050)

Not listed.

#### SARA 304 Emergency release notification

Not regulated.

### Superfund Amendments and Reauthorization Act of 1986 (SARA)

#### Hazard categories

Immediate Hazard - Yes  
Delayed Hazard - Yes  
Fire Hazard - No  
Pressure Hazard - No  
Reactivity Hazard - No

#### SARA 302 Extremely hazardous substance

No

#### SARA 311/312 Hazardous chemical

No

### NFPA ratings

Health: 2  
Flammability: 0  
Instability: 0

### HMIS® ratings

Health: 2  
Flammability: 0  
Physical hazard: 0

### Other federal regulations

#### Clean Air Act (CAA) Section 112 Hazardous Air Pollutants (HAPs) List

Not regulated.

#### Clean Air Act (CAA) Section 112(r) Accidental Release Prevention (40 CFR 68.130)

Not regulated.

#### Safe Drinking Water Act (SDWA)

Not regulated.

#### Food and Drug Administration (FDA)

Not regulated.

### US state regulations

#### US. Massachusetts RTK - Substance List

Not regulated.

#### US. New Jersey Worker and Community Right-to-Know Act

Not regulated.

#### US. Pennsylvania RTK - Hazardous Substances

Not regulated.

#### US. Rhode Island RTK

Not regulated.

#### US. California Proposition 65

California Safe Drinking Water and Toxic Enforcement Act of 1986 (Proposition 65): This material is not known to contain any chemicals currently listed as carcinogens or reproductive toxins.

### International Inventories

Country(s) or region	Inventory name	On inventory (yes/no)*
Australia	Australian Inventory of Chemical Substances (AICS)	No
Canada	Domestic Substances List (DSL)	No
Canada	Non-Domestic Substances List (NDSL)	No
China	Inventory of Existing Chemical Substances in China (IECSC)	No

Country(s) or region	Inventory name	On inventory (yes/no)*
Europe	European Inventory of Existing Commercial Chemical Substances (EINECS)	Yes
Europe	European List of Notified Chemical Substances (ELINCS)	No
Japan	Inventory of Existing and New Chemical Substances (ENCS)	No
Korea	Existing Chemicals List (ECL)	No
New Zealand	New Zealand Inventory	Yes
Philippines	Philippine Inventory of Chemicals and Chemical Substances (PICCS)	No
United States & Puerto Rico	Toxic Substances Control Act (TSCA) Inventory	No

\*A "Yes" indicates that all components of this product comply with the inventory requirements administered by the governing country(s)  
A "No" indicates that one or more components of the product are not listed or exempt from listing on the inventory administered by the governing country(s).

## 16. Other information, including date of preparation or last revision

<b>Issue date</b>	11-29-2013
<b>Revision date</b>	11-29-2013
<b>Version #</b>	13
<b>Further information</b>	HMIS® is a registered trade and service mark of the NPCA.
<b>HMIS® ratings</b>	Health: 2 Flammability: 0 Physical hazard: 0
<b>NFPA ratings</b>	Health: 2 Flammability: 0 Instability: 0
<b>References</b>	GSK Hazard Determination
<b>Disclaimer</b>	The information and recommendations in this safety data sheet are, to the best of our knowledge, accurate as of the date of issue. Nothing herein shall be deemed to create any warranty, express or implied. It is the responsibility of the user to determine the applicability of this information and the suitability of the material or product for any particular purpose.
<b>Revision Information</b>	Other information, including date of preparation or last revision: Further information