Safety Data Sheet (SDS) NITRIC ACID

Section 1:	Product and Company Ident	ification		
Product Name:	NITRIC ACID	Index Number:	007-004-00-1	
Product Number(s):				
	S050101, S050102, S050106, S050110, S050101-SSNQ03, S050101-SSNQ09, S050102-SSEH03, S050102-SSEH11, S050102-SSNH03, S050102-SSNH11, S050106-SSEJ05, S050106-SSNJ05, S050110-SSEF01, S050110-SSEF02, S050110-SSEF03, S050110-SSEF04, S050110-			
SSEF05, S050110-SSEF06,	S050110-SSEF07, S050110-SSEF08, S05011			
S050110-SSNF05, S050110-	SSNF06, S050110-SSNF07, S050110-SSNF08			
Synonyms:	Aqua fortis; Azotic acid; Hydrogen nitr			
Chemical names:	DE Salpetersäure; ES Ácido nítrico; FR	Acide nitrique; IT Acid	o nitric; NL Salpeterzuur	
Supplier:	SEASTAR CHEMICALS Inc.			
Address:	10005 McDonald Park Road, Sidney, B		1	
Phone Number:	250-655-5880	Fax Number:	250-655-5888	
CANUTEC (CAN):	613-996-6666			
Section 2:	Hazards Identification			
	Emergency O	verview		
Appearance:	Clear, colourless or yellowish liquid			
Target Organs:	Eyes, skin, respiratory system, teeth.			
	on in accordance with 29 CFR 1910 (OSH/	,	regulation (EC) No 1272/2008	
Classification:	Skin corrosion – Category 1B	Pictograms:		
Signal Word:	Corrosive to metals – Category 1		GHS05	
Hazard Statements:	Danger			
	kin burns and eye damage.			
H290: May be corrosive	5 6			
Precautionary Statem	nents:			
P234: Keep only in orig				
P260: Do not breathe fu P264: Wash thoroughly	umes/gas/mist/vapours/spray.			
	gloves/protective clothing/eye protection	n/face protection.		
	SWALLOWED: Rinse mouth. Do NOT ir			
	ON SKIN (or hair): Take off immediate	5	5	
P304+P340: IF INHALED: Remove victim to fresh air and keep at rest in a position comfortable for breathing.				
P305+P351+P338: IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing.				
P310: Immediately call a POISON CENTER or doctor.				
P363: Wash contaminated clothing before reuse.				
P390: Absorb spillage to prevent material damage.				
P405: Store locked up.				
P406: Store in corrosion resistant container with a resistant inner liner. P501: Dispose of contents/container according to federal, regional and local government requirements.				
	inscontainer according to rederal, regio	inai anu iocal governme		
Section 3:	Composition/Information on	Ingredients		

Section 3:	composition/informat	tion on ingredients	
CAS No.	Chemical Name	Percent	EINECS / ELINCS No.
7697-37-2	Nitric acid	5 – 20%	231-714-2
7732-18-5	Water	Balance	231-791-2



Section 4:	First Aid Measures
In case of contact:	
Inhalation:	Take proper precautions to ensure your own safety before attempting rescue (e.g. wear appropriate protective equipment, use the "buddy" system). Remove source of contamination or move victim to fresh air. If breathing is difficult, oxygen may be beneficial if administered by trained personnel, preferably on a doctor's advice. Do not allow victim to move about unnecessarily. Symptoms of pulmonary edema can be delayed up to 48 hours after exposure. Immediately transport victim to an emergency care facility.
Skin:	Avoid direct contact with this chemical. Wear chemical protective clothing, if necessary. Immediately remove contaminated clothing, shoes, and leather goods. As quickly as possible, flush contaminated area with lukewarm, gently running water for at least 20-30 minutes, by the clock. DO NOT INTERRUPT FLUSHING. If necessary, keep emergency vehicle waiting. Immediately transport victim to an emergency care facility. Discard contaminated clothing, shoes and leather goods. Keep contaminated clothing under water in a closed container until it can be safely discarded.
Eye:	Avoid direct contact. Wear chemical protective gloves, if necessary. Immediately flush the contaminated eye(s) with lukewarm, gently flowing water for at least 20-30 minutes, by the clock, holding the eyelid(s) open. Neutral saline may be used as soon as it is available. DO NOT INTERRUPT FLUSHING. If necessary, keep emergency vehicle waiting. Take care not to rinse contaminated water into the unaffected eye or onto the face. If irritation persists, repeat flushing. Quickly transport victim to an emergency care facility. Do NOT allow victim to rub or keep eyes closed.
Ingestion:	NEVER give anything by mouth if victim is rapidly losing consciousness, or is unconscious or convulsing. Have victim rinse mouth thoroughly with water. DO NOT INDUCE VOMITING. Have victim drink 240 to 300 mL (8 to 10 oz.) of water. If milk is available, it may be administered after the water has been given. If vomiting occurs naturally, repeat administration of water. Quickly transport victim to an emergency facility.
Notes to Physician/Doctor:	Provide general supportive measures (comfort, warmth, rest). Consult a doctor and/or the nearest Poison Control Centre for all exposures except minor instances of inhalation or skin contact. All first aid procedures should be periodically reviewed by a doctor familiar with the material and its conditions of use in the workplace.

Section 5: Fire Fighting Measures

Fire Hazard Summary:

Nitric acid does not burn. Can react with many metals, particularly in powdered form, to form extremely flammable hydrogen gas. Generates heat when mixed with water. During a fire, nitric acid decomposes with the release of corrosive nitrogen oxide gases. Explosive decomposition may occur under fire conditions and closed containers may rupture violently due to rapid decomposition, if exposed to fire or excessive heat for a sufficient period of time. Firefighter's normal protective equipment (Bunker Gear) will not provide adequate protection. Chemical protective clothing (e.g. chemical splash suit) and positive pressure self-contained breathing apparatus (NIOSH approved or equivalent) may be necessary.

Extinguishing	Nitric acid does not burn. Extinguish fire using extinguishing agent suitable for the surrounding	
Media:	fire and not contraindicated for use with nitric acid. Nitric acid is an oxidizer. Therefore, flooding quantities of water spray or fog should be used to fight fires involving nitric acid.	
Extinguishing Media to be Avoided:	DO NOT use dry chemical powders containing sodium bicarbonate, potassium bicarbonate, sodium carbonate, calcium carbonate, ammonium phosphate, or ammonium sulfate. Nitric acid may react violently with these extinguishing agents.	
Flash Point:	Not combustible (does not burn).	
Lower Flammable (Explosiv	e) Limit (LFL/LEL):	Not applicable
Upper Flammable (Explosive) Limit (UFL/UEL):		Not applicable
Autoignition Temperature:		Not applicable
Sensitivity to Mechanical Impact:		Probably not sensitive. Normally stable.
Sensitivity to Static Charge:		Will not accumulate static charge or be ignited by a static discharge.
Electrical Conductivity:		3.77×10 ¹² pS/m (0°C, 100% nitric acid)



Minimum Ignition Energy:	Not applicable
Combustion and Thermal	Liquid decomposes to a limited extent when heated, producing corrosive
Decomposition Products:	nitrogen oxides.
Decomposition Products:	Thill ogen oxides.

NATIONAL FIRE PROTECTION ASSOCIATION (NFPA) HAZARD IDENTIFICATION

Nitric acid	Less than 40%
Health:	3 – Short exposure could cause serious temporary or residual injury.
Flammability:	0 – Will not burn under typical fire conditions.
Reactivity:	0 – Normally stable, even under fire conditions, and not reactive with water.
Special Hazard:	Not applicable

Section 6: Accidental Release Measures

Spill Precautions:

Restrict access to area until completion of clean-up. Ensure clean-up is conducted by trained personnel only. Remove or isolate flammable and combustible materials. Ventilate area. Dangerous levels of nitrogen oxides may form during spills of nitric acid. Consider evacuation of down-wind areas. Wear adequate personal protective equipment. Extinguish or remove all ignition sources. Notify government occupational health and safety and environmental authorities.

Clean-up:

Do not touch spilled material. Prevent material from entering sewers, waterways or confined spaces. Keep materials that can burn away from spilled material. Stop or reduce leak if safe to do so. Contain spill with earth, sand, or absorbent material that does not react with spilled material. Do not use sawdust or other organic materials, which will react with nitric acid creating a fire or health hazard.

<u>SMALL SPILLS</u>: Soak up spill with absorbent material that does not react with spilled chemical. Put material in suitable, covered, labelled containers. Flush area with large quantities of water. Contaminated absorbent material will pose the same hazards as the spilled product. Only trained personnel should attempt to neutralize spills. Neutralizing spill with sodium bicarbonate, sodium carbonate or calcium carbonate will produce large amounts of carbon dioxide gas. Ensure adequate ventilation.

LARGE SPILLS: Evacuate area. Contact fire and emergency services and supplier for advice.

Section 7: Handling and Storage

Handling:

This material is CORROSIVE (to eyes, skin and respiratory tract). Before handling, it is extremely important that engineering controls are operating and that protective equipment requirements and personal hygiene measures are being followed. People working with this chemical should be properly trained regarding its hazards and its safe use. Maintenance and emergency personnel should be advised of potential hazards.

Unprotected persons should avoid all contact with this chemical including contaminated equipment. Immediately report leaks, spills or failures of the engineering controls. If nitric acid is released, immediately put on a suitable respirator and leave the area until the severity of the release is determined. In case of leaks or spills, escape-type respiratory protective equipment should be available in the work area.

Be aware of typical signs and symptoms of poisoning and first aid procedures. Any signs of illness should be reported immediately to supervisory personnel. Seek medical attention for all exposures even if an exposure did not seem excessive. Symptoms of a severe exposure can be delayed.

Avoid contact with all incompatible materials. Nitric acid is a strong acid and is very reactive. It is not combustible but it enhances the combustion of other substances (nitric acid greater than 40%). Nitric acid may react violently or explosively with many organic and inorganic chemicals. Flammable hydrogen gas is released on contact with many common metals, particularly metal in powdered form. Significant heat is generated upon contact with water. See Section 10 for more information.

Avoid generating vapours or mists. Prevent the release of vapours or mists into the workplace. If possible, use closed handling systems for processes involving this material. If a closed handling system is not possible, use the smallest possible amounts in a well-ventilated area, separate from the storage area. Ensure that handling systems are corrosion-



resistant.

Inspect containers for damage or leaks before handling. Label containers. Handle containers carefully to avoid damage. Keep containers tightly closed when not in use to avoid spillage, vapour release or contamination of the contents. Never return unused or contaminated material to its original container.

Cautiously, dispense into sturdy containers made of compatible materials. Use corrosion-resistant transfer equipment when dispensing. Secondary protective containers must be used when this material is being carried. When diluting, always add acid to cold water slowly and in small amounts. Never use hot water and never add water to the acid. Water added to acid can cause uncontrolled boiling and splashing.

Always assume that empty containers contain hazardous residues. Never reuse empty containers, even if they appear to be clean. Do not perform any welding, cutting, soldering, drilling or other hot work on an empty vessel, container or piping until all material has been cleared. Have suitable emergency equipment for fires, spills and leaks readily available. Practice good housekeeping. Maintain handling equipment. Comply with applicable regulations.

Storage:

Store in a cool, dry, well-ventilated area, out of direct sunlight and away from heat. Keep quantities stored as small as possible. Store away from incompatible materials such as flammable materials, oxidizing materials, reducing materials, and strong bases. See Section 10 for more information.

Use corrosion-resistant structural materials, lighting and ventilation systems in the storage area. Wood and other organic/combustible materials should not be used on floors, structural materials and ventilation systems in the storage area. Storage area should be clearly identified, clear of obstruction and accessible only to trained and authorized personnel.

Keep storage area separate from work areas. Post warning signs. Inspect all incoming containers to make sure they are properly labelled and not damaged. Always store in original labelled container. Suitable storage may include glass bottles and carboys. Protect the label and keep it visible. Keep containers tightly closed when not in use and when empty. Protect from damage. Store containers at a convenient height for handling, below eye level if possible. Inspect storage area regularly for deficiencies, including damaged or leaking containers, signs of corrosion or poor housekeeping.

Keep absorbents for leaks and spills readily available. Contain spills or leaks by storing in trays made from compatible materials. Provide raised sills or ramps at doorways or create a trench which drains to a safe location. Keep empty containers in separate storage area. Assume that empty containers contain hazardous residues. Keep tightly closed. Avoid bulk storage indoors. Storage tanks should be above ground and surrounded with a dike capable of holding entire contents.

Store according to the occupational health and safety regulations and fire and building codes which will describe the kind of storage area and the type of storage containers for a specified amount of the material. Have appropriate fire extinguishers available in and near the storage area.

Section 8: Exposure Controls/Personal Protection

General Exposure Precautions:

NOTE: Exposure to this material can be controlled in many ways. The measures appropriate for a particular worksite depend on how this material is used and on the extent of exposure. This general information can be used to help develop specific control measures. Ensure that control systems are properly designed and maintained. Comply with occupational, environmental, fire, and other applicable regulations.

Engineering Controls:

Engineering methods to control hazardous conditions are preferred. Methods include mechanical ventilation (dilution and local exhaust), process or personnel enclosure, control of process conditions, and process modification (e.g., substitution of a less hazardous material). Administrative controls and personal protective equipment may also be required.

Because of the high potential hazard associated with this substance, stringent control measures such as enclosure or isolation may be necessary. Use a corrosion-resistant local exhaust ventilation system separate from other exhaust ventilation systems. Exhaust directly to the outside. Supply sufficient replacement air to make up for air removed by exhaust system. Treatment of exhaust emissions to prevent environmental contamination may be required.

NOTE: Do not use organic or combustible materials such as wood in the construction of ventilation or control systems.



Personal Protective Equipment:

If engineering controls and work practices are not effective in controlling exposure to this material, then wear suitable personal protective equipment including approved respiratory protection. Have appropriate equipment available for use in emergencies such as spills or fire.

If respiratory protection is required, institute a complete respiratory protection program including selection, fit testing, training, maintenance and inspection. Refer to the CSA Standard Z94.4-11, "Selection, Use and Care of Respirators," available from the Canadian Standards Association.

Eye / Face protection:	Wear chemical safety goggles as described by OSHA's eye and face protection regulations in 29 CFR 1910.133 or European Standard EN166. A face shield may also be necessary.			
Skin protection:	Wear impervious gloves and appropriate protective clothing. Choose body protection according to the amount and concentration of the substance at the work place. Have a safety shower/eye-wash readily available in the immediate work area.			
Resistance of Materials for Protective Clothing:	Guidelines for Nitric acid, less than 30 percent: RECOMMENDED (resistance to breakthrough longer than 8 hours): Butyl rubber; Natural rubber; Neoprene rubber; Polyvinyl chloride; Viton [™] ; Viton [™] /Butyl rubber; Barrier (PE/PA/PE); Trellchem [™] HPS and VPS; Tychem [™] SL (Saranex [™]), CPF 3, F, BR/LV, Responder [™] , and TK. RECOMMENDED (resistance to breakthrough longer than 4 hours): Polyethylene; Silver Shield/4H [™] (polyethylene/ethylene vinyl alcohol). CAUTION, use for short periods only (resistance to breakthrough within 1 to 4 hours): Nitrile rubber. NOT RECOMMENDED for use (resistance to breakthrough less than 1 hour): Polyvinyl alcohol.			
Inhalation / Ventilation:	NIOSH/OSHA RECOMMENDATIONS FOR NITRIC ACID CONCENTRATIONS IN AIR: UP TO 25 ppm: SAR operated in a continuous-flow mode; or full face piece chemical cartridge respirator with cartridge(s) to protect against nitric acid; or gas mask with canister to protect against nitric acid; or full face piece SCBA; or full face piece SAR. EMERGENCY OR PLANNED ENTRY INTO UNKNOWN CONCENTRATION OR IDLH CONDITIONS: Positive pressure, full face piece SCBA; or positive pressure, full face piece SAR with an auxiliary positive pressure SCBA. ESCAPE: Gas mask with canister to protect against nitric acid; or escape-type SCBA.			
Personal Hygiene:	Remove contaminated clothing immediately. Keep contaminated clothing thoroughly wet or immersed in water in closed containers. Discard or launder before rewearing. Inform laundry personnel of contaminant's hazards. Do not eat or drink in work areas. Wash hands thoroughly after handling this material. Maintain good housekeeping.			
EXPOSURE G	JIDEL	INES – Listed under Nitric acid	l, as HNO₃	
NIOSH:			25 ppm	
ACGIH:	TLV-T	WA: 2 ppm; TLV-STEL: 4 ppm		
OSHA PELs:	PEL-T	WA: 2 ppm; PEL-T-TWA: 2 ppm	n (5 mg/m³); PEL-STEL: 4 ppm	
Section 9:		Physical and Chemica	al Properties	
Form:		Liquid	Melting/Freezing Point:	50% w/w: -20 °C (-4 °F)
Colour:		Clear to yellow	Boiling Point:	10% w/w: 102.2 °C (215.96 °F) 20% w/w: 104.4 °C (219.92 °F)
Odour:		Strong – acrid odour	pH:	1.0 (0.1 M solution)
Odour Thresh	old:	0.29 - 0.98 ppm (detection)	Density: (at 20 °C)	5% w/w: 1.0256 g/mL
Chemical For		HNO ₃		10% w/w: 1.0543 g/mL
Formula Weig	ght:	63.01 g/mol		20% w/w: 1.115 g/mL
Vapour Densi	ty:	2.17 (air=1) (calculated)	Solubility in Water:	Soluble in all proportions.

Form:	Liquid	Melting/Freezing Point:	50% w/w: -20 °C (-4 °F)
Colour:	Clear to yellow	Boiling Point:	10% w/w: 102.2 °C (215.96 °F)
			20% w/w: 104.4 °C (219.92 °F)
Odour:	Strong – acrid odour	pH:	1.0 (0.1 M solution)
Odour Threshold:	0.29 - 0.98 ppm (detection)	Density: (at 20 °C)	5% w/w: 1.0256 g/mL
Chemical Formula:	HNO ₃		10% w/w: 1.0543 g/mL
Formula Weight:	63.01 g/mol		20% w/w: 1.115 g/mL
Vapour Density:	2.17 (air=1) (calculated)	Solubility in Water:	Soluble in all proportions.
Vapour Pressure: (Partial pressure, 25 °C)	40% w/w: 0.12 mmHg	Solubility in Other Liquids:	Reacts with many organic solvents (e.g. alcohols, ketones, ethers, esters)



Section 10: Stability and Reactivity

Normally stable. Nitric acid has a tendency to slowly decompose at room temperature to form nitrogen oxides, which may colour the acid yellow or red. The decomposition is accelerated by exposure to light and increases in temperature.

Oxidizing Properties:

The NFPA lists nitric acid (40% or less) as a Class 1 oxidizer. A Class 1 oxidizer meets the definition of an oxidizer (any material that readily yields oxygen or other oxidizing gas, or that readily reacts to promote or initiate combustion of combustible materials) and does not moderately increase the burning rate of combustible materials with which it comes into contact.

Incompatibility – Materials to be Avoided:

NOTE: Chemical reactions that could result in a hazardous situation (e.g. generation of flammable or toxic chemicals, fire or detonation) are listed here. Many of these reactions can be done safely if specific control measures (e.g. cooling of the reaction) are in place. Although not intended to be complete, an overview of important reactions involving common chemicals is provided to assist in the development of safe work practices.

Nitric acid is a strong acid and a strong oxidizing agent and is very reactive. Nitric acid is frequently involved in reactive incidents.

MOST METALS particularly POWDERED METALS (e.g. antimony, bismuth, germanium, manganese or titanium), ALKALI METALS (e.g. lithium or sodium) or ALKALINE EARTH METALS (e.g. magnesium or calcium) - may react violently or explosively, and/or cause fire, with generation of extremely flammable hydrogen gas.

ORGANIC CHEMICALS (e.g., alcohols, acids, anhydrides, aldehydes, ketones, amines, ethers, hydrocarbons, alkanethiols, nitriles, nitroalkanes and nitroaromatics) – may react violently or explosively, and/or ignite spontaneously.

NON-METALS (e.g. arsenic, boron, finely divided carbon, phosphorus or silicon), NON-METAL HYDRIDES (e.g. arsine, phosphine, stibine or tetraborane) or REDUCING AGENTS (e.g. potassium phosphinate) - may react violently or explosively and ignite.

CROTONALDEHYDE, HYDRAZINE, DIMETHYLHYDRAZINE, DIVINYL ETHER, PYROCATECHOL - ignite spontaneously (hypergolic) on contact with concentrated nitric acid, the ignition delay being 1 millisecond (ms).

AMMONIA, ANILINE, DIBORANE, FURFURYL ALCOHOL or TERPENES - mixtures are self-igniting.

SULFIDES (e.g. sodium or potassium sulfide) - toxic and flammable hydrogen sulfide gas and toxic sulfur dioxide gas may be generated.

CARBIDES (e.g. cesium carbide), FLUORINE, PHOSPHORUS HALIDES (e.g. phosphorus trichloride) or OTHER PHOSPHORUS COMPOUNDS (e.g. cadmium phosphide) – may ignite and/or explode.

METAL CYANIDES (e.g. sodium cyanide, potassium cyanide or calcium cyanide) - mixture produces a violent reaction, with formation of very toxic and flammable hydrogen cyanide.

SULFUR HALIDES (e.g. sulfur dichloride or disulfur dibromide) - interaction is violent, with generation of the corresponding hydrogen halide.

Conditions to avoid:	Air, light, high temperatures.
Hazardous Decomposition Products:	Nitrogen oxides.
Hazardous Polymerization:	None reported.

Corrosivity to Metals:

Nitric acid (5-70%) is corrosive (corrosion rate greater than 1.25 mm/year) to most metals at 21 °C (70 °F), including carbon steel (e.g. types 1010, 1020, 1075 and 1095), type 3003 aluminum, cast iron (e.g. gray, ductile and high nickel cast iron), nickel, nickel-base alloys (e.g. Monel and Hastelloy B and D), copper, copper-nickel, bronze (unspecified), aluminum bronze, silicon bronze, brass (unspecified), admiralty brass, naval brass and lead. It attacks (corrosion rate less than or equal to 0.5 mm/year to greater than or equal to 1.27 mm/year) some 400 series stainless steels at 21 °C. Stainless steels with high chromium content (most 300 series) exhibit excellent or good resistance to nitric acid concentrations ranging from 0-65% up to the boiling point. Types 304, 304L, R12S, 2RE10 (high chromium and nickel content) and 7-Mo duplex stainless steel are particularly recommended for use with nitric acid. High-silicon cast iron (Duriron) and high-chromium cast iron, higher chromium nickel alloys (e.g. G-30 and 690), nickel-base alloys (e.g. Hastelloy C and Incoloy 825), tantalum, titanium, zirconium, gold and platinum-type metals also have excellent resistance to nitric acid. Aluminum alloys (types 1100 and 3003) can only be used for very high concentrations of nitric acid (80-100%).

Corrosivity to Non-Metals:

Nitric acid (5-100%) at 21 °C attacks plastics, such as acrylonitrile-butadienestyrene (ABS), nylon, styrene-acrylonitrile, polystyrene and polyurethane; and elastomers, such as nitrile Buna N (NBR), natural rubber, isoprene, neoprene, chloroprene, polyester, styrene butadiene (SBR), polyurethane, chlorinated polyethylene, ethylene-propylene, ethylene-propylene terpolymer and low density polyethylene. Nitric acid (5-100%) does not attack plastics, such as Teflon, and other fluorocarbons; and elastomers such as Viton and other fluorocarbons (e.g. Kalrez and Chemraz (up to 50%)). Nitric acid (concentrations up to 50%) does not attack plastics, e.g. polyvinyl chloride (PVC), chlorinated polyvinyl chloride (CPVC), polypropylene, thermoset polyester (Bisphenol A-fumarate and Isophthalic acid), high density polyethylene and ultra high molecular weight polyethylene and elastomers e.g. butyl rubber.



Section 1	Section 11: Toxicological Information			
	Potential Health Effects			
Inhalation:	May be fatal if inhaled. Nitric acid readily forms a high vapour concentration at room temperature. It is very corrosive and can release other corrosive and toxic gases upon contact with air, organic materials or metals, thus posing a very serious inhalation hazard. Symptoms of exposure include dryness and/or burning pain in the nose and throat, cough, chest pain, shortness of breath and difficulty breathing. A severe exposure can result in a potentially fatal accumulation of fluid in the lungs (pulmonary edema). Symptoms of pulmonary edema (chest pain and shortness of breath) can be delayed for up to 24 or 48 hours after exposure. The presence of nitrogen oxide gases contributes significantly to the toxic effects observed following nitric acid exposure. However, the strongly corrosive nitric acid is expected to be a severe respiratory irritant on its own.			
Skin:	Nitric acid is corrosive. Corrosive materials are capable of producing severe burns, blisters, ulcers and permanent scarring, depending on the concentration of the solution and the duration of contact. Concentrated solutions produce burns, lower concentrations cause a change in skin colour from yellow to brown, and dilute solutions cause mild irritation and hardening of the skin.			
Eye:	Nitric acid is corrosive. Corrosive materials are capable of producing severe eye burns and permanent injury, including blindness, depending on the concentration of the solutions and duration of contact.			
Ingestion:	Nitric acid is corrosive and can cause burns to the lips, tongue, throat and stomach, abdominal pain, nausea, vomiting, diarrhea, kidney damage, and death if ingested. May cause perforation of the digestive tract. Because of immediate pain when taken into the mouth, strong mineral acids are not often swallowed.			
Chronic:	Long-term exposure may cause skin and respiratory irritation, with possible development of lung injury (e.g. chronic bronchitis). Exposure to nitric acid vapours, mists or aerosols may cause dental erosion and jaw necrosis. To the best of our knowledge, the chronic toxicity of this substance has not been fully investigated.			
	Effects of Long Term (Chronic) Exposure			

	Effects of Long-Term (Chronic) Exposure	
RTECS#:	QU5775000	
LD50/LC50:	Inhalation LC50: 260 mg/m ³ /30M (rat)	
Epidemiology:	No information available.	
Teratogenicity:	No information available.	
Reproductive Effects:	No information available.	
Neurotoxicity:	No information available.	
Mutagenicity:	No information available.	
Carcinogenicity:	Not listed as a carcinogen by ACGIH, IARC, NTP, or CA Prop 65.	
Section 12: Ecological Information		
Ecotoxicity:	Mosquito fish (<i>Gambusia affinis</i>): LC50 = 72 mg/L/96H	
Persistence and degradability:	Nitric acid will be gradually neutralized by hardness minerals (calcium and magnesium) in water. The nitrate ion may persist longer but will ultimately be consumed as a plant nutrient.	
Bioaccumulative _potential:	Bioaccumulation is not anticipated for inorganic compounds that are miscible with water.	
Mobility in soil:	No information available.	
Results of PBT and vPvB assessment:	Not applicable for inorganic substances.	

Section 13: Disposal Considerations

Review federal, regional and local government requirements prior to disposal. Authorities may not permit disposal of waste nitric acid until certain neutralization standards have been achieved. Store material for disposal as indicated in Section 7. Disposal by secure landfill may be acceptable.



Section 14: Transport	Information		
	t Information		
	CANADIAN TRANSPORTATION OF DANGEROUS GOODS (TDG) SHIPPING INFORMATION		
	CID, other than red fuming, with less than 65% nitric acid		
UN Number: UN2031 Class			
	HAZARDOUS MATERIALS SHIPPING INFORMATION (49 CFR)		
	CID other than red fuming with not more than 20 percent nitric acid Codes: 8 Packing Group: II		
	national Carriage of Dangerous Goods by Road (ADR)		
Proper Shipping Name / Description: NITR UN No.: 2031 Label	IC ACID, other than red fuming, with less than 65% nitric acid		
International Maritime Dangerous Goods (I			
	IC ACID other than red fuming, with less than 65% nitric acid or Division (Sub Risk): 8 Packing Group: II EMS: F-A, S-B		
International Air Transport Association (IA			
•	: acid other than red fuming, with 20% or less nitric acid		
	or Division (Sub Risk): 8 Packing Group: II		
Section 15: Regulator	ry Information		
OSHA Hazards:	CAS #7697-37-2 meets criteria for hazardous material, as defined by 29 CFR 1910.1200.		
SARA Title III: Section 302:	This material contains Nitric acid (CAS# 7697-37-2), which is subject to the reporting requirement of 1,000 lbs RQ.		
SARA Title III: Section 313:	This material contains Nitric acid (CAS# 7697-37-2), which is subject to the reporting requirements of Section 313 of SARA Title III.		
Massachusetts Right To Know	CAS# 7697-37-2 is listed, 50 lbs RQ.		
Pennsylvania Right To Know	CAS# 7697-37-2 is listed, E (environmental hazard).		
New Jersey Right To Know	CAS# 7697-37-2 is listed, RTK# 1356.		
California Prop. 65	CAS# 7697-37-2 is not subject to this act. CAS# 10024-97-2 (Nitrous oxide, decomposition product)		
·	is subject to this act, type of toxicity: developmental, female. CAS# 7732-18-5 is not subject to this		
	act.		
Inventory Status:	Canada DSL/NDSL Inventory List: CAS# 7697-37-2 is listed. CAS# 7732-18-5 is listed.		
	US TSCA Inventory List: CAS# 7697-37-2 is listed. CAS# 7732-18-5 is listed.		
Section 16: Other Inf	ormation		

The statements contained herein are offered for informational purposes only and are based upon technical data. SEASTAR CHEMICALS Inc. believes them to be accurate but does not purport to be all-inclusive. The above-stated product is intended for use only by persons having the necessary technical skills and facilities for handling the product at their discretion and risk. Since conditions and manner of use are outside our control, we (SEASTAR CHEMICALS Inc) make no warranty of merchantability or any such warranty, express or implied with respect to information and we assume no liability resulting from the above product or its use. Users should make their own investigations to determine suitability of information and product for their particular purposes.