

Balchem Corporation

Part Number: **08727** Version No: **11.15**

Safety Data Sheet according to OSHA HazCom Standard (2012) requirements

SECTION 1 Identification

Product Identifier

Product name	e Metalosate Multimineral Amino Acid Soluble Powder	
Synonyms	Multimineral Polvo Soluble (08727)	
Other means of identification	08727	

Recommended use of the chemical and restrictions on use

Relevant identified uses Organic Mineral Foliar Supplement

Name, address, and telephone number of the chemical manufacturer, importer, or other responsible party

Registered company name	Balchem Corporation
Address	52 Sunrise Park Road New Hampton, NY 10958 United States
Telephone	1.845.326.5600
Website	www.balchem.com
Email	sds@balchem.com

Emergency phone number

Emergene		
Asso	ociation / Organisation	Chemtrec CCN#2275
I	Emergency telephone numbers	1.800.424.9300 (USA)
Other	emergency telephone numbers	+1.703.527.3887 (International)

SECTION 2 Hazard(s) identification

Classification of the substance	or mixture
Classification	Serious Eye Damage/Eye Irritation Category 2A, Specific Target Organ Toxicity - Repeated Exposure Category 2, Combustible Dust

Label elements

Hazard pictogram(s)	
Signal word	Warning

Hazard statement(s)

H319	Causes serious eye irritation.
H373	May cause damage to organs through prolonged or repeated exposure. (Organ) (Oral, Inhalation)
	May form combustible dust concentrations in air

Hazard(s) not otherwise classified

Not Applicable

Precautionary statement(s) Prevention

P260	Do not breathe dust/fume.	
P280	Wear protective gloves, protective clothing, eye protection and face protection.	
P264	P264 Wash all exposed external body areas thoroughly after handling.	

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Precautionary statement(s) Response

P305+P351+P338	IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing.	
P314	Get medical advice/attention if you feel unwell.	
P337+P313	If eye irritation persists: Get medical advice/attention.	

Precautionary statement(s) Storage

Not Applicable

Precautionary statement(s) Disposal

P501 Dispose of contents/container to authorised hazardous or special waste collection point in accordance with any local regulation.

Not Applicable

SECTION 3 Composition / information on ingredients

Substances

See section below for composition of Mixtures

Mixtures

CAS No	%[weight]	Name
7782-63-0	1-15	ferrous sulfate heptahydrate
10043-52-4	1-15	calcium chloride
10034-99-8	20-40	magnesium sulfate, heptahydrate
7446-19-7	1-15	zinc sulfate heptahydrate
10034-96-5	1-15	manganese sulfate, hydrate
1344-73-6	1-15	copper sulfate, basic

SECTION 4 First-aid measures

Description of first aid measur	es
Eye Contact	 If this product comes in contact with the eyes: Wash out immediately with fresh running water. Ensure complete irrigation of the eye by keeping eyelids apart and away from eye and moving the eyelids by occasionally lifting the upper and lower lids. Seek medical attention without delay; if pain persists or recurs seek medical attention. Removal of contact lenses after an eye injury should only be undertaken by skilled personnel.
Skin Contact	If skin contact occurs: Immediately remove all contaminated clothing, including footwear. Flush skin and hair with running water (and soap if available). Seek medical attention in event of irritation.
Inhalation	 If fumes, aerosols or combustion products are inhaled remove from contaminated area. Other measures are usually unnecessary.
Ingestion	 IF SWALLOWED, REFER FOR MEDICAL ATTENTION, WHERE POSSIBLE, WITHOUT DELAY. For advice, contact a Poisons Information Centre or a doctor. Urgent hospital treatment is likely to be needed. In the mean time, qualified first-aid personnel should treat the patient following observation and employing supportive measures as indicated by the patient's condition. If the services of a medical officer or medical doctor are readily available, the patient should be placed in his/her care and a copy of the SDS should be provided. Further action will be the responsibility of the medical specialist. If medical attention is not available on the worksite or surroundings send the patient to a hospital together with a copy of the SDS. Where medical attention is not immediately available or where the patient is more than 15 minutes from a hospital or unless instructed otherwise: INDUCE vomiting with fingers down the back of the throat,ONLY IF CONSCIOUS. Lean patient forward or place on left side (head-down position, if possible) to maintain open airway and prevent aspiration. NOTE: Wear a protective glove when inducing vomiting by mechanical means.

Most important symptoms and effects, both acute and delayed

See Section 11

Indication of any immediate medical attention and special treatment needed

Magnesium is present in the blood, as a normal constituent, at concentrations between 1.6 to 2.2 meq/L. Some 30% is plasma bound. At serum magnesium levels of 3-4 meq/L, signs of CNS depression, loss of reflexes, muscular tone and power, and bradycardia occur. Cardiac arrest (sometimes fatal) and/or respiratory paralysis can occur at plasma levels of 10-15 meq/L. For acute or short term repeated exposures to magnesium:

- Symptomatic hypermagnesaemia appears rarely in the absence of intestinal or renal disease.
- Elevated magnesium levels may cause hypocalcaemia because of decreased parathyroid hormone activity and decreased end-organ responsiveness. ۶
- ٠ Patients with severe hypermagnesemia may develop sudden respiratory arrest and must be watched closely for apnoea.
- ۲ Use fluids, then vasopressors for hypotension. Frequently hypotension responds to calcium administration.
- Induce emesis or administer lavage if patient presents within 4 hours of ingestion. Use sodium cathartics, with caution, in presence of cardiac or renal failure. ٠
- Activated charcoal is not useful.
- Calcium is an antagonist of magnesium action and is an effective antidote when serum levels exceed 5 meq/L and the patient exhibits symptoms. The adult dose of calcium

gluconate is 10 mlL of a 10% solution over several minutes. [Ellenhorn and Barceloux: Medical Toxicology]

for copper intoxication:

- Unless extensive vomiting has occurred empty the stomach by lavage with water, milk, sodium bicarbonate solution or a 0.1% solution of potassium ferrocyanide (the resulting copper ferrocyanide is insoluble).
- Administer egg white and other demulcents.
- Maintain electrolyte and fluid balances.
- Morphine or meperidine (Demerol) may be necessary for control of pain.
- If symptoms persist or intensify (especially circulatory collapse or cerebral disturbances, try BAL intramuscularly or penicillamine in accordance with the supplier's
- recommendations
- Treat shock vigorously with blood transfusions and perhaps vasopressor amines.
- If intravascular haemolysis becomes evident protect the kidneys by maintaining a diuresis with mannitol and perhaps by alkalinising the urine with sodium bicarbonate. ۶
- It is unlikely that methylene blue would be effective against the occassional methaemoglobinemia and it might exacerbate the subsequent haemolytic episode.
- Institute measures for impending renal and hepatic failure. [GOSSELIN, SMITH & HODGE: Commercial Toxicology of Commercial Products]
- A role for activated charcoals for emesis is, as yet, unproven.
- In severe poisoning CaNa2EDTA has been proposed. [ELLENHORN & BARCELOUX: Medical Toxicology]

For acute or short term repeated exposures to iron and its derivatives:

- Always treat symptoms rather than history
- In general, however, toxic doses exceed 20 mg/kg of ingested material (as elemental iron) with lethal doses exceeding 180 mg/kg.
- Control of iron stores depend on variation in absorption rather than excretion. Absorption occurs through aspiration, ingestion and burned skin.
- Hepatic damage may progress to failure with hypoprothrombinaemia and hypoglycaemia. Hepatorenal syndrome may occur.
- Iron intoxication may also result in decreased cardiac output and increased cardiac pooling which subsequently produces hypotension.
- Serum iron should be analysed in symptomatic patients. Serum iron levels (2-4 hours post-ingestion) greater that 100 ug/dL indicate poisoning with levels, in excess of 350 ug/dL,
- being potentially serious. Emesis or lavage (for obtunded patients with no gag reflex)are the usual means of decontamination
- Activated charcoal does not effectively bind iron.
- Catharsis (using sodium sulfate or magnesium sulfate) may only be used if the patient already has diarrhoea.
- Deferoxamine is a specific chelator of ferric (3+) iron and is currently the antidote of choice. It should be administered parenterally. [Ellenhorn and Barceloux: Medical Toxicology]

Both dermal and oral toxicity of manganese salts is low because of limited solubility of manganese. No known permanent pulmonary sequelae develop after acute manganese exposure. Treatment is supportive.

[Ellenhorn and Barceloux: Medical Toxicology]

In clinical trials with miners exposed to manganese-containing dusts, L-dopa relieved extrapyramidal symptoms of both hypo kinetic and dystonic patients. For short periods of time symptoms could also be controlled with scopolamine and amphetamine. BAL and calcium EDTA prove ineffective. [Gosselin et al: Clinical Toxicology of Commercial Products.]

SECTION 5 Fire-fighting measures

Extinguishing media

- There is no restriction on the type of extinguisher which may be used.
- Use extinguishing media suitable for surrounding area

Special hazards arising from the substrate or mixture

Fire Incompatibility None known. Special protective equipment and precautions for fire-fighters

Fire Fighting	 Alert Fire Brigade and tell them location and nature of hazard. Wear breathing apparatus plus protective gloves in the event of a fire. Prevent spillage from entering drains or water courses. Use fire fighting procedures suitable for surrounding area. DO NOT approach containers suspected to be hot. Cool fire exposed containers with water spray from a protected location. If safe to do so, remove containers from path of fire.
	This product may present a dust explosion risk in the form it is manufactured and sold. This conclusion is based on the product composition and on results of Kst testing of similar products. Further processing of the product, such as manipulation of particle size, and the environment in which it is used (e.g., humidity effects) may increase or decrease the risk of dust explosions.
	Dusts fall into one of three Kst* classes. Class 1 dusts; Kst 1-200 m3/sec; Class 2 dusts; 201-299 m3/sec. Class 3 dusts; Kst 300 or more. Most agricultural dusts (grains, flour etc.) are Class 1; pharmaceuticals and other speciality chemicals are typically Class 1 or 2; most unoxidised metallic dusts are Class 3. The higher the Kst, the more energetically the dust will burn and the greater is the explosion risk and the greater is the speed of the explosion.
	Standard test conditions, used to derive the Kst, are representative of industrial conditions, but do not represent an absolute worst case. Increased levels of turbulence increase the speed of the explosion dramatically. * Kst - a normalised expression of the burning dust pressure rise rate over time.
Fire/Explosion Hazard	Dusts with Minimum Ignition Energies (MIEs) ranging between 20 and 100 mJ may be sensitive to ignition. They require that: · plant is grounded · personnel might also need to be grounded
	the use of high resistivity materials (such as plastics) should be restricted or avoided during handling or in packaging
	The majority of ignition accidents occur within or below this range. The MIE of a dust/air mix depends on the particle size the water content and the temperature of the dust. The finer and the dryer the dust the lower the MIE. Higher temperatures cause lower MIE and an increased risk of dust explosion.
	Quoted values for MIE generally are only representative. Characteristics may change depending upon the process and conditions of use or any changes made to the dust during use, including further grinding or mixing with other products. In order to obtain more specific data for dust, as used, it is recommended that further characterisation testing is performed.
	 Combustible solid which burns but propagates flame with difficulty; it is estimated that most organic dusts are combustible (circa 70%) - according to the circumstances under which the combustion process occurs, such materials may cause fires and / or dust explosions. Organic powders when finely divided over a range of concentrations regardless of particulate size or shape and suspended in air or some
	Continuación

 other oxidizing medium may form explosive dust-air mixtures and result in a fire or dust explosion (including secondary explosions). Avoid generating dust, particularly clouds of dust in a confined or unventilated space as dusts may form an explosive mixture with air, and any source of ignition, i.e., flame or spark, will cause fire or explosion. Dust clouds generated by the fine grinding of the solid are a particular hazard; accumulations of fine dust (420 micron or less) may burn rapidly and fiercely if ignited - particles exceeding this limit will generally not form flammable dust clouds; once initiated, however, larger particles up to 1400 microns diameter will contribute to the propagation of an
 explosion. In the same way as gases and vapours, dusts in the form of a cloud are only ignitable over a range of concentrations; in principle, the concepts of lower explosive limit (LEL) and upper explosive limit (UEL) are applicable to dust clouds but only the LEL is of practical use; - this is because of the inherent difficulty of achieving homogeneous dust clouds at high temperatures (for dusts the LEL is often called the 'Minimum Explosible Concentration', MEC).
When processed with flammable liquids/vapors/mists, ignitable (hybrid) mixtures may be formed with combustible dusts. Ignitable mixtures will increase the rate of explosion pressure rise and the Minimum Ignition Energy (the minimum amount of energy required to ignite dust clouds - MIE) will be lower than the pure dust in air mixture. The Lower Explosive Limit (LEL) of the vapour/dust mixture will be lower than the individual LELs for the vapors/mists or dusts.
A dust explosion may release of large quantities of gaseous products; this in turn creates a subsequent pressure rise of explosive force capable of damaging plant and buildings and injuring people.
Usually the initial or primary explosion takes place in a confined space such as plant or machinery, and can be of sufficient force to damage or rupture the plant. If the shock wave from the primary explosion enters the surrounding area, it will disturb any settled dust layers, forming a second dust cloud, and often initiate a much larger secondary explosion. All large scale explosions have resulted from chain reactions of this
 type. Dry dust can be charged electrostatically by turbulence, pneumatic transport, pouring, in exhaust ducts and during transport.
 Build-up of electrostatic charge may be prevented by bonding and grounding. Powder handling equipment such as dust collectors, dryers and mills may require additional protection measures such as explosion venting. All movable parts coming in contact with this material should have a speed of less than 1-meter/sec.
A sudden release of statically charged materials from storage or process equipment, particularly at elevated temperatures and/ or pressure, may result in ignition especially in the absence of an apparent ignition source.
 One important effect of the particulate nature of powders is that the surface area and surface structure (and often moisture content) can vary widely from sample to sample, depending of how the powder was manufactured and handled; this means that it is virtually impossible to use flammability data published in the literature for dusts (in contrast to that published for gases and vapours).
 Autoignition temperatures are often quoted for dust clouds (minimum ignition temperature (MIT)) and dust layers (layer ignition temperature (LIT)); LIT generally falls as the thickness of the layer increases. Decomposition may produce toxic fumes of:
sulfur oxides (SOx)
metal oxides
May emit poisonous fumes.
May emit corrosive fumes.

Personal precautions, protective equipment and emergency procedures

See section 8

Environmental precautions

See section 12

Methods and material for containment and cleaning up

Minor Spills	 Remove all ignition sources. Clean up all spills immediately. Avoid contact with skin and eyes. Control personal contact with the substance, by using protective equipment. Use dry clean up procedures and avoid generating dust. Place in a suitable, labelled container for waste disposal. Environmental hazard - contain spillage.
Major Spills	 Environmental hazard - contain spillage. Moderate hazard. CAUTION: Advise personnel in area. Control personal contact by wearing protective clothing. Prevent spillage from entering drains or water courses. Recover product wherever possible. IF DRY: Use dry clean up procedures and avoid generating dust. Collect residues and place in sealed plastic bags or other containers for disposal. IF WET: Vacuum/shovel up and place in labelled containers for disposal. ALWAYS: Wash area down with large amounts of water and prevent runoff into drains.

Personal Protective Equipment advice is contained in Section 8 of the SDS.

SECTION 7 Handling and storage

Precautions for safe handling

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Safe handling

	 Atmosphere should be regularly checked against established exposure standards to ensure safe working conditions are maintained. Organic powders when finely divided over a range ofconcentrations regardless of particulate size or shape and suspended in air or some other oxidizing medium may form explosive dust-air mixtures and result in a fire or dust explosion (including secondary explosions). Minimise airborne dust and eliminate all ignition sources. Keep away from heat, hot surfaces, sparks, and flame. Establish good housekeeping practices. Remove dust accumulations on a regular basis by vacuuming or gentle sweeping to avoid creating dust clouds. Use continuous suction at points of dust generation to capture and minimise the probability of a 'secondary' explosion. According to NFPA Standard 654, dust layers 1/32 inch (0.8 mm) thick can be sufficient to warrant immediate cleaning of the area. Do not use air hoses for cleaning. Minimise dry sweeping to avoid generation of dust clouds. Vacuum dust-accumulating surfaces and remove to a chemical disposal area. Vacuums with explosion-proof motors should be used. Control sources of static electricity. Dusts or their packages may accumulate static charges, and static discharge can be a source of ignition. Solids handling systems must be designed in accordance with applicable standards (e.g., NFPA including 654 and 77) and other national guidance. Do not empty directly into flammable solvents or in the presence of flammable vapors. The operator, the packaging container and all equipment must be grounded with electrical bonding and grounding systems. Plastic bags and plastics cannot be grounded, and antistatic bags do not completely protect against development of static charges. Empty containers may contain residual dust which has the potential to accumulate following setting. Such dusts may explode in the presence of an appropriate ignition source.
Other information	 Store in original containers. Keep containers securely sealed. Store in a cool, dry area protected from environmental extremes. Store away from incompatible materials and foodstuff containers. Protect containers against physical damage and check regularly for leaks. Observe manufacturer's storage and handling recommendations contained within this SDS.

Conditions for safe storage, including any incompatibilities

Suitable container	 Polyethylene or polypropylene container. Check all containers are clearly labelled and free from leaks.
Storage incompatibility	Inorganic alkaline earth metal derivative. Derivative of very electropositive metal. Inorganic derivative of Group 11 metal. Calcium chloride (and its hydrates): * are incompatible with boric acid, calcium oxide, bromine trifluoride, 2-furan, percarboxylic acid * may produce explosive hydrogen gas on contact with zinc * catalyse exothermic polymerisation of methyl vinyl ether * produce heat on contact with water * attack metals * WARNING: Avoid or control reaction with peroxides. All <i>transition metal</i> peroxides should be considered as potentially explosive. For example transition metal complexes of alkyl hydroperoxides may decompose explosively. * The pi-complexes formed between chromium(0), vanadium(0) and other transition metals (haloarene-metal complexes) and mono-or poly-fluorobenzene show extreme sensitivity to heat and are explosive. * Avoid reaction with borohydrides or cyanoborohydrides * Metals and their oxides or salts may react violently with chlorine trifluoride and bromine trifluoride. * These trifluorides are hypergolic oxidisers. They ignite on contact (without external source of heat or ignition) with recognised fuels - contact with these materials, following an ambient or slightly elevated temperature, is often violent and may produce ignition. * The state of subdivision may affect the results. Copper sulfate: * reacts violently with strong bases, hydroxylamine (with ignition), magnesium (producing hydrogen gas) * in contact with potassium chlorate is potentially explosive * solutions are acidic and can react with metals to evolve flammable hydrogen gas corrosive to some metals including steel. * is incompatible with sulfuric acid, caustics, ammonia, aliphatic amines, alkanolamines, anides, alkylene oxides, epichlorohydrin, organic anhydrides, isocyanates, vinyl acetate. * dusts or mists may react with acetylene to form shock-sensitive copper acetylides.

SECTION 8 Exposure controls / personal protection

Control parameters

Occupational Exposure Limits (OEL)

INGREDIENT DATA Source Ingredient Material name TWA STEL Peak Notes US NIOSH Recommended ferrous sulfate Not Not Iron salts (soluble, as Fe) 1 mg/m3 Not Available Exposure Limits (RELs) heptahydrate Available Available US ACGIH Threshold Limit ferrous sulfate Not Not Iron salts, soluble, as Fe 1 mg/m3 Not Available Available Available Values (TLV) heptahydrate US OSHA Permissible Exposure manganese Manganese compounds (as Not Not 5 mg/m3 Not Available Limits (PELs) Table Z-1 sulfate, hydrate Mn) Available Available [*Note: Also see specific listings for US NIOSH Recommended manganese Manganese compounds and Manganese cyclopentadienyl tricarbonyl, Not 1 mg/m3 3 mg/m3 Available Methyl cyclopentadienyl manganese Exposure Limits (RELs) sulfate, hydrate fume (as Mn) tricarbonyl, and Manganese tetroxide.] Manganese, elemental and US ACGIH Threshold Limit 0.1 manganese Not Not inorganic compounds, as Mn A4 sulfate, hydrate Available Available Values (TLV) mg/m3 (Inhalable particulate matter)

Source	Ingredient	Material name	TWA	STEL	Peak	Notes	
US ACGIH Threshold Limit Values (TLV)	manganese sulfate, hydrate	Manganese, elemental and inorganic compounds, as N (Respirable particulate matter)		Not Available	Not Available	A4	
Emergency Limits							
Ingredient	TEEL-1	т	EEL-2			TEEL-3	
ferrous sulfate heptahydrate	8.2 mg/m3	4	1 mg/m3			250 mg/m3	
ferrous sulfate heptahydrate	15 mg/m3	1	70 mg/m3			990 mg/m3	
calcium chloride	16 mg/m3	1	70 mg/m3			1,100 mg/m3	
calcium chloride	12 mg/m3	1	30 mg/m3			790 mg/m3	
calcium chloride	13 mg/m3	1.	40 mg/m3			850 mg/m3	
calcium chloride	24 mg/m3	2	60 mg/m3			1,600 mg/m3	
magnesium sulfate, heptahydrate	33 mg/m3	3	70 mg/m3			2,300 mg/m3	
magnesium sulfate, heptahydrate	20 mg/m3	2	20 mg/m3			1,300 mg/m3	
zinc sulfate heptahydrate	27 mg/m3		170 mg/m3			1,000 mg/m3	
zinc sulfate heptahydrate	15 mg/m3		97 mg/m3			580 mg/m3	
manganese sulfate, hydrate	9.2 mg/m3	1	15 mg/m3			90 mg/m3	
manganese sulfate, hydrate	8.2 mg/m3	8.2 mg/m3 14 mg/m3				430 mg/m3	
Ingredient	Original IDLH	Original IDLH		Revis	sed IDLH		
ferrous sulfate heptahydrate	Not Available		Not Available		vailable		
calcium chloride	Not Available		Not Available		vailable		
magnesium sulfate, heptahydrate	Not Available		Not Available		vailable		
zinc sulfate heptahydrate	Not Available	Not Available		Not Available			
manganese sulfate, hydrate	500 mg/m3		Not Available		vailable		
copper sulfate, basic	Not Available			Not A	Not Available		

Exposure controls

Appropriate engineering controls	Assess operations based upon available dust explosion inforprecautionary measures against possible dust explosions. I suppression of dust handling equipment. Where explosions is should preferably be calculated based on Kst rather than ar with an oxygen level below the limiting oxygen concentratio event of excessive oxygen being detected. The maximum surface temperature of enclosures potentiall minimum ignition temperature (MIE) of the dust cloud. The An isolated (insulated) human body can readily produce elee be highly effective in protecting workers and will typically be The basic types of engineering controls are: Process controls which involve changing the way a job actific Enclosure and/or isolation of emission source which keeps 'adds' and 'removes' air in the work environment. Ventilation ventilation system must match the particular process and cliemployers may need to use multiple types of controls to protection might consist of: (a): particle dust respirators, if necessary, combined with ar (b): filter respirators, with absorption cartridge or canister of (c): fresh-air hoods or masks. Air contaminants generated in the workplace possess varyi circulating air required to effectively remove the contaminar	f prevention is not possible, consi venting is considered to be the mo- n St value. If nitrogen purging is or n. The system should include an or y exposed to this material should effect of dust layers should be rev ectrostatic discharges in excess of a barrier between the worker and be independent of worker interaction vity or process is done to reduce t a selected hazard 'physically' awa n can remove or dilute an air conta hemical or contaminant in use. event employee overexposure. andled as powders or crystals; ev the substance in air could occur, r in absorption cartridge; the right type; ng 'escape' velocities which, in tur	der protection by use of co ost appropriate method of p onsidered as the protective oxygen monitoring and shu be based on values obtain iewed. 50 mJ, but have been rec the hazard. Well-designed ns to provide this high leve he risk. ay from the worker and ver aminant if designed proper	entainment, venting or protection, vent areas a system, it must operate ut-down facility in the ed by taking 2/3 of the orded up to 100 mJ. d engineering controls can al of protection. Initiation that strategically ity. The design of a elatively large, a certain Id be considered.
	Type of Contaminant:			Air Speed:
direct spray, spray painting in shallow booths, drum filling, conveyer loading, crusher dusts, gas discharge (active generation into zone of rapid air motion)				1-2.5 m/s (200-500 ft/min)
	grinding, abrasive blasting, tumbling, high speed wheel ge very high rapid air motion).	nerated dusts (released at high in	itial velocity into zone of	2.5-10 m/s (500-2000 ft/min)
	Within each range the appropriate value depends on: Lower end of the range	Upper end of the range		
	1: Room air currents minimal or favourable to capture	1: Disturbing room air currents		

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Metalosate Multimineral Amino Acid Soluble Powder

	2: Contaminants of low toxicity or of nuisance value only.	2: Contaminants of high toxicity			
	3: Intermittent, low production.	3: High production, heavy use			
	4: Large hood or large air mass in motion	4: Small hood-local control only			
Personal protection	Simple theory shows that air velocity falls rapidly with dista with the square of distance from the extraction point (in sim accordingly, after reference to distance from the contamina 4-10 m/s (800-2000 ft/min) for extraction of crusher dusts g producing performance deficits within the extraction appare more when extraction systems are installed or used.	nple cases). Therefore the air speed at ating source. The air velocity at the extr generated 2 metres distant from the ex	t the extraction point should be adjusted, raction fan, for example, should be a minimum traction point. Other mechanical considerations		
Eye and face protection	 Safety glasses with side shields. Chemical goggles. Contact lenses may pose a special hazard; soft contact lenses may absorb and concentrate irritants. A written policy document, describing the wearing of lenses or restrictions on use, should be created for each workplace or task. This should include a review of lens absorption and adsorption for the class of chemicals in use and an account of injury experience. Medical and first-aid personnel should be trained in their removal and suitable equipment should be readily available. In the event of chemical exposure, begin eye irrigation immediately and remove contact lens as soon as practicable. Lens should be removed at the first signs of eye redness or irritation - lens should be removed in a clean environment only after workers have washed hands thoroughly. [CDC NIOSH Current Intelligence Bulletin 59], [AS/NZS 1336 or national equivalent] 				
Skin protection	See Hand protection below				
Hands/feet protection	 NOTE: The material may produce skin sensitisation in predisp equipment, to avoid all possible skin contact. Contaminated leather items, such as shoes, belts and The selection of suitable gloves does not only depend on the manufacturer. Where the chemical is a preparation of sever and has therefore to be checked prior to the application. The exact break through time for substances has to be obtomaking a final choice. Personal hygiene is a key element of effective hand care. Of washed and dried thoroughly. Application of a non-perfume Suitability and durability of glove type is dependent on usary frequency and duration of contact, chemical resistance of glove material, glove thickness and dexterity Select gloves tested to a relevant standard (e.g., Europe E When prolonged or frequently repeated contact may occuminutes according to EN 374, AS/NZS 2161.10.1 or national equivalent) is recommer. Some glove polymer types are less affected by movement. Contaminated gloves should be replaced. As defined in ASTM F-739-96 in any application, gloves and Excellent when breakthrough time < 20 min. Fair when breakthrough time < 20 min. Fair when breakthrough time < 20 min. For general applications, gloves with a thickness typically of It should be emphasised that glove thickness is not necess efficiency of the glove will be dependent on the exact comp consideration of the task requirements and knowledge of b Glove thickness may also vary depending on the glove and data should always be taken into account to ensure selecti Note: Depending on the activity being conducted, gloves of the ploves (down to 0.1 mm or less) may be required the purceture potential. Gloves must only be worn on clean hands. After using glove moisturiser is recommended. Experience indicates that the following polymers are suitab particles are not present. polychloroprene. hutyl rubber.<!--</th--><th>watch-bands should be removed and he material, but also on further marks i ral substances, the resistance of the g tained from the manufacturer of the pro- Gloves must only be worn on clean ha ad moisturiser is recommended. ge. Important factors in the selection o 3. 374, US F739, AS/NZS 2161.1 or n: ur, a glove with a protection class of 5 of al equivalent) is recommended. ction class of 3 or higher (breakthrough ended. tt and this should be taken into account e rated as: greater than 0.35 mm, are recommend sarily a good predictor of glove resistar position of the glove material. Therefor reakthrough times. nufacturer, the glove type and the glov ion of the most appropriate glove for th f varying thickness may be required fo where a high degree of manual dexte be just for single use applications, there ere there is a mechanical (as well as a ves, hands should be washed and drie ole as glove materials for protection ag</th><th>destroyed. of quality which vary from manufacturer to love material can not be calculated in advance otective gloves and has to be observed when inds. After using gloves, hands should be if gloves include: ational equivalent). or higher (breakthrough time greater than 240 in time greater than 60 minutes according to EN it when considering gloves for long-term use. ed. ince to a specific chemical, as the permeation e, glove selection should also be based on we model. Therefore, the manufacturers technic ic tasks. r specific tasks. For example: rity is needed. However, these gloves are only in disposed. in chemical) risk, i.e., where there is abrasion or d thoroughly. Application of a non-perfumed</th>	watch-bands should be removed and he material, but also on further marks i ral substances, the resistance of the g tained from the manufacturer of the pro- Gloves must only be worn on clean ha ad moisturiser is recommended. ge. Important factors in the selection o 3. 374, US F739, AS/NZS 2161.1 or n: ur, a glove with a protection class of 5 of al equivalent) is recommended. ction class of 3 or higher (breakthrough ended. tt and this should be taken into account e rated as: greater than 0.35 mm, are recommend sarily a good predictor of glove resistar position of the glove material. Therefor reakthrough times. nufacturer, the glove type and the glov ion of the most appropriate glove for th f varying thickness may be required fo where a high degree of manual dexte be just for single use applications, there ere there is a mechanical (as well as a ves, hands should be washed and drie ole as glove materials for protection ag	destroyed. of quality which vary from manufacturer to love material can not be calculated in advance otective gloves and has to be observed when inds. After using gloves, hands should be if gloves include: ational equivalent). or higher (breakthrough time greater than 240 in time greater than 60 minutes according to EN it when considering gloves for long-term use. ed. ince to a specific chemical, as the permeation e, glove selection should also be based on we model. Therefore, the manufacturers technic ic tasks. r specific tasks. For example: rity is needed. However, these gloves are only in disposed. in chemical) risk, i.e., where there is abrasion or d thoroughly. Application of a non-perfumed		
Body protection	See Other protection below				
Other protection	 Overalls. P.V.C apron. Barrier cream. Skin cleansing cream. Eye wash unit. 				

Respiratory protection

Particulate. (AS/NZS 1716 & 1715, EN 143:2000 & 149:001, ANSI Z88 or national equivalent)

· Respirators may be necessary when engineering and administrative controls do not adequately prevent exposures.

The decision to use respiratory protection should be based on professional judgment that takes into account toxicity information, exposure measurement data, and frequency and likelihood of the worker's exposure - ensure users are not subject to high thermal loads which may result in heat stress or distress due to personal protective equipment (powered, positive flow, full face apparatus may be an option).

Published occupational exposure limits, where they exist, will assist in determining the adequacy of the selected respiratory protection. These may be government mandated or vendor recommended.

Certified respirators will be useful for protecting workers from inhalation of particulates when properly selected and fit tested as part of a complete respiratory protection program.
 Where protection from nuisance levels of dusts are desired, use type N95 (US) or type P1 (EN143) dust masks. Use respirators and components tested and approved under appropriate government standards such as NIOSH (US) or CEN (EU)

· Use approved positive flow mask if significant quantities of dust becomes airborne.

· Avoid creating dust conditions.

Class P2 particulate filters are used for protection against mechanically and thermally generated particulates or both.

P2 is a respiratory filter rating under various international standards, Filters at least 94% of airborne particles

Suitable for:

· Relatively small particles generated by mechanical processes, e.g., grinding, cutting, sanding, drilling, sawing.

· Sub-micron thermally generated particles, e.g., welding fumes, fertilizer and bushfire smoke.

· Biologically active airborne particles under specified infection control applications, e.g., viruses, bacteria, COVID-19, SARS.

SECTION 9 Physical and chemical properties

Information on basic physical and chemical properties

Appearance	Tan to Brown Powder		
Physical state	Divided Solid Powder	Relative density (Water = 1)	Not Available
Odour	Not Available	Partition coefficient n-octanol / water	Not Available
Odour threshold	Not Available	Auto-ignition temperature (°C)	Not Available
pH (as supplied)	Not Applicable	Decomposition temperature	Not Available
Melting point / freezing point (°C)	Not Available	Viscosity (cSt)	Not Available
Initial boiling point and boiling range (°C)	Not Available	Molecular weight (g/mol)	Not Available
Flash point (°C)	Not Available	Taste	Not Available
Evaporation rate	Not Available	Explosive properties	Not Available
Flammability	Not Available	Oxidising properties	Not Available
Upper Explosive Limit (%)	Not Available	Surface Tension (dyn/cm or mN/m)	Not Applicable
Lower Explosive Limit (%)	Not Available	Volatile Component (%vol)	Not Available
Vapour pressure (kPa)	Not Available	Gas group	Not Available
Solubility in water	Miscible	pH as a solution (1%)	4.0 - 4.5
Vapour density (Air = 1)	Not Available	VOC g/L	Not Available

SECTION 10 Stability and reactivity

Reactivity	See section 7
Chemical stability	 Unstable in the presence of incompatible materials. Product is considered stable. Hazardous polymerisation will not occur.
Possibility of hazardous reactions	See section 7
Conditions to avoid	See section 7
Incompatible materials	See section 7
Hazardous decomposition products	See section 5

SECTION 11 Toxicological information

Information on toxicological effects

Inhaled

The material is not thought to produce either adverse health effects or irritation of the respiratory tract following inhalation (as classified by EC Directives using animal models). Nevertheless, adverse systemic effects have been produced following exposure of animals by at least one other

	route and good hygiene practice requires that exposure be kept to a minimum and that suitable control measures be used in an occupational setting. Levels above 10 micrograms per cubic metre of suspended inorganic sulfates in the air may cause an excess risk of asthmatic attacks in susceptible people. Persons with impaired respiratory function, airway diseases and conditions such as emphysema or chronic bronchitis, may incur further disability if excessive concentrations of particulate are inhaled. If prior damage to the circulatory or nervous systems has occurred or if kidney damage has been sustained, proper screenings should be conducted on individuals who may be exposed to further risk if handling and use of the material result in excessive exposures. Manganese fume is toxic and produces nervous system effects characterised by tiredness. Acute poisoning is rare although acute inflammation of the lungs may occur. A chemical pneumonia may also result from frequent exposure. Inhalation of freshly formed metal oxide particles sized below 1.5 microns and generally between 0.02 to 0.05 microns may result in 'metal fume fever'. Symptoms may be delayed for up to 12 hours and begin with the sudden onset of thirst, and a sweet, metallic or foul taste in the mouth. Other symptoms usually subside within 24-36 hours following removal from exposure. Copper poisoning following exposure to copper dusts and fume may result in headache, cold sweat and weak pulse. Capillary, kidney, liver and brain damage are the longer term manifestations of such poisoning. Inhalation of freshly formed metal oxide particles sized below 1.5 microns and generally between 0.02 to 0.05 microns may result in metal fume fever'. Symptoms may be delayed for up to 12 hours and begin with the sudden onset of thirst, and a sweet, metallic or foul taste in the mouth. Other symptoms usually subside within 24-36 hours following removal from exposure.
Ingestion	Accidental ingestion of the material may be harmful; animal experiments indicate that ingestion of less than 150 gram may be fatal or may produce serious damage to the health of the individual. Compared with other metals, the calcium ion and most calcium compounds have low toxicity. Acute calcium poisoning is rare, and occurs only when calcium compounds are taken in high doses over a long period, or given through a vein. Excessive consumption of calcium carbonate antacids or pills over a period of weeks or months can cause milk-alkali syndrome, with high blood calcium and potentially fatal kidney failure. Excessive calcium supplementation can be detrimental to cardiovascular health, especially in men. Calcium supplementation given by mouth may reduce the absorption of thyroxine, when taken within 4-6 hours of each other. This may lead to inadequate thyroid hormone replacement and then hypothyroidism. Magnesium salts are generally absorbed so slowly that swallowing these cause few toxic effects, with purging being the most significant. If it cannot be removed (for example in bowel obstruction or paralysis), it may irritate the gut lining and be absorbed into the body. Side effects of magnesium salts include upset stomach, dry mouth, dry nose, dry throat, drowsiness, nausea, heartburn, and thickening of the lining of the throat and nose. The magnesium ion causes salt disturbances, central nervous system depression, involvement of the heart, loss of reflexes and death from paralysis of breathing; these effects, however, are rare without pre-existing kidney or bowel disorders. Early signs and symptoms of magnesium poisoning include nausea, vomiting, general unwellness and confusion. There may be low blood pressure due to dilation of blood vessels. A slow heart beat is common, which may eventually lead to stoppage of the heart. Sulfates are not well absorbed orally, but can cause diarrhoea. A metallic taste, nausea, vomiting and burning feeling in the upper stomach region occur after ingestion of copper and its deri
Skin Contact	This material can cause inflammation of the skin on contact in some persons. The material may accentuate any pre-existing dermatitis condition Exposure to copper, by skin, has come from its use in pigments, ointments, ornaments, jewellery, dental amalgams and IUDs (intra-uterine devices), and in killing fungi and algae. Although copper is used in the treatment of water in swimming pools and reservoirs, there are no reports of toxicity from these applications. Open cuts, abraded or irritated skin should not be exposed to this material Entry into the blood-stream, through, for example, cuts, abrasions or lesions, may produce systemic injury with harmful effects. Examine the skin prior to the use of the material and ensure that any external damage is suitably protected.
Eye	Although the material is not thought to be an irritant (as classified by EC Directives), direct contact with the eye may cause transient discomfort characterised by tearing or conjunctival redness (as with windburn). Slight abrasive damage may also result. Copper salts, in contact with the eye, may produce inflammation of the conjunctiva, or even ulceration and cloudiness of the cornea.
Chronic	Repeated or long-term occupational exposure is likely to produce cumulative health effects involving organs or biochemical systems. Strong evidence exists that this substance may cause irreversible mutations (though not lethal) even following a single exposure. Skin contact with the material is more likely to cause a sensitisation reaction in some persons compared to the general population. Ample evidence exists from experimentation that reduced human fertility is directly caused by exposure to the material. In a case of chronic abuse of magnesium citrate, symptoms seen included tiredness and severe low blood pressure which did not respond to treatment. Blood tests revealed extremely high levels of magnesium, and the patient was found to have a perforated ulcer of the duodenum. Kidney failure and death followed. A patient with normal kidney function developed stoppage of breathing and slow heart rate after receiving 90 grams of magnesium sulfate over 18 hours. Animal testing suggests that magnesium sulfate may reduce both fertility and the weight of offspring. Overexposure to the breathable dust may cause coughing, wheezing, difficulty in breathing and impaired lung function. Chronic symptoms may include decreased vital lung capacity and chest infections. Repeated exposures in the workplace to high levels of fine-divided dusts may produce a condition known as pneumoconiosis, which is the lodgement of any inhaled dusts in the lung, irrespective of the effect. This is particularly true when a significant number of particles less than 0.5 microns (1/50000 inch) are present. Lung shadows are seen in the X-ray. Symptoms of pneumoconiosis may include a progressive dry cough, shortness of breath on exertion, increased chest expansion, weakness and weight loss. As the disease progresses, the cough produces stringy phlegm, vital capacity decreases further, and shortness of breath becomes more severe. Other signs or symptoms include changed breath sounds, reduced oxygen uptake during exercise, emphysema an

	concentrations in vitro. Cancer-causing potential: There was insufficient information to evaluate the cancer-causing activity of copper monochloride. Manganese is an essential trace element. Chronic exposure to low levels of manganese can include a mask-like facial expression, spastic gait, tremors, slurred speech, disordered muscle tone, fatigue, anorexia, loss of strength and energy, apathy and poor concentration.			
Metalosate Multimineral	ΤΟΧΙΟΙΤΥ	IRRITATION		
Amino Acid Soluble Powder	Not Available	Not Available		
	TOXICITY		IRRITATION	
ferrous sulfate heptahydrate	Dermal (rat) LD50: >881 mg/kg ^[1]		Not Available	
	Oral (rat) LD50: 13 mg/kg ^[1]			
calcium chloride			IRRITATION Not Available	
calcium chioride	Dermal (rat) LD50: >2000 mg/kg ^[1] Oral (rat) LD50: >2000 mg/kg ^[1]			
	ΤΟΧΙCΙΤΥ		IRRITATION	
magnesium sulfate,	Dermal (rat) LD50: >2000 mg/kg ^[1]		Not Available	
heptahydrate	Oral (rat) LD50: >1000-2000 mg/kg ^[1]			
	ΤΟΧΙΟΙΤΥ		IRRITATION	
zinc sulfate heptahydrate	Oral (mouse) LD50: 200 mg/kg ^[2]		Not Available	
	TOXICITY IRRITATION			
manganese sulfate, hydrate	Oral (rat) LD50: 2150 mg/kg ^[2]	1	lot Available	
	ΤΟΧΙCITY	IRRITATION		
copper sulfate, basic	Not Available	Not Available		
Legend:	1. Value obtained from Europe ECHA Registered Substances - Acute to	xicity 2 * Value obtained from	m manufacturer's SDS Unless otherwise	
	specified data extracted from RTECS - Register of Toxic Effect of chemi			
	Laboratory (in vitro) and animal studies show, exposure to the material may result in a possible risk of irreversible effects, with the possibility producing mutation. The following information refers to contact allergens as a group and may not be specific to this product. Contact allergies quickly manifest themselves as contact eczema, more rarely as urticaria or Quincke's oedema. The pathogenesis of contact eczema involves a cell-mediated (T lymphocytes) immune reaction of the delayed type. Other allergic skin reactions, e.g., contact urticaria, involve antibody-mediated immune reactions. The significance of the contact allergen is not simply determined by its sensitisation potential: t distribution of the substance and the opportunities for contact with it are equally important. A weakly sensitising substance which is widely distributed can be a more important allergen than one with stronger sensitising potential with which few individuals come into contact. From a clinical point of view, substances are noteworthy if they produce an allergic test reaction in more than 1% of the persons tested. For copper and its compounds (typically copper chloride): Acute toxicity: There are no reliable acute oral toxicity results available. In an acute dermal toxicity study (OECD TG 402), one group of 5 m		uct. te's oedema. The pathogenesis of contact ic skin reactions, e.g., contact urticaria, determined by its sensitisation potential: the sensitising substance which is widely few individuals come into contact. From a n 1% of the persons tested. study (OECD TG 402), one group of 5 male	
Metalosate Multimineral Amino Acid Soluble Powder	rats and 5 groups of 5 female rats received doses of 1000, 1500 and 200 copper monochloride were 2,000 mg/kg bw or greater for male (no death 1500 and 2000 mg/kg bw, and one at 1,000 mg/kg bw. Symptom of the I reddish changes were observed on application sites in all treated animal black urine was observed in females at 2,000, 1,500 and 1,000 mg/kg bw mortality and clinical signs. No reliable skin/eye irritation study was available. The acute dermal stud irritation. Repeat dose toxicity: In repeated dose toxicity study performed accord Sprague-Dawley rats for 30 days to males and for 39 - 51 days to female value was 5 and 1.3 mg/kg bw/day for male and female rats, respectivel observed in female rats in the high dose group. Erythropoietic toxicity (a of squamous cell hyperplasia of the forestomach was increased in a dos and was statistically significant in males at doses of 20 mg/kg bw/day ar are considered to be local, non-systemic effect on the forestomach whic Genotoxicity: An in vitro genotoxicity study with copper monochlorides <i>Salmonella typhimurium</i> strains (TA 98, TA 100, TA 1535, and TA 1537) vitro test for chromosome aberration in Chinese hamster lung (CHL) cell aberrations at the concentration of 50, 70 and 100 ug/mL without S9 mis	n observed) and 1,224 mg/k hardness of skin, an exudati is. Skin inflammation and inj w. Female rats appeared to l dy with copper monochloride ling to OECD TG 422, coppe es at concentrations of 0, 1.2 y. No death was observed in naemia) was seen in both so se-dependent manner in mal di formales at doses of 5 n h result from organize results in a with and without S9 mix at o s showed that copper mono	g bw for female. Four females died at both ion of hardness site, the formation of scar and jury were also noted. In addition, a reddish or be more sensitive than male based on a suggests that it has a potential to cause skin er monochloride was given orally (gavage) to 3, 5.0, 20, and 80 mg/kg bw/day. The NOAEL n male rats. One treatment-related death was exes at the 80 mg/kg bw/day. The frequency le and female rats at all treatment groups, mg/kg bw/day doses. The observed effects administration of copper monochloride. bacterial reverse mutation test with concentrations of up to 1,000 ug/plate. An in bochloride induced structural and numerical	

in vivo mammalian erythrocyte micronucleus assay, all animals dosed (15 - 60 mg/kg bw) with copper monochloride exhibited similar PCE/(PCE+NCE) ratios and MNPCE frequencies compared to those of the negative control animals. Therefore copper monochloride is not an in vivo mutagen.

Carcinogenicity: there was insufficient information to evaluate the carcinogenic activity of copper monochloride.

Reproductive and developmental toxicity: In the combined repeated dose toxicity study with the reproduction/developmental toxicity screening test (OECD TG 422), copper monochloride was given orally (gavage) to Sprague-Dawley rats for 30 days to males and for 39-51 days to females

	at concentrations of 0, 1.3, 5.0, 20, and 80 mg/kg bw/day. The NOAEL of copper monochloride for fertility toxicity was 80 mg/kg bw/day for the parental animals. No treatment-related effect was observed on the reproductive organs and the fertility parameters assessed. For developmental toxicity the NOAEL was 20 mg/kg bw/day. Three of 120 pups appeared to have icterus at birth; 4 of 120 pups appeared runted at the highest dose tested (80 mg/kg bw/day).
CALCIUM CHLORIDE	For calcium: Toxicity from calcium is not common, because the gastrointestinal tract normally limits the amount of calcium absorbed. Therefore, short-term intake of large amounts of calcium does not generally produce any ill effects aside from constipation and an increased risk of kidney stones. However, more severe toxicity can occur when excess calcium is ingested over long periods, or when calcium is combined with increased amounts of vitamin D, which increases calcium absorption. Calcium toxicity is also found sometimes after excessive administration of calcium via a vein. Toxicity shows as abnormall deposition of calcium in tissues and by elevated blood calcium levels. However, high blood calcium is often due to other causes, such as abnormally high amounts of parathyroid hormone (PTH). Usually, under these circumstances, bone density is lost, and the resulting high blood calcium can cause kidney stones and abdominal pain. Some cancers can also cause high blood calcium, either by secreting abnormal proteins that act like PTH or by invading and killing bone cells causing them to release calcium. Very high levels of calcium can result in appetite loss, nausea, vomiting, abdominal pain, confusion, seizures, and even coma. For calcium chloride: Acute toxicity: The acute oral toxicity of calcium chloride is low. It is attributed to the severe irritating property to the gastrointestinal tract. In humans, acute oral toxicity is rare because large single doses cause nausea and vomiting. There is very little toxicity by skin contact. High blood calcium generally occurs only when there are other factors that affect calcium balance, such as kidney inefficiency and primary thyroid overactivity. Animal testing indicates that calcium chloride is at most slightly irritating to skin, but severely irritation. Repeat dose toxicity: Animal testing did not show evidence of chronic toxicity. Calcium and chloride are both essential nutrients and a daily intake has been recommended. Genetic toxicity: Test results for genetic tox
MAGNESIUM SULFATE, HEPTAHYDRATE	Oral (man) TDLo: 183 mg/kg/4h-L Nil reported
ZINC SULFATE HEPTAHYDRATE	Exposure may produce irreversible effects*. NOTE: Substance has been shown to be mutagenic in at least one assay, or belongs to a family of chemicals producing damage or change to cellular DNA. Oral (human) TDLo: 45 mg/kg/7d-C Eye (rabbit): 0.42 mg moderate Oral (man) TDLo: 180 mg/kg/6w-I Equivocal tumorigenic agent by RTECS criteria. for zinc sulfate heptahydrate Sleep, ataxia, respiratory stimulation, somnolence, coma, diarrhoea, changes in endocrine pancreas recorded.
MANGANESE SULFATE, HYDRATE	Not available.
COPPER SULFATE, BASIC	No significant acute toxicological datum was identified in literature search.
Metalosate Multimineral Amino Acid Soluble Powder & COPPER SULFATE, BASIC	For copper sulfate Copper sulfate is corrosive. Side effects are diverse and multi-systemic, and include severe gastrointestinal symptoms and signs, metallic taste in the mouth, burning pain in the chest, headache, sweating, shock and damage to brain, liver and kidneys. It has been reported as a cause of human suicide. On exposure, it can cause dose dependent damage to the skin and eye, also, eczema and allergic reactions. Long term effects can lead to anaemia and degenerative changes and are more likely in individuals with Wilson's disease, a condition which causes excessive absorption and storage of copper. It has adverse effects on reproduction and fertility as well as cancer and embryo toxic effects. Although it is excreted in the faeces, there is residual accumulation the liver, brain, heart, kidney and muscles.

SECTION 12 Ecological information

Metalosate Multimineral	Endpoint	Test Duration (hr)		Species	Value	Sou	irce
mino Acid Soluble Powder	Not Available	Not Available		Not Available	Not Available	e Not	Available
	Endpoint	Test Duration (hr)		Species	Value		Source
	EC50 (ECx)	48		Crustacea	12.35-16.72 m	ng/L	4
errous sulfate heptahydrate	LC50	96		Fish	6.27-50.35 mg	ı/L	4
	EC50	48		Crustacea	12.35-16.72 m	35-16.72 mg/L	
calcium chloride	Endpoint NOEC (ECx) LC50 EC50 EC50 EC50	Test Duration (hr) 0 96 72 48 96	Crusta	or other aquatic pla		Value 8.879 mg/L 3 mg/L 2900 mg/L 52 mg/L 1109.9 mg/L	Source 4 1 2 1 4
	En du ciut	Test Duration (In)	O meniae		Valu		Source
magnesium sulfate, heptahydrate	Endpoint EC0 (ECx)	Test Duration (hr)	Species	er aquatic plants		ng/L	1
		12	Aigae of our	iei aquatic piants	220	mg/∟	1

	EC50	48		Crustacea			266.4-417.3 mg/L		4
	Endpoint		Test Duration (hr)		Species	Valu	le	So	urce
zinc sulfate heptahydrate	EC50 (ECx)		120		Fish		0.001 mg/L		4100
	EC50		48		Crustacea		0.04 mg/L		
	Endpoint	Tes	t Duration (hr)	Species			Value		Source
	LC50	96		Fish			130.465 mg/L		4
manganese sulfate, hydrate	NOEC (ECx)	96		Fish	Fish		84 mg/L		5
	NOEC (ECx)	1440		Crustacea		0.01 mg/L		2	
	LC50	96		Fish		0.19-12.49 mg/L		4	
	EC50	72		Algae or other aquatic plants		61 mg/L		2	
	EC50	48		Crustacea		7.09-9.36 mg/L		4	
	Endpoint		Test Duration (hr)		Species	Value		6	ource
	-		. ,		•		45	4	burce
copper sulfate, basic	EC50 (ECx)		48 96	Crustacea			0.07-0.115 mg/L		
					Fish 0.47 mg				
	EC50		48		Crustacea	0.07-0.1	15 mg/L	4	
Legend:		Aquatic T	oxicity Data 2. Europe E Toxicity Data 5. ECETO /endor Data						

For Copper Sulfate:

Terrestrial Fate: Soil - If released to soil, copper sulfate may leach to groundwater and may partially oxidize or bind to humic materials, clay or hydrous oxides of iron and manganese. Since copper is an element, it will persist indefinitely. Copper is bound or adsorbed, to organic materials, and to clay and mineral surfaces. The degree of adsorption to soils depends on the acidity or alkalinity of the soil. Copper sulfate is one of the more mobile metals in soil; however, its leaching potential is low in all but sandy soils. When applied with irrigation water, copper sulfate does not accumulate in the surrounding soils; however, some 60% is deposited in the sediments at the bottom of the irrigation ditch, where it becomes adsorbed to clay, mineral, and organic particles. Copper compounds also settle out of solution. Plants – Copper sulfate is toxic to plants and kills by photosynthesis disruption. Blue-green algae have been shown to become increasingly resistant to the algaecide after 26 years of use.

Aquatic Fate: In water, copper sulfate will bind to carbonates as well as humic materials, clay and hydrous oxides of iron and manganese. As an element, copper can persist indefinitely.

Ecotoxicity: Copper is accumulated by plants and animals, but it does not appear to biomagnify from plants to animals. Copper sulphate is practically non-toxic to birds and poses less of a threat to birds than to other animals. Copper sulfate is highly toxic to fish and *Daphnia magna* water fleas. Even at recommended rates of application, this material may be poisonous to trout and other fish, especially in soft or acid waters. Its toxicity to fish generally decreases as water hardness increases. Fish eggs are more resistant than young fish fry to the toxic effects of copper sulfate. Copper sulfate is toxic to aquatic invertebrates, such as crab, shrimp, and oysters. Higher concentrations of the material caused some behavioral changes, such as secretion of mucous, and discharge of eggs and embryos. Bees are endangered by Bordeaux mixtures of copper(II) sulfate and hydrated lime. Copper sulfate may be poisonous to sheep and chickens at normal application rates. Most animal life in soil, including large earthworms, have been eliminated by the extensive use of copper containing fungicides in orchards.

For magnesium compounds in general:

Fish LC50: 100-400 mg/L

For copper:

Atmospheric Fate - Copper is unlikely to accumulate in the atmosphere due to a short residence time for airborne copper aerosols. Airborne coppers, however, may be transported over large distances. Air Quality Standards: no datum available.

Aquatic Fate: Toxicity of copper is affected by pH and hardness of water. Total copper is rarely useful as a predictor of toxicity. In natural sea water, more than 98% of copper is organically bound and in river waters a high percentage is often organically bound, but the actual percentage depends on the river water and its pH.

Ecotoxicity: Copper accumulates significantly in the food chain. The toxic effect of copper in the aquatic biota depends on the bio-availability of copper in water which, in turn, depends on its physico-chemical form (i.e., speciation). Bioavailability is decreased by complexation and adsorption of copper by natural organic matter, iron and manganese hydrated oxides, and chelating agents excreted by algae and other aquatic organisms. Copper exhibits significant toxicity in some aquatic organisms. Some algal species are very sensitive to copper. Silicate, iron, manganese and EDTA may reduce bioavailability.

For copper: Ecotoxicity - Significant effects are expected on various species of microalgae, some species of macroalgae, and a range of invertebrates, including crustaceans, gastropods and sea urchins. Copper is moderately toxic to crab and their larvae and is highly toxic to gastropods (mollusks, including oysters, mussels and clams). In fish, the acute lethal concentrations of copper depends both on test species and exposure conditions. Waters with high concentrations of copper can have significant effects on diatoms and sensitive invertebrates, notably cladocerans (water fleas). Most taxonomic groups of macroalgae and invertebrates will be severely affected. For Inorganic Sulfate:

Environmental Fate - Sulfates can produce a laxative effect at concentrations of 1000 - 1200 mg/liter, but no increase in diarrhea, dehydration or weight loss. The presence of sulfate in drinking-water can also result in a noticeable taste. Sulfate may also contribute to the corrosion of distribution systems. No health-based guideline value for sulfate in drinking water is proposed.

Atmospheric Fate: Sulfates are removed from the air by both dry and wet deposition processes. Wet deposition processes including rain-out (a process that occurs within the clouds) and washout (removal by precipitation below the clouds) which contribute to the removal of sulfate from the atmosphere.

Terrestrial Fate: Soil - In soil, the inorganic sulfates can adsorb to soil particles or leach into surface water and groundwater. Plants - Sodium sulfate is not very toxic to terrestrial plants however; sulfates can be taken up by plants and be incorporated into the parenchyma of the plant. Some plants (e.g., corn and *Kochia scoparia*) are capable of accumulating sulfate to concentrations that are potentially toxic to ruminants. Jack pine are the most sensitive plant species.

Aquatic Fate: Sulfate in water can also be reduced by sulfate bacteria (Thiobacilli) which use them as a source of energy. In anaerobic environments sulfate is biologically reduced to (hydrogen) sulfide by sulfate reducing bacteria, or incorporated into living organisms as source of sulfur. Sodium sulfate is not reactive in aqueous solution at room temperature. Sodium sulfate will completely dissolve, ionize and distribute across the entire planetary 'aquasphere'. Some sulfates may eventually be deposited with the majority of sulfates participating in the sulfur cycle in which natural and industrial sodium sulfates are not distinguishable.

Ecotoxicity: Significant bioconcentration or bioaccumulation is not expected. Algae are the most sensitive to sodium sulfate and toxicity occurs in bacteria from 2500mg/L. Sulfates are not acutely toxic to fish or invertebrates. *Daphnia magna* water fleas and fathead minnow appear to be the least sensitive species. Activated sludge showed a very low sensitivity to sodium sulfate. Overall it can be concluded that sodium sulfate has no acute adverse effect on aquatic and sediment dwelling organisms. No datum was found for long term toxicity. For Copper: Typical foliar levels of copper are: Uncontaminated soils (0.3-250 mg/kg); Contaminated soils (150-450 mg/kg); Mining/smelting soils (6.1-25 mg/kg, 80 mg/kg, 300 mg/kg).

Terrestrial Fate: Plants - Generally, vegetation reflects soil copper levels in its foliage. This is dependent upon the bioavailability of copper and the physiological requirements of species concerned. Crops are often more sensitive to copper than the native flora. Soil: In soil, copper levels are raised by application of fertilizer, fungicides, from deposition of highway dusts and from urban, mining and industrial sources. Chronic and or acute effects on sensitive species occur as a result of human activities such as copper fertilizer addition and addition of sludge. When soil levels exceed 150 mg Cu/kg, native and agricultural species show chronic effects. Soils in the range 500-1000 mg Cu/kg are largely devoid of vegetation cover. The organic content of the soil appears to be a key factor affecting the bioavailability of copper. On normal forest soils, non-rooted plants such as mosses and lichens show

higher copper concentrations. The fruiting bodies and mycorrhizal sheaths of soil fungi associated with higher plants in forests often accumulate copper to much higher levels than plants at the same site.

For Manganese and its Compounds:

Environmental Fate: Manganese is a naturally occurring element in the environment occurring as a result of weathering of geological material. It also occurs from its use in steel manufacture/ coal mining. The most commonly occurring of 11 possible oxidation states are +2 (e.g., manganese chloride or sulfate); +4 (e.g., manganese dioxide); and +7 (e.g., potassium permanganate); although the latter is unstable in the environment.

Atmospheric Fate: Elemental/inorganic manganese compounds may exist in air as suspended particulates from industrial emissions or soil erosion. Manganese-containing particles are mainly removed from the atmosphere by gravitational settling - large particles tend to fall out faster than small particles. The half-life of airborne particles is usually on the order of days, depending on the size of the particle and atmospheric conditions. Some removal by washout mechanisms such as rain may also occur, although it is of minor significance in comparison to dry deposition.

Terrestrial Fate: Manganese in soil can migrate as particulate matter to air or water and soluble manganese compounds can be leached from the soil. High soil pH reduces manganese availability while low soil pH will increase availability, even to the point of toxicity. Soils high in organic matter tie up manganese such that high organic matter soils can be manganese deficient. Fertilization with materials containing chlorine, nitrate, and/or sulfate, can also enhance manganese uptake, (termed the anion effect). Adsorption of soluble manganese to soil/sediments increases as positive ions increase, (cation), and organic matter increases. In some cases, adsorption of manganese to soils may not be a readily reversible process. At low concentrations, manganese may be fixed by clays and will not be released into solution readily. Bacteria and microflora can increase the mobility of manganese.

Aquatic Fate: Most manganese salts, with the exception of phosphates, carbonates, and oxides, are soluble in water. Solubility is controlled by the precipitation of insoluble forms, (species). In most oxygenated waters, the most common form is insoluble manganese oxide. Manganese chloride is the dominant form at pH 4-7, but may oxidize at pH>8 or 9. Ecotoxicity: While lower organisms (plankton, aquatic plants, and some fish) can significantly bioconcentrate manganese, higher organisms (including humans) tend to maintain manganese balance. Manganese in water may be significantly concentrated at lower levels of the food chain.

Uptake of manganese by aquatic invertebrates and fish increases with temperature and decreases with pH. Fish and crustaceans appear to be the most sensitive to acute and chronic exposures. The substance has low toxicity to trout but, is moderately toxic to Coho salmon. The substance is toxic to *Daphnia* water fleas and moderately toxic to freshwater algae *Pseudomonas putida* and *Photobacterium phosphoreum* bacteria.

DO NOT discharge into sewer or waterways.

Persistence and degradability

Ingredient	Persistence: Water/Soil	Persistence: Air
ferrous sulfate heptahydrate	HIGH	HIGH
magnesium sulfate, heptahydrate	HIGH	HIGH
zinc sulfate heptahydrate	HIGH	HIGH

Bioaccumulative potential

Ingredient	Bioaccumulation
ferrous sulfate heptahydrate	LOW (BCF = 52)
magnesium sulfate, heptahydrate	LOW (LogKOW = -2.2002)
zinc sulfate heptahydrate	LOW (BCF = 112)

Mobility in soil

Ingredient	Mobility
ferrous sulfate heptahydrate	LOW (KOC = 6.124)
magnesium sulfate, heptahydrate	LOW (KOC = 6.124)
zinc sulfate heptahydrate	LOW (KOC = 6.124)

SECTION 13 Disposal considerations

Waste treatment methods	
Product / Packaging disposal	 Containers may still present a chemical hazard/ danger when empty. If container can not be cleaned sufficiently well to ensure that residuals do not remain or if the container cannot be used to store the same product, then puncture containers, to prevent re-use, and bury at an authorised landfill. Where possible retain label warnings and SDS and observe all notices pertaining to the product. Legislation addressing waste disposal requirements may differ by country, state and/ or territory. Each user must refer to laws operating in their area. In some areas, certain wastes must be tracked. A Hierarchy of Controls seems to be common - the user should investigate: Reduction Reuse Recycling Disposal (if all else fails) This material may be recycled if unused, or if it has not been contaminated so as to make it unsuitable for its intended use. Shelf life considerations should also be applied in making decisions of this type. Note that properties of a material may change in use, and recycling or reuse may not always be appropriate. In most instances the supplier of the material should be consulted. It may be necessary to collect all wash water for treatment before disposal. In all cases disposal to sewer may be subject to local laws and regulations and these should be considered first. Where in doubt contact the responsible authority. Recycle wherever possible. Consult State Land Waste Management Authority for disposal. Bury residue in an authorised landfill. Recycle containers if possible, or dispose of in an authorised landfill.

SECTION 14 Transport information

Labels Required

Marine Pollutant No

Land transport (DOT): NOT REGULATED FOR TRANSPORT OF DANGEROUS GOODS

Air transport (ICAO-IATA / DGR): NOT REGULATED FOR TRANSPORT OF DANGEROUS GOODS

Sea transport (IMDG-Code / GGVSee): NOT REGULATED FOR TRANSPORT OF DANGEROUS GOODS

Transport in bulk according to Annex II of MARPOL and the IBC code

Not Applicable

Transport in bulk in accordance with MARPOL Annex V and the IMSBC Code

Product name	Group
ferrous sulfate heptahydrate	Not Available
calcium chloride	Not Available
magnesium sulfate, heptahydrate	Not Available
zinc sulfate heptahydrate	Not Available
manganese sulfate, hydrate	Not Available
copper sulfate, basic	Not Available

Transport in bulk in accordance with the ICG Code

Product name	Ship Type
ferrous sulfate heptahydrate	Not Available
calcium chloride	Not Available
magnesium sulfate, heptahydrate	Not Available
zinc sulfate heptahydrate	Not Available
manganese sulfate, hydrate	Not Available
copper sulfate, basic	Not Available

SECTION 15 Regulatory information

Safety, health and environmental regulations / legislation specific for the substance or mixture

ferrous sulfate heptahydrate is found on the following regulatory lists

US - Massachusetts - Right To Know Listed Chemicals

US ACGIH Threshold Limit Values (TLV)

US CWA (Clean Water Act) - List of Hazardous Substances

US DOE Temporary Emergency Exposure Limits (TEELs)

calcium chloride is found on the following regulatory lists

US DOE Temporary Emergency Exposure Limits (TEELs)

US Toxic Substances Control Act (TSCA) - Chemical Substance Inventory

magnesium sulfate, heptahydrate is found on the following regulatory lists

FEI Equine Prohibited Substances List - Controlled Medication

- FEI Equine Prohibited Substances List (EPSL)
- US DOE Temporary Emergency Exposure Limits (TEELs)

zinc sulfate heptahydrate is found on the following regulatory lists

US - Massachusetts - Right To Know Listed Chemicals

- US CWA (Clean Water Act) List of Hazardous Substances
- US CWA (Clean Water Act) Priority Pollutants
- US CWA (Clean Water Act) Toxic Pollutants
- US DOE Temporary Emergency Exposure Limits (TEELs)

manganese sulfate, hydrate is found on the following regulatory lists

US - California Hazardous Air Pollutants Identified as Toxic Air Contaminants

- US ACGIH Threshold Limit Values (TLV)
- US ACGIH Threshold Limit Values (TLV) Carcinogens
- US Clean Air Act Hazardous Air Pollutants
- US DOE Temporary Emergency Exposure Limits (TEELs)

copper sulfate, basic is found on the following regulatory lists

US CWA (Clean Water Act) - Priority Pollutants

US CWA (Clean Water Act) - Toxic Pollutants

US EPCRA Section 313 Chemical List

Federal Regulations

Superfund Amendments and Reauthorization Act of 1986 (SARA)

US List of Active Substances Exempt from the TSCA Inventory Notifications (Active-Inactive) Rule

US NIOSH Recommended Exposure Limits (RELs)

US Toxic Substances Control Act (TSCA) - Chemical Substance Inventory US TSCA Chemical Substance Inventory - Interim List of Active Substances

US TSCA Chemical Substance Inventory - Interim List of Active Substances

US Toxic Substances Control Act (TSCA) - Chemical Substance Inventory US TSCA Chemical Substance Inventory - Interim List of Active Substances

US EPA Integrated Risk Information System (IRIS)

- US EPCRA Section 313 Chemical List
- US Toxic Substances Control Act (TSCA) Chemical Substance Inventory US TSCA Chemical Substance Inventory - Interim List of Active Substances

US EPCRA Section 313 Chemical List

- US NIOSH Recommended Exposure Limits (RELs)
- US OSHA Permissible Exposure Limits (PELs) Table Z-1
- US Toxic Substances Control Act (TSCA) Chemical Substance Inventory

US TSCA Chemical Substance Inventory - Interim List of Active Substances

US List of Active Substances Exempt from the TSCA Inventory Notifications (Active-Inactive) Rule

Continuación...

US Toxic Substances Control Act (TSCA) - Chemical Substance Inventory

Continuación...

Metalosate Multimineral Amino Acid Soluble Powder

Section 311/312 hazard categories

Flammable (Gases, Aerosols, Liquids, or Solids)	No
Gas under pressure	No
Explosive	No
Self-heating	No
Pyrophoric (Liquid or Solid)	No
Pyrophoric Gas	No
Corrosive to metal	No
Oxidizer (Liquid, Solid or Gas)	No
Organic Peroxide	No
Self-reactive	No
In contact with water emits flammable gas	No
Combustible Dust	Yes
Carcinogenicity	No
Acute toxicity (any route of exposure)	No
Reproductive toxicity	No
Skin Corrosion or Irritation	No
Respiratory or Skin Sensitization	No
Serious eye damage or eye irritation	Yes
Specific target organ toxicity (single or repeated exposure)	Yes
Aspiration Hazard	No
Germ cell mutagenicity	No
Simple Asphyxiant	No
Hazards Not Otherwise Classified	No

Name	Reportable Quantity in Pounds (Ib)	Reportable Quantity in kg
ferrous sulfate heptahydrate	1000	454
zinc sulfate heptahydrate	1000	454

National Inventory Status

National Inventory	Status	
Australia - AIIC / Australia Non-Industrial Use	Yes	
Canada - DSL	Yes	
Canada - NDSL	o (ferrous sulfate heptahydrate; calcium chloride; magnesium sulfate, heptahydrate; zinc sulfate heptahydrate; manganese sulfate, hydrate; opper sulfate, basic)	
China - IECSC	Yes	
Europe - EINEC / ELINCS / NLP	Yes	
Japan - ENCS	No (ferrous sulfate heptahydrate; copper sulfate, basic)	
Korea - KECI	Yes	
New Zealand - NZIoC	Yes	
Philippines - PICCS	Yes	
USA - TSCA	Yes	
Taiwan - TCSI	Yes	
Mexico - INSQ	No (copper sulfate, basic)	
Vietnam - NCI	Yes	
Russia - FBEPH	No (copper sulfate, basic)	
Legend:	Yes = All CAS declared ingredients are on the inventory No = One or more of the CAS listed ingredients are not on the inventory. These ingredients may be exempt or will require registration.	

SECTION 16 Other information

Revision Date	09/05/2022
Initial Date	01/07/2018

SDS Version Summary

Version	Date of Update	Sections Updated
10.15	08/05/2022	Acute Health (inhaled), Chronic Health, Exposure Standard, Personal Protection (Respirator)

Other information

Classification of the preparation and its individual components has drawn on official and authoritative sources as well as independent review by the Chemwatch Classification committee using available literature references.

The SDS is a Hazard Communication tool and should be used to assist in the Risk Assessment. Many factors determine whether the reported Hazards are Risks in the workplace or other settings. Risks may be determined by reference to Exposures Scenarios. Scale of use, frequency of use and current or available engineering controls must be considered.

Definitions and abbreviations

PC-TWA: Permissible Concentration-Time Weighted Average PC-STEL: Permissible Concentration-Short Term Exposure Limit IARC: International Agency for Research on Cancer ACGIH: American Conference of Governmental Industrial Hygienists STEL: Short Term Exposure Limit TEEL: Temporary Emergency Exposure Limit. IDLH: Immediately Dangerous to Life or Health Concentrations ES: Exposure Standard OSF: Odour Safety Factor NOAEL :No Observed Adverse Effect Level LOAEL: Lowest Observed Adverse Effect Level TLV: Threshold Limit Value LOD: Limit Of Detection OTV: Odour Threshold Value BCF: BioConcentration Factors **BEI: Biological Exposure Index** AIIC: Australian Inventory of Industrial Chemicals DSL: Domestic Substances List NDSL: Non-Domestic Substances List IECSC: Inventory of Existing Chemical Substance in China EINECS: European INventory of Existing Commercial chemical Substances ELINCS: European List of Notified Chemical Substances NLP: No-Longer Polymers ENCS: Existing and New Chemical Substances Inventory KECI: Korea Existing Chemicals Inventory NZIoC: New Zealand Inventory of Chemicals PICCS: Philippine Inventory of Chemicals and Chemical Substances TSCA: Toxic Substances Control Act TCSI: Taiwan Chemical Substance Inventory INSQ: Inventario Nacional de Sustancias Químicas NCI: National Chemical Inventory FBEPH: Russian Register of Potentially Hazardous Chemical and Biological Substances

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