Balchem Corporation

Catalogue number: Various Version No: 16.22 Safety Data Sheet according to OSHA HazCom Standard (2012) requirements

Issue Date: 4/8/2020 S.GHS.USA.EN

SECTION 1 IDENTIFICATION

Product Identifier

Product name	Choline Chloride, Dry, Feed Grade
Synonyms	50% Choline Chloride Dry, on Silica (F3040035, F3040130); 60% Choline Chloride Dry (F3070030); 60% Choline Chloride with Carrier (F3070530); 70% Choline Chloride Dry (F3080030); 70% Choline Chloride with Carrier (F3080530); Choline Chloride 60% Vegetable Carrier (F3070330); Choline Chloride 70% Vegetable Carrier (F3080330); PetShure® 60% Choline Chloride Dry (F3070730); PetShure® 70% Choline Chloride Dry (F3080930); PetShure® 97% Choline Chloride Dry (F3524120).
Other means of identification	Various

Recommended use of the chemical and restrictions on use

Relevant identified uses	Feed Additive
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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	
Website	www.balchem.com	
Email	sds@balchem.com	

Emergency phone number

Association / Organisation	Chemtrec CCN#2275
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



Note: The hazard category numbers found in GHS classification in section 2 of this SDSs are NOT to be used to fill in the NFPA 704 diamond. Blue = Health Red = Fire Yellow = Reactivity White = Special (Oxidizer or water reactive substances)

Classification Not classified for physical or health hazards according to 2012 OSHA HazCom Standard (29 CFR 1910.1200)

Label elements

Hazard pictogram(s)	Not Applicable

SIGNAL WORD NOT APPLICABLE

Hazard statement(s)

Not Applicable

Hazard(s) not otherwise classified

May form combustible dust

Precautionary statement(s) Prevention
Not Applicable
Precautionary statement(s) Response
Not Applicable
Precautionary statement(s) Storage
Not Applicable
Precautionary statement(s) Disposal

Not Applicable

SECTION 3 COMPOSITION / INFORMATION ON INGREDIENTS

Substances

See section below for composition of Mixtures

Mixtures

CAS No	%[weight]	Name
Not Applicable	Not applicable	Not classified for physical or health hazards according to 2012 OSHA HazCom Standard (29 CFR 1910.1200)

SECTION 4 FIRST-AID MEASURES

Description of first aid measures

Eye Contact	 If in eyes, hold eyelids apart and flush the eye continuously with running water. Continue flushing for at least 15 minutes. 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. 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 DO NOT induce vomiting. If vomiting occurs, lean patient forward or place on left side (head-down position, if possible) to maintain open airway and prevent aspiration. Observe the patient carefully. Never give liquid to a person showing signs of being sleepy or with reduced awareness; i.e., becoming unconscious. Give water to rinse out mouth, then provide liquid slowly and as much as casualty can comfortably drink. If poisoning occurs, contact a doctor or Poisons Information Centre.

Most important symptoms and effects, both acute and delayed

See Section 11

Indication of any immediate medical attention and special treatment needed

For exposures to quaternary ammonium compounds;

- For ingestion of concentrated solutions (10% or higher): Swallow promptly a large quantity of milk, egg whites / gelatin solution. If not readily available, a slurry of activated charcoal may be useful. Avoid alcohol. Because of probable mucosal damage omit gastric lavage and emetic drugs.
- For dilute solutions (2% or less): If little or no emesis appears spontaneously, administer syrup of Ipecac or perform gastric lavage.
- If hypotension becomes severe, institute measures against circulatory shock.
- If respiration laboured, administer oxygen and support breathing mechanically. Oropharyngeal airway may be inserted in absence of gag reflex. Epiglottic or laryngeal edema may necessitate a tracheotomy.
- Persistent convulsions may be controlled by cautious intravenous injection of diazepam or short-acting barbiturate drugs. [Gosselin et al, Clinical Toxicology of Commercial Products]

SECTION 5 FIRE-FIGHTING MEASURES

Extinguishing media

- Foam.
- Dry chemical powder.BCF (where regulations permit).
- Carbon dioxide.
- Water spray or fog Large fires only.

Special hazards arising from the substrate or mixture

Fire Incompatibility Avoid contamination with oxidising agent, e.g., nitrates, oxidising acids, chlorine bleaches, pool chlorine etc., as ignition may result.	
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Special protective equipment and precautions for fire-fighters

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		Fire Fighting	 Alert Fire Brigade and tell them location and nature of hazard. Wear breathing apparatus plus protective gloves. Prevent spillage from entering drains or water courses. Use water delivered as a fine spray to control fire and cool adjacent 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.
	Fire	e/Explosion Hazard	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

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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.
* Kst - a normalised expression of the burning dust pressure rise rate over time.
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 junition 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.
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.
70% dry choline chloride is an ST1 dust.
Pmax = 6.8 bar Kst = 137 bar-m/s
Rate of Pressure Rise = 505 bar/s
 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 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 ginted - particles exceeding this limit will generally not form flammable dust duvide: according or particular particular particular by the particular by the propagation of an explosion of accumulations of fine dust (420 micron or less) may burn rapidly and fiercely if ginted - particles exceeding this limit will generally not form flammable dust duvide: according to the particular particular by the part
 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 'Winimum 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 studien release of statically charged materials from storage of process equipment, particularly at elevated temperatures and/ of 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 or now 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. Combustion products include:
carbon monoxide (CO) carbon dioxide (CO2)
hydrogen chloride
phosgene
nitrogen oxides (NUX) other pyrolysis products typical of burning organic material.
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	 Clean up waste regularly and abnormal spills immediately. Avoid breathing dust and contact with skin and eyes. Wear protective clothing, gloves, safety glasses and dust respirator. Use dry clean up procedures and avoid generating dust. Vacuum up or sweep up. NOTE: Vacuum cleaner must be fitted with an exhaust micro filter (HEPA type) (consider explosion-proof machines)
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designed to be grounded during storage and use). > Dampen with water to prevent dusting before sweeping. > Place in suitable containers for disposal. 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

Safe handling	 Avoid all personal contact, including inhalation. Wear protective clothing when risk of exposure occurs. Use in a well-ventilated area. DO NOT enter confined spaces until atmosphere has been checked. Avoid contact with incompatible materials. When handing, DO NOT est, drink or smoke. Keep containers securely sealed when not in use. Avoid physical damage to containers. Always wash hands with scap and water after handing. Work clothes should be laundered separately. Launder contaminated clothing before re-use. Use good occupational work practice. Observe manufacturer's storage and handling recommendations contained within this SDS. Atmosphere should be regularly checked against established exposure standards to ensure safe working conditions are maintained. Organic powders when finely divided over a range of concentrations 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 accumulation of dusts. Particular attention should be given to overhead and hidden horizontal surfaces to 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. Vaccuums with explosion-proor motors
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. Protect containers against physical damage and check regularly for leaks. Observe manufacturer's storage and handling recommendations contained within this SDS. For major quantities: Consider storage in bunded areas - ensure storage areas are isolated from sources of community water (including stormwater, ground water, lakes and streams). Ensure that accidental discharge to air or water is the subject of a contingency disaster management plan; this may require consultation with local authorities.

Conditions for safe storage, including any incompatibilities

Suitable container	 Glass container is suitable for laboratory quantities Polyethylene or polypropylene container. Check all containers are clearly labelled and free from leaks.
Storage incompatibility	 Quaternary ammonium cations are unreactive toward even strong electrophiles, oxidants, and acids. They also are stable toward most nucleophiles. The latter is indicated by the stability of the hydroxide salts such as tetramethylammonium hydroxide and tetrabutylammonium hydroxide. Quaternary ammonium compounds are deactivated by anionic detergents (including common soaps). With exceptionally strong bases, quat cations degrade. They undergo Sommelet–Hauser rearrangement and Stevens rearrangement, as well as dealkylation under harsh conditions. Quaternary ammonium cations containing N–C–C–H units can also undergo the Hofmann elimination and Emde degradation. Avoid reaction with oxidising agents

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SECTION 8 EXPOSURE CONTROLS / PERSONAL PROTECTION

Control parameters

OCCUPATIONAL EXPOSURE LIMITS (OEL)

INGREDIENT DATA

Not Available

EMERGENCY LIMITS

Ingredient	Material name	TEEL-1	TEEL-2	TEEL-3
Choline Chloride, Dry, Feed Grade	Not Available	Not Available	Not Available	Not Available
Ingredient	Original IDLH		Revised IDLH	
choline chloride	Not Available		Not Available	

OCCUPATIONAL EXPOSURE BANDING

Ingredient	Occupational Exposure Band Rating	Occupational Exposure Band Limit
choline chloride	E	≤ 0.01 mg/m³
Notes:	Occupational exposure banding is a process of assigning chemicals into specific categories or bands based on a chemical's potency and the adverse health outcomes associated with exposure. The output of this process is an occupational exposure band (OEB), which corresponds to range of exposure concentrations that are expected to protect worker health.	

Exposure controls

	Enclosed local exhaust ventilation is required at points of d HEPA terminated local exhaust ventilation should be consid Barrier protection or laminar flow cabinets should be consid A fume hood or vented balance enclosure is recommended When handling quantities up to 500 grams in either a stanc preferred. Quantities up to 1 kilogram may require a design enclosures. Quantities exceeding 1 kilogram should be har containment technology. Manufacturing and pilot plant operations require barrier/ co Barrier/ containment technology and direct coupling (totally typically use double or split butterfly valves and hybrid unid Glove bags, isolator glove box systems are optional. HEPA Fume-hoods and other open-face containment technolog non-routine emergencies maximum local and general exha 'escape' velocities which, in turn, determine the 'capture ve	ust, fume or vapour generation. dered at point of generation of dust dered for laboratory scale handling. I for weighing/ transferring quantitie and laboratory with general dilution hated laboratory using fume hood, b ddled in a designated laboratory or ntainment and direct coupling tech e enclosed processes that create a lirectional airflow/ local exhaust ver filtration of exhaust from dry produ acceptable when face velocities of gies are required to prevent migrati- uust are necessary. Air contaminant locities' of fresh circulating air requ	, fumes or vapours. es exceeding 500 mg. o ventilation (e.g., 6-12 air oiological safety cabinet, o containment laboratory us nologies. barrier between the equipp tillation solutions (e.g. pow titlation solutions (e.g. pow tit handling areas is requi i at least 1 m/s (200 feet/m on of the material to uncor s generated in the workpla ired to effectively remove 1	changes per hour) is r approved vented ing appropriate barrier/ ment and the room) vder containment booths). red. ninute) are achieved. ntrolled areas. For ace possess varying the contaminant.
	Type of Contaminant:			Air Speed:
	solvent, vapours, etc. evaporating from tank (in still air)			0.25-0.5 m/s (50-100 ft/min.)
	aerosols, fumes from pouring operations, intermittent convelocity into zone of active generation)	tainer filling, low speed conveyer tra	ansfers (released at low	0.5-1 m/s (100-200 ft/min.)
Appropriate engineering direct spray, drum filling, conveyer loading, crusher dusts, gas discharge (active generation motion)			nto zone of rapid air	1-2.5 m/s (200-500 ft/min.)
controls	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		
	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		
	Simple theory shows that air velocity falls rapidly with distance away from the opening of a simple extraction pipe. Velocity generally decreases with the square of distance from the extraction point (in simple cases). Therefore the air speed at the extraction point should be adjusted, accordingly, after reference to distance from the contaminating source. The air velocity at the extraction point. Other mechanical considerations, producing performance deficits within the extraction apparatus, make it essential that theoretical air velocities are multiplied by factors of 10 or more when extraction systems are installed or used. The need for respiratory protection should also be assessed where incidental or accidental exposure is anticipated: Dependent on levels of contamination, PAPR, full face air purifying devices with P2 or P3 filters or air supplied respirators should be evaluated. The following protective devices are recommended where exposures exceed the recommended exposure control guidelines by factors of: 10; high efficiency particulate (HEPA) filters or cartridges 10-25; loose-fitting (Tyvek or helmet type) HEPA powered-air purifying respirator. 25-50; a full face-piece negative pressure respirator with HEPA filters 50-100; tight-fitting, full face-piece HEPA PAPR 100-1000; a hood-shroud HEPA PAPR or full face-piece supplied air respirator operated in pressure demand or other positive pressure mode.			ity generally decreases ould be adjusted, e, should be a minimum anical considerations, lied by factors of 10 or rendent on levels of lines by factors of:



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Eye and face protection	 When handling very small quantities of the material eye protection may not be required. For laboratory, larger scale or bulk handling or where regular exposure in an occupational setting occurs: Chemical goggles. Face shield. Full face shield may be required for supplementary but never for primary protection of eyes. 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	The selection of suitable gloves does not only depend on the material, but also on further marks of quality which way from manufacturer to manufacturer. Where the chemical is a preparation of several substances, the resistance of the glove material can not be calculated in advance and has therefore to be checked prior to the application. The exact threak through time for substances has to be obtained from the manufacturer of the protective gloves and has to be observed when making a final choice. Personal hygines is a key element of effective hand care. Gloves must only be wom on clean hands. After using gloves, hands should be washed and dired thoroughly. Application of a non-perfumed molisturiaer is recommended. Suitability and duratition of contrad. - denical resistance of glove material, - glove thickness and - devicetify Select gloves tested to a relevant standard (e.g., Europe EN 374, US F739, ASNZS 2161.1 or national equivalent). - Mhen ordingetor (frequently repeated contain two y court, a glove with a protection class of 5 or higher (freakthrough time greater than 240 minutes according to EN 374, ASNZS 2161.10 or national equivalent) is recommended. - When ordingetor fortunes in the protection class of 3 or higher (freakthrough time greater than 240 minutes according to EN 374, ASNZS 2161.10 or national equivalent) is recommended. - Some glove polymer types are less affected by movement and this should be taken into account when considering gloves for long-term use. - Contaminated gloves should be reglaced. - Some glove polymer types are less affected by movement and this should be taken into account when considering gloves for long-term use. - Fordinet in the take regulave thinking a good predictor of glove resistance to a specific chamical, as the permeation efficiency of the glove thickness is no holesadows - Fordinet motions, gloves with a thickness typ polyling greater than 0.35 mm, are recommended. - Thore withing type manal diagrades - Proving theore material diagrades t
Body protection	See Other protection below
Other protection	 For quantities up to 500 grams a laboratory coat may be suitable. For quantities up to 1 kilogram a disposable laboratory coat or coverall of low permeability is recommended. Coveralls should be buttoned at collar and cuffs. For quantities over 1 kilogram and manufacturing operations, wear disposable coverall of low permeability and disposable shoe covers. For manufacturing operations, air-supplied full body suits may be required for the provision of advanced respiratory protection. Eye wash unit. Ensure there is ready access to an emergency shower. For Emergencies: vinyl suit

Respiratory protection

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

Required Minimum Protection Factor	Half-Face Respirator	Full-Face Respirator	Powered Air Respirator
up to 10 x ES	P1	-	PAPR-P1
up to 50 x ES	Air-line*	- P2	- PAPR-P2
up to 100 x ES	-	P3	-

		Air-line*	-
100+ x ES	-	Air-line**	PAPR-P3
* Negative pressure demand ** Continuous flow			

* - Negative pressure demand ** - Continuous flow

A(All classes) = Organic vapours, B AUS or B1 = Acid gasses, B2 = Acid gas or hydrogen cyanide(HCN), B3 = Acid gas or hydrogen cyanide(HCN), E = Sulfur dioxide(SO2), G = Agricultural chemicals, K = Ammonia(NH3), Hg = Mercury, NO = Oxides of nitrogen, MB = Methyl bromide, AX = Low boiling point organic compounds(below 65 degC)

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.
- + Use approved positive flow mask if significant quantities of dust becomes airborne.
- Try to avoid creating dust conditions.

SECTION 9 PHYSICAL AND CHEMICAL PROPERTIES

Information on basic physical and chemical properties

Appearance	White / Yellow / Brown granules or powder		
Physical state	Solid	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 Applicable
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	Immiscible	pH as a solution (10%)	6
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.
Ingestion	Accidental ingestion of the material may be damaging to the health of the individual. Nausea, vomiting, abdominal discomfort and diarrhoea can occur after large doses of choline. Other side effects reported include incontinence or worsening of incontinence, depression or worsening of depression and an unpleasant fishy odour, restlessness, fast breathing, and convulsions. Sluggish movement, tremor, and rigidity as well as liver damage have been observed at high doses. Very high doses may be lethal.

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	Concentrated solutions of many cationics may cause cor (sometimes bloody) may follow ingestion.	rosive damage	to mucous membranes and the oesophagus. Nausea and vomiting	
Skin Contact	Skin contact is not thought to have harmful health effects (as classified under EC Directives); the material may still produce health damage following entry through wounds, lesions or abrasions. 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. This material can cause inflammation of the skin on contact in some persons.			
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.			
Chronic	Long-term exposure to the product is not thought to produce chronic effects adverse to the health (as classified by EC Directives using animal models); nevertheless exposure by all routes should be minimised as a matter of course. Prolonged or repeated skin contact may cause degreasing, followed by drying, cracking and skin inflammation.			
Choline Chloride, Dry, Feed	TOXICITY IRRITATION			
Grade	Not Available Not Available			
	TOXICITY	IRRITATION		
choline chloride	Oral (rat) LD50: 3400 mg/kg ^[2] Eye: no adverse effect observed (not irritating) ^[1]			

 Legend:
 1. Value obtained from Europe ECHA Registered Substances - Acute toxicity 2.* Value obtained from manufacturer's SDS. Unless otherwise specified data extracted from RTECS - Register of Toxic Effect of chemical Substances

Skin: no adverse effect observed (not irritating)^[1]

Acute Toxicity	×	Carcinogenicity	×
Skin Irritation/Corrosion	×	Reproductivity	×
Serious Eye Damage/Irritation	×	STOT - Single Exposure	×
Respiratory or Skin sensitisation	×	STOT - Repeated Exposure	×
Mutagenicity	×	Aspiration Hazard	×



Data either not available or does not fill the criteria for classification
 Data available to make classification

SECTION 12 ECOLOGICAL INFORMATION

Toxicity	
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Choline Chloride, Dry, Feed Grade		ADPOINT TEST DURATION (HR)		SPECIES	VALUE		SOURCE
	NOT AValiable NOT AValiable NOT AValiable NOT AValiable NOT AValiable NOT AValiable						
	ENDPOINT	TEST DURATION (HR)	SPEC	IES		VALUE	SOURCE
choline chloride	LC50	96	Fish	Fish :		>10 mg/L	2
	EC50	48	Crusta	Crustacea		>320 mg/L	4
	EC50	72	Algae	or other aquatic plants		>500 mg/L	2
	NOEC	96	Fish	Fish 10		10 mg/L	2
Legend:	Extracted from 1. IUCLID Toxicity Data 2. Europe ECHA Registered Substances - Ecotoxicological Information - Aquatic Toxicity 3. EPIWIN Suite V3.12 (QSAR) - Aquatic Toxicity Data (Estimated) 4. US EPA, Ecotox database - Aquatic Toxicity Data 5. ECETOC Aquatic Hazard Assessment Data 6. NITE (Japan) - Bioconcentration Data 7. METI (Japan) - Bioconcentration Data 8. Vendor Data						

For choline and its salts (typically choline chloride):

Environmental Fate: Choline chloride is a quaternary amine salt, it dissociates in water into the corresponding positively charged quaternary hydroxyl alkylammonium ion and the negatively charged chloride ion. Chlorine chloride is a weak acid with a lowest pH value of 4 being measured at different concentrations.

Photodegradation: In the air, models predict that the substance will be rapidly degraded with a calculated t1/2 of about 6.9 hours for OH-radicals.

Stability in Water: Choline chloride is a quaternary ammonium salt and dissociates in water. Hydrolysis at environmental pH conditions is not to be expected due to the chemical structure of choline chloride.

Transport between Environmental Compartments: Releases into the environment may occur during production, further processing and use of choline chloride. Models indicate water to be the main target compartment (100% for pure choline chloride), with amounts partitioning to other compartments (air, soil, sediment, suspended sediments, fish and aerosol) considered negligible (< 0.0001%).

Choline chloride will not adsorb on soil and sediments or suspended solids.

Choline chloride will not rapidly evaporate into the atmosphere.

Biodegradation: Choline chloride was shown to be readily biodegradable.

Bioaccumulation: No measured datum on bioaccumulationis available, however model predictions show that bioaccumulation is not likely in aquatic organisms. The bioaccumulation potential of choline chloride is considered to be low.

Ecotoxicity: Fish LC50 (96 h): Oryzias latipes >100 mg/L (nominal and measured, flow through OECD 203); Leuciscus idus >10,000 mg/L (nominal, static, DIN 38412, part 15); Limanda limandu >1000 mg/L (nominal, semistatic, OECD 203)

Daphnia magna EC50 (48 h): >500 mg/L (nominal); NOEC 125 (Directive 79/831 EEC, C2, static)

Algae ErC50 (72 h): Pseudokirchneriella subcapitata >1000 mg/L (nominal and measured); NOEC 32 mg/L (growth rate) (OECD 201); EbC50 Scenedesmus subspicatus >500 mg/L (nominal); NOEC >500 mg/L (growth rate and biomass) (DIN 38412, part 9, static)

Bacterial EC10 (17 h): Pseudomonas putida 113 mg/L; EC50 (17 h): 133 mg/L; EC90 (17 h): 278 mg/L (all nominal) (DIN 38412, part 8 static)

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Choline Chloride, Dry, Feed Grade

Based on these data choline chloride is considered unlikely to be harmful to aquatic organisms.

NOEC of 30.2 mg/L could also be considered, resulting in a PNECaqua of 3.02 mg/L, which is close to the other value obtained using the acute data.

For Quaternary Ammonium Compounds (QACs): QACs are white, crystalline powders. Low molecular weight QACs are very soluble in water, but slightly or not at all soluble in solvents such as ether, petrol and benzene. As the molecular weight and chain lengths increases, the solubility in polar solvents (e.g., water) decreases and the solubility in non-polar solvents increases.

Environmental Fate: A major part of the QACs is discharged into wastewater and removed in the biological processes of sewage treatment, however; the aerobic and anaerobic biodegradability of QACs is not well investigated. Only sparse data are available concerning stability, solubility and biodegradability. In general, it seems that the biodegradability decreases with increasing numbers of alkyl chains. Within each category the biodegradability seems inversely proportional to the alkyl chain length. Heterocyclic QACs are less degradable than the non-cyclic.

Ecotoxicity: Significant bioaccumulation is not expected.

Aquatic Fate: The toxicity of QACs is known to be greatly reduced in the environment because of preferential binding to dissolved organics in surface water.

For Chloride: Although inorganic chloride ions are not normally considered toxic they can exist in effluents at acutely toxic levels. Incidental exposure to inorganic chloride may occur in occupational settings where chemicals management policies are improperly applied. The toxicity of chloride salts depends on the counter-ion (cation) present; that of chloride itself is unknown. Chloride toxicity has not been observed in humans except in the special case of impaired sodium chloride metabolism, e.g., in congestive heart failure. Healthy individuals can tolerate the intake of large quantities of chloride provided that there is an intake of fresh water following ingestion. Although excessive intake of rinking-water containing sodium chloride at concentrations above 2.5 g/L has been reported to produce hypertension, this effect is believed to be related to the sodium ion concentration. Chloride concentrations in excess of about 250 mg/L can give rise to detectable taste in water. Consumers can, however, become accustomed to concentrations in excess of 250 mg/L. No health-based guideline value is proposed for chloride in drinking-water. Chloride is almost completely absorbed in normal individuals. In metal pipes, chloride reacts with metal ions to form soluble salts thus increasing levels of metals in drinking-water. Chloride enhances galvanic corrosion in lead pipes and can also increase the rate of pitting corrosion of metal pipes. Aquatic Fate: Inorganic chlorine eventually finds its way into aquatic systems and becomes bio-available. Chloride increases the electrical conductivity of water and thus increases its corrosivity.

Ecotoxicity: When excessive inorganic chloride ions are introduced to aquatic environments, the resulting salinity can exceed the tolerances of most freshwater organisms.

Persistence and degradability

Ingredient	Persistence: Water/Soil	Persistence: Air
choline chloride	LOW	LOW

Bioaccumulative potential

Ingredient	Bioaccumulation	
choline chloride	LOW (LogKOW = -3.1555)	
Mobility in soil		
Ingredient	Mobility	

SECTION 13 DISPOSAL CONSIDERATIONS

Waste treatment methods

choline chloride

Product / Packaging disposal	 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.
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SECTION 14 TRANSPORT INFORMATION

Labels Required

Marine Pollutant

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

HIGH (KOC = 1.717)

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

No

Not Applicable

SECTION 15 REGULATORY INFORMATION

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

CHOLINE CHLORIDE IS FOUND ON THE FOLLOWING REGULATORY LISTS

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

US TSCA Chemical Substance Inventory - Interim List of Active Substances

Federal Regulations

Superfund Amendments and Reauthorization Act of 1986 (SARA)

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	No
Specific target organ toxicity (single or repeated exposure)	No
Aspiration Hazard	No
Germ cell mutagenicity	No
Simple Asphyxiant	No
Hazards Not Otherwise Classified	No

US. EPA CERCLA HAZARDOUS SUBSTANCES AND REPORTABLE QUANTITIES (40 CFR 302.4) None Reported

State Regulations

None Reported

National Inventory Status

National Inventory	Status
Australia - AICS	Yes
Canada - DSL	Yes
Canada - NDSL	Yes
China - IECSC	Yes
Europe - EINEC / ELINCS / NLP	Yes
Japan - ENCS	Yes
Korea - KECI	Yes
New Zealand - NZIoC	Yes
Philippines - PICCS	Yes
USA - TSCA	Yes
Taiwan - TCSI	Yes
Mexico - INSQ	Yes
Vietnam - NCI	Yes
Russia - ARIPS	Yes
Legend:	Yes = All CAS declared ingredients are on the inventory No = One or more of the CAS listed ingredients are not on the inventory and are not exempt from listing(see specific ingredients in brackets)

SECTION 16 OTHER INFORMATION

Revision Date	25/03/2020
Initial Date	04/04/2019

SDS Version Summary

Version	Issue Date	Sections Updated
15.22.1.1.1	23/03/2020	Ingredients, Synonyms

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

Catalogue number: Various Version No: 16.22

Choline Chloride, Dry, Feed Grade

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 OSF: Odour Safety Factor NOAEL :No Observed Adverse Effect Level LOAEL: Lowest 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

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