

TALARC NiCrMo TALARC

Chemwatch: **5256-76** Version No: **3.1.1.1** Safety Data Sheet according to WHS and ADG requirements Chemwatch Hazard Alert Code: 4

Issue Date: 01/11/2019 Print Date: 11/11/2020 L.GHS.AUS.EN

SECTION 1 Identification of the substance / mixture and of the company / undertaking

Product Identifier

Product name	TALARC NICrMo
Synonyms	E10018M; 3.2 mm, VAC Pack 2 Kg, Part no. INE80B32
Other means of identification	Not Available

Relevant identified uses of the substance or mixture and uses advised against

Relevant identified uses	Low-alloy basic-coated electrode with Ni-Cr-Mo additions designed for welding high yield strength steels with minimum tensile strength higher than 690 MPa. Good impact strength at low temperatures. Suitable for the metal working industry, offshore fabrication, chemical and petrochemical industry. It also has applications in fabrications of HSLA (high-strength low-alloy) steels, which may be used for industrial machinery construction, cranes and other highly stressed structural components. for welding fume: In addition to complying with any individual exposure standards for specific contaminants, where current manual welding processes are used, the fume concentration inside the welder's helmet should not exceed 5 mg/m3, when collected in accordance with the appropriate standard (AS 3640, for example). ES* TWA: 5 mg/m3 TLV* TWA: 5 mg/m3 Most welding, even with primitive ventilation, does not produce exposures inside the welding helmet above 5 mg/m3. That which does should be controlled (ACGIH). Inspirable dust concentrations in a worker's breathing zone shall be collected and measured in accordance with AS 3640, for example. Metal content can be analytically determined by OSHA Method ID25 (ICP-AES) after total digestion of filters and dissolution of captured metals. Sampling of the Respirable Dust fraction requires cyclone separator devices (elutriators) and procedures to comply with AS 2985 (for example).
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Details of the supplier of the safety data sheet

Registered company name	TALARC
Address	10-16 Syme Street Brunswick VIC 3056 Australia
Telephone	+61 3 9388 0588
Fax	+61 3 9388 0710
Website	www.talarc.com.au
Email	sales@talarc.com

Emergency telephone number

Association / Organisation	TALARC
Emergency telephone numbers	+61 3 9388 0588 (Hours 9am-5pm AEST)
Other emergency telephone numbers	Not Available

SECTION 2 Hazards identification

Classification of the substance or mixture

Poisons Schedule	Not Applicable
Classification ^[1]	Acute Toxicity (Inhalation) Category 4, Carcinogenicity Category 1A

Legend:	1. Classified by Chemwatch; 2. Classification drawn from HCIS; 3. Classification drawn from Regulation (EU) No 1272/2008 - Annex VI
Label elements	
Hazard pictogram(s)	
Signal word	Danger
Hazard statement(s)	
H332	Harmful if inhaled.
H350	May cause cancer.
Precautionary statement	(s) Prevention
P201	Obtain special instructions before use.
P271	Use only outdoors or in a well-ventilated area.

Precautionary statement(s) Response

P281

P261

P308+P313	IF exposed or concerned: Get medical advice/attention.
P312	Call a POISON CENTER or doctor/physician if you feel unwell.
P304+P340	IF INHALED: Remove victim to fresh air and keep at rest in a position comfortable for breathing.

Use personal protective equipment as required.

Avoid breathing dust/fumes.

Precautionary statement(s) Storage

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Precautionary statement(s) Disposal

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

SECTION 3 Composition / information on ingredients

Substances

See section below for composition of Mixtures

Mixtures

CAS No	%[weight]	Name
Not Available		low alloy steel welding electrode
Not Available		which in use generates:
Not Available	>60	welding fumes
Not Available		as
1309-37-1.		iron oxide fume
7439-96-5.		manganese fume
7440-02-0		nickel fume
69012-64-2		silica welding fumes
7440-47-3		chromium fume
7439-98-7		molybdenum fume

SECTION 4 First aid measures

Description of first aid measures

Eye Contact

	 DO NOT attempt to remove particles attached to or embedded in eye. Lay victim down, on stretcher if available and pad BOTH eyes, make sure dressing does not press on the injured eye by placing thick pads under dressing, above and below the eye. Seek urgent medical assistance, or transport to hospital. For "arc eye", i.e. welding flash or UV light burns to the eye: Place eye pads or light clean dressings over both eyes. Seek medical assistance. For THERMAL burns: Do NOT remove contact lens Lay victim down, on stretcher if available and pad BOTH eyes, make sure dressing does not press on the injured eye by placing thick pads under dressing, above and below the eye. Seek urgent medical assistance, or transport to hospital.
Skin Contact	 If skin or hair contact occurs: Flush skin and hair with running water (and soap if available). Seek medical attention in event of irritation. For thermal burms: Decontaminate area around burn. Consider the use of cold packs and topical antibiotics. For first-degree burns (affecting top layer of skin) Hold burned skin under cool (not cold) running water or immerse in cool water until pain subsides. Use compresses if running water is not available. Cover with sterile non-adhesive bandage or clean cloth. Do NOT apply butter or ointments; this may cause infection. Give over-the counter pain relievers if pain increases or swelling, redness, fever occur. For second-degree burns (affecting top two layers of skin) Cool the burn by immerse in cold running water for 10-15 minutes. Use compresses if running water is not available. Do NOT apply butter or ointments; this may cause infection. Brower these burns if the sterile, nonstick bandage and secure in place with gauze or tape. To prevent shock: (unless the person has a head, neck, or leg injury, or it would cause discomfort): Lay the person flat. Elevate feet about 12 inches. Elevate burns rea above heart level, if possible. Cover the person with coat or blanket. Seek medical assistance. For third-degree burns Seek medical assistance. For third-degree burns Seek medical assistance. In the mean time: Protect burn area acover loosely with sterile, nonstick bandage or, for large areas, a sheet or other material that will not leave lint in wound. Separate burned toes and fingers with dry, sterile dressings. Do not soak burn in water or apply ointments or butter; this may cause infection. To prevent shock see above. For an ainway burn, do not place pillow under the person's head wh
Inhalation	 If fumes or combustion products are inhaled remove from contaminated area. Lay patient down. Keep warm and rested. Prostheses such as false teeth, which may block airway, should be removed, where possible, prior to initiating first aid procedures. Apply artificial respiration if not breathing, preferably with a demand valve resuscitator, bag-valve mask device, or pocket mask as trained. Perform CPR if necessary. Transport to hospital, or doctor.
Ingestion	Not normally a hazard due to physical form of product.
ingeetion	

Indication of any immediate medical attention and special treatment needed

Copper, magnesium, aluminium, antimony, iron, manganese, nickel, zinc (and their compounds) in welding, brazing, galvanising or smelting operations all give rise to thermally produced particulates of smaller dimension than may be produced if the metals are divided mechanically. Where insufficient ventilation or respiratory protection is available these particulates may produce "metal fume fever" in workers from an acute or long term exposure.

- Onset occurs in 4-6 hours generally on the evening following exposure. Tolerance develops in workers but may be lost over the weekend. (Monday Morning Fever)
- Pulmonary function tests may indicate reduced lung volumes, small airway obstruction and decreased carbon monoxide diffusing capacity but these abnormalities resolve after several months.
- Although mildly elevated urinary levels of heavy metal may occur they do not correlate with clinical effects.
- The general approach to treatment is recognition of the disease, supportive care and prevention of exposure.
- Seriously symptomatic patients should receive chest x-rays, have arterial blood gases determined and be observed for the development of tracheobronchitis and pulmonary edema.

[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.]

For carbon monoxide intoxications:

- Administer pure oxygen by the best means possible. An oro-nasal mask is usually best. Artificial respiration is necessary wherever breathing is inadequate. Apnoeic patients have often been saved by persistent and efficient artificial ventilation. A patent airway must be carefully maintained. Patients with 40% carboxyhaemoglobin or more and an uncompensated metabolic acidosis (arterial pH less than 7.4) should be managed aggressively with ventilatory support/ hyperbaric oxygenation.
- Gastric aspiration and lavage early in the course of therapy may prevent aspiration pneumonitis and reveal the presence of ingested intoxicants.
- Avoid stimulant drugs including carbon dioxide. DO NOT inject methylene blue.
- Hypothermia has been employed to reduce the patient's oxygen requirement.
- Consider antibiotics as prophylaxis against pulmonary infection.
- A whole blood transfusion may be useful if it can be given early in the treatment program.
- Infuse sodium bicarbonate and balanced electrolyte solutions if blood analyses indicate a significant metabolic acidosis.
- Ancillary therapy for brain oedema may be necessary if hypoxia has been severe.
- Ensure absolute rest in bed for at least 48 hours; in severe poisonings, 2 to 4 weeks in bed may prevent sequelae.
- Watch for late neurological, psychiatric and cardiac complications. GOSSELIN, SMITH HODGE: Clinical Toxicology of Commercial Products 5th Ed. BIOLOGICAL EXPOSURE INDEX (BEI)

These represent the determinants observed in specimens collected from a healthy worker exposed at the Exposure Standard (ES or TLV):

Determinant	Sampling time	Index	Comments
Carboxyhaemoglobin in blood	end of shift	3.5% of haemoglobin	B, NS
Carbon monoxide in end-exhaled air	end of shift	20 ppm	B, NS
B: Background levels occur in specimens collected from subjects N	IOT exposed		

NS: Non-specific determinant; also observed after exposure to other material

SECTION 5 Firefighting 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	Welding electrodes should not be allowed to come into contact with strong acids or other substances which are corrosive to metals.
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Advice for firefighters

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, by any means available, 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. Equipment should be thoroughly decontaminated after use. Slight hazard when exposed to heat, flame and oxidisers.
Fire/Explosion Hazard	 Welding arc and metal sparks can ignite combustibles. Non combustible. Not considered to be a significant fire risk, however containers may burn. In a fire may decompose on heating and produce toxic / corrosive fumes.
HAZCHEM	Not Applicable

SECTION 6 Accidental release measures

Personal precautions, protective equipment and emergency procedures

Environmental precautions

See section 12

Methods and material for containment and cleaning up

Minor Spills	Clean up all spills immediately. Wear impervious gloves and safety glasses. Use dry clean up procedures and avoid generating dust. Place in suitable containers for disposal.
Major Spills	 Minor hazard. Clear area of personnel. Alert Fire Brigade and tell them location and nature of hazard. Control personal contact with the substance, by using protective equipment if risk of overexposure exists. Prevent, by any means available, spillage from entering drains or water courses. Contain spill/secure load if safe to do so. Bundle/collect recoverable product and label for recycling. Collect remaining product and place in appropriate containers for disposal. Clean up/sweep up area. Water may be required. If contamination of drains or waterways occurs, advise emergency services.

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. Prevent concentration in hollows and sumps. DO NOT enter confined spaces until atmosphere has been checked. DO NOT allow material to contact humans, exposed food or food utensils. Avoid contact with incompatible materials. When handling, DO NOT eat, drink or smoke. Keep containers securely sealed when not in use. Avoid physical damage to containers. Always wash hands with soap and water after handling. 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
Other information	maintained. Keep dry. Store under cover.
Other Information	 Protect containers against physical damage. Observe manufacturer's storage and handling recommendations contained within this SDS.

Conditions for safe storage, including any incompatibilities

Suitable container	 Packaging as recommended by manufacturer. Check that containers are clearly labelled
Storage incompatibility	Welding electrodes should not be allowed to come into contact with strong acids or other substances which are corrosive to metals.

SECTION 8 Exposure controls / personal protection

Control parameters

Occupational Exposure Limits (OEL)

INGREDIENT DATA

Source	Ingredient	Material name	TWA	STEL	Peak	Notes
Australia Exposure Standards	welding fumes	Welding fumes (not otherwise classified)	5 mg/m3	Not Available	Not Available	Not Available
Australia Exposure Standards	iron oxide fume	Iron oxide fume (Fe2O3) (as Fe)	5 mg/m3	Not Available	Not Available	Not Available

Source	Ingredient	Material name		TWA	STEL	Peak	Notes
Australia Exposure Standards	manganese fume	Manganese, fume (as Mn))	1 mg/m3	3 mg/m3	Not Available	Not Availabl
Australia Exposure Standards	nickel fume	Nickel, metal		1 mg/m3	Not Available	Not Available	Not Availabl
Australia Exposure Standards	nickel fume	Nickel, powder		1 mg/m3	Not Available	Not Available	Not Availabl
Australia Exposure Standards	chromium fume	Chromium (metal)		0.5 mg/m3	Not Available	Not Available	Not Availabl
Emergency Limits							
Emergency Limits Ingredient	Material name		TEEL-1	TE	EL-2	TEEL-3	
	Material name Iron oxide; (Ferric o	oxide)	TEEL-1 15 mg/m3		EL-2 0 mg/m3	TEEL-3 2,200 mg/r	m3
Ingredient		oxide)		36			
Ingredient iron oxide fume	Iron oxide; (Ferric o	oxide)	15 mg/m3	36 5 I	i0 mg/m3	2,200 mg/i	m3
Ingredient iron oxide fume manganese fume	Iron oxide; (Ferric of Manganese	·	15 mg/m3 3 mg/m3	36 5 1 50	:0 mg/m3 mg/m3	2,200 mg/r 1,800 mg/r	m3
Ingredient iron oxide fume manganese fume nickel fume	Iron oxide; (Ferric o Manganese Nickel	·	15 mg/m3 3 mg/m3 4.5 mg/m3	36 5 I 50 50	0 mg/m3 ng/m3 mg/m3	2,200 mg/r 1,800 mg/r 99 mg/m3	m3 m3

Ingredient	Original IDLH	Revised IDLH
welding fumes	Not Available	Not Available
iron oxide fume	2,500 mg/m3	Not Available
manganese fume	500 mg/m3	Not Available
nickel fume	10 mg/m3	Not Available
silica welding fumes	Not Available	Not Available
chromium fume	250 mg/m3	Not Available
molybdenum fume	Not Available	Not Available

Occupational Exposure Banding

Ingredient	Occupational Exposure Band Rating	Occupational Exposure Band Limit
molybdenum fume	E	≤ 0.01 mg/m³
Notes:	Occupational exposure banding is a process of assigning chemica potency and the adverse health outcomes associated with exposu band (OEB), which corresponds to a range of exposure concentra	ire. The output of this process is an occupational exposure

MATERIAL DATA

Exposure controls

Appropriate engineering controls	 Engineering controls are used to remove a hazard or place a barrier between the wool engineering controls can be highly effective in protecting workers and will typically be provide this high level of protection. The basic types of engineering controls are: Process controls which involve changing the way a job activity or process is done to a Enclosure and/or isolation of emission source which keeps a selected hazard "physic that strategically "adds" and "removes" air in the work environment. Ventilation can redesigned properly. The design of a ventilation system must match the particular proce Employers may need to use multiple types of controls to prevent employee overexposes. Special ventilation requirements apply for processes which result in the generation of zinc fume. For work conducted outdoors and in open work spaces, the use of mechanical (grequired as a minimum. (Open work spaces exceed 300 cubic meters per welder For indoor work, conducted in limited or confined work spaces, use of mechanical mandatory. (In confined spaces always check that oxygen has not been depleted corrosion of aluminium) Local exhaust systems must be designed to provide a minimum capture velocity at the metre/sec. Air contaminants generated in the workplace possess varying "escape" vertication effectively remove the contamination of the space work is a provide a minimum capture velocity at the metre/sec. Air contaminants generated in the workplace possess varying "escape" vertication vertications of the space of the contamination of the space of the spac	e independent of worker interactions to reduce the risk. ally" away from the worker and ventilation emove or dilute an air contaminant if ess and chemical or contaminant in use. sure. aluminium, copper, fluoride, manganese or eneral exhaust or plenum) ventilation is) I ventilation by local exhaust systems is by excessive rusting of steel or snowflake the fume source, away from the worker, of 0.5 elocities which, in turn, determine the nant.
	Type of Contaminant:	Air Speed:
	welding, brazing fumes (released at relatively low velocity into moderately still air)	0.5-1.0 m/s (100-200 f/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
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 fan, for example, should be a minimum of 1-2 m/s (200-400 f/min.) for extraction of welding or brazing fumes generated 2 meters distant from 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.

Articles or manufactured items, in their original condition, generally don't require engineering controls during handling or in normal use.

Exceptions may arise following extensive use and subsequent wear, during recycling or disposal operations where substances, found in the article, may be released to the environment.

Engineering controls are used to remove a hazard or place a barrier between the worker and the hazard. Well-designed engineering controls can be highly effective in protecting workers and will typically be independent of worker interactions to provide this high level of protection.

The basic types of engineering controls are:

Process controls which involve changing the way a job activity or process is done to reduce the risk.

Enclosure and/or isolation of emission source which keeps a selected hazard "physically" away from the worker and ventilation that strategically "adds" and "removes" air in the work environment. Ventilation can remove or dilute an air contaminant if designed properly. The design of a ventilation system must match the particular process and chemical or contaminant in use. Employers may need to use multiple types of controls to prevent employee overexposure.

Special ventilation requirements apply for processes which result in the generation of barium, chromium, lead, or nickel fume and in those processes which generate ozone.

The use of mechanical ventilation by local exhaust systems is required as a minimum in all circumstances (including outdoor work). (In confined spaces always check that oxygen has not been depleted by excessive rusting of steel or snowflake corrosion of aluminium)

Local exhaust systems must be designed to provide a minimum capture velocity at the fume source, away from the worker, of 0.5 metre/sec. Air contaminants generated in the workplace possess varying "escape" velocities which, in turn, determine the "capture velocities" of fresh circulating air required to effectively remove the contaminant.

Type of Contaminant:	Air Speed:
welding broking fumor (released at relatively law velocity into mederately still air)	0.5-1.0 m/s
welding, brazing fumes (released at relatively low velocity into moderately still air)	(100-200 f/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
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 fan, for example, should be a minimum of 1-2 m/s (200-400 f/min.) for extraction of welding or brazing fumes generated 2 meters distant from 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.

For manual arc welding operations the nature of ventilation is determined by the location of the work.

- ▶ For outdoor work, natural ventilation is generally sufficient.
- For indoor work, conducted in open spaces, use mechanical (general exhaust or plenum) ventilation. (Open work spaces exceed 300 cubic metres per welder)
- For work conducted in limited or confined spaces, mechanical ventilation, using local exhaust systems, is required. (In confined spaces always check that oxygen has not been depleted by excessive rusting of steel or snowflake corrosion of aluminium)

Mechanical or local exhaust ventilation may not be required where the process working time does not exceed 24 mins. (in an 8 hr. shift) provided the work is intermittent (a maximum of 5 mins. every hour). Local exhaust systems must be designed to provide a minimum capture velocity at the fume source, away from the worker, of 0.5 metre/sec. Air contaminants generated in the workplace possess varying "escape" velocities which, in turn, determine the "capture velocities" of fresh circulating air required to effectively remove the contaminant.

Type of Contaminant:

Air Speed:

welding, brazing fumes (released at relatively low velocity into moderately still air) 0.5-1.0 m/s (100-200 f/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
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 fan, for example, should be a minimum of 1-2.5 m/s (200-500 f/min.) for extraction of gases discharged 2 meters distant from 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.

Engineering controls are used to remove a hazard or place a barrier between the worker and the hazard. Well-designed engineering controls can be highly effective in protecting workers and will typically be independent of worker interactions to provide this high level of protection.

The basic types of engineering controls are:

Process controls which involve changing the way a job activity or process is done to reduce the risk.

Enclosure and/or isolation of emission source which keeps a selected hazard "physically" away from the worker and ventilation that strategically "adds" and "removes" air in the work environment. Ventilation can remove or dilute an air contaminant if designed properly. The design of a ventilation system must match the particular process and chemical or contaminant in use. Employers may need to use multiple types of controls to prevent employee overexposure.

Local exhaust ventilation usually required. If risk of overexposure exists, wear approved respirator. Correct fit is essential to obtain adequate protection. Supplied-air type respirator may be required in special circumstances. Correct fit is essential to ensure adequate protection.

An approved self contained breathing apparatus (SCBA) may be required in some situations.

Provide adequate ventilation in warehouse or closed storage area. Air contaminants generated in the workplace possess varying "escape" velocities which, in turn, determine the "capture velocities" of fresh circulating air required to effectively remove the contaminant.

Type of Contaminant:	Air Speed:
solvent, vapours, degreasing etc., evaporating from tank (in still air).	0.25-0.5 m/s (50-100 f/min.)
aerosols, fumes from pouring operations, intermittent container filling, low speed conveyer transfers, welding, spray drift, plating acid fumes, pickling (released at low velocity into zone of active generation)	0.5-1 m/s (100-200 f/min.)
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 f/min.)
grinding, abrasive blasting, tumbling, high speed wheel generated dusts (released at high initial velocity into zone of very high rapid air motion).	2.5-10 m/s (500-2000 f/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
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 fan, for example, should be a minimum of 1-2 m/s (200-400 f/min) for extraction of solvents generated in a tank 2 meters distant from 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.

Personal protection



Welding helmet with suitable filter. Welding hand shield with suitable filter.

Eye and face protection

Goggles or other suitable eye protection shall be used during all gas welding or oxygen cutting operations. Spectacles without side shields, with suitable filter lenses are permitted for use during gas welding operations on light work, for torch brazing or for inspection.

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	 For most open welding/brazing operations, goggles, even with appropriate filters, will not afford sufficient facial protection for operators. Where possible use welding helmets or handshields corresponding to EN 175, ANSI Z49:12005, AS 1336 and AS 1338 which provide the maximum possible facial protection from flying particles and fragments. [WRIA-WTIA Technical Note 7] An approved face shield or welding helmet can also have filters for optical radiation protection, and offer additional protection against debris and sparks. UV blocking protective spectacles with side shields or welding goggles are considered primary protection, with the face shield or welding helmet considered secondary protection. The optical filter in welding goggles, face mask or helmet must be a type which is suitable for the sort of work being done. A filter suitable for gas welding, for instance, should not be used for arc welding. Face masks which are self dimming are available for arc welding, MIG, TIG and plasma cutting, and allow better vision before the arc is struck and after it is extinguished. For submerged arc welding use a lens shade which gives just sufficient arc brightness to allow weld pool control.
Skin protection	See Hand protection below
Hands/feet protection	 Welding Gloves Safety footwear Welding gloves conforming to Standards such as EN 12477:2001, ANSI Z49.1, AS/NZS 2161:2008 produced from leather, rubber, treated cotton, or alumininised These gloves protect against mechanical risk caused by abrasion, blade cut, tear and puncture Other gloves which protect against thermal risks (heat and fire) might also be considered - these comply with different standards to those mentioned above. One pair of gloves may not be suitable for all processes. For example, gloves that are suitable for low current Gas Tungsten Arc Welding (GTAW) (thin and flexible) would not be proper for high-current Air Carbon Arc Cutting (CAC-A) (insulated, tough, and durable)
Body protection	See Other protection below
Other protection	Before starting; consider that protection should be provided for all personnel within 10 metres of any open arc welding operation. Welding sites must be adequately shielded with screens of non flammable materials. Screens should permit ventilation at floor and ceiling levels. Overalls

Respiratory protection

Welding of powder coated metal requires good general area ventilation, and ventilated mask as local heat causes minor coating decomposition releasing highly discomforting fume which may be harmful if exposure is regular.

Welding or flame cutting of metals with chromate pigmented primers or coatings may result in inhalation of highly toxic chromate fumes. Exposures may be significant in enclosed or poorly ventilated areas

Respiratory protection not normally required due to the physical form of the product.

SECTION 9 Physical and chemical properties

Information on basic physical and chemical properties

Appearance	Copper-coloured solid welding rod with no odour; insoluble in water.		
Physical state	Manufactured	Relative density (Water = 1)	~7.8
Odour	Not Available	Partition coefficient n-octanol / water	Not Available
Odour threshold	Not Available	Auto-ignition temperature (°C)	Not Applicable
pH (as supplied)	Not Applicable	Decomposition temperature	Not Available
Melting point / freezing point (°C)	~1500	Viscosity (cSt)	Not Applicable
Initial boiling point and boiling range (°C)	Not Applicable	Molecular weight (g/mol)	Not Applicable
Flash point (°C)	Not Applicable	Taste	Not Available
Evaporation rate	Not Applicable	Explosive properties	Not Available
Flammability	Not Applicable	Oxidising properties	Not Available
Upper Explosive Limit (%)	Not Applicable	Surface Tension (dyn/cm or mN/m)	Not Applicable
Lower Explosive Limit (%)	Not Applicable	Volatile Component (%vol)	Not Applicable
Vapour pressure (kPa)	Not Applicable	Gas group	Not Available

Chemwatch: 5256-76		Page 10 of 19	Issue Date: 01/11/2019
Version No: 3.1.1.1		TALARC NiCrMo	Print Date: 11/11/2020
Solubility in water Immiscib	ble	pH as a solution ((1%) Not Applicable

VOC g/L

Not Applicable

SECTION 10 Stability and reactivity

Not Available

Vapour density (Air = 1)

Reactivity	See section 7	
Chemical stability	Product is considered stable and 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

- -	
	Fumes evolved during welding operations may be irritating to the upper-respiratory tract and may be harmful if inhaled.
	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 include upper respiratory tract irritation accompanied by coughing and a dryness of the mucous membranes, lassitude and a generalised feeling of malaise. Mild to severe headache, nausea, occasional vomiting, fever or chills, exaggerated mental activity, profuse sweating, diarrhoea, excessive urination and prostration may also occur. Tolerance to the fumes develops rapidly, but is quickly lost. All symptoms usually subside within 24-36 hours following removal from exposure.
	Bronchial and alveolar exudate are apparent in animals exposed to molybdenum by inhalation. Molybdenum fume may produce
	bronchial irritation and moderate fatty changes in liver and kidney.
	Acute carbon monoxide exposure can mimic acute gastroenteritis or food poisoning with accompanying nausea and vomiting. Rapidly fatal cases of poisoning are characterised by congestion and hemorrhages in all organs. The extent of the tissue and organ damage is related to the duration of the post-hypoxic unconsciousness. Exposure to carbon monoxide can result in immediate effects and, depending on the severity of the exposure, delayed effects. These delayed effects may occur days to weeks after the initial exposure. Signs of brain or nerve injury may appear at any time within three weeks following an acute exposure. Characteristically, those patients manifesting delayed neuropathology are middle aged or older. Most of the neurological symptoms associated with carbon monoxide exposure can resolve within a year but memory deficits and gait
	disturbances may remain
Inhaled	Symptoms of poisoning resulting from carbon monoxide exposure include respiratory disorders, diarrhoea and shock. Carbon monoxide competes with oxygen for haemoglobin binding sites and has a 240-fold affinity for these sites compared to oxygen. In addition to oxygen deficiency further disability is produced by the formation of carboxymyoglobin (COHb) in muscles, to produce disturbances in muscle metabolism, particularly that of the heart.
Index	The tissues most affected by carbon monoxide are those which are most sensitive to oxygen deprivation such as the brain and the heart. The overt lesion in these tissues is mostly haemorrhage. The severe headache associated with exposure is believed to be caused by cerebral oedema and increased intracranial pressure resulting from excessive transudate leakage of fluids through the hypoxic capillaries.
	Carbon monoxide induced hypoxia in the cochlea and brain stem leads to central hearing loss and vestibular dysfunction
	(vertigo, nausea, vomiting) with the vestibular symptoms usually more prominent than the hearing loss
	At low levels carbon monoxide may cause poor concentration, memory and vision problems, vertigo, muscular weakness and loss of muscle coordination, rapid and stretorous breathing, intermittent heart beat, loss of sphincter control and rarely coma and death. At higher levels (200 ppm for 2-3 hours), it may cause headaches, fatigue and nausea. At very high levels (400 ppm) the symptoms intensify and will be life-threatening after three hours. Exposure to levels of 1200 ppm or greater are immediately dangerous to life. When carbon monoxide levels in air exceed 3% (30,000 ppm), death occurs almost at once. Carbon monoxide is not a cumulative poison since COHb is fully dissociable and once exposure has ceased, the hemoglobin will revert to oxyhemoglobin. The biological half life of carbon monoxide in the blood in sedentary adults is 2- 5 hours and the elimination becomes slower as the concentration decreases.
	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 include upper respiratory tract irritation accompanied by coughing and a dryness of the mucous membranes, lassitude and a generalised feeling of malaise. Mild to severe headache, nausea, occasional vomiting, fever or chills, exaggerated mental activity, profuse sweating, diarrhoea, excessive urination and prostration may also occur. Tolerance to the fumes develops rapidly, but is quickly lost. All symptoms usually subside within 24-36 hours following removal from exposure.
Ingestion	Not normally a hazard due to physical form of product.
	Skin contact does not normally present a hazard, though it is always possible that occasionally individuals may be found who
Skin Contact	react to substances usually regarded as inert. Ultraviolet radiation (UV) is generated by the electric arc in the welding process. Skin exposure to UV can result in severe burns, in many cases without prior warning.

	Exposure to infrared radiation (IR), produced by the electric arc and other flame cutting equipment may heat the skin surface and the tissues immediately below the surface. Except for this effect, which can progress to thermal burns in some situations, infrared radiation is not dangerous to welders. Most welders protect themselves from IR (and UV) with a welder's helmet (or glasses) and
	protective clothing.
Eye	Fumes from welding/brazing operations may be irritating to the eyes. Ultraviolet (UV) radiation can also damage the lens of the eye. Many arc welders are aware of the condition known as "arc-eye," a sensation of sand in the eyes. This condition is caused by excessive eye exposure to UV. Exposure to ultraviolet rays may also increase the skin effects of some industrial chemicals (coal tar and cresol compounds, for example). Exposure of the human eye to intense visible light can produce adaptation, pupillary reflex, and shading of the eyes. Such actions are protective mechanisms to prevent excessive light from being focused on the retina. In the arc welding process, eye exposure to intense visible light is prevented for the most part by the welder's helmet. However, some individuals have sustained
	retinal damage due to careless "viewing" of the arc. At no time should the arc be observed without eye protection.
Chronic	On the basis, primarily, of animal experiments, concern has been expressed that the material may produce carcinogenic or mutagenic diffectory assessment. Repeated or prolonged exposure may also damage the liver and may cause a decrease in the heart rate. Systemic poinoing may result from inhibition or chronic ingestion of manganese containing substances. Progressive and permanent disability can occur from chronic magnatese poisoining if it is not treated, but it is not total. Chronic exposure has been associated with two major effects; bronchilis/pneumonitis following inhibition cellular damage to the paralin northcalation appears to substain cellular damage to the paralin. Symptoms appear bolice any pathology is evident and may include a mask-like facial expression, spatial cellular, damage to the ganglion. Symptoms appear bolice any pathology is evident and may include a mask-like facial expression, spatial cellular, depending on exposure levids weakness, headache and sparse. Manganese psychosis following with certain definite features: ancountable laubyles, eucloris, impulsive acts, absentimidentes, smental confusion, aggressiveness and hallucinations. The final stage is characterised by speech difficulties, muscular thetic, spassic gait and other nervous system effects. Symptoms resemble those of Parkinson's disease. Rat studies include the gradual accumulation of brain maganese to produce lesions mimicking these found in Parkinsonism. If the disease is diagnoed whilst still in the early taskes and the patient is removed from exposure, he course may be reversed.

lesions were not present in the lungs. Lesions of the nose and larynx were similar to those observed in rats.

Molybdenum trioxide was not mutagenic in any of five strains of Salmonella typhimurium and did not induce sister chromatid

exchanges or chro S9 metabolic activ	prosomal aberrations in cultured Chinese hamster ovary cells in vitro. All tests were conducted with or withor ation enzymes
	as been described in experimental animals exposed sub-chronically to molybdenum trioxide.
	f molybdenum trioxide action in lung carcinogenicity is not known; the material is not mutagenic.
	sions of the nose and larynx of rats and in the nose, larynx and lungs of mice were apparently due to the
•	
-	more durable epithelium in response to chronic exposure. nt of Health and Human Services (1) concluded that there was equivocal evidence of carcinogenic activity in
•	based on a marginally significant positive trend of alveolar/ bronchiolar adenoma or carcinogenic activity in
	cinogenic activity in female F344/N rats; that there was some evidence of carcinogenic activity in male
	that there was evidence of carcinogenic activity in female B6C3F1 mice
	gy Program: Technical Report Series 462, April 1997
Principal route of	exposure is inhalation of welding fumes from electrodes and workpiece. Reaction products arising from
electrode core and	flux appear as welding fume depending on welding conditions, relative volatilities of metal oxides and any
coatings on the w	orkpiece. Studies of lung cancer among welders indicate that they may experience a 30-40% increased risk
compared to the g	eneral population. Since smoking and exposure to other cancer-causing agents, such as asbestos fibre, may
influence these re	sults, it is not clear whether welding, in fact, represents a significant lung cancer risk. Whilst mild steel weldin
represents little ris	k, the stainless steel welder, exposed to chromium and nickel fume, may be at risk and it is this factor which
may account for the	e overall increase in lung cancer incidence among welders. Cold isolated electrodes are relatively harmless.
Long-term (chroni	c) exposure to low levels of carbon monoxide may produce heart disease and damage to the nervous system
Exposure of pregr	ant animals to carbon monoxide may cause low birthweight, increased foetal mortality and nervous system
damage to the off	pring.
	is a common cause of fatal poisoning in industry and homes. Non fatal poisoning may result in permanent
-	amage. Carbon monoxide reduces the oxygen carrying capacity of the blood. Effects on the body are
	eversible as long as brain cell damage or heart failure has not occurred. Avoid prolonged exposure, even to
	ns. A well-established and probably causal relationship exists between maternal smoking (resulting in
	bin levels of 2-7% in the foetus) and low birth weight. There also appears to be a dose-related increase in
•	nd a retardation of mental ability in infants born to smoking mothers.
	wborn infant are considered to be very susceptible to CO exposure for several reasons:
•	obin has a greater affinity for CO than maternal hemoglobin.
	ices in uptake and elimination of CO, the fetal circulation is likely to have COHb levels higher (up to 2.5 times
	e maternal circulation.
	COHb in fetal blood is 3 times longer than that of maternal blood.
	s has a comparatively high rate of O2 consumption, and a lower O2 tension in the blood than adults, a
	O2 transport has the potential to produce a serious hypoxia.
	gas readily crosses the placenta and CO exposure during pregnancy can be teratogenic.
	low levels may initiate or enhance deleterious myocardial alterations in individuals with restricted coronary
	Ind decreased myocardial lactate production Linde
-	rated by industrial processes such as welding, give rise to a number of potential health problems. Particles
	ron (respirables) articles may cause lung deterioration. Particles of less than 1.5 micron can be trapped in th
lungs and, depend	lent on the nature of the particle, may give rise to further serious health consequences.
Exposure to fume	containing high concentrations of water-soluble chromium (VI) during the welding of stainless steels in
-	as been reported to result in chronic chrome intoxication, dermatitis and asthma. Certain insoluble chromium

Exposure to turne containing high concentrations of water-soluble chromium (VI) during the welding of stainless steels in confined spaces has been reported to result in chronic chrome intoxication, dermatitis and asthma. Certain insoluble chromium (VI) compounds have been named as carcinogens (by the ACGIH) in other work environments. Chromium may also appear in welding fumes as Cr2O3 or double oxides with iron. These chromium (III) compounds are generally biologically inert. Welding fume with high levels of ferrous materials may lead to particle deposition in the lungs (siderosis) after long exposure. This clears up when exposure stops. Chronic exposure to iron dusts may lead to eye disorders.

Silica and silicates in welding fumes are non-crystalline and believed to be non-harmful.

Other welding process exposures can arise from radiant energy UV flash burns, thermal burns or electric shock The welding arc emits ultraviolet radiation at wavelengths that have the potential to produce skin tumours in animals and in over-exposed individuals, however, no confirmatory studies of this effect in welders have been reported.

TALARC NiCrMo	ΤΟΧΙΟΙΤΥ	IRRITATION
	Not Available	Not Available
	тохісіту	IRRITATION
welding fumes	Not Available	Not Available
	тохісіту	IRRITATION
iron oxide fume	5500 mg/kg ^[2]	Not Available
	Oral (rat) LD50: >10000 mg/kg ^[2]	
	ΤΟΧΙΟΙΤΥ	IRRITATION
	2.3 mg/kg ^[2]	Eye (rabbit) 500mg/24H Mild
manganese fume	Oral (rat) LD50: >2000 mg/kg ^[1]	Eye: no adverse effect observed (not irritating) ^[1]
		Skin (rabbit) 500mg/24H Mild

WELDING FUMES

		Skin: no adverse effect observed (not irritating) ^[1]
	ΤΟΧΙCΙΤΥ	IRRITATION
	0.1 mg/kg ^[2]	Eye: no adverse effect observed (not irritating) ^[1]
nickel fume	500 mg/kg ^[2]	Skin: no adverse effect observed (not irritating) ^[1]
	Oral (rat) LD50: >9000 mg/kg ^[2]	
	Oral (rat) LD50: 5000 mg/kg ^[2]	
	TOXICITY	IRRITATION
silica welding fumes	Not Available	Eye: no adverse effect observed (not irritating) ^[1]
		Skin: no adverse effect observed (not irritating) ^[1]
	TOXICITY	IRRITATION
chromium fume	Not Available	Not Available
	TOXICITY	IRRITATION
	Oral (rat) LD50: >2000 mg/kg ^[1]	Not Available
molybdenum fume	Oral (rat) LD50: 2689 mg/kg ^[1]	
	Oral (rat) LD50: 4040 mg/kg ^[1]	
Legend:	1. Value obtained from Europe ECHA Registered Substances - Acute toxicity 2.* Value obtained from manufacturer's SDS	

Most welding is performed using electric arc processes - manual metal arc, metal inert gas (MIG) and tungsten inert gas welding (TIG) – and most welding is on mild steel.

In 2017, an IARC working group has determined that "sufficient evidence exists that welding fume is a human lung carcinogen (Group 1).

A complicating factor in classifying welding fumes is its complexity. Generally, welding fume is a mixture of metal fumes (i.e., iron, manganese, chromium, nickel, silicon, titanium) and gases (i.e., carbon monoxide, ozone, argon, carbon dioxide). Welding fume can contain varying concentrations of individual components that are classified as human carcinogens, including hexavalent chrome and nickel. However the presence of such metals and the intensity of exposure to each differ significantly according to a number of variables, including the type of welding technique used and the composition of the base metal and consumable. Nonetheless, IARC did not differentiate between these variables in its decision.

There has been considerable evidence over several decades regarding cancer risks in relation to welding activities. Several case-control studies reported excess risks of ocular melanoma in welders. This association may be due to the presence in some welding environments of fumes of thorium-232, which is used in tungsten welding rods

Different welding environments may present different and complex profiles of exposures. In one study to characterise welding fume aerosol nanoparticles in mild steel metal active gas welding showed a mass median diameter (MMMD) of 200-300 nm. A widespread consensus seems to have formed to the effect that some welding environments, notably in stainless steel welding, do carry risks of lung cancer. This widespread consensus is in part based on empirical evidence regarding risks among stainless steel welders and in part on the fact that stainless steel welding entails moderately high exposure to nickel and chromium VI compounds, which are recognised lung carcinogens. The corollary is that welding without the presence of nickel and chromium VI compounds, namely mild-steel welding, should not carry risk. But it appears that this line of reasoning in not supported by the accumulated body of epidemiologic evidence. While there remained some uncertainty about possible confounding by smoking and by asbestos, and some possible publication bias, the overwhelming evidence is that there has been an excess risk of lung cancer among welders as a whole in the order of 20%-40%. The most begrudging explanation is that there is an as-yet unexplained common reason for excess lung cancer risks that applies to all types of welders. It has been have proposed that iron fumes may play such a role, and some Finnish data appear to support this hypothesis, though not conclusively. This hypothesis would also imply that excess lung cancer risks among welders are not unique to welders, but rather may be shared among many

types of metal working occupations. Welders are exposed to a range of fumes and gases (evaporated metal, metal oxides, hydrocarbons, nanoparticles, ozone, oxides of nitrogen (NOx)) depending on the electrodes, filler wire and flux materials used in the process, but also physical exposures such as electric and magnetic fields (EMF) and ultraviolet (UV) radiation. Fume particles contain a wide variety of oxides and salts of metals and other compounds, which are produced mainly from electrodes, filler wire and flux materials. Fumes from the welding of stainless-steel and other alloys contain nickel compounds and chromium[VI] and [III]. Ozone is formed during most electric arc welding, and exposures can be high in comparison to the exposure limit, particularly during metal inert gas welding of aluminium. Oxides of nitrogen are found during manual metal arc welding and particularly during gas welding. Welders who weld painted mild steel can also be exposed to a range of organic compounds produced by pyrolysis.

In one study particle elemental composition was mainly iron and manganese. Ni and Cr exposures were very low in the vicinity of mild steel welders, but much higher in the background in the workshop where there presumably was some stainless steel welding.

Personal exposures to manganese ranged from 0.01-4.93 mg/m3 and to iron ranged from 0.04-16.29 mg/m3 in eight Canadian welding companies. Types of welding identified were mostly (90%) MIG mild steel, MIG stainless steel, and TIG aluminum. Carbon monoxide levels were less than 5.0 ppm (at source) and ozone levels varied from 0.4-0.6 ppm (at source).

	 Welders, especially in shipyards, may also be exposed to asbestos dust. Physical exposures such as electric and magnetic fields (EMF) and ultraviolet (UV) radiation are also common. In all, the in vivo studies suggest that different welding fumes cause varied responses in rat lungs in vivo , and the toxic effects typically correlate with the metal composition of the fumes and their ability to produce free radicals. In many studies both soluble and insoluble fractions of the stainless steel welding fumes were required to produce most types of effects, indicating that the responses are not dependent exclusively on the soluble metals Lung tumourigenicity of welding fumes was investigated in lung tumour susceptible (A/J) strain of mice. Male mice were exposed by pharyngeal aspiration four times (once every 3 days) to 85 ug of gas metal arc-mild steel (GMA-MS), GMA-SS, or manual metal arc-SS (MMA-SS) fume. At 48 weeks post-exposure, GMA-SS caused the greatest increase in tumour multiplicity and incidence, but did not differ from sham exposure. Tumour incidence in the GMA-SS group versus sham control was close to significance at 78 weeks post exposure. Histopathological analysis of the ungs of these mice showed the GMA-SS group having an increase in preneoplasia/tumour multiplicity and incidence compared to the GMA-MS and sham groups at 48 weeks. The increase in incidence in the GMA-SS exposed mice was significant compared to the GMA-MS group but not to the sham-exposed animals, and the difference in incidence between the GMA-SS and MMA-SS groups was of border-line significante (p = 0.06). At 78 week s post-exposure, no statistically significant differences A significantly higher frequency of micronuclei in peripheral blood lymphocytes (binucleated cell assay) and higher mean levels of both centromere-positive and centromere-negative micronuclei was observed in welders (n=27) who worked without protective device compared to controls (n=30). The rate of micronucleated cells
	Not available. Refer to individual constituents.
NICKEL FUME	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: the 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.
	WARNING: This substance has been classified by the IARC as Group 2B: Possibly Carcinogenic to Humans. Tenth Annual Report on Carcinogens: Substance anticipated to be Carcinogen [National Toxicology Program: U.S. Dep. of Health & Human Services 2002]
SILICA WELDING FUMES	For silica amorphous: Derived No Adverse Effects Level (NOAEL) in the range of 1000 mg/kg/d. In humans, synthetic amorphous silica (SAS) is essentially non-toxic by mouth, skin or eyes, and by inhalation. Epidemiology studies show little evidence of adverse health effects due to SAS. Repeated exposure (without personal protection) may cause mechanical irritation of the eye and drying/cracking of the skin. When experimental animals inhale synthetic amorphous silica (SAS) dust, it dissolves in the lung fluid and is rapidly eliminated. If swallowed, the vast majority of SAS is excreted in the faeces and there is little accumulation in the body. Following absorption across the gut, SAS is eliminated via urine without modification in animals and humans. SAS is not expected to be broken down (metabolised) in mammals. After ingestion, there is limited accumulation of SAS in body tissues and rapid elimination occurs. Intestinal absorption has not been calculated, but appears to be insignificant in animals and humans. SASs injected subcutaneously are subjected to rapid dissolution and removal. There is no indication of metabolism of SAS in animals or humans based on chemical structure and available data. In contrast to crystalline silica, SAS is soluble in physiological media and the soluble chemical species that are formed are eliminated via the urinary tract without modification. Both the mammalian and environmental toxicology of SASs are significantly influenced by the physical and chemical properties, particularly those of solubility and particle size. SAS has no acute intrinsic toxicity by inhalation. Adverse effects, including suffocation, that have been reported were caused by the presence of high numbers of respirable particles generated to meet the required test atmosphere. These results are not representative of exposure to commercial SASs and should not be used for human risk assessment. Though repeated exposure of the skin may cause dryness and cracking, SAS is not a skin or eye irritant, and it is not a sen
	Inhalation (rat), 13 weeks, Lowest Observed Effect Level (LOEL) =1.3 mg/m3 based on mild reversible effects in the lungs. Inhalation (rat), 90 days, LOEL = 1 mg/m3 based on reversible effects in the lungs and effects in the nasal cavity. For silane treated synthetic amorphous silica: Repeated dose toxicity: oral (rat), 28-d, diet, no significant treatment-related adverse effects at the doses tested.

	manufacture of SAS. Respiratory symptoms in S exposure, while serial pulmonary function value SAS. Reports indicate high/prolonged exposures to a experiments these effects were reversible. [PAT	s and chest radiographs are not a morphous silicas induced lung fib	dversely affected by long-term exposure to
CHROMIUM FUME	For chrome(III) and other valence states (excep For inhalation exposure, all trivalent and other of The mechanisms of chromium toxicity are very deal of uncertainty about how chromium exerts chromium toxicity than trivalent chromium toxicit chromium compounds and on the genotoxicity at consensus from various reviews and agencies is compounds is lacking. Epidemiological studies of production and use, and chrome plating) conclu associated with an increased risk of respiratory exposure studies to mixtures that were mainly eleather tanners, who were exposed to trivalent or carcinogenicity of trivalent or elemental chromium The lesser potency of trivalent chromium relative hexavalent chromium and its greater ability to et The general inability of trivalent chromium are n divalent, or trivalent chromium compounds can absorption is simply less efficient in comparison compounds exist as tetrahedral chromate anion are permeable across nonselective membranes though these channels, instead being absorbed absorbed than hexavalent chromium, workers e urine at the end of a workday. Absorbed chromi the foetus. Although there is ample in vivo evide gastrointestinal tract and can be reduced to the trivalent chromium is converted to hexavalent chromi the foetus. Although there is ample in vivo evide gastrointestinal tract and can be reduced to the trivalent chromium is converted to hexavalent chromi chromium molecule appears to be chromodulin, four chromic ions. Chromodulin may facilitate in metabolism. Inorganic trivalent chromium comp being converted into biologically active forms by Chromium can be a potent sensitiser in a small The most sensitive endpoint identified in animal	t hexavalent): hromium compounds are treated complex, and although many stud its toxic influence. Much more is k ty. There is an abundance of infor and mutagenicity of chromium com s that evidence of carcinogenicity of workers in a number of industri- de that while occupational expost system cancers (primarily bronch elemental and trivalent (ferrochrom chromium were consistently negat im and its compounds, the genoto e to hexavalent chromium is likely net cells. enter cells verse membranes and thus be ab cplanation for the overall absence not traverse membranes and reac to absorption of hexavalent chror s, resembling the forms of other r . Trivalent chromium forms octaine via passive diffusion and phagoc xposed to trivalent chromium is trivalent form by ascorbate and g momium in biological systems. In from the tissues. Although not full also referred to as (GTF). Chrom teractions of insulin with its recep- iounds, which do not appear to ha r humans and animals minority of humans, both from de studies of acute exposure to triva- to trivalent chromium is associate romium in the human body, its poi other than hexavalent exhibit a recep- ion the tast exavalent exhibit a recep- tion the the hexavalent exhibit a recep- tion the the human body, its poi other than hexavalent exhibit a recep-	ies on chromium are available, there is a great known about the mechanisms of hexavalent mation available on the carcinogenic potential of npounds in experimental systems. The of elemental, divalent, or trivalent chromium es (chromate production, chromate pigment ure to hexavalent chromium compounds is ogenic and nasal), results from occupational nium alloy worker) were inconclusive. Studies in rive. In addition to the lack of direct evidence of exic evidence is overwhelmingly negative. In related to the higher redox potential of esorbed or reach peripheral tissue in significant of systemic trivalent chromium toxicity. eadily either. This is not to say that elemental, h peripheral tissue, the mechanism of mium compounds. Hexavalent chromium natural anions like sulfate and phosphate which edral complexes which cannot easily enter ytosis. Although trivalent chromium is less well ave had detectable levels of chromium in the ut the body via the bloodstream, and can reach efficiently reduced to trivalent chromium in the lutathione in the lungs, there is no evidence that general, trivalent chromium compounds are y characterized, the biologically active trivalent todulin is an oligopeptide complex containing tor site, influencing protein, glucose, and lipid ve insulin-potentiating properties, are capable of rmal and inhalation exposures. Nent chromium appears to involve the ed with impaired lung function and lung damage. tential mechanism of action in cells, and elative lack of toxicity the toxicity of elemental
MOLYBDENUM FUME	Asthma-like symptoms may continue for months non-allergenic condition known as reactive airwa levels of highly irritating compound. Key criteria in a non-atopic individual, with abrupt onset of p exposure to the irritant. A reversible airflow patter hyperreactivity on methacholine challenge testin also been included in the criteria for diagnosis of disorder with rates related to the concentration of other hand, is a disorder that occurs as result of nature) and is completely reversible after exposi- production.	ays dysfunction syndrome (RADS for the diagnosis of RADS include ersistent asthma-like symptoms v ern, on spirometry, with the presen ing and the lack of minimal lympho of RADS. RADS (or asthma) follow of and duration of exposure to the exposure due to high concentration	 which can occur following exposure to high the absence of preceding respiratory disease, within minutes to hours of a documented nce of moderate to severe bronchial cytic inflammation, without eosinophilia, have ving an irritating inhalation is an infrequent irritating substance. Industrial bronchitis, on the ions of irritating substance (often particulate in
SILICA WELDING FUMES & CHROMIUM FUME	The substance is classified by IARC as Group 3 NOT classifiable as to its carcinogenicity to hum Evidence of carcinogenicity may be inadequate	nans.	
CHROMIUM FUME & MOLYBDENUM FUME	No significant acute toxicological data identified	in literature search.	
Acute Toxicity	✓	Carcinogenicity	✓
Skin Irritation/Corrosion	×	Reproductivity	×
Serious Eye Damage/Irritation	×	STOT - Single Exposure	×
Respiratory or Skin sensitisation	×	STOT - Repeated Exposure	×
Mutagenicity	×	Aspiration Hazard	×

Legend:

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

SECTION 12 Ecological information

_		
IOX	IC	utv.
107		

	Endpoint	Test Duration (hr)		Species		Value	Source
TALARC NiCrMo	Not Available	Not Available		Not Available		Not Available	Not Availabl
	Endpoint	Test Duration (hr)		Species		Value	Source
welding fumes	Not Available	Not Available		Not Available		Not Available	Not Availabl
	Endpoint	Test Duration (hr)		Species		Value	Sourc
	LC50	96		Fish		0.05mg/L	2
iron oxide fume	EC50	48		Crustacea		5.11mg/L	2
	EC50	72		Algae or other aquatic plants		18mg/L	2
	NOEC	504		Fish		0.52mg/L	2
	Endpoint	Test Duration (hr)		Species		Value	Sourc
	LC50	96		Fish		>3.6mg/L	2
	EC50	48		Crustacea		>1.6mg/L	2
manganese fume	EC50	72		Algae or other aquatic plants		2.8mg/L	2
	EC10	72		Algae or other aquatic plants		2.6mg/L	2
	NOEC	48		Crustacea		1.6mg/L	2
	Endpoint	Test Duration (hr)	Sp	pecies	Valu	e	Sourc
	LC50	96	Fi	sh	0.00	3-0.1mg/L	2
nickel fume	EC50	48	Cr	rustacea	0.00	1-0.576mg/L	2
	EC50	72	Al	gae or other aquatic plants	0.00	1-0.43mg/L	2
	NOEC	240	Cr	rustacea	>0.0	01-0.715mg/L	2
	Endpoint	Test Duration (hr)		Species		Value	Sourc
oilion welding fumoo	LC50	96		Fish		>100mg/L	2
silica welding fumes	EC50	72		Algae or other aquatic plants		4-200mg/L	2
	NOEL	72		Algae or other aquatic plants		10-mg/L	2
	Endpoint	Test Duration (hr)		Species		Value	Source
chromium fume	Not Available	Not Available		Not Available		Not Available	Not Availab
molybdenum fume	Endpoint	Test Duration (hr)		Species		Value	Sourc
	LC50	96		Fish		1-339mg/L	2
	EC50	48		Crustacea		1-472.6mg/L	2
	EC50	72		Algae or other aquatic plants		1-568.9mg/L	2
	NOEC	672		Crustacea		0.67mg/L	2
Legend:	3. EPIWIN St	uite V3.12 (QSAR) - Aquatic To:	xicity Data (Es	egistered Substances - Ecotoxic timated) 4. US EPA, Ecotox data n) - Bioconcentration Data 7. ME	base - Aqu	atic Toxicity Da	nta 5.

Vendor Data

DO NOT discharge into sewer or waterways.

Persistence and degradability

Ingredient

Persistence: Water/Soil

Persistence: Air

No Data available for all ingredients No Data available for all ingredients	Ingredient	Persistence: Water/Soil	Persistence: Air
		No Data available for all ingredients	No Data available for all ingredients

Bioaccumulative potential

Ingredient	Bioaccumulation
	No Data available for all ingredients
Mehilityin eeil	

Mobility in soil

Ingredient	Mobility
	No Data available for all ingredients

SECTION 13 Disposal considerations

Waste treatment methods	
Product / Packaging disposal	 Recycle wherever possible or consult manufacturer for recycling options. 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
HAZCHEM	Not Applicable

Land transport (ADG): 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

SECTION 15 Regulatory information

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

welding fumes is found on the following regulatory lists	
International Agency for Research on Cancer (IARC) - Agents Classified by the IARC Monographs	International Agency for Research on Cancer (IARC) - Agents Classified by the IARC Monographs - Group 1 : Carcinogenic to humans
iron oxide fume is found on the following regulatory lists	
Australia Standard for the Uniform Scheduling of Medicines and Poisons	Australian Inventory of Industrial Chemicals (AIIC)
(SUSMP) - Schedule 4	International Agency for Research on Cancer (IARC) - Agents Classified by
Australia Standard for the Uniform Scheduling of Medicines and Poisons (SUSMP) - Schedule 5	the IARC Monographs
Australia Standard for the Uniform Scheduling of Medicines and Poisons (SUSMP) - Schedule 6	
manganese fume is found on the following regulatory lists	
Australia Hazardous Chemical Information System (HCIS) - Hazardous	Australian Inventory of Industrial Chemicals (AIIC)
Chemicals	
1	

nickel fume is found on the following regulatory lists

Australia Hazardous Chemical Information System (HCIS) - Hazardous Chemicals	International Agency for Research on Cancer (IARC) - Agents Classified by the IARC Monographs
Chemicale	
Australian Inventory of Industrial Chemicals (AIIC)	International Agency for Research on Cancer (IARC) - Agents Classified by
Chemical Footprint Project - Chemicals of High Concern List	the IARC Monographs - Group 2B : Possibly carcinogenic to humans
silica welding fumes is found on the following regulatory lists	
Australia Hazardous Chemical Information System (HCIS) - Hazardous	Australian Inventory of Industrial Chemicals (AIIC)
Chemicals	
chromium fume is found on the following regulatory lists	
Australian Inventory of Industrial Chemicals (AIIC)	International Agency for Research on Cancer (IARC) - Agents Classified by
	the IARC Monographs

National Inventory Status

molybdenum fume is found on the following regulatory lists

Australian Inventory of Industrial Chemicals (AIIC)

National Inventory	Status
Australia - AIIC	Yes
Australia - Non-Industrial Use	No (iron oxide fume; manganese fume; nickel fume; silica welding fumes; chromium fume; molybdenum fume)
Canada - DSL	Yes
Canada - NDSL	No (iron oxide fume; manganese fume; nickel fume; silica welding fumes; chromium fume; molybdenum fume)
China - IECSC	Yes
Europe - EINEC / ELINCS / NLP	Yes
Japan - ENCS	No (manganese fume; nickel fume; chromium fume; molybdenum fume)
Korea - KECI	Yes
New Zealand - NZIoC	Yes
Philippines - PICCS	Yes
USA - TSCA	Yes
Taiwan - TCSI	Yes
Mexico - INSQ	No (silica welding fumes)
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	01/11/2019
Initial Date	01/08/2017

SDS Version Summary

Version	Issue Date	Sections Updated
2.1.1.1	01/08/2017	Spills (major)
3.1.1.1	01/11/2019	One-off system update. NOTE: This may or may not change the GHS classification

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_\circ
- IDLH: Immediately Dangerous to Life or Health Concentrations
- 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
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