

Version No: A-3.00 Safety data sheet according to REACH Regulation (EC) No 1907/2006, Directive 2020/878

Issue Date: 01/04/2022

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

Product name	EM-Tec C20 Carbon Conductive Micro-Tip Pen
Synonyms	
Other means of identification	15-001120

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

Relevant identified uses	Electrically conductive coating and EMI/RFI shield
Uses advised against	Not Applicable

1.3. Details of the supplier of the safety data sheet

Registered company name	Rave Scientific	
Address	100 Franklin Square Dr. Suite 101 Somerset, NJ 08873	
Telephone	1-732-898-3828	
Fax	Not Available	
Website	https://www.ravescientific.com/	
Email	sales@ravescientific.com	info@ravescientific.com

1.4. Emergency telephone number

Association / Organisation	National Emergency Telephone
Emergency telephone numbers	911
Other emergency telephone numbers	911

SECTION 2 Hazards identification

2.1. Classification of the substance or mixture

Classified according to EU Regulation Nr.1272/2008-VI [1]	
Legend:	1. Classified by according to EU Regulation NR 1272/2008-VI

2.2. Label elements

Hazard pictogram(s)	

Hazard	statement(s)
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H336	May cause drowsiness or dizziness.
H225	Highly flammable liquid and vapour.
H318	Causes serious eye damage.
H317	May cause an allergic skin reaction.
H351	Suspected of causing cancer.

EUH066	Repeated exposure may cause skin dryness or cracking.
Dressutionary statement(s) Dre	
Precautionary statement(s) Pre	vention
P201	Obtain special instructions before use.
P210	Keep away from heat, hot surfaces, sparks, open flames and other ignition sources. No smoking.
P271	Use only outdoors or in a well-ventilated area.
P280	Wear protective gloves, protective clothing, eye protection and face protection.
P240	Ground and bond container and receiving equipment.
P241	Use explosion-proof electrical/ventilating/lighting/intrinsically safe equipment.
P242	Use non-sparking tools.
P243	Take action to prevent static discharges.
P261	Avoid breathing mist/vapours/spray.

P272 Contaminated work clothing should not be allowed out of the workplace.

Precautionary statement(s) Response

P305+P351+P338	IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing.
P308+P313	IF exposed or concerned: Get medical advice/ attention.
P310	Immediately call a POISON CENTER/doctor/physician/first aider.
P370+P378	In case of fire: Use alcohol resistant foam or normal protein foam to extinguish.
P302+P352	IF ON SKIN: Wash with plenty of water.
P333+P313	If skin irritation or rash occurs: Get medical advice/attention.
P362+P364	Take off contaminated clothing and wash it before reuse.
P303+P361+P353	IF ON SKIN (or hair): Take off immediately all contaminated clothing. Rinse skin with water [or shower].
P304+P340	IF INHALED: Remove person to fresh air and keep comfortable for breathing.

Precautionary statement(s) Storage

P403+P235	Store in a well-ventilated place. Keep cool.
P405	Store locked up.

Precautionary statement(s) Disposal

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

2.3. Other hazards

Inhalation, skin contact and/or ingestion may produce health damage*.

Cumulative effects may result following exposure*.

May produce discomfort of the respiratory system and skin*.

HARMFUL: may cause lung damage if swallowed

SECTION 3 Composition / information on ingredients

3.1.Substances

See 'Composition on ingredients' in Section 3.2

3.2.Mixtures

1.CAS No 2.EC No 3.Index No 4.REACH No	%[weight]	Name	Classified according to EU Regulation Nr. 1272/2008/CLP Plus Amendments	SCL / M-Factor	Nanoform Particle Characteristics
1.67-64-1 2.200-662-2 3.606-001-00-8 4.Not Available	36	<u>acetone</u> * -	Flammable Liquids Category 2, Serious Eye Damage/Eye Irritation Category 2, Specific Target Organ Toxicity - Single Exposure (Narcotic Effects) Category 3; H225, H319, H336 [2]	Not Available	Not Available
1.110-19-0 2.203-745-1 429-360-0 3.607-026-00-7 4.Not Available	30	isobutyl acetate * -	Flammable Liquids Category 2; H225 ^[2]	Not Available	Not Available
1.71-36-3 2.200-751-6 3.603-004-00-6 4.Not Available	10	n-butanol	Flammable Liquids Category 3, Acute Toxicity (Oral) Category 4, Skin Corrosion/Irritation Category 2, Serious Eye Damage/Eye Irritation Category 1, Specific Target Organ Toxicity - Single Exposure (Respiratory Tract Irritation) Category 3, Specific Target Organ Toxicity - Single Exposure (Narcotic Effects) Category 3; H226, H302, H315, H318, H335, H336 ^[2]	Not Available	Not Available
1.1333-86-4 2.215-609-9 422-130-0 435-640-3 3.Not Available 4.Not Available	6	carbon black	Carcinogenicity Category 2; H351 ^[1]	Not Available	Not Available

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1.CAS No 2.EC No 3.Index No 4.REACH No		%[weight]	Name	Classified according to EU Regulation Nr. 1272/2008/CLP Plus Amendments	SCL / M-Factor	Nanoform Particle Characteristics
1.108-65-6 2.203-603-9 3.607-195-00-7 4.Not Available		4	propylene glycol monomethyl ether acetate, alpha-isomer *	Flammable Liquids Category 3; H226 [2]	Not Available	Not Available
1.25619-56-1 2.247-132-7 3.Not Available 4.Not Available		0.5	<u>barium dinonyl</u> naphthalenesulfonate	Acute Toxicity (Oral and Inhalation) Category 4, Skin Corrosion/Irritation Category 2, Serious Eye Damage/Eye Irritation Category 1, Hazardous to the Aquatic Environment Long-Term Hazard Category 2; H302+H332, H315, H318, H411 ^[1]	Not Available	Not Available
Le	egend:			n according to EU Regulation Nr.1272/2008-VI; 3. Classification o endocrine disrupting properties	Irawn from C&L	.; * EU IOELVs

SECTION 4 First aid measures

4.1. Description of first aid measures

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Eye Contact	 If this product comes in contact with the eyes: Immediately hold eyelids apart and flush the eye continuously with running water. Ensure complete irrigation of the eye by keeping eyelids apart and away from eye and moving the eyelids by occasionally lifting the upper and lower lids. Continue flushing until advised to stop by the Poisons Information Centre or a doctor, or for at least 15 minutes. Transport to hospital or doctor without delay. 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 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, without delay.
Ingestion	 Immediately give a glass of water. First aid is not generally required. If in doubt, contact a Poisons Information Centre or a doctor. If spontaneous vomiting appears imminent or occurs, hold patient's head down, lower than their hips to help avoid possible aspiration of vomitus.

4.2 Most important symptoms and effects, both acute and delayed

See Section 11

4.3. Indication of any immediate medical attention and special treatment needed

Any material aspirated during vomiting may produce lung injury. Therefore emesis should not be induced mechanically or pharmacologically. Mechanical means should be used if it is considered necessary to evacuate the stomach contents; these include gastric lavage after endotracheal intubation. If spontaneous vomiting has occurred after ingestion, the patient should be monitored for difficult breathing, as adverse effects of aspiration into the lungs may be delayed up to 48 hours. Treat symptomatically.

for simple esters:

BASIC TREATMENT

- _____
- Establish a patent airway with suction where necessary.
- Watch for signs of respiratory insufficiency and assist ventilation as necessary.
- Administer oxygen by non-rebreather mask at 10 to 15 l/min.
- Monitor and treat, where necessary, for pulmonary oedema .
- Monitor and treat, where necessary, for shock.
- DO NOT use emetics. Where ingestion is suspected rinse mouth and give up to 200 ml water (5 ml/kg recommended) for dilution where patient is able to swallow, has a strong gag reflex and does not drool.
- Give activated charcoal.

ADVANCED TREATMENT

- + Consider orotracheal or nasotracheal intubation for airway control in unconscious patient or where respiratory arrest has occurred.
- Positive-pressure ventilation using a bag-valve mask might be of use.
- Monitor and treat, where necessary, for arrhythmias.
- Start an IV D5W TKO. If signs of hypovolaemia are present use lactated Ringers solution. Fluid overload might create complications.
- Drug therapy should be considered for pulmonary oedema.
- + Hypotension with signs of hypovolaemia requires the cautious administration of fluids. Fluid overload might create complications.
- Treat seizures with diazepam.
- Proparacaine hydrochloride should be used to assist eye irrigation.

EMERGENCY DEPARTMENT

- Laboratory analysis of complete blood count, serum electrolytes, BUN, creatinine, glucose, urinalysis, baseline for serum aminotransferases (ALT and AST), calcium, phosphorus and magnesium, may assist in establishing a treatment regime. Other useful analyses include anion and osmolar gaps, arterial blood gases (ABGs), chest radiographs and electrocardiograph.
- Positive end-expiratory pressure (PEEP)-assisted ventilation may be required for acute parenchymal injury or adult respiratory distress syndrome.

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Consult a toxicologist as necessary.

BRONSTEIN, A.C. and CURRANCE, P.L. EMERGENCY CARE FOR HAZARDOUS MATERIALS EXPOSURE: 2nd Ed. 1994

To treat poisoning by the higher aliphatic alcohols (up to C7):

- Gastric lavage with copious amounts of water.
- It may be beneficial to instill 60 ml of mineral oil into the stomach.
- Oxygen and artificial respiration as needed.
- Electrolyte balance: it may be useful to start 500 ml. M/6 sodium bicarbonate intravenously but maintain a cautious and conservative attitude toward electrolyte replacement unless shock or severe acidosis threatens.
- ▶ To protect the liver, maintain carbohydrate intake by intravenous infusions of glucose.
- Haemodialysis if coma is deep and persistent. [GOSSELIN, SMITH HODGE: Clinical Toxicology of Commercial Products, Ed 5]

BASIC TREATMENT

Establish a patent airway with suction where necessary.

- Watch for signs of respiratory insufficiency and assist ventilation as necessary.
- Administer oxygen by non-rebreather mask at 10 to 15 l/min.
- Monitor and treat, where necessary, for shock.
- Monitor and treat, where necessary, for pulmonary oedema.
- Anticipate and treat, where necessary, for seizures.
- DO NOT use emetics. Where ingestion is suspected rinse mouth and give up to 200 ml water (5 ml/kg recommended) for dilution where patient is able to swallow, has a strong gag reflex and does not drool.
- Give activated charcoal.
- _____

ADVANCED TREATMENT

- Consider orotracheal or nasotracheal intubation for airway control in unconscious patient or where respiratory arrest has occurred.
- Positive-pressure ventilation using a bag-valve mask might be of use.
- Monitor and treat, where necessary, for arrhythmias.
- Start an IV D5W TKO. If signs of hypovolaemia are present use lactated Ringers solution. Fluid overload might create complications.
- If the patient is hypoglycaemic (decreased or loss of consciousness, tachycardia, pallor, dilated pupils, diaphoresis and/or dextrose strip or glucometer readings below 50 mg), give 50% dextrose.
- + Hypotension with signs of hypovolaemia requires the cautious administration of fluids. Fluid overload might create complications.
- Drug therapy should be considered for pulmonary oedema.
- Treat seizures with diazepam.
- Proparacaine hydrochloride should be used to assist eye irrigation.
- -----
- EMERGENCY DEPARTMENT
- Laboratory analysis of complete blood count, serum electrolytes, BUN, creatinine, glucose, urinalysis, baseline for serum aminotransferases (ALT and AST), calcium, phosphorus and magnesium, may assist in establishing a treatment regime. Other useful analyses include anion and osmolar gaps, arterial blood gases (ABGs), chest radiographs and electrocardiograph.
- Positive end-expiratory pressure (PEEP)-assisted ventilation may be required for acute parenchymal injury or adult respiratory distress syndrome.
- Acidosis may respond to hyperventilation and bicarbonate therapy.
 Haemodialysis might be considered in patients with severe intoxication.
- Consult a toxicologist as necessary. BRONSTEIN, A.C. and CURRANCE, P.L. EMERGENCY CARE FOR HAZARDOUS MATERIALS EXPOSURE: 2nd Ed. 1994

For C8 alcohols and above.

Symptomatic and supportive therapy is advised in managing patients.

For acute or short term repeated exposures to acetone:

- Symptoms of acetone exposure approximate ethanol intoxication.
- About 20% is expired by the lungs and the rest is metabolised. Alveolar air half-life is about 4 hours following two hour inhalation at levels near the Exposure Standard; in overdose, saturable metabolism and limited clearance, prolong the elimination half-life to 25-30 hours.
- There are no known antidotes and treatment should involve the usual methods of decontamination followed by supportive care.

[Ellenhorn and Barceloux: Medical Toxicology]

Management:

Measurement of serum and urine acetone concentrations may be useful to monitor the severity of ingestion or inhalation.

Inhalation Management:

- Maintain a clear airway, give humidified oxygen and ventilate if necessary.
- If respiratory irritation occurs, assess respiratory function and, if necessary, perform chest X-rays to check for chemical pneumonitis.
- Consider the use of steroids to reduce the inflammatory response.
- Treat pulmonary oedema with PEEP or CPAP ventilation.

Dermal Management:

- P Remove any remaining contaminated clothing, place in double sealed, clear bags, label and store in secure area away from patients and staff.
- Irrigate with copious amounts of water.
- An emollient may be required.
- Eye Management:

Irrigate thoroughly with running water or saline for 15 minutes.

- Stain with fluorescein and refer to an ophthalmologist if there is any uptake of the stain.
- Oral Management:

▶ No GASTRIC LAVAGE OR EMETIC

Encourage oral fluids.

Systemic Management:

- Monitor blood glucose and arterial pH.
- Ventilate if respiratory depression occurs.
- If patient unconscious, monitor renal function.
- Symptomatic and supportive care

The Chemical Incident Management Handbook:

Guy's and St. Thomas' Hospital Trust, 2000

BIOLOGICAL EXPOSURE INDEX

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

End of shift

Acetone in urine

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

Comments

NS

50 mg/L

5.1. Extinguishing media

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

5.2. Special hazards arising from the substrate or mixture

eine opeena nanaa ae antenig ne	
Fire Incompatibility	Avoid contamination with oxidising agents i.e. nitrates, oxidising acids, chlorine bleaches, pool chlorine etc. as ignition may result
5.3. Advice for firefighters	
Fire Fighting	
Fire/Explosion Hazard	 Liquid and vapour are highly flammable. Severe fire hazard when exposed to heat, flame and/or oxidisers. Vapour may travel a considerable distance to source of ignition. Heating may cause expansion or decomposition leading to violent rupture of containers. On combustion, may emit toxic fumes of carbon monoxide (CO). Combustion products include: carbon dioxide (CO2) metal oxides other pyrolysis products typical of burning organic material. Contains low boiling substance: Closed containers may rupture due to pressure buildup under fire conditions.

SECTION 6 Accidental release measures

6.1. Personal precautions, protective equipment and emergency procedures

See section 8

6.2. Environmental precautions

See section 12

6.3. Methods and material for containment and cleaning up

6.3. Methods and material for c	containment and cleaning up								
Minor Spills	 Remove all ignition sources. Clean up all spills immediately. Avoid breathing vapours and contact with skin and eyes. Control personal contact with the substance, by using protective equipment. Contain and absorb small quantities with vermiculite or other absorbent material. Wipe up. Collect residues in a flammable waste container. 								
	Chemical Class: ester and ethers For release onto land: recommende SORBENT TYPE RANK APPLICATIO				rder of priority.				
	LAND SPILL - SMALL								
	cross-linked polymer - particulate	1	shovel	shovel	R, W, SS				
	cross-linked polymer - pillow	1	throw	pitchfor	k R, DGC, RT				
	sorbent clay - particulate	2	shovel	shovel	R,I, P				
	wood fiber - particulate	3	shovel	shovel	R, W, P, DGC				
	wood fiber - pillow	3	throw	pitchfor	k R, P, DGC, RT				
	treated wood fiber - pillow	3	throw	pitchfor	k DGC, RT				
	LAND SPILL - MEDIUM								
Major Spills	cross-linked polymer - particulate	1	blower	skiploa	der R,W, SS				
	cross-linked polymer - pillow	2	throw	skiploa	der R, DGC, RT				
	sorbent clay - particulate	3	blower	skiploa	der R, I, P				
	polypropylene - particulate	3	blower	skiploa	der W, SS, DGC				
	expanded mineral - particulate	4	blower	skiploa	der R, I, W, P, DGC				
	wood fiber - particulate	4	blower	skiploa	der R, W, P, DGC				
	wood fiber - particulate 4 blower skiploader R, W, P, DGC Legend DGC: Not effective where ground cover is dense R; Not reusable I: l: Not incinerable P: Effectiveness reduced when rainy RT:Not effective where terrain is rugged SS: Not for use within environmentally sensitive sites W: Effectiveness reduced when windy Reference: Sorbents for Liquid Hazardous Substance Cleanup and Control; R.W Melvold et al: Pollution Technology Review No. 150: Noyes Data Corporation 1988								

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SORBENT TYPE RANK	APPLICATIO	ΟN	COLLE		IMITATIONS	
LAND SPILL - SMALL						
cross-linked polymer	- particulate	1	shovel	shovel	R, W, SS	
cross-linked polymer	- pillow	1	throw	pitchfork	R, DGC, RT	
sorbent clay - particul	late	2	shovel	shovel	R,I, P	
wood fiber - pillow		3	throw	pitchfork	R, P, DGC, RT	
treated wood fiber - p	illow	3	throw	pitchfork	DGC, RT	
foamed glass - pillow		4	throw	pichfork	R, P, DGC, RT	
polypropylene - partic sorbent clay - particul		2 2	blower blower	skiploade skiploade	er R, I, W, P, DG	
polypropylene - mat		3	throw	skiploade		
expanded mineral - p polyurethane - mat	articulate	3	blower	skipload		
egend DGC: Not effective whe R; Not reusable Not incinerable E: Effectiveness reduc RT:Not effective where SS: Not for use within W: Effectiveness reduc Reference: Sorbents fc	ed when rain terrain is rug environmenta ced when win	y Igeo ally dy	d sensitive	sites	anup and Control;	

6.4. Reference to other sections

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

SECTION 7 Handling and storage

7.1. Precaution	is for safe	e handling	

Safe handling	 Containers, even those that have been emptied, may contain explosive vapours. Do NOT cut, drill, grind, weld or perform similar operations on or near containers. Contains low boiling substance: Storage in sealed containers may result in pressure buildup causing violent rupture of containers not rated appropriately. Check for bulging containers. Vent periodically Always release caps or seals slowly to ensure slow dissipation of vapours 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. Avoid smoking, naked lights, heat or ignition sources. When handling, DO NOT eat, drink or smoke. Yapour may ignite on pumping or pouring due to static electricity. DO NOT use plastic buckets. Earth and secure metal containers when dispensing or pouring product. Use spark-free tools when handling. Avoid ontact with incompatible materials. Keep containers securely sealed. Avoid physical damage to containers. Work clothes should be laundered separately. 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. DO NOT allow clothing wet with material to stay in contact with skin
Fire and explosion protection	See section 5
Other information	 Store in original containers in approved flame-proof area. No smoking, naked lights, heat or ignition sources. DO NOT store in pits, depressions, basements or areas where vapours may be trapped. Keep containers securely sealed. Store away from incompatible materials in a cool, dry well ventilated area. Protect containers against physical damage and check regularly for leaks. Observe manufacturer's storage and handling recommendations contained within this SDS.

Continued...

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7.2. Conditions for safe storage, including any incompatibilities

Suitable container	 Packing as supplied by manufacturer. Plastic containers may only be used if approved for flammable liquid. Check that containers are clearly labelled and free from leaks. For low viscosity materials (i) : Drums and jerry cans must be of the non-removable head type. (ii) : Where a can is to be used as an inner package, the can must have a screwed enclosure. For materials with a viscosity of at least 2680 cSt. (23 deg. C) For manufactured product having a viscosity of at least 250 cSt. (23 deg. C) Manufactured product that requires stirring before use and having a viscosity of at least 20 cSt (25 deg. C): (i) Removable head packaging; (ii) Cans with friction closures and (iii) low pressure tubes and cartridges may be used. Where combination packages are used, and the inner packages are of glass, there must be sufficient inert cushioning material in contact with inner and outer packages In addition, where inner packagings are glass and contain liquids of packing group I there must be sufficient inert absorbent to absorb any spillage, unless the outer packaging is a close fitting moulded plastic box and the substances are not incompatible with the plastic.
Storage incompatibility	 Acetone: may react violently with chloroform, activated charcoal, aliphatic amines, bromine, bromine trifluoride, chlorotriazine, chromic(IV) acid, chromic(VI) acid, chromium trioxide, chromy chloride, hexachloromelamine, iodine heptalluoride, iodoform, liquid oxygen, nitrosyl chloride, nitrosyl perchlorate, percenter and explosive percivides in contact with strong oxidisers, fluorine, hydrogen percvide (90%), sodium perchlorate, 2-methyl-1.3-butadiene can increase the explosive sensitivity of nitromethane on contact flow or agitation may generate electrostatic charges due to low conductivity disolves or attacks most rubber, resins, and plastics (polyethylenes, polyester, vinyl ester, PVC, Neoprene, Viton) Alcohols are incompatible with strong acids, acid chlorides, acid anhydrides, oxidising and reducing agents. react with strong acids, strong causics, aliphatic amines, isocyanates, acetaldehyde, benzoyl percvide, chromic acid, chromium oxide, dialkytics, dichlorine oxide, ethylene oxide, hypochlorous acid, sloporoyl chloroccarbonate, lithium tetrahydroaluminate, nitrogen dioxide, pentafluoroguanidine, phosphorus halides, phosphorus pentasulfide, tangerine oil, triethylaluminium, triisobutylaluminum should not be heated above 49 deg. C. when in contact with aluminium equipment Esters react with acids to liberate heat along with alcohols and acids. Strong oxidising acids may cause a vigorous reaction with esters that is sufficiently exothermic to ignite the reaction products. <li< th=""></li<>

7.3. Specific end use(s)

See section 1.2

SECTION 8 Exposure controls / personal protection

8.1. Control parameters

Ingredient	DNELs Exposure Pattern Worker	PNECs Compartment
acetone	Dermal 186 mg/kg bw/day (Systemic, Chronic) Inhalation 1 210 mg/m ³ (Systemic, Chronic) Inhalation 2 420 mg/m ³ (Local, Acute) Dermal 62 mg/kg bw/day (Systemic, Chronic) * Inhalation 200 mg/m ³ (Systemic, Chronic) * Oral 62 mg/kg bw/day (Systemic, Chronic) *	 10.6 mg/L (Water (Fresh)) 1.06 mg/L (Water - Intermittent release) 21 mg/L (Water (Marine)) 30.4 mg/kg sediment dw (Sediment (Fresh Water)) 3.04 mg/kg sediment dw (Sediment (Marine)) 29.5 mg/kg soil dw (Soil) 100 mg/L (STP)
isobutyl acetate	Dermal 10 mg/kg bw/day (Systemic, Chronic) Inhalation 300 mg/m ³ (Systemic, Chronic) Inhalation 300 mg/m ³ (Local, Chronic) Dermal 10 mg/kg bw/day (Systemic, Acute) Inhalation 600 mg/m ³ (Systemic, Acute) Inhalation 600 mg/m ³ (Local, Acute) Dermal 5 mg/kg bw/day (Systemic, Chronic) * Inhalation 35.7 mg/m ³ (Systemic, Chronic) * Oral 5 mg/kg bw/day (Systemic, Chronic) * Inhalation 35.7 mg/m ³ (Local, Chronic) * Dermal 5 mg/kg bw/day (Systemic, Acute) * Inhalation 300 mg/m ³ (Systemic, Acute) * Inhalation 300 mg/m ³ (Local, Acute) *	0.17 mg/L (Water (Fresh)) 0.017 mg/L (Water - Intermittent release) 0.34 mg/L (Water (Marine)) 0.877 mg/kg sediment dw (Sediment (Fresh Water)) 0.088 mg/kg sediment dw (Sediment (Marine)) 0.075 mg/kg soil dw (Soil) 200 mg/L (STP)
n-butanol	Inhalation 310 mg/m ³ (Local, Chronic) Dermal 3.125 mg/kg bw/day (Systemic, Chronic) * Inhalation 55.357 mg/m ³ (Systemic, Chronic) * Oral 1.562 mg/kg bw/day (Systemic, Chronic) * Inhalation 155 mg/m ³ (Local, Chronic) *	0.082 mg/L (Water (Fresh)) 0.008 mg/L (Water - Intermittent release) 2.25 mg/L (Water (Marine)) 0.324 mg/kg sediment dw (Sediment (Fresh Water)) 0.032 mg/kg sediment dw (Sediment (Marine))

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Ingredient	DNELs Exposure Pattern Worker	PNECs Compartment		
		0.017 mg/kg soil dw (Soil) 2476 mg/L (STP)		
	Inhalation 1 mg/m3 (Systemic, Chronic)	1 mg/L (Water (Fresh))		
carbon black	Inhalation 0.5 mg/m ³ (Local, Chronic)	0.1 mg/L (Water - Intermittent release)		
	Inhalation 0.06 mg/m ³ (Systemic, Chronic) *	10 mg/L (Water (Marine))		
	Dermal 796 mg/kg bw/day (Systemic, Chronic)	0.635 mg/L (Water (Fresh))		
	Inhalation 275 mg/m ³ (Systemic, Chronic)	0.064 mg/L (Water - Intermittent release)		
and the state of t	Inhalation 550 mg/m ³ (Local, Acute)	6.35 mg/L (Water (Marine))		
propylene glycol monomethyl	Dermal 320 mg/kg bw/day (Systemic, Chronic) *	3.29 mg/kg sediment dw (Sediment (Fresh Water))		
ether acetate, alpha-isomer	Inhalation 33 mg/m ³ (Systemic, Chronic) *	0.329 mg/kg sediment dw (Sediment (Marine))		
	Oral 36 mg/kg bw/day (Systemic, Chronic) *	0.29 mg/kg soil dw (Soil)		
	Inhalation 33 mg/m³ (Local, Chronic) *	100 mg/L (STP)		

* Values for General Population

Occupational Exposure Limits (OEL)

INGREDIENT DATA

Source	Ingredient	Material name	TWA	STEL	Peak	Notes
EU Consolidated List of Indicative Occupational Exposure Limit Values (IOELVs)	acetone	Acetone	500 ppm / 1210 mg/m3	Not Available	Not Available	Not Available
UK Workplace Exposure Limits (WELs)	acetone	Acetone	500 ppm / 1210 mg/m3	3620 mg/m3 / 1500 ppm	Not Available	Not Available
EU Consolidated List of Indicative Occupational Exposure Limit Values (IOELVs)	isobutyl acetate	Isobutyl acetate	50 ppm / 241 mg/m3	723 mg/m3 / 150 ppm	Not Available	Not Available
UK Workplace Exposure Limits (WELs)	isobutyl acetate	Isobutyl acetate	150 ppm / 724 mg/m3	903 mg/m3 / 187 ppm	Not Available	Not Available
UK Workplace Exposure Limits (WELs)	n-butanol	Butan-1-ol	Not Available	154 mg/m3 / 50 ppm	Not Available	Sk
UK Workplace Exposure Limits (WELs)	carbon black	Carbon black	3.5 mg/m3	7 mg/m3	Not Available	Not Available
EU Consolidated List of Indicative Occupational Exposure Limit Values (IOELVs)	propylene glycol monomethyl ether acetate, alpha-isomer	1-Methoxypropyl- 2-acetate	50 ppm / 275 mg/m3	550 mg/m3 / 100 ppm	Not Available	Skin
UK Workplace Exposure Limits (WELs)	propylene glycol monomethyl ether acetate, alpha-isomer	1-Methoxypropyl acetate	50 ppm / 274 mg/m3	548 mg/m3 / 100 ppm	Not Available	Sk

Emergency Limits

Ingredient	TEEL-1	TEEL-2	TEEL-3
acetone	Not Available	Not Available	Not Available
isobutyl acetate	450 ppm	1300* ppm	7500** ppm
n-butanol	60 ppm	800 ppm	8000** ppm
carbon black	9 mg/m3	99 mg/m3	590 mg/m3
propylene glycol monomethyl ether acetate, alpha-isomer	Not Available	Not Available	Not Available

Ingredient	Original IDLH	Revised IDLH	
acetone	2,500 ppm	Not Available	
isobutyl acetate	1,300 ppm	Not Available	
n-butanol	1,400 ppm	Not Available	
carbon black	1,750 mg/m3	Not Available	
propylene glycol monomethyl ether acetate, alpha-isomer	Not Available	Not Available	
barium dinonyl naphthalenesulfonate	Not Available	Not Available	
Occupational Exposure Banding			
Ingredient	Occupational Exposure Band Rating	Occupational Exposure Band Limit	
barium dinonyl naphthalenesulfonate	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 a
	range of exposure concentrations that are expected to protect worker health.

I MATERIAL DATA

for isobutyl acetate:

Odour Threshold Value: 0.40-0.44 ppm (recognition) The TLV-TWA is identical with that of n-butyl acetate and is thought to minimise the potential for ocular and upper respiratory tract irritation.

Odour Threshold Value: 3.6 ppm (detection), 699 ppm (recognition)

Saturation vapour concentration: 237000 ppm @ 20 C

NOTE: Detector tubes measuring in excess of 40 ppm, are available.

Exposure at or below the recommended TLV-TWA is thought to protect the worker against mild irritation associated with brief exposures and the bioaccumulation, chronic irritation of the respiratory tract and headaches associated with long-term acetone exposures. The NIOSH REL-TWA is substantially lower and has taken into account slight irritation experienced by volunteer subjects at 300 ppm. Mild irritation to acclimatised workers begins at about 750 ppm - unacclimatised subjects will experience irritation at about 350-500 ppm but acclimatisation can occur rapidly. Disagreement between the peak bodies is based largely on the view by ACGIH that widespread use of acetone, without evidence of significant adverse health effects at higher concentrations, allows acceptance of a higher limit.

Half-life of acetone in blood is 3 hours which means that no adjustment for shift-length has to be made with reference to the standard 8 hour/day, 40 hours per week because body clearance occurs within any shift with low potential for accumulation.

A STEL has been established to prevent excursions of acetone vapours that could cause depression of the central nervous system.

Odour Safety Factor(OSF)

OSF=38 (ACETONE)

for propylene glycol monomethyl ether acetate (PGMEA)

Saturated vapour concentration: 4868 ppm at 20 C.

A two-week inhalation study found nasal effects to the nasal mucosa in animals at concentrations up to 3000 ppm. Differences in the teratogenic potential of the alpha (commercial grade) and beta isomers of PGMEA may be explained by the formation of different metabolites. The beta-isomer is thought to be oxidised to methoxypropionic acid, a homologue to methoxyacetic acid which is a known teratogen. The alpha- form is conjugated and excreted. PGMEA mixture (containing 2% to 5% beta isomer) is a mild skin and eye irritant, produces mild central nervous system effects in animals at 3000 ppm and produces mild CNS impairment and upper respiratory tract and eye irritation in humans at 1000 ppm. In rats exposed to 3000 ppm PGMEA produced slight foetotoxic effects (delayed sternabral ossification) - no effects on foetal development were seen in rabbits exposed at 3000 ppm.

For n-butanol:

Odour Threshold Value: 0.12-3.4 ppm (detection), 1.0-3.5 ppm (recognition)

NOTE: Detector tubes for n-butanol, measuring in excess of 5 ppm are commercially available.

Exposure at or below the TLV-TWA is thought to provide protection against hearing loss due to vestibular and auditory nerve damage in younger workers and to protect against the significant risk of headache and irritation.

25 ppm may produce mild irritation of the respiratory tract 50 ppm may produce headache and vertigo.

Higher concentrations may produce marked irritation, sore throat, coughing, nausea, shortness of breath, pulmonary injury and central nervous system depression characterised by headache, dizziness, dullness and drowsiness.

6000 ppm may produce giddiness, prostration, narcosis, ataxia, and death.

Odour Safety Factor (OSF)

OSF=60 (n-BUTANOL)

8.2. Exposure controls

	be highly effective in protecting workers and will typically be The basic types of engineering controls are: Process controls which involve changing the way a job acti Enclosure and/or isolation of emission source which keeps 'adds' and 'removes' air in the work environment. Ventilatio ventilation system must match the particular process and c Employers may need to use multiple types of controls to pr For flammable liquids and flammable gases, local exhaust equipment should be explosion-resistant.	a selected hazard 'physically' away from the worker and ventilation the n can remove or dilute an air contaminant if designed properly. The d hemical or contaminant in use. event employee overexposure. ventilation or a process enclosure ventilation system may be required ing 'escape' velocities which, in turn, determine the 'capture velocities	nat strategically esign of a I. Ventilation
	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 com plating acid fumes, pickling (released at low velocity into z	tainer filling, low speed conveyer transfers, welding, spray drift, cone of active generation)	0.5-1 m/s (100-200 f/min.)
	direct spray, spray painting in shallow booths, drum filling, into zone of rapid air motion)	conveyer loading, crusher dusts, gas discharge (active generation	1-2.5 m/s (200-500 f/min.)
controls	with the square of distance from the extraction point (in sim accordingly, after reference to distance from the contamina 1-2 m/s (200-400 f/min.) for extraction of solvents generate considerations, producing performance deficits within the effactors of 10 or more when extraction systems are installed Adequate ventilation is typically taken to be that white room or enclosure containing the dangerous substance. Ventilation for plant and machinery is normally consi- might potentially be present to no more than 25% of the LE additional safeguards are provided to prevent the formation emergency shutdown of the process might be used together and gas turbine enclosures.	Upper end of the range 1: Disturbing room air currents 2: Contaminants of high toxicity 3: High production, heavy use 4: Small hood-local control only nce away from the opening of a simple extraction pipe. Velocity gene ple cases). Therefore the air speed at the extraction point should be ting source. The air velocity at the extraction point should be d in a tank 2 meters distant from the extraction point. Other mechanic xtraction apparatus, make it essential that theoretical air velocities ar l or used. ch limits the average concentration to no more than 25% of the LEL w dered adequate if it limits the average concentration of any dangerou L. However, an increase up to a maximum 50% LEL can be acceptal of a hazardous explosive atmosphere. For example, gas detectors I er with maintaining or increasing the exhaust ventilation on solvent ev	adjusted, I be a minimum of cal e multiplied by within the building, s substance that ole where inked to aporating ovens maintenance in

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	space, the ventilation should ensure that the concentration of the dangerous substance does not exceed 10% of the LEL (irrespective of the provision of suitable breathing apparatus)
8.2.2. Personal protection	
Eye and face protection	 Safety glasses with side shields. Chemical goggles. Contact lenses may pose a special hazard; soft contact lenses may absorb and concentrate irritants. A written policy document, describing the wearing of lenses or restrictions on use, should be created for each workplace or task. This should include a review of lens absorption and adsorption for the class of chemicals in use and an account of injury experience. Medical and first-aid personnel should be trained in their removal and suitable equipment should be readily available. In the event of chemical exposure, begin eye irrigation immediately and remove contact lens as soon as practicable. Lens should be removed at the first signs of eye redness or irritation - lens should be removed 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	 Wear adent/ forover or safety gunboots, e.g. PVC. Wear adent/ forover or safety gunboots, e.g. Rubber NOTE: The material may produce skin sensitisation in predisposed individuals. Care must be taken, when removing gloves and other protective equipment, to avoid all possible skin contact. Contaminated leather items, such as shoes, bets and watch-bands should be removed and destroyed. For sets: Do NOT use natural rubber, butyl rubber, EPDM or polystyrene-containing materials. The selection of suitable gloves does not only depend on the material, but takes on further marks of quality which vary 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 selection of suitable gloves does not only depend on the material, but takes the glove material can not be calculated in advance and has therefore to be checked prior to the application. The avact break 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 hygiene is a key element of effective hand care. Gloves must only be worn on clean hands. After using gloves, hands should be washed and dired thoroughly. Application of a non-perfumed moisturiser is recommended. Suitability and durability of glove type is dependent on usage. Important factors in the selection of gloves include: - frequency and duration of contact, - deventry Select gloves tested to a relevant standard (e.g. Europe EN 374, US F739, AS/M25 2161.1 or national equivalent). - When nory brief contact is expected, a glove with a protection class of 5 or higher (breakthrough time greater tha
Body protection	See Other protection below
Other protection	 Overalls. PVC Apron. PVC protective suit may be required if exposure severe. Eyewash unit. Ensure there is ready access to a safety shower. Some plastic personal protective equipment (PPE) (e.g. gloves, aprons, overshoes) are not recommended as they may produce static electricity. For large scale or continuous use wear tight-weave non-static clothing (no metallic fasteners, cuffs or pockets). Non sparking safety or conductive footwear should be considered. Conductive footwear describes a boot or shoe with a sole made from a conductive compound chemically bound to the bottom components, for permanent control to electrically ground the foot an shall dissipate static electricity from the body to reduce the possibility of ignition of volatile compounds. Electrical resistance must range between 0 to 500,000 ohms. Conductive shoes should be stored in lockers close to their nom in which they are won. Personnel who have been issued conductive footwear should not wear them from their place of work to their homes and return.

Recommended material(s)

GLOVE SELECTION INDEX

Glove selection is based on a modified presentation of the: 'Forsberg Clothing Performance Index'. Type AX Filter of sufficient capacity. (AS/NZS 1716 & 1715, EN 143:2000 & 149:2001, ANSI Z88 or national equivalent)

Where the concentration of gas/particulates in the breathing zone, approaches or

Respiratory protection

The effect(s) of the following substance(s) are taken into account in the *computer-generated* selection: EM-Tec C20 Carbon Conductive Micro-Tip Pen

Material	CPI
PE/EVAL/PE	А
TEFLON	В
BUTYL	С
BUTYL/NEOPRENE	С
CPE	С
HYPALON	С
NATURAL RUBBER	С
NATURAL+NEOPRENE	С
NEOPRENE	С
NITRILE	С
NITRILE+PVC	С
PE	С
PVA	С
PVC	С
PVDC/PE/PVDC	С
SARANEX-23	С
SARANEX-23 2-PLY	С
VITON/NEOPRENE	С

* CPI - Chemwatch Performance Index

A: Best Selection

B: Satisfactory; may degrade after 4 hours continuous immersion

C: Poor to Dangerous Choice for other than short term immersion NOTE: As a series of factors will influence the actual performance of the glove, a final

selection must be based on detailed observation. -

* Where the glove is to be used on a short term, casual or infrequent basis, factors such as 'feel' or convenience (e.g. disposability), may dictate a choice of gloves which might otherwise be unsuitable following long-term or frequent use. A qualified practitioner should be consulted. exceeds the 'Exposure Standard' (or ES), respiratory protection is required. Degree of protection varies with both face-piece and Class of filter; the nature of protection varies with Type of filter.

Required Minimum Protection Factor	Half-Face Respirator	Full-Face Respirator	Powered Air Respirator
up to 10 x ES	AX-AUS	-	AX-PAPR-AUS / Class 1
up to 50 x ES	-	AX-AUS / Class 1	-
up to 100 x ES	-	AX-2	AX-PAPR-2 ^

^ - Full-face

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)

- Cartridge respirators should never be used for emergency ingress or in areas of unknown vapour concentrations or oxygen content.
- The wearer must be warned to leave the contaminated area immediately on detecting any odours through the respirator. The odour may indicate that the mask is not functioning properly, that the vapour concentration is too high, or that the mask is not properly fitted. Because of these limitations, only restricted use of cartridge respirators is considered appropriate.
- Cartridge performance is affected by humidity. Cartridges should be changed after 2 hr of continuous use unless it is determined that the humidity is less than 75%, in which case, cartridges can be used for 4 hr. Used cartridges should be discarded daily, regardless of the length of time used

Selection of the Class and Type of respirator will depend upon the level of breathing zone contaminant and the chemical nature of the contaminant. Protection Factors (defined as the ratio of contaminant outside and inside the mask) may also be important.

Required minimum protection factor	Maximum gas/vapour concentration present in air p.p.m. (by volume)	Half-face Respirator	Full-Face Respirator
up to 10	1000	AX-AUS / Class 1	-
up to 50	1000	-	AX-AUS / Class 1
up to 50	5000	Airline *	-
up to 100	5000	-	AX-2
up to 100	10000	-	AX-3
100+		-	Airline**

** - Continuous-flow or positive pressure demand.

 $\begin{array}{l} \mathsf{A}(\mathsf{AII}\ classes) = \mathsf{Organic}\ vapours, \mathsf{B}\ \mathsf{AUS}\ or\ \mathsf{B1} = \mathsf{Acid}\ gases, \ \mathsf{B2} = \mathsf{Acid}\ gas\ or\ hydrogen\ cyanide(\mathsf{HCN}), \ \mathsf{B3} = \mathsf{Acid}\ gas\ or\ hydrogen\ cyanide(\mathsf{HCN}), \ \mathsf{E} = \mathsf{Sulfur}\ dioxide(\mathsf{SO2}), \ \mathsf{G} = \\ \mathsf{Agricultural}\ chemicals, \ \mathsf{K} = \mathsf{Ammonia}(\mathsf{NH3}), \ \mathsf{Hg} = \mathsf{Mercury}, \ \mathsf{NO} = \mathsf{Oxides}\ of\ nitrogen, \ \mathsf{MB} \\ = \ \mathsf{Methyl}\ bromide, \ \mathsf{AX} = \mathsf{Low}\ boiling\ point\ organic\ compounds(below\ 65\ deg\ C) \\ \end{array}$

8.2.3. Environmental exposure controls

See section 12

SECTION 9 Physical and chemical properties

9.1. Information on basic physical and chemical properties

Appearance	Black		
Physical state	Liquid	Relative density (Water = 1)	0.89
Odour	Not Available	Partition coefficient n-octanol / water	Not Available
Odour threshold	Not Available	Auto-ignition temperature (°C)	465
pH (as supplied)	Not Available	Decomposition temperature	Not Available
Melting point / freezing point (°C)	Not Available	Viscosity (cSt)	128.090
Initial boiling point and boiling range (°C)	56	Molecular weight (g/mol)	Not Available
Flash point (°C)	-17	Taste	Not Available
Evaporation rate	Not Available	Explosive properties	Not Available
Flammability	HIGHLY FLAMMABLE.	Oxidising properties	Not Available

Upper Explosive Limit (%)	12	Surface Tension (dyn/cm or mN/m)	Not Available
Lower Explosive Limit (%)	2	Volatile Component (%vol)	Not Available
Vapour pressure (kPa)	Not Available	Gas group	Not Available
Solubility in water	Partly miscible	pH as a solution (Not Available%)	Not Available
Vapour density (Air = 1)	>2	VOC g/L	Not Available
Nanoform Solubility	Not Available	Nanoform Particle Characteristics	Not Available
Particle Size	Not Available		

9.2. Other information

Not Available

SECTION 10 Stability and reactivity

10.1.Reactivity	See section 7.2
10.2. Chemical stability	 Unstable in the presence of incompatible materials. Product is considered stable. Hazardous polymerisation will not occur.
10.3. Possibility of hazardous reactions	See section 7.2
10.4. Conditions to avoid	See section 7.2
10.5. Incompatible materials	See section 7.2
10.6. Hazardous decomposition products	See section 5.3

SECTION 11 Toxicological information

11.1. Information on toxicological effects

Inhaled	 Evidence shows, or practical experience predicts, that the material produces irritation of the respiratory system, in a substantial number of individuals, following inhaltation. In contrast to most organs, the lung is able to respond to a chemical insult by first removing or neutralising the irritant and then repairing the damage. The repair process, which initially evolved to protect mammalian lungs from foreign matter and antigens, may however, produce further lung damage resulting in the impairment of gas exchange, the primary function of the lungs. Respiratory tract irritation often results in an inflammatory response involving the recruitment and activation of many cell types, mainly derived from the vascular system. Inhalation of vapours may cause drowsiness and dizziness. This may be accompanied by narcosis, reduced alerthess, loss of reflexes, lack of coordination and vertigo. The main effects of simple aliphatic esters are narcosis and irritation and anaesthesia at higher concentrations. These effects become greater as the molecular weights and boiling points increase. Central nervous system depression, headche, drowsiness, dizziness, coma and neurobehavioral changes may also be symptomatic of overexposure. Respiratory tract involvement may produce mucous membrane irritation, dyspnea, and tachypnea, pharyngtis, bronchtis, pneumonitis and, in massive exposures, pulmonary oedema (which may be delayed). Gastrointestinal effects include nausea, vomiting, diarrhoee and abdominal cramps. Liver and kidney damage may result from massive exposures by mice to 6600 ppm produced signs of marked central nervous system (CNS) depression, including prostration after 2 hours, narcosis after 3 hours and sone deaths. Although n-butanol is odourous and generally possesses adequate warning properies, the offactor senses may become faigued. Exposure to aliphatic alcohols with more than 3 carbons may produce central nervous system effects such as headache, dixizness, drevinim. CNS depressi
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Systemic effects of acetone inhalation exposure include central nervous system depression, light-headedness, incoherent speech, ataxia, stupor, hypotension, tachycardia, metabolic acidosis, hyperglycaemia and ketosis. Rarely, convulsions and tubular necrosis may be evident. Other symptoms of exposure may include restlessness, headache, vomiting, low blood-pressure and rapid and irregular pulse, eye and throat irritation, weakness of the legs and dizziness. Inhalation of high concentrations may produce dryness of the mouth and throat, nausea, uncoordinated movement, loss of coordinated speech, drowsiness and, in severe cases, coma. Inhalation of acetone vapours over long periods causes irritation of the respiratory tract, coughing and headache. Rats exposed to 52200 ppm vapour for 1 hour showed clear signs of narcosis; fatalities occurred at 126600 ppm.
Effects on the nervous system characterise over-exposure to higher aliphatic alcohols. These include headache, muscle weakness, giddiness, ataxia, (loss of muscle coordination), confusion, delirium and coma. Gastrointestinal effects may include nausea, vomiting and diarrhoea. In the absence of effective treatment, respiratory arrest is the most common cause of death in animals acutely poisoned by the higher alcohols. Aspiration of liquid alcohols produces an especially toxic response as they are able to penetrate deeply in the lung where they are absorbed and may produce pulmonary injury. Those possessing lower viscosity elicit a greater response. The result is a high blood level and prompt death at doses otherwise tolerated by ingestion without aspiration. In general the secondary alcohols are less toxic than the corresponding primary isomers. As a general observation, alcohols are more powerful central nervous system depressants than their aliphatic analogues. In sequence of decreasing depressant potential, lertiary alcohols, increased. Within the homologous series of aliphatic alcohols, narcotic potency may increase even faster than lethality of the vater solubility is diminished and lipophilicity is increased. Within the homologous series of aliphatic alcohols, narcotic potency may increase even faster than lethality of a cortionue to increase with increasing chain length. Aliphatic alcohols series (greater than C7) but animal data establish that lethality (0.2 mi) of these behaves like a hydrocarbon solven in causing death from pulmonary oedema. Primary alcohols are metabolised to corresponding aldehydes and acids; a significant metabolic acidosis may occur. Secondary alcohols are converted to ketones, which are also certral nervous system depressants and which, in the case of the higher homologues persist. The beaterial nervous system depressants and which, in the case of the higher homologues persist. Swallowing of n-butanol may course system depressants and which, in the case of the higher homologues pe
 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. Repeated exposure may cause skin cracking, flaking or drying following normal handling and use. Most liquid alcohols appear to act as primary skin irritants in humans. Significant percutaneous absorption occurs in rabbits but not apparently in man. Open cuts, abraded or irritated skin should not be exposed to this material Entry into the blood-stream through, for example, cuts, abrasions, puncture wounds 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. Skin contact with the material may damage the health of the individual; systemic effects may result following absorption. The material may produce moderate skin irritation; limited evidence or practical experience suggests, that the material either: produces significant, but moderate, inflammation when applied to the healthy intact skin of animals (for up to four hours), such inflammation being present twenty-four hours or more after the end of the exposure period. Skin irritation may also be present after prolonged or repeated exposure; this may result in a form of contact dermatitis (nonallergic). The dermatitis is often characterised by skin redness (erythema) and swelling (oedema) which may progress to blistering (vesiculation), scaling and thickening of the epidermis.
When applied to the eye(s) of animals, the material produces severe ocular lesions which are present twenty-four hours or more after instillation. Workers exposed to 200 ppm n-butanol showed ocular symptoms including corneal inflammation, burning sensation, blurring of vision, lachrymation, and photophobia. 100 ppm produced no systemic effects and reports of irritation of the eyes was rare.
On the basis, primarily, of animal experiments, concern has been expressed that the material may produce carcinogenic or mutagenic effects; in respect of the available information, however, there presently exists inadequate data for making a satisfactory assessment. Long-term exposure to respiratory irritants may result in disease of the airways involving difficult breathing and related systemic problems. Practical experience shows that skin contact with the material is capable either of inducing a sensitisation reaction in a substantial number of individuals, and/or of producing a positive response in experimental animals. Substances that can cause occupational asthma (also known as asthmagens and respiratory sensitisers) can induce a state of specific airway hyper-responsiveness via an immunological, irritant or other mechanism. Once the airways have become hyper-responsive, further exposure to the substance, sometimes even to tiny quantities, may cause respiratory symptoms. These symptoms can range in severity from a runny nose to asthma. Not all workers who are exposed to a sensitiser will become hyper-responsive and it is impossible to identify in advance who are likely to become hyper-responsive. Substances than can cuase occupational asthma should be distinguished from substances which may trigger the symptoms of asthma in people with pre-existing air-way hyper-responsiveness. The latter substances are not classified as asthmagens or respiratory sensitisers Wherever it is reasonably practicable, exposure to substances that can cuase occupational asthma should be diverse from becoming hyper-responsive. Activities giving rise to short-term peak concentrations should receive particular attention when risk management is being considered. Health surveillance is appropriate for all employees exposed or liable to be exposed to a substance which may cause occupational asthma and there should be appropriate consultation with an occupational health professional over the degree of risk and level of surveillance. To

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Serious damage (clear functional disturbance or morphological change which may have toxicological significance) is likely to be caused by repeated or prolonged exposure. As a rule the material produces, or contains a substance which produces severe lesions. Such damage may become apparent following direct application in subchronic (90 day) toxicity studies or following sub-acute (28 day) or chronic (two-year) toxicity tests. Exposure to the material may cause concerns for human fertility, generally on the basis that results in animal studies provide sufficient evidence to cause a strong suspicion of impaired fertility in the absence of toxic effects. or evidence of impaired fertility occurring at around the same dose

levels as other toxic effects, but which are not a secondary non-specific consequence of other toxic effects. Prolonged or repeated skin contact may cause drying with cracking, irritation and possible dermatitis following.

Limited evidence suggests that repeated or long-term occupational exposure may produce cumulative health effects involving organs or biochemical systems.

Serious systemic effects from exposure to n-butanol in the form of auditory and vestibular nerve damage have been reported amongst workers in France and Mexico. Audiologic impairment was produced in workers exposed to 80 ppm n-butanol with unprotected noise exposure. Workers exposed over a 15 year period (1929-1944) exhibited severe vertigo and vertiges gravis. Workers exposed from 3-11 years without personal protective equipment from noise experienced greater hearing loss (hypoacusia) in direct relation to exposure time when compared to a control group exposed to industrial noise of 90-100 dB but with n-butanol exposure. Average hearing loss was not large but the workers had central frequencies of 21.98 dB (11.59 dB minimum and 32.30 dB maximum) with a mean widening of the break between 3000 and 4000 Hz of 42.22 dB. There was a tendency of the averages to decrease as the frequencies moved away from the central zone. Affected workers were aged from 20-39 years. [ACGIH Documentation of TLVs] Workers exposed to 700 ppm acetone for 3 hours/day for 7-15 years showed inflammation of the respiratory tract, stomach and duodenum,

Workers exposed to 700 ppm acetone for 3 hours/day for 7-15 years showed inflammation of the respiratory tract, stomach and duodenum, attacks of giddiness and loss of strength. Exposure to acetone may enhance liver toxicity of chlorinated solvents.

EM-Tec C20 Carbon	TOXICITY	IRF	RITATION	
Conductive Micro-Tip Pen	Not Available	Not	Available	
	ΤΟΧΙΟΙΤΥ	IRRITATION	1	
	Dermal (rabbit) LD50: 20000 mg/kg ^[2]		•): 500 ppm - irritant	
	Inhalation(Mouse) LC50; 44 mg/L4h ^[2]		: 20mg/24hr -moderate	
	Oral (Rat) LD50; 5800 mg/kg ^[2]		: 3.95 mg - SEVERE	
acetone			e effect observed (irritating) ^[1]	
			: 500 mg/24hr - mild	
			:395mg (open) - mild	
			verse effect observed (not irritating) ^[1]	
	ΤΟΧΙΟΙΤΥ		IRRITATION	
isobutyl acetate	Dermal (rabbit) LD50: >5000 mg/kg ^[1]		Skin(rabbit): 500 mg open mild	
	Inhalation(Rat) LC50; >23.4 mg/l4h ^[1]			
	Oral (Rabbit) LD50; 4763 mg/kg ^[2]			
	ΤΟΧΙΟΙΤΥ	IRRITATION		
	Dermal (rabbit) LD50: 3400 mg/kg ^[2]	Eye (human): 50) ppm - irritant	
	Inhalation(Rat) LC50; 8000 ppm4h ^[2]	Eye (rabbit): 1.6		
n-butanol	Oral (Rat) LD50; 790 mg/kg ^[2]	Eye (rabbit): 24	mg/24h-SEVERE	
		Eye: adverse ef	fect observed (irreversible damage) ^[1]	
			5 mg/24h-moderate	
		Skin: adverse e	ifect observed (irritating) ^[1]	
	ΤΟΧΙΟΙΤΥ	IRRITATION		
carbon black	Dermal (rabbit) LD50: >3000 mg/kg ^[2]		erse effect observed (not irritating) ^[1]	
carbon black	Oral (Rat) LD50; >8000 mg/kg ^[1]		erse effect observed (not irritating) ^[1]	
propylene glycol monomethyl		IRRITATION	1	
ether acetate, alpha-isomer	dermal (rat) LD50: >2000 mg/kg ^[1]		e effect observed (not irritating) ^[1]	
	Oral (Rat) LD50; 3739 mg/kg ^[2]	Skin: no advers	e effect observed (not irritating) ^[1]	
	TOXICITY		IRRITATION	
barium dinonyl	Dermal (rabbit) LD50: >2000 mg/kg ^[2]		Eye (rabbit): 250 mg/5d mild	
naphthalenesulfonate	Inhalation(Rat) LC50; >5.25 mg/L4h ^[2]			
	Oral (Rat) LD50; 3000 mg/kg ^[2]			

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EM-Tec C20 Carbon Conductive Micro-Tip Pen	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. Generally,linear and branched-chain alkyl esters are hydrolysed to their component alcohols and carboxylic acids in the intestinal tract, blood and most tissues throughout the body. Following hydrolysis the component alcohols and carboxylic acids are metabolized Oral acute toxicity studies have been reported for 51 of the 67 esters of aliphatic acyclic primary alcohols and aliphatic linear saturated carboxylic acids. The very low oral acute toxicity of this group of esters is demonstrated by oral LD50 values greater than 1850 mg/kg bw Genotoxicity studies have been performed in vitro using the following esters of aliphatic acyclic primary alcohols and aliphatic linear saturated carboxylic acids: methyl acetate, butyl acetate, butyl aterate and the structurally related isoamyl formate and demonstrates that these substances are not genotoxic. The JEFCA Committee concluded that the substances in this group would not present safety concerns at the current levels of intake the esters of aliphatic acyclic primary alcohols and aliphatic linear saturated cardoxylic acids are generally used
ACETONE	The material may cause skin irritation after prolonged or repeated exposure and may produce a contact dermatitis (nonallergic). This form of dermatitis is often characterised by skin redness (erythema) and swelling epidermis. Histologically there may be intercellular oedema of the spongy layer (spongiosis) and intracellular oedema of the epidermis.
ISOBUTYL ACETATE	The material may produce moderate eye irritation leading to inflammation. Repeated or prolonged exposure to irritants may produce conjunctivitis. Inhalation (rat): 8000ppm/4h Skin(rabbit): 500 mg/24hr moderate
N-BUTANOL	The material may produce severe irritation to the eye causing pronounced inflammation. Repeated or prolonged exposure to irritants may produce conjunctivitis. for n-butanol Acute toxicity: n-Butanol (BA) was only slightly toxic to experimental animals following acute oral, dermal, or inhalation exposure. The acute oral LD50 values for female rats ranged from 790 to 4360 mg/kg. Different strains of rat were used in each of four studies, which may account for the variability. Oral LD50 values for mice, rabbits, hamsters, dogs, and male rats all fell within the same range. The rat inhalation LC0 of 8000 ppm (24000 mg/m3) indicates very low inhalation toxicity (no lethality at 8000 ppm). The rabbit dermal LD50 was 3402 mg/kg, indicating that BA can penetrate the skin, but not very readily. Animal experiments and human experience indicate that BA is, at most, moderately irritating to the skin, but it is a severe eye irritant. These effects are most likely due to BA s localised defatting and drying characteristics. Although no animal data are available, human studies and experience show that BA is not likely to be a skin sensitiser. The median dort threshold for BA (0.17 ppm) is well below the lowest nasal irritation threshold in humans (289 ppm), allowing warning of possible chemical exposure prior to nasal irritation courring. Human studies are complicated by the odor characteristics of the material, as the odor threshold is well below the levels at which irritation is observed. Repeat dose toxicity: An in vivo toxicokinetics study confirmed the rapid metabolism of n-butyl acetate (BAc) to BA. Hydrolysis of BAc in blood and brain was estimated to be 99 percent complete within 2.7 minutes (elimination to: exposure to BAc, the metabolic precursor of BA, produced transient hypoactivity (during exposure only) at 1500 and 3000 ppm (7185 and 14370 mg/m3) along with decreased body weight and food consumption, but no post exposure neurotoxicity even at 3000 ppm. A concurrent subchronic neurotoxicity study under the
CARBON BLACK	Inhalation (rat) TCLo: 50 mg/m3/6h/90D-I Nil reported WARNING: This substance has been classified by the IARC as Group 2B: Possibly Carcinogenic to Humans.
PROPYLENE GLYCOL MONOMETHYL ETHER ACETATE, ALPHA-ISOMER	A BASF report (in ECETOC) showed that inhalation exposure to 545 ppm PGMEA (beta isomer) was associated with a teratogenic response in rabbits; but exposure to 145 ppm and 36 ppm had no adverse effects. The beta isomer of PGMEA comprises only 10% of the commercial material, the remaining 90% is alpha isomer. Hazard appears low but emphasizes the need for care in handling this chemical. [I.C.I] *Shin-Etsu SDS for propylene glycol ethers include propylene glycol n-butyl ether (PnB); dipropylene glycol n-butyl ether (DPnB); dipropylene glycol methyl ether acetate (DPMA); tripropylene glycol methyl ether (TPM). Testing of a wide variety of propylene glycol ethers are less toxic than some ethers of the ethylene series. The common toxicities associated with the lower molecular weight homologues of the ethylene series, such as adverse effects on reproductive organs, the developing embryo and fetus, blood (haemolytic effects), or thymus, are not seen with the commercial-grade propylene glycol ethers. In the ethylene series, metabolism of the terminal hydroxyl group produces an alkoxyacetic acid. The reproductive and developmental toxicities of the lower molecular weight homologues in the ethylene series are on associated with the reproductive toxicity but can cause haemolysis in sensitive species, also through formation of an alkoxyacetic acid. The predominant alpha isomer of all the PGEs (thermodynamically favored during manufacture of PGEs) is a secondary alcohol incapable of forming an alkoxypropionic acid. In contrast beta-isomers are able to form the alkoxypropionic acids and these are linked to teratogenic effects (and possibly haemolytic effects). This alpha isomer cannot form an alkoxypropionic acid, this is the most likely reason for the lack of toxicity shown by the PGEs as distinct from the lower molecular weight ethylene glycol ethers. More importantly, however, very extensive empirical test data show that this class of commercial-grade glycol ethers. More importantly, however, very extensive empirical tes

	alcohol group), show a very similar pattern of low to non-detectable toxicity of any type at doses or exposure levels greatly exceeding those showing pronounced effects from the ethylene series. One of the primary metabolites of the propylene glycol ethers is propylene glycol, which is of low toxicity and completely metabolised in the body. As a class, the propylene glycol ethers are rapidly absorbed and distribution is rapid. Most excretion for PGEs is via the urine and expired air. A small portion is excreted in the faces. As a group PGEs exhibits low acute toxicity by the oral, darmal, and inhalation routes. Rat oral LDS0s range from >3,000 mg/kg (PHB) to >5,000 mg/kg (PHB). Do >5,000 mg/kg (PHB), Do >5,000 mg/kg (PHB), Do >5,000 mg/kg (PHB), Do >5,000 mg/kg (PHB), Do >5,010 mg/kg (PHB), DO
BARIUM DINONYL NAPHTHALENESULFONATE	 Tokichy information for barium sulfonates (barium salts of various alkyl and aryl sulfonic acids in oil solution): For dinonylnaphthalenes: The chemicals exhibit a very low order of toxicity to rats or rabbits by the oral, inhalation, or dermal routes. Human sensitisation study results are available for two members of the category (dinonylnaphthalene sulfonic acid, calcium salt; dinonylnaphthalene sulfonic acid, barium salt appears to be the most biologically active member of the category. For altaryl sulfonate petroleum additives: Mammalian Toxicology - Acute. Existing data on acute mammalian toxicity indicates a low concern for acute toxicity. Acute oral toxicity: In all but one studies, there were no deaths that could be attributed to treatment with the test material when administered at the limit dose of 2000 or 5000 mg/kg. In some studies, the primary clinical observations were darhea and reduced food consumption (without a change in body weight). These effects are consistent with the gastrointestinal irritant properties of detergents in an oil-based vehicle. In other studies, decreased body weight gain or ruffled fur was observed. In one study where deaths occurred, animals were administered dose levels well above the 2000 mg/kg limit dose. Overall, the acute oral LD50 for these substances was greater than the 2000 mg/kg limit dose indicating a relatively low order of toxicity. Acute dermal toxicity: No mortality was observed for any tested substance were agrease and reduced food consumption (withing included dry, flaky skin, desquamation and hyperkeratosis. Overall, the acute dermal application. In some cases, the cutaneous findings included dry, flaky skin, desquamation and hyperkeratosis. Overall, the acute dermal application. In some cases, the cutaneous findings included dry, flaky skin, desquamation and hyperkeratosis. Overall, the acute dermal coccurred to was greater than the 2000 mg/kg limit do

For dinonyInaphthalenesulfonic acid (DNNSA) and its salts: In general, a compound needs to be dissolved before it can be taken up from the gastro-intestinal tract after oral administration. Calcium bis(di

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C8-C10, branched, C9 rich, alkylnaphthalene sulphonate) (CaDNNSA) has a measured water solubility of 0.266 mg/L and therefore it is expected to dissolve into the gastrointestinal fluids to a very limited extent. Uptake by passive diffusion is possible, but limited due to the high molecular weight of the salt (average MW 959) and its dissociation product DNNSA (MW 461). CaDNNSA has a high log Pow 6.6), which makes the compound relatively hydrophobic. This characteristic will enable micellular solubilisation by bile salts in the gastro-intestinal tract which allows some crossing of lipid biomembranes. The structure contains an ionizable group (SO3H), which might hamper diffusion across biological membranes. In addition, the molecular size of the molecule of 19 Å does not favor uptake across the biological membranes. In the 90-day study on CaDNNSA in the highest dose group 6/10 females died showing alterations in the gastro-intestinal tract, a small thymus and bone marrow atrophy. The surviving females at 1000 mg/kg bw showed similar effects and a reduced body weight (gain). The effects on the gastro-intestinal tract also became apparent in males at 300 and 1000 mg/kg bw. These animals also had a reduced body weight (gain). Other effects included changes in numbers of white blood cells, lymphocytes, platelets as well as effects on several biochemical parameters. Macroscopy and histopathology indicated that next to the GI-tract mainly the thymus and bone marrow could be considered as potentially affected in males at 300 mg/kg bw and above and in females at 1000 mg/kg bw. The effects on blood and blood forming organs as well as on the immune system are indicative for some absorption of the substance. This absorption may be enhanced due to the effects on the gastro intestinal fraction intestinal fraction in the substance integritient of the substance.

The metabolism of DNNSA salts is mainly contingent on both the nature of the alkyl groups and the nature and extent of naphthalene ring substitutions. There are currently no metabolism studies of CaDNNSA, however, the US EPA has evaluated the metabolism of analogs in in the sodium alkyl naphthalenesulfonate cluster (SANS), a group of sodium salts of naphthalenesulfonic acids . In a US EPA final rule for SANS, it was stated that "the 1- or 2-sulfonic acid sodium salt moieties on the naphthalene ring may provide a handle by which these compounds can be readily conjugated and eliminated." Though the available information is not definitive for CaDNNSA, where the alkyl chains are much larger than for the naphthalenesulfonic acids evaluated by EPA, it is expected that the metabolism of the substance will be a factor, enhancing elimination. If absorbed, wide distribution of the CaDNNSA throughout the body is not expected based on its molecular size (18 Å). In general, molecules of this size do not pass readily through cell membranes, thus limiting wide distribution. Excretion of CaDNNSA and its potential metabolites will occur via the bile (high molecular weight) or the urine (low molecular weight). Irritation:

Calcium bis(di-C8-10, branched, C9 rich, alkylnaphthalenesulphonate) is irritating to skin and eyes. It is not corrosive. Sensitisation:

In the Buehler assay the substance was shown to be a weak skin sensitiser, while a human patch test showed no sensitization in human volunteers.

Genetic toxicity:

The Barium analog was found to be non-mutagenic in the Ames bacterial reverse mutation assay and the mouse lymphoma test (MLA). The substance was did not cause chromosomal aberrations in human peripheral lymphocytes.

Reproductive toxicity

DNNSA (di C8-C10, branched, C9 rich, alkylnaphthalene sulphonic acid) is the major structural component of Calcium bis(di c8-c10, branched, c9 rich, alkylnaphthalene sulphonate). The OECD 422 repeat dose and reproduction/development study with DNNSA provides reliable read-across for developmental endpoints for Calcium bis(di c8-c10, branched, c9 rich, alkylnaphthalene sulphonate).

A second OECD 422 study conducted with another analog, Barium bis(di c8-c10, branched, c9 rich, alkylnaphthalene sulphonate), showed no effects on development at the highest dose in the study of 150 mg/kg/day. Together these studies show that Calcium bis(di c8-c10, branched, c9 rich, alkylnaphthalene sulphonate) is not a developmental toxin.

*REACh Dossier

Animal studies show that calcium sulfonates with a TBN greater than 300 are not skin sensitisers while the results in animals at a TBN (Total Base Number) of 300 exhibit a mixed skin sensitisation response. However, human repeat insult patch tests clearly show that high TBN overbased calcium sulfonates (TBN = 300) are not sensitisers and that low TBN calcium sulfonates do not cause sensitisation in a substantial number of persons at concentrations of 10% or lower within the definition of sensitisation under EU Regulation (EC) No. 1272/2008. The weight-of-evidence indicates that low TBN sodium and calcium sulfonates (TBN < 300) are skin sensitisers with a specific concentration limit (SCL) of 10% and that high TBN sodium and calcium sulfonates (TBN = 300) are not skin sensitisers. Studies in guinea pigs show that low TBN benzenesulfonic acid, mono-C20-24 (even)-sec-alkyl derivs., para-, sodium salts (EC No. None; CAS No. None; TBN = 3) is a skin sensitizer while benzenesulfonic acid, mono-C20-24 (even)-sec-alkyl derivs., para-, sodium salts TBN = 448) is not a skin sensitiser. Studies in guinea pigs and human volunteers show that low TBN benzenesulfonic acid, 4-(mono-C15 -36 branched alkyl derivs., C24 rich) and benzenesulfonic acid, 4-octadecyl, calcium salts (EC 939-141-9; TBN = 13) are skin sensitisers. Numerous well-conducted, reliable, controlled human (HRIPT) studies with benzene, polypropene derivs., sulfonated, calcium salts (EC 616-278-7; TBN values ranging from 13 to 85), sulfonic acids, petroleum, calcium salts (EC 263-093-9; TBN = 30 to 100), and benzenesulfonic acid, 4-(mono-C15-36 branched alkyl derivs., C24 rich) and benzenesulfonic acid, 4-octadecyl, calcium salts (EC 939-141-6; TBN = 13) show that low TBN calcium sulfonates do not cause sensitisation in a substantial number of subjects at 10% and lower. High TBN calcium sulfonates, sulfonic acids, petroleum, calcium salts (EC 263-093-9; TBN = 375 and 400) do not cause skin sensitisation in guinea pigs. Results of guinea pigs studies at TBN = 300 are mixed; two studies of sulfonic acids, petroleum, calcium salts, (EC 263-093-9) report no skin sensitisation while one study of sulfonic acids, petroleum, calcium salts (EC 263-093-9) and one study of benzene, polypropene derivs., sulfonated, calcium salts (EC 616-278-7) report skin sensitisation, However, numerous well-conducted, reliable, controlled human (HRIPT) studies with benzene, polypropene derivs., sulfonated, calcium salts (EC 616-278-7; TBN = 300) and sulfonic acids, petroleum, calcium salts (EC 263-093-9; TBN = 300) also show that high TBN (TBN = 300) do not cause skin sensitisation. In accordance with EU CLP Regulation (EC) No. 1272/2008, classification is required for low TBN sodium and calcium sulfonates (TBN < 300) with a specific concentration limit of 10% and classification is not required for high TBN calcium sulfonates (TBN = 300). Linear alkylbenzene sulfonates (LAS) are classified as Irritant (Xi) with the risk phrases R38 (Irritating to skin) and R41 (Risk of serious damage to eyes) according to CESIO (CESIO 2000). LAS are not included in Annex 1 of list of dangerous substances of Council Directive 67/548/EEC. Linear alkylbenzene sulfonic acids (LABS) are strong acids (pKa<2) are classified as corrosive (R34) Branched materials exhibit comparable toxicity to linear species.

Acute toxicity: The available data indicate minimal to moderate toxicity, with LD50 values ranging from 500 to 2000 mg/kg body weight (bw). Acute inhalation data also indicate a lack of significant toxicity. Available dermal exposure data also shows a lack of significant toxicity. LAS are readily absorbed by the gastrointestinal tract after oral administration in animals. LAS are not readily absorbed through the skin. The bulk is metabolised in the liver to sulfophenylic carboxyl acids. The metabolites are excreted primarily via the urine and faeces. The main urinary metabolites in rats are sulfophenyl butanoic acid and sulfophenyl pentanoic acid. Accumulation of LAS or its main metabolites has not been established in any organ after repeated oral ingestion.

No serious injuries or fatalities in man have been reported following accidental ingestion of LAS-containing detergent. The main clinical signs observed after oral administration to rats of doses near or greater than the LD50 values consisted of reduced voluntary activity, diarrhoea, weakness etc. Death usually occurred within 24 hours of administration. Rats appear to be more sensitive to LAS than mice.

LAS and branched alkylbenzene sulfonates may cause irritation of the eyes, skin and mucous membranes. LAS are relatively more irritating to the skin than the corresponding branched alkylbenzene sulfonates. The potential of LAS to irritate the skin depends on the concentration applied. LAS have been classified as irritating to skin at concentrations above 20% according to EU-criteria. Human skin can tolerate contact with solution of up to 1% LAS for 24 hours resulting in only mild irritation. Application of > 5% LAS to the eyes of rabbits produced irritation. Concentration of < 0.1% LAS produced mild to no irritation.

Skin sensitization was not seen in 2,294 volunteers exposed to LAS or in 17,887 exposed to formulations of LAS.

Repeat dose toxicity: A feeding study indicated that LAS, when administered for 2 years at extremely high levels (0.5%) in the diets to rats, produced no adverse effects on growth, health or feed efficiency.

Genotoxicity: The mutagenic potential of LAS was tested using Salmonella typhimurium strains, using Ames test. In these studies, LAS was not mutagenic. The available long-term studies are inadequate for evaluating the carcinogenic potential of LAS in laboratory animals. The studies available (oral administration to rats and mice) do not show any evidence of carcinogenicity.

Reproductive toxicity: In general no specific effect of LAS on reproductive processes has been seen, although dosages causing maternal toxicity may also induce some effects on reproduction. No teratogenic effects attributed to LAS exposure have been observed. Environmental and Health Assessment of Substances in Household Detergents and Cosmetic Detergent Products, Environment Project, 615, 2001. Torben Madsen et al: Miljoministeriet (Danish Environmental Protection Agency)

	Sensitisation: There is a single, key study for sensitization of the arr documented, GLP guinea pig sensitization study with non-sensitizer in guinea pigs as none of the test anim topical challenge. Repeat dose toxicity: A GLP guideline study with p-toluenesulphonic acid (f 28 days. The highest dose was 500 mg/kg bw/day (>- mg/kg bw/day. Toxicity to reproduction: No fertility studies are reported for the aromatic sulph looked at reproductive organs and development of off read-across for this endpoint. The 90-day oral rat and related compound sodium xylene sulfonate (CAS No. on reproductive organs were reported at doses rough foetal toxicity was the highest dose tested - 3000 mg/ day. The conclusion of the study was no indications of Genetic toxicity: There is a fully documented, GLP Guideline (OECD 4 Aberration Test for one of the aromatic sulphonic acid metabolic activation. The Ames test exposed up to 50 per liter of the test substance. These studies concludd There is an additional, published report of an Ames T Exposures up to 10,000 micrograms/plate were done p-toluenesulphonic acid; that is, not mutagenic and nor There are no in vivo mutagenicity studies for the aromatic conducted under GLP. Up to 240 mg (rats) and 727 m week for 104 weeks. There were no treatment related and other organs. The increased incidence of epiderr reported as 240 mg/kg bw/day for rats and 727 m/kg Elimination: The US EPA has evaluated the metabolism of analog naphthalenesulfonic acids . In a US EPA final rule for	e, including manufacturing plant book i.e., a 65% aqueous solution with < 5 omatic sulphonic acids. None of the te toluene-4-sulphonic acid (CAS No. 10 hals showed a positive response to cor CAS No. 104-15-4) reported no advers 490 mg/kg bw/day based on >98% acid ionic acids. There are however studies fspring. Hydrotropes are the salt form of a oral mouse studies and the 2-year ch 1300-72-7) included examination of s ally equivalent to those in the developm /kg bw/day which is equivalent to 936 of developmental toxicity including tera 471) Ames Test and a fully documente is, p-toluenesulphonic acid (CAS No. 200 micrograms/plate and the chromose e the substance is neither mutagenic r set for another of the aromatic sulphorie with and without metabolic activation. ot cytotoxic. natic sulphonic acids, but there are two rest for another of the aromatic sulphorie with and without metabolic activation. ot cytotoxic. sulphonic acids Two hydrotrope studied n. sulphonic acids Two hydrotrope studied incidences of mononuclear cell leuked mal hyperplasia may have been related g bw/day for mice. spin in the sodium alkyl naphthalenesulf SANS, it was stated that "the 1- or 2-s	of injuries and a physician report, show toluene- % free sulphuric acid) is an irritant to the eye and skin. sted animals showed positive responses in a, well 04-15-4). The test substance can be considered a mbined intradermal and topical induction followed by se effects to male and female rats exposed orally for tive ingredient). Therefore the NOAEL was set at 500 a for the chemically related hydrotrope substances that of the sulphonic acids and therefore are used as ironic dermal rat and mouse studies with the closely ex organs of both sexes. No treatment related effects ental toxicity study. he NOAEL for both maternal and mg active ingredient per kilogram body weight per togenesis. d, GLP Guideline (OECD 473) Chromosome 104-15-4). Both tests were conducted with and without some aberration test exposed up to 1902 micrograms iorcytotoxic. hic acids, benzenesulfonic acid (CAS No. 98-11-3). The conclusion is the same as for the o in vivo mouse micronucleus studies for the related CAS 28088-63-3). Both are GLP-compliant Guideline nees were not mutagenic in these assays.
EM-Tec C20 Carbon Conductive Micro-Tip Pen	ring may provide a handle by which these compounds Asthma-like symptoms may continue for months or ex- condition known as reactive airways dysfunction sync compound. Key criteria for the diagnosis of RADS inco- onset of persistent asthma-like symptoms within minu spirometry, with the presence of moderate to severe to lymphocytic inflammation, without eosinophilia, have irritating inhalation is an infrequent disorder with rates Industrial bronchitis, on the other hand, is a disorder to particulate in nature) and is completely reversible after	ven years after exposure to the materi drome (RADS) which can occur followi clude the absence of preceding respira ites to hours of a documented exposu bronchial hyperreactivity on methachoi also been included in the criteria for di s related to the concentration of and du that occurs as result of exposure due to	al ceases. This may be due to a non-allergenic ng exposure to high levels of highly irritating tory disease, in a non-atopic individual, with abrupt re to the irritant. A reversible airflow pattern, on line challenge testing and the lack of minimal agnosis of RADS. RADS (or asthma) following an irrition of exposure to the irritating substance. o high concentrations of irritating substance (often
EM-Tec C20 Carbon Conductive Micro-Tip Pen	by oral gavage. Acetone-induced increases in relative study. Acetone treatment caused increases in the rela- effects and the effects may have been associated wit were also noted in male rats along with hyperpigment decreased spleen weights. Overall, the no-observed- (2258 mg/kg/d), 2% for female mice (5945 mg/kg/d), reduction in foetal weight, and a slight, but statistically 15,665 mg/m3 and in rats at 26,100 mg/m3. The no- rats and mice. Teratogenic effects were not observed in rats and mic in mice treated with up to 0.2 mL of acetone did not re The scientific literature contains many different studie	mice and rats that were administered a kidney weight changes were observe ative liver weight in male and female ra h microsomal enzyme induction. Haer tation in the spleen. The most notable effect-levels in the drinking water stud and 5% for female rats (3100 mg/kg/d) y significant increase in the percent inc observable-effect level for developmen ce tested at 26,110 and 15,665 mg/m3 eveal any increase in organ tumor inci is that have measured either the neuror is ranging from about 600 to greater th tty shown that 8-hr exposures in excer digit span scores. Clinical case studies	acetone in the drinking water and again in rats treated ad in male and female rats used in the oral 13-week ats that were not associated with histopathologic natologic effects consistent with macrocytic anaemia findings in the mice were increased liver and y were 1% for male rats (900 mg/kg/d) and male mice). For developmental effects, a statistically significant cidence of later resorptions were seen in mice at tal toxicity was determined to be 5220 mg/m3 for both , respectively. Lifetime dermal carcinogenicity studies dence relative to untreated control animals. bbehavioural performance or neurophysiological an 2375 mg/m3 were not associated with any s, controlled human volunteer studies, animal
ISOBUTYL ACETATE & N-BUTANOL	The material may cause skin irritation after prolonged dermatitis is often characterised by skin redness (ervi spongy layer (spongiosis) and intracellular oedema o	thema) and swelling the epidermis. His	
CARBON BLACK & BARIUM			
DINONYL	No significant acute toxicological data identified in lite	erature search.	
NAPHTHALENESULFONATE			
NAPHTHALENESULFONATE Acute Toxicity	×	Carcinogenicity	~
	× × ×	Carcinogenicity Reproductivity	× ×

Respiratory or Skin sensitisation	~	STOT - Re	epeated Exposure	×
Mutagenicity	×		Aspiration Hazard	×
		Legend:		ot available or does not fill the criteria for classification le to make classification

11.2.1. Endocrine Disruption Properties

Not Available

SECTION 12 Ecological information

12.1. Toxicity Value EM-Tec C20 Carbon Endpoint Test Duration (hr) Species Source **Conductive Micro-Tip Pen** Not Available Not Available Not Available Not Available Not Available Test Duration (hr) Endpoint Species Value Source 0.001mg/L NOEC(ECx) 12h Fish 4 LC50 Fish 3744.6-5000.7mg/L 96h 4 acetone EC50 48h Crustacea 6098.4mg/L 5 FC50 96h 9.873-27.684mg/l 4 Algae or other aquatic plants Endpoint Test Duration (hr) Species Value Source EC0(ECx) >15.5mg/l 2 48h Crustacea isobutyl acetate LC50 96h Fish 16.6mg/l 2 246mg/l 2 EC50 72h Algae or other aquatic plants EC50 48h Crustacea 24.6mg/l 2 Value Endpoint Test Duration (hr) Species Source NOEC(ECx) 504h Crustacea 4.1mg/l 2 96h 100-500mg/l LC50 Fish 4 n-butanol EC50 72h Algae or other aquatic plants >500mg/l 1 EC50 48h 1 Crustacea >500mg/l 2 EC50 96h Algae or other aquatic plants 225mg/l Endpoint Test Duration (hr) Species Value Source NOEC(ECx) 24h Crustacea 3200mg/l 1 carbon black LC50 96h Fish >100mg/l 2 EC50 72h >0.2mg/l 2 Algae or other aquatic plants 48h 33.076-41.968mg/l EC50 Crustacea 4 Value Endpoint Test Duration (hr) Species Source NOEC(ECx) 336h Fish 47.5mg/l 2 I C50 96h Fish >100ma/l 2 propylene glycol monomethyl ether acetate, alpha-isomer EC50 72h Algae or other aquatic plants 2 >1000mg/l EC50 48h 373mg/l 2 Crustacea EC50 Algae or other aquatic plants >1000mg/l 2 96h Endpoint Test Duration (hr) Species Value Source barium dinonyl naphthalenesulfonate Not Available Not Available Not Available Not Available Not Available Extracted from 1. IUCLID Toxicity Data 2. Europe ECHA Registered Substances - Ecotoxicological Information - Aquatic Toxicity 4. US EPA, Legend: Ecotox database - Aquatic Toxicity Data 5. ECETOC Aquatic Hazard Assessment Data 6. NITE (Japan) - Bioconcentration Data 7. METI (Japan) - Bioconcentration Data 8. Vendor Data

Harmful to aquatic organisms.

For ketones

Ketones, unless they are alpha, beta--unsaturated ketones, can be considered as narcosis or baseline toxicity compounds

Hydrolysis may also involve the addition of water to ketones to yield ketals under mild acid conditions. However, this addition of water is thermodynamically favorable only for low molecular weight ketones. This addition is an equilibrium reaction that is reversible upon a change of water concentration and the reaction ultimately leads to no permanent change in the structure of the ketone substrateThe higher molecular weight ketones do no form stable ketals. Therefore, the ketones are stable to water under ambient environmental conditions Another possible reaction of ketones in water involves the enolic hydrogen on the carbony bonded to the carbonyl function. Under conditions of high pH (pH greater than 10), the enolic proton is abstracted by base (OH-) forming a carbanion intermediate that may react with other organic substrates (e.g., ketones, esters, aldehydes) containing a center for nucleophilic attack. The reactions, commonly recognized as condensation reactions, produce higher molecular weight products. Under ambient conditions of temperature, pH, and low

concentration, these condensation reactions are unfavorable.

Based on its reactions in air, it seems likely that ketones undergo photolysis in water. It is probable that ketones will be biodegraded to an appreciable degree by micro-organisms in soil and water. They are unlikely to bioconcentrate or biomagnify.

for acetone log Kow: -0.24 Half-life (hr) air: 312-1896 Half-life (hr) H2O surface water: 20 Henry's atm m3 /mol: 3.67E-05 BOD 5: 0.31-1.76,46-55% COD: 1.12-2.07 ThOD: 2.2 BCF: 0.69 Environmental fate:

Acetone preferentially locates in the air compartment when released to the environment. A substantial amount of acetone can also be found in water, which is consistent with the high water to air partition coefficient and its small, but detectable, presence in rain water, sea water, and lake water samples. Very little acetone is expected to reside in soil, biota, or suspended solids. This is entirely consistent with the physical and chemical properties of acetone and with measurements showing a low propensity for soil absorption and a high preference for moving through the soil and into the ground water

In air, acetone is lost by photolysis and reaction with photochemically produced hydroxyl radicals; the estimated half-life of these combined processes is about 22 days. The relatively long half-life allows acetone to be transported long distances from its emission source.

Acetone is highly soluble and slightly persistent in water, with a half-life of about 20 hours; it is minimally toxic to aquatic life.

Acetone released to soil volatilises although some may leach into the ground where it rapidly biodegrades.

Acetone does not concentrate in the food chain. Acetone meets the OECD definition of readily biodegradable which requires that the biological oxygen demand (BOD) is at least 70% of the theoretical oxygen demand (THOD) within the 28-day test period

Drinking Water Standard: none available

Soil Guidelines: none available Air Quality Standards: none available.

Ecotoxicity:

Testing shows that acetone exhibits a low order of toxicity

Fish LC50: brook trout 6070 mg/l; fathead minnow 15000 mg/l

Bird LC0 (5 day): Japanese quail, ring-neck pheasant 40,000 mg/l Daphnia magna LC50 (48 h): 15800 mg/l; NOEC 8500 mg/l

Aquatic invertebrate 2100 - 16700 mg/l

Aquatic plant NOEC: 5400-7500 mg/l

Daphnia magna chronic NOEC 1660 mg/l

Acetone vapors were shown to be relatively toxic to two types insects and their eggs. The time to 50% lethality (LT50) was found to be 51.2 hr and 67.9 hr when the flour beetle (*Tribolium confusum*) and the flour moth (*Ephestia kuehniella*) were exposed to an airborne acetone concentration of 61.5 mg/m3. The LT50 values for the eggs were 30-50% lower than for the adult. The direct application of acetone liquid to the body of the insects or surface of the eggs did not, however, cause any mortality.

The ability of acetone to inhibit cell multiplication has been examined in a wide variety of microorganisms. The results have generally indicated mild to minimal toxicity with NOECs greater than 1700 mg/L for exposures lasting from 6 hr to 4 days. Longer exposure periods of 7 to 8 days with bacteria produced mixed results; but overall the data indicate a low degree of toxicity for acetone. The only exception to these findings were the results obtained with the flagellated protozoa (Entosiphon sulcatum) which yielded a 3-day NOEC of 28 ma/L.

DO NOT discharge into sewer or waterways.

12.2. Persistence and degradability

Ingredient	Persistence: Water/Soil	Persistence: Air
acetone	LOW (Half-life = 14 days)	MEDIUM (Half-life = 116.25 days)
isobutyl acetate	LOW	LOW
n-butanol	LOW (Half-life = 54 days)	LOW (Half-life = 3.65 days)
propylene glycol monomethyl ether acetate, alpha-isomer	LOW	LOW

12.3. Bioaccumulative potential

Ingredient	Bioaccumulation
acetone	LOW (BCF = 0.69)
isobutyl acetate	LOW (LogKOW = 1.78)
n-butanol	LOW (BCF = 0.64)
propylene glycol monomethyl ether acetate, alpha-isomer	LOW (LogKOW = 0.56)

12.4. Mobility in soil

Ingredient	Mobility
acetone	HIGH (KOC = 1.981)
isobutyl acetate	LOW (KOC = 17.48)
n-butanol	MEDIUM (KOC = 2.443)
propylene glycol monomethyl ether acetate, alpha-isomer	HIGH (KOC = 1.838)

12.5. Results of PBT and vPvB assessment

	Р	В	т
Relevant available data	Not Available	Not Available	Not Available
PBT	X	×	×
vPvB	X	×	×
PBT Criteria fulfilled?	· · · · · · · · · · · · · · · · · · ·	· · · · · · · · · · · · · · · · · · ·	No

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15-001120 EM-Tec C20 Carbon Conductive Micro-Tip Pen

vPvB

12.6. Endocrine Disruption Properties

Not Available

12.7. Other adverse effects

Not Available

SECTION 13 Disposal considerations

13.1. Waste treatment methods	
Product / Packaging disposal	 Containers may still present a chemical hazard/ danger when empty. Return to supplier for reuse/ recycling if possible. Otherwise: If container can not be cleaned sufficiently well to ensure that residuals do not remain or if the container cannot be used to store the same product, then puncture containers, to prevent re-use, and bury at an authorised landfill. Where possible retain label warnings and SDS and observe all notices pertaining to the product. Legislation addressing waste disposal requirements may differ by country, state and/ or territory. Each user must refer to laws operating in their area. In some areas, certain wastes must be tracked. A Hierarchy of Controls seems to be common - the user should investigate: Reduction Recycling Disposal (if all else fails) This material may be recycled if unused, or if it has not been contaminated so as to make it unsuitable for its intended use. If it has been contaminated, it may be possible to reclaim the product by filtration, distillation or some other means. Shelf life considerations should also be appropriate. Do NOT allow wash water form cleaning or process equipment to enter drains. It may be necessary to collect all wash water for treatment before disposal. In all cases disposal to sever may be subject to local laws and regulations and these should be considered first. Where in doubt contact the responsible. Consult manufacturer for recycling options or consult local or regional waste management authority for disposal if no suitable treatment or disposal facility can be identified. Dispose of by: burial in a land-fill specifically licensed to accept chemical and / or pharmaceutical wastes or Incineration in a licensed appropriate. Decontaminate empty containers. Observe all label safeguards until containers are cleaned and destroyed.
Waste treatment options	Not Available
Sewage disposal options	Not Available

SECTION 14 Transport information

Labels Required



Land transport (ADR-RID)

14.1. UN number	1263	
14.2. UN proper shipping name	PAINT or PAINT RELATED MATE	RIAL
14.3. Transport hazard class(es)	Class 3 Subrisk Not Applicable	
14.4. Packing group	11	
14.5. Environmental hazard	Not Applicable	
	Hazard identification (Kemler)	33
	Classification code	F1
14.6. Special precautions for	Hazard Label	3
user	Special provisions	163 367 640C 650 640D
	Limited quantity	5 L
	Tunnel Restriction Code	2 (D/E)

Air transport (ICAO-IATA / DGR)

14.1. UN number	1263
14.2. UN proper shipping name	PAINT or PAINT RELATED MATERIAL

No

14.3. Transport hazard class(es)	ICAO/IATA Class	3	
	ICAO / IATA Subrisk	Not Applicable	
	ERG Code	3L	
14.4. Packing group	I		
14.5. Environmental hazard	Not Applicable		
14.6. Special precautions for user	Special provisions		A3 A72 A192
	Cargo Only Packing Instructions		364
	Cargo Only Maximum Qty / Pack		60 L
	Passenger and Cargo Packing Instructions		353
	Passenger and Cargo Maximum Qty / Pack		5 L
	Passenger and Cargo Limited Quantity Packing Instructions		Y341
	Passenger and Cargo Limited Maximum Qty / Pack		1 L

Sea transport (IMDG-Code / GGVSee)

1263		
PAINT or PAINT RELATED MATERIAL		
IMDG Class 3 IMDG Subrisk Not Applicable		
1		
Not Applicable		
EMS NumberF-E, S-ESpecial provisions163 367Limited Quantities5 L		

Inland waterways transport (ADN)

	,		
14.1. UN number	1263		
14.2. UN proper shipping name	PAINT or PAINT RELATED MATERIAL		
14.3. Transport hazard class(es)	3 Not Applicable		
14.4. Packing group	1		
14.5. Environmental hazard	Not Applicable		
	Classification code	F1	
	Special provisions	163; 367; 640C; 640D; 650	
14.6. Special precautions for user	Limited quantity	5L	
	Equipment required	PP, EX, A	
	Fire cones number	1	
	Limited quantity Equipment required	5L	

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

Not Applicable

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

Product name	Group
acetone	Not Available
isobutyl acetate	Not Available
n-butanol	Not Available
carbon black	Not Available
propylene glycol monomethyl ether acetate, alpha-isomer	Not Available
barium dinonyl naphthalenesulfonate	Not Available

14.9. Transport in bulk in accordance with the ICG Code

Product name	Ship Type
acetone	Not Available
isobutyl acetate	Not Available
n-butanol	Not Available
carbon black	Not Available

Product name	Ship Type
propylene glycol monomethyl ether acetate, alpha-isomer	Not Available
barium dinonyl naphthalenesulfonate	Not Available

SECTION 15 Regulatory information

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

acetone is found on the following regulatory lists		
EU Consolidated List of Indicative Occupational Exposure Limit Values (IOELVs)	European Union - European Inventory of Existing Commercial Chemical Substances	
EU REACH Regulation (EC) No 1907/2006 - Annex XVII - Restrictions on the	(EINECS)	
manufacture, placing on the market and use of certain dangerous substances, mixtures and articles	European Union (EU) Regulation (EC) No 1272/2008 on Classification, Labelling and Packaging of Substances and Mixtures - Annex VI	
Europe EC Inventory	r ackaging of Substances and Mixtures - Annex Vi	
isobutyl acetate is found on the following regulatory lists		
EU Consolidated List of Indicative Occupational Exposure Limit Values (IOELVs)	European Union - European Inventory of Existing Commercial Chemical Substances	
EU REACH Regulation (EC) No 1907/2006 - Annex XVII - Restrictions on the	(EINECS)	
manufacture, placing on the market and use of certain dangerous substances, mixtures and articles	European Union (EU) Regulation (EC) No 1272/2008 on Classification, Labelling and Packaging of Substances and Mixtures - Annex VI	
Europe EC Inventory		
n-butanol is found on the following regulatory lists		
EU European Chemicals Agency (ECHA) Community Rolling Action Plan (CoRAP) List of Substances	European Union - European Inventory of Existing Commercial Chemical Substances (EINECS)	
EU REACH Regulation (EC) No 1907/2006 - Annex XVII - Restrictions on the	European Union (EU) Regulation (EC) No 1272/2008 on Classification, Labelling and	
manufacture, placing on the market and use of certain dangerous substances, mixtures and articles	Packaging of Substances and Mixtures - Annex VI	
Europe EC Inventory		
carbon black is found on the following regulatory lists		
Chemical Footprint Project - Chemicals of High Concern List	International Agency for Research on Cancer (IARC) - Agents Classified by the IARC	
EU European Chemicals Agency (ECHA) Community Rolling Action Plan (CoRAP) List	Monographs	
of Substances	International Agency for Research on Cancer (IARC) - Agents Classified by the IARC	
Europe EC Inventory	Monographs - Group 2B: Possibly carcinogenic to humans International WHO List of Proposed Occupational Exposure Limit (OEL) Values for	
European Union - European Inventory of Existing Commercial Chemical Substances (EINECS)	Manufactured Nanomaterials (MNMS)	
(======================================		
propylene glycol monomethyl ether acetate, alpha-isomer is found on the following re-	egulatory lists	
EU Consolidated List of Indicative Occupational Exposure Limit Values (IOELVs)	European Union - European Inventory of Existing Commercial Chemical Substances	
EU REACH Regulation (EC) No 1907/2006 - Annex XVII - Restrictions on the	(EINECS)	
manufacture, placing on the market and use of certain dangerous substances, mixtures and articles	European Union (EU) Regulation (EC) No 1272/2008 on Classification, Labelling and Packaging of Substances and Mixtures - Annex VI	
Europe EC Inventory		
barium dinonyl naphthalenesulfonate is found on the following regulatory lists		
Europe EC Inventory	European Union - European Inventory of Existing Commercial Chemical Substances	
	(EINECS)	

This safety data sheet is in compliance with the following EU legislation and its adaptations - as far as applicable - : Directives 98/24/EC, - 92/85/EEC, - 94/33/EC, - 2008/98/EC, - 2010/75/EU; Commission Regulation (EU) 2020/878; Regulation (EC) No 1272/2008 as updated through ATPs.

15.2. Chemical safety assessment

No Chemical Safety Assessment has been carried out for this substance/mixture by the supplier.

National Inventory Status

National Inventory	Status
Australia - AIIC / Australia Non-Industrial Use	Yes
Canada - DSL	Yes
Canada - NDSL	No (acetone; isobutyl acetate; n-butanol; carbon black; propylene glycol monomethyl ether acetate, alpha-isomer; barium dinonyl naphthalenesulfonate)
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 - FBEPH	Yes

National Inventory	Status
Legend:	Yes = All CAS declared ingredients are on the inventory No = One or more of the CAS listed ingredients are not on the inventory. These ingredients may be exempt or will require registration.

SECTION 16 Other information

Revision Date	01/04/2022
Initial Date	04/04/2017

Full text Risk and Hazard codes		
H226	Flammable liquid and vapour.	
H302	Harmful if swallowed.	
H302+H332	Harmful if swallowed or if inhaled.	
H315	Causes skin irritation.	
H319	Causes serious eye irritation.	
H335	May cause respiratory irritation.	
H411	Toxic to aquatic life with long lasting effects.	

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. For detailed advice on Personal Protective Equipment, refer to the following EU CEN Standards:

EN 166 Personal eye-protection

EN 340 Protective clothing

EN 374 Protective gloves against chemicals and micro-organisms

EN 13832 Footwear protecting against chemicals

EN 133 Respiratory protective devices

Definitions and abbreviations

PC-TWA: Permissible Concentration-Time Weighted Average

PC-STEL: Permissible Concentration-Short Term Exposure Limit

IARC: International Agency for Research on Cancer

ACGIH: American Conference of Governmental Industrial Hygienists

STEL: Short Term Exposure Limit

TEEL: Temporary Emergency Exposure Limit.

IDLH: Immediately Dangerous to Life or Health Concentrations

ES: Exposure Standard

OSF: Odour Safety Factor

NOAEL :No Observed Adverse Effect Level LOAEL: Lowest Observed Adverse Effect Level

TLV: Threshold Limit Value LOD: Limit Of Detection

OTV: Odour Threshold Value

BCF: BioConcentration Factors

BEI: Biological Exposure Index

AIIC: Australian Inventory of Industrial Chemicals

- DSL: Domestic Substances List
- NDSL: Non-Domestic Substances List

IECSC: Inventory of Existing Chemical Substance in China

EINECS: European INventory of Existing Commercial chemical Substances

ELINCS: European List of Notified Chemical Substances

NLP: No-Longer Polymers

ENCS: Existing and New Chemical Substances Inventory

KECI: Korea Existing Chemicals Inventory

NZIoC: New Zealand Inventory of Chemicals PICCS: Philippine Inventory of Chemicals and Chemical Substances

TSCA: Toxic Substances Control Act

TCSI: Taiwan Chemical Substance Inventory

INSQ: Inventario Nacional de Sustancias Químicas

NCI: National Chemical Inventory

FBEPH: Russian Register of Potentially Hazardous Chemical and Biological Substances

Reason For Change

A-3.00 - Added UFI number and modifications to the safety data sheet