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PRODUCT: MONO PROPYLENE GLYCOL (MPG) REVISION:3 DATED: 30/06/18 PAGE 1 OF 13

PRODUCT SPECIFICATION

Product Name Alternative Name Specification Reference Mono Propylene Glycol Propane-1-2,-diol, 1,2-propanediol MPG/2 (09/07)

SALES SPECIFICATION

Exclusion of Liability

NOTES

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Health and Safety

A Material Safety Data Sheet has been issued describing the health, safety and environmental properties of this product, identifying the potential hazards and giving advice on the handling precautions and emergency procedures. This must be consulted fully before handling, storage and use.

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SAFETY DATA SHEET IDENTIFICATION OF THE SUBSTANCE/PREPARATION AND COMPANY

1.1 Product Identifier

1.

Chemical Name (EINECS)Monopropylene GlycolChemical FormulaC3H8O2SynonymsPropane-1,2-diol1,2-propanediolMPGCAS Number57-55-6EINECS Number200-338-0REACH Registration Number01-2119456809-23-XXXX

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

Identified use(s)

Exposure Scenario title	Exposure Scenario Group	Sector of Use	Applicable Use Descriptors (PROC or PC)	Applicable Use Descriptors
: Agrochemical uses	Consumer		PC 12, PC 27	ERC 8d
	Professional		PROC 4, PROC 8a, PROC 8b, PROC 11, PROC 13	ERC 8a
	Professional		PROC 4, PROC 8a, PROC 8b, PROC 11, PROC 13	ERC 8d
: Distribution of substance	Industrial	SU 9	PROC 1, PROC 2, PROC 3, PROC 4, PROC 8a, PROC 8b, PROC 9, PROC 15	ERC 1
	Industrial	SU 9	PROC 1, PROC 2, PROC 3, PROC 4, PROC 8a, PROC 8b, PROC 9, PROC 15	ERC 2
: Formulation & (re)packing of substances and mixtures	Industrial	SU 10	PROC 1, PROC 2, PROC 3, PROC 4, PROC 5, PROC 8a, PROC 8b, PROC 9, PROC 14, PROC 15	ERC 2
: Functional Fluids	Consumer		PC 16, PC 17	ERC 9a
	Consumer		PC 16, PC 17	ERC 9b
	Industrial		PROC 1, PROC 2, PROC 3, PROC 4, PROC 8a, PROC 8b, PROC 9, PROC 5	ERC 7
	Professional		PROC 1, PROC 2, PROC 3, PROC 8a, PROC 9, PROC 20	ERC 9a
	Professional		PROC 1, PROC 2, PROC 3, PROC 8a, PROC 9, PROC 20	ERC 9b
: Laboratory agents	Industrial		PROC 10, PROC 15	ERC 4
	Professional		PROC 10, PROC 15	ERC 8a
: Manufacture of substance	Industrial	SU 9	PROC 1, PROC 2, PROC 3, PROC 4, PROC 8a, PROC 8b, PROC 15	ERC 1
: Mining chemicals	Industrial		PROC 1, PROC 2, PROC 3, PROC 4, PROC 8a, PROC 8b, PROC 9, PROC 23	ERC 8d
: Other Consumer Uses	Consumer		PC 28, PC 29, PC 39	ERC 8a
	Consumer		PC 28, PC 29, PC 39	ERC 8d
: Polymer processing	Industrial	SU 10	PROC 1, PROC 2, PROC 3, PROC	ERC 3
			4, PROC 5, PROC 6, PROC 8a, PROC 8b, PROC 14, PROC 21	
	Industrial	SU 10	PROC 1, PROC 2, PROC 3, PROC 4, PROC 8a, PROC 8b, PROC 14, PROC 21	ERC 6c
: Use as binders and release agents	Industrial		PROC 1, PROC 2, PROC 3, PROC 4, PROC 5, PROC 6, PROC 7, PROC 8b, PROC 9, PROC 12, PROC 13, PROC 14, PROC 15, PROC 10	ERC 4

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	Professional	PROC 1, PROC 2, PROC 3, PROC 4, PROC 5, PROC 6, PROC 8a, PROC 8b, PROC 10, PROC 11, PROC 14, PROC 9, PROC 13, PROC 19	ERC 8c
: Use in Cleaning Agents	Consumer	PC 3, PC 4, PC 9c, PC 9b, PC 9a, PC 24, PC 35	ERC 8a
	Consumer	PC 3, PC 4, PC 9c, PC 9b, PC 9a, PC 24, PC 35	ERC 8d
	Industrial	PROC 1, PROC 2, PROC 3, PROC 4, PROC 7, PROC 8a, PROC 8b, PROC 9, PROC 13	ERC 4
	Professional	PROC 1, PROC 2, PROC 3, PROC 4, PROC 8a, PROC 8b, PROC 10, PROC 11, PROC 13, PROC 9	ERC 8a
	Professional	PROC 1, PROC 2, PROC 3, PROC 4, PROC 8a, PROC 8b, PROC 10, PROC 11, PROC 13, PROC 9	ERC 8d
: Use in/as de-icing/anti-icing applications/agents (consumer use)	Consumer	PC 4	ERC 8d
Use in/as de-icing/anti-icing applications/agents (professional)	Professional	PROC 2, PROC 8b, PROC 2	ERC 8d
Uses in Coatings	Consumer	PC 1, PC 4, PC 9a, PC 9b, PC 9c, PC 18, PC 23, PC 24, PC 31	ERC 8a
	Consumer	PC 1, PC 4, PC 9a, PC 9b, PC 9c, PC 18, PC 23, PC 24, PC 31	ERC 8d
	Industrial	PROC 1, PROC 2, PROC 3, PROC 4, PROC 5, PROC 7, PROC 8a, PROC 8b, PROC 10, PROC 13, PROC 9, PROC 15	ERC 4
	Professional	PROC 1, PROC 2, PROC 3, PROC 4, PROC 5, PROC 8a, PROC 8b, PROC 9, PROC 10, PROC 11, PROC 13, PROC 15, PROC 19	ERC 4
	Professional	PROC 1, PROC 2, PROC 3, PROC 4, PROC 5, PROC 8a, PROC 8b, PROC 9, PROC 10, PROC 11, PROC 13, PROC 15, PROC 19	ERC 8a
	Professional	PROC 1, PROC 2, PROC 3, PROC 4, PROC 5, PROC 8a, PROC 8b, PROC 9, PROC 10, PROC 11, PROC 13, PROC 15, PROC 19	ERC 8b
Water treatment chemicals	Industrial	PROC 1, PROC 2, PROC 3, PROC 4, PROC 5, PROC 8a, PROC 8b	ERC 4
	Professional	PROC 1, PROC 2, PROC 3, PROC 4, PROC 8b	ERC 8a
	Professional	PROC 1, PROC 2, PROC 3, PROC 4, PROC 8b	ERC 8b
	Professional	PROC 1, PROC 2, PROC 3, PROC 4, PROC 8b	ERC 8d
	Professional	PROC 1, PROC 2, PROC 3, PROC 4, PROC 8b	ERC 8e

Uses advised against

Group	Use advised against	Use descriptors	Environmental release category (ERC)
Consumer	No uses advised against		
Industrial	No uses advised against		
Professional	No uses advised against		
Group	Use advice against	Use descriptors	Article (AC)
Consumer	No uses advised against		
Industrial	No uses advised against		
Professional	No uses advised against		

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Email: <u>iı</u>	Email: <u>info@lucemill.com</u>										
1.4 Emergency telephone number Tel: 44(0) 141 776 7237											
2. HAZARDS IDENTIFICATION											
2.1 Cla	assification	of the substance or mixtu	re								
211 Ba	mulation 12	72/2008 (CT P)									
Not clas	sified as dan	gerous according to the cri	teria of Regulation	(EC) No 1272/2	008						
212FF	Silicu as dan	67/548/FFC & Directive	1000/15/FC	(EC) 10 1272727	008						
Not clas	sified as dan	gerous according to the cri	teria of directive(s)	67/548/FFC and	1/or 1999/45	/FC					
3	since as can	COMPOSITION/IN	FORMATION		DIFNTS						
Substan	0005										
Propan	e-1.2-diol										
CAS	EINECS	REACH registration	Classification	Classification	Content	Note					
Number	Number	number	according to	according to							
			Directive	Regulation							
57-55-6	200-338-0	01-2119456809-23-XXXX	67/548/EEC	1272/2008	>99%	Substance with a community					
57 55 0	200 550 0	01 211) 10000) 20 Millin			22270	workplace exposure limit.					
See sect	ion 16 for th	e full text of the R, H- and	EUH-phrases decl	ared above							
Occupat	ional exposu	re limits, if available, are l	isted in section 8								
4.		FIRST AID MEAS	URES								
4.1 Desc	ription of fi	rst aid measures									
General	Advice										
Check th	e vital functi	ions. Unconscious: maintai	n adequate airway	and respiration. F	Respiratory a	arrest: artificial respiration					
or oxyge	n: Victim in	shock: on his back with leg	gs slightly raised. V	omiting: prevent	asphyxia/as	spiration pneumonia.					
Prevent c	cooling by co	overing the victim (no warr	ning up). Cardiac a	rrest: perform res	suscitation.	Victim conscious with					
labored b	reathing: ha	lf-seated. Keep watching th	ne victim. Give psy	chological aid. K	eep the vict	im calm, avoid physical					
strain. De	epending on	the victim's condition: doc	tor/hospital. Alcoh	ol consumption in	ncreases the	toxicity.					
Inhalatio)n	(. C 1 'n D 'n (11	L (/ 1 ² 1							
Remove	te vicum in	tto fresh air. Respiratory pr	oblems: consult a c	loctor/medical se	rvice.						
SKIN CON	tim to a doct	or if irritation persists Rin	se with water. Do r	ot apply (chemic	al) noutraliz	ing agents					
Eve conf	act	of it initiation persists. Kin	se with water. Do i	iot appry (enemie	ai) incutianz	ang agents.					
Rinse wi	th water. Do	not apply neutralizing age	nts. Take victim to	an ophthalmolog	ist if irritati	on persists.					
Ingestion	1					F i i i i i i i i i i					
Rinse mo	outh with wa	ter. Consult a doctor/medic	cal service if you fe	el unwell.							
4.2 Most	import syn	ptoms and effects, both a	acute and delayed								
4.2.1 Ac	ute symptor	ns									
If applica	able and avai	lable it will be listed below	v.								
After inl	nalation:										
Dry /sore	throat. EXF	OSURE TO HIGH CONC	ENTRATIONS.								
Alter ski	in contact:	altin Dury altin ON CONTR									
Slight irr	nation. Red	skin. Dry skin. ON CONT	UNUUSEAPUSUR	E/CONTACT.							
Slight im	tation Dode	ness of the eve tissue									
After inc	restion:	icss of the eye ussue.									
Nausea.	Abdominal r	ain. AFTER ABSORPTIO	N OF HIGH OUA	NTITIES:							

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4.2.2 Delayed symptoms
If applicable and available it will be listed below.
4.3 Indication of any immediate medical attention and special treatment
needed If applicable and available it will be listed below.
5. FIRE FIGHTING MEASURES
5.1 Extinguishing Media
Suitable extinguishing media: Carbon dioxide. Water spray, Polyvalent foam. BC powder: Preferably: alcohol-resistant
foam.
Unsuitable extinguishing media: Container may slop over if solid jet(water/foam) is applied.
5.2 Special hazards arising from the substance or mixture
Upon combustion CO and C02 are formed.
5.3 Advice for fire-fighters
5.3.1 Instructions:
Cool tanks/drums with water spray/remove them into safety.
5.3.2 Special protective equipment for fire-fighters:
Gloves. Protective clothing. Heat/fire exposure: compressed air/oxygen apparatus.
6. ACCIDENTAL RELEASE MEASURES
6.1 Personal precautions, protective equipment and emergency procedures
6.1.1 For non-emergency personnel
See heading 8.2
6.1.2 For emergency responders
General
Gloves
Protective clothing
Suitable protective clothing
butyl rubber, natural rubber, polyethylene, PVC,
polyethylene/ethylenevinylalcohol Unsuitable protective clothing
6.2 Environmental precautions
Contain released substance, pump into suitable containers. Plug the leak, cut off the supply.
6.3 Methods and material for containment and cleaning up
Take up liquid spill into a non combustible material e.g.: sand, earth, vermiculite. Scoop absorbed substance into
closing containers. Clean contaminated surfaces with an excess of water. Wash clothing and equipment after handling.
6.4 Reference to other sections
See heading 13
7. HANDLING AND STORAGE
7.1 Precautions for safe handling
Keep away from naked flames/heat. At temp>flashpoint: use spark-/explosion proof appliances. Finely divided: spark-
and explosion proof appliances. Finely divided: keep away from ignition sources/sparks. Observe normal
hygiene standards. Keep container tightly closed.
7.2 Conditions for safe storage, including any incompatibilities
7.2.1 Safe storage requirements:
Store in a dry area. Ventilation at floor level. Store at ambient temperature. Keep out of direct sunlight. Meet the
legal requirements.
7.2.2 Keep away from:
Oxidizing agents, reducing agents, (strong) acids, water/moisture.
7.2.3 Suitable packaging material:
Stainless steel, carbon steel, aluminium, copper, nickel, bronze, steel with plastic inner lining.
7.3 Specific end use(s)
For relevant identified uses, see exposure scenarios attached in annex. See information supplied by the manufacturer.

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8. **EXPOSURE CONTROLS/PERSONAL PROTECTION** 8.1 Control parameters 8.1.1 Occupational exposure If limit values are applicable and available these will be listed below. a) Occupational exposure limit values Limit Value (UK) Propane-1,2-diol (total(vapour and part.) and particulates Short time value -ppm - mg/m³ Time-weighted average 10 P/474 T mg/m3 - P/150 T exposure limit ppm b) National biographical limit values If limit values are applicable and available these will be listed below. 8.1.2 Sampling Methods Product Name Number Remarks Sampling method Test Propylene Glycol OSHA CSI Propylene Glycol NIOSH 5523 Applicable limit values when using the substance or mixture as intended adsorption tubes 8.1.3 If limit values are applicable and available these will be listed below. 8.1.4 DNEL/PNEC values Acute: systemic/local effects workers Effect level Value Type Remark (DNEL/DMEL) DNEL Acute systemic effects dermal Not quantifiable Acute systemic effects inhalation Not quantifiable Acute local effects dermal Not quantifiable Acute local effects inhalation Not quantifiable Long-term systemic effects dermal Not quantifiable Long-term systemic effects inhalation 186mg/m³ Long-term local effects dermal Not quantifiable Long-term local effects inhalation 10 mg/m^3 Acute: systemic/local effect general population Value Remark Effect level Type (DNEL/DMEL) DNEL Acute systemic effects dermal Not quantifiable Acute systemic effects inhalation Not quantifiable Acute -systemic effects oral Not quantifiable Not quantifiable Acute local effects dermal Acute local effects inhalation Not quantifiable Long-term systemic effects dermal Not quantifiable Long-term systemic effects inhalation 50 mg/m^3 Long term -systemic effects oral Not quantifiable Long-term local effects dermal Not quantifiable Long-term local effects inhalation 10mg/m^3 PNEC Compartments Value Remark 206 mg/l FRESH WATER Marine water 26 mg/l Fresh water sediment 572 mg/kg sediment dw 57.2 mg/kg sediment dw Marine water sediment SOIL 50 mg/kg soil dw STP 2000 mg/l 8.1.5 Control banding

If applicable and available it will be listed below:

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8.2 Exposure controls									
The information in this section is a general description	Always use the relevant exposur	e scenarios that correspond							
to your identified use. For relevant identified uses see	exposure scenarios attached in an	nex							
A ppropriate engineering controls	exposure seenarios attached in an								
Observe normal hygiene standards. Keen container tig	atly closed. Do not est, drink or su	moke during work Keen away							
from naked flames/heat									
At tempsflashpoint: use snark-/explosion proof appliat	Irom naked Hames/neat.								
Finely divided: spark- and explosion proof appliances	ices.								
Finely divided: keen away from ignition sources/snark	C								
Recipitatory notection									
Respiratory protection not required in normal condition	18.								
Hand protection									
Wear suitable gloves									
Gloves material									
Materials for protective clothing (excellent resistance)									
Materials for protective clothing (good resistance)									
Butyl rubber, natural rubber, polyethylene, PVC, polye	thylene/ethylenevinylalcohol.								
Eye protection	• • •								
safety glasses									
Skin protection									
Protective clothing									
Environmental exposure controls See 6.2,6.3 and 13									
9. PHYSICAL AND CHEM	AICAL PROPERTIES								
9.1 Information on basic physical and chemical pro	perties								
Appearance	Liquid								
Colour	Colourlass								
Odour	Almost adourlass								
	Almost odouriess								
Malting point/frogging point	<u>0.3-7.3</u>	Test data							
Deiling point/heeling ronge	<-20 C 1002.2 kD ₂	Test data							
Flash point	104 °C	Test data							
	$\frac{104}{0.2 \text{ kp}_{2}} \otimes 20^{9} \text{C}$	Test data							
Palating regions density	0.2 HPa @ 20 C	Test data							
Relative vapour density	$\frac{2.0}{1.02 \times 20^{9} \text{C}}$								
Relative density	1.03 @20 °C								
Ethanol	Complete								
	Complete								
Ether	12a/100m1								
	1.07	Trat data							
Log POW	-1.07	Test data							
Dynamia Viscosity	>400 C								
Explosive properties	0.0434 Pa.s @25 C	with avalaging manantics							
Explosive properties	No chemical group associated	with explosive properties							
9.2 Other information	No chemical group associated	with oxidising properties							
Minimum ignition energy									
SADT									
Specific conductivity	440000pS/m								
Surface tension	$0.0716 \text{ N/m} @ 21.5^{\circ}\text{C}$								
Solidification (freezing) point									
Softening point									
Critical temperature									
Critical pressure	_								
Relative density saturated vapour/air mixture									
Saturation concentration									

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STABILITY AND REACTIVITY

10.1 Reactivity

10.

11.

Temperature above flashpoint: higher fire/explosion hazard.

10.2 Chemical stability

Hygroscopic.

10.3 Possibility of hazardous reactions

Reacts violently with (strong) oxidizers: (increased) risk of fire. Violent to explosive reaction with (strong) acids.

10.4 Conditions to avoid

Keep away from naked flames/heat. At temp>flashpoint: use spark-/explosion proof appliances. Finely divided: spark-and explosion proof appliances. Finely divided: keep away from ignition sources/sparks.

10.5 Incompatible materials

Oxidizing agents, reducing agents, (strong) acids, water/moisture.

10.6 Hazardous decomposition products

Upon combustion CO and C02 are formed.

TOXICOLOGICAL INFORMATION

Toxicokinetics: summary

Oral absorption: Toxicokinetic behavior of monopropylene glycol and its structural homologue tripropylene glycol upon oral administration to rats was investigated in a well-conducted and well-reported study (The Dow Chemical Company, 1995). In this study, two groups of 5 male rats were administered a single oral dose of either radiolabeled (14C) tripropylene glycol or non-radiolabeled monopropylene glycol by gavage in water at target concentrations 40 mg/kg bw and 50 mg/kg bw, respectively. The excreta were collected for ca. 24 hours postdosing. After sacrifice 24 hours postdosing the remaining radioactivity in tissues was determined for the first group and urine was analyzed for free and acidabile conjugates of mono-, di- and tripropylene glycol for both groups. While the absorption of monopropylene glycol has not been specifically investigated in the study, the data on tripropylene glycol indicate that it is rapidly adsorbed if administered by gavage, based on the average recovery of ca. 91% of the 14C label administered from excreta, C02, skin, tissues and carcass after ca. 24 hours postdosing sacrifice. The absorption of tripropylene glycol via oral route was calculated to amount to at least 86%, based on 5% of the administered dose recovered in faeces. As monopropylene glycol has a significantly lower molecular weight, its absorption from the gut is expected to occur even faster. Toxicokinetic behavior of monopropylene glycol in humans and experimental animals was also evaluated by the NTP CERHR expert panel (National Toxicology Program, 2004a), which concluded that available data indicate rapid and extensive absorption. Therefore a value of 100% for oral absorption shall be used for risk assessment for monopropylene glycol.

Distribution: No data on the distribution of monopropylene glycol were reported in the study; however, in case of tripropylene glycol, approximately 10% of the radiolabeled dose was recovered in tissues and carcass, with the liver and kidney having the greatest amount of radiolabel per gram of tissue 24 hours after dosing (ca. 0.2 and 0.1%, respectively). The 14C concentration in blood was approximately 6.4 and 2.8 -fold lower than in liver and kidney, respectively. The expert panel of NTP CERHR (National Toxicology Program, 2004a) concluded that monopropylene glycol is rapidly distributed into total body water.

Metabolism and excretion: In the study with rats administered monopropylene glycol and radiolabeled monopropylene glycol, the data on the animals indicate that approximately 11% of the monopropylene glycol administered was recovered in the urine as free monopropylene glycol (with < 1% of the dose recovered as acid-labile conjugates). In the study with radiolabeld tripropylene glycol, twenty-four hours after administration of a single oral dose of 40 mg/kg bw to male rats, only 5.8% of the dose was recovered as unmetabolized parent compound in the urine, while 7.2% was recovered as acid-labile conjugates of tripropylene glycol, 5.1% and 3.3% as free and acid-labile conjugates of dipropylene glycol and 3.3% and 0.6% as free and acid-labile conjugates of monopropylene glycol, respectively. A large fraction (21%) of the 14C-tripropylene glycol dose was catabolized all the way to 14C02, indicating considerable breakdown of tripropylene glycol. According to the NTP CERHR expert panel report (National Toxicology Program, 2004a), the rate-determining step in the metabolism is alcohol dehydrogenase which, when saturated, switches from a first order process into a zero order process. Saturation of metabolism appears to occur in rats and rabbits at a dose of about 1600 to 2000 mg/kg bw, whereas in humans this seems to happen at a dose of about 200 mg/kg bw. In accordance with a zero order process, the half-life of monopropylene glycol in humans and rats increases from about 1.5 hours to more than 5 hours with increasing doses above metabolic saturation. By a NAD-dependent reaction, alcohol dehydrogenase converts monopropylene glycol to lactate.

Since monopropylene glycol has a chiral center, technical grade monopropylene glycol results in the formation of 50/50 D, Llactate. L-lactate is indistinguishable from endogenous lactate, which is a good substrate for gluconeogenesis. D-lactate is less readily converted to glucose than L- lactate, which prolongs its half-life leading, under conditions of prolonged exposure, to D-lactic acidosis. It is difficult to cause L-lactic acidosis even with very high doses of

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monopropylene glycol because of its efficient detoxification via gluconeogenesis. The second reason for lack of development of L-lactic acidosis is the saturation of alcohol dehydrogenase, which results in a constant rate of lactate production. Due to removal of L-lactate by gluconeogenesis, a further increase in lactate levels is not possible after saturation of metabolism. The excretion of monopropylene glycol is species-dependent. Humans clear about 45% of monopropylene glycol via kidney, and in dogs, up to 88%. In rats and rabbits, very little of the parent compound is excreted by the kidney until saturation of metabolism occurs. Inhibition of alcohol dehydrogenase by pyrazole increases urinary excretion of monopropylene glycol to 75% in rats, as expected. Since monopropylene glycol has very low intrinsic toxicity, saturation of metabolism plays a protective role in its toxicity since the conversion of monopropylene glycol to the more toxic lactate (particularly D-lactate) is slowed.

Inhalation route of exposure: Only limited data addressing the absorption of monopropylene glycol by inhalation are available. Bau et al. (1971) reported that less than 5% of a technetium-labeled aerosol containing 10% monopropylene glycol in deionized water was taken up by human volunteers after inhalation for 1 hour in a mist tent. The authors measured the aerosol mass median diameter to be 4.8 - 5.4 microns, a size small enough to have enabled penetration to the deep lung. Ninety percent of the dose was found in the nasopharynx and it rapidly entered the stomach with very little entering the lungs. Monopropylene glycol was not directly measured, not allowing the determination of absorption through the nasal mucosa. However, the low dose rate from inhalation exposure and the small surface area would not lead to significant absorption of monopropylene glycol

Dermal route of exposure: An in vitro skin penetration study (El du Pont de Nemours and Company, 2007) with the monopropylene glycol using human cadaver skin and performed under infinite dose conditions, was available for assessment. A nominal dose of 1200 pL/cm2 (ca. 1.2 g/cm2) of the neat substance was applied for 24 hours under occlusive conditions to 6 skin replicates representing 5 human subjects. By the conclusion of the 24-hour exposure interval, only a negligible portion of the applied dose of neat monopropylene glycol (0.14%) had penetrated through the skin into the receptor fluid. The integrity of human skin, as determined by electrical impendance (El), was affected by continuous exposure to monopropylene glycol under occlusive conditions. The ratio of the post-El values was 0.33, confirming that the barrier properties of the stratum corneum were altered by monopropylene glycol.

In general, monopropylene glycol was detected in receptor fluid within about an hour of application (lag time ~ 6 hours); steady-state penetration, which was represented by no less than 4 data points, was determined to be 95.4 |ig/cm2/h (r2]0.999). This represents the maximum flux for neat monopropylene glycol. Based on the slope at steady-state (95.4 ng/cm2/h) and the concentration of monopropylene glycol in the applied solution, taken as its density (1,036,000 pg/cm3), the permeability coefficient for neat monopropylene glycol calculated to be 9.21x10-5cm/h. Based on the results of the study, a value of 40% for dermal absorption has been chosen by expert judgment to be used in the risk assessment. This value has been chosen as an average value between the percentage of dermal absorption obtained in the study and the maximal oral absorption (corresponding to 100%), and is considered to represent a worst-case approach, Propane-1,2-diol

	Parameter	Method	Value	Exposure time	Species	
Acute toxicity: oral	LD50	Equivalent or similar to	22000mg/kg bw		Rat	Experimental value
		UECD 401			(Male/Telliale)	
Acute toxicity: dermal	LD50	Equivalent or similar to OECD 402	>2000mg/kg bw	2h	Rabbit	Experimental value
Acute toxicity: inhalation	LC50	Equivalent or similar to OECD 403	317042 mg/l	2h	Rabbit	Experimental value

Propane-1,2-diol

		Results	Method	Exposure time	Time point	Species	
Corre	osion/irritation: eye	Not irritating	OECD 405:AcuteEye Irritation /Corrosion		24;48;72 hours	Rabbit	Experimental value
Corre	osion/irritation: skin	Not irritating	Equivalent or similar to OECD 404		24;48;72 hours	Rabbit	Experimental value
		Slightly irritating	Patch test	24h	24 hours	Human	Experimental value
A Co inhal	orrosion/irritation: ation	No data available					

Propane-1,2-diol

	Result	Method	Exposure	Observation time	Species	
			time			
Sensitisation: skin	Not sensitising	OECD 429: Skin Sensitization;	-		Mouse	Experimental value
		Local Lymph Node Assay				
	Not sensitising	Patch test			Human	Experimental value
					(male/female)	
Sensitisation: inhalation						Not relevant expert
						judgement

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Propane-1,2-diol											
	Parameter	Method	Value	Organ	Effect	Exposure time	Species				
Specific target organ toxicity:	NOAEL	Other	1700 mg/kg		No effects	102 weeks	Rat	Experimental			
oral			bw/day			(daily, 5	(Male/female)	value			
						days/week)					
Specific target organ toxicity:	NOAEL	Other	0.02ml		No effects	10 weeks (daily,	Mouse	Experimental			
dermal			(twice a			5days/week)	(Male/female)	value			
			week)								
Specific target organ toxicity:	LOAEC	Other	160mg/m ³	Nose	No effects	90days	Rat	Experimental			
inhalation							(Male/female)	value			

Propane-1,2-diol

.								
	Result	Method		Exposure time	Test substrate	Organ	Effect	
Germ cell mutagenicity	Negative	OECD 471: Bacterial			Bacteria			Experimental
		Reverse Mutat	ion Test		(S.typhimurium))		value
	Negative	OECD 473: in Vitro mammalian Chromosome Aberration Test			Human lymphocytes			Experimental value
	Negative	OECD 475: Mammalian Bone marrow Chromosome Aberration test			Rat (Male)			Experimental value
Propane-1,2-diol								
-	Parameter	Value	Method	Exposure times	Species Or	gan	Effect]
Carcinogenicity	NOAEL	1700mg/kg	other	102 weeks	Rat			Experimental
		Bw/day		(daily,5 days/week)	Male/female)			value

Propane-1,2-diol

	Parameter	Method	Value	Exposure time	Species	Organ I	Effect	
Developmental toxicity	NOAEL	Equivalent or similar	10400 mg/kg	9 days	Mouse		No effect	Experimental
		to OECD 414	bw/day		(Male/female)			value
Effects on fertility	NOAEL	OECD 416: two-	10100 mg/kg		Mouse		No effect	Experimental
		generation	bw/day		(Male/female)			value
		reproduction toxicity	-					
		study						

Conclusion

12.

Low acute toxicity by the oral route.

Low acute toxicity by the dermal route.

Low acute toxicity by the inhalation route.

Not classified as irritating to the skin.

Not classified as irritating to the eye.

Not sensitizing for skin.

No respiratory sensitization data available.

Low sub-chronic toxicity by the oral route.

Low sub-chronic toxicity by the dermal route.

Low sub-chronic toxicity by inhalation route.

Not classified for carcinogenicity

Not classified for mutagenic or genotoxic toxicity (negative result).

Not classified for reprotoxic or developmental toxicity

ECOLOGICAL INFORMATION

12.1 Toxici	ity						
LC50 fishe	s						
Parameter	Method	Value	Duration	Species	Test design	Fresh/salt water	
LC50	other	40613mg/l	96h	ONCORHYNCHUS	STATIC	FRESH	Experimental value
				MYKISS	SYSTEM	WATER	
EC50 Daph	nia						
Parameter	Method	Value	Duration	Species	Test design	Fresh/salt water	
LC50	EPA600/4-90/027	18340 mg/l	48h	CERIODAPHNIA	STATIC	FRESH	Experimental value
				DUBIA	SYSTEM	WATER	
LC50	FIFRA 72-3	18800 mg/l	96h	Americamysis bahia	STATIC	SALT WATER	Experimental value
					SYSTEM		

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EC50 other	aquatic organisms								
Parameter	Method	Value	Duration	Species	Test des	sign Fresh/salt water			
Threshold limit	t algae	1							
Parameter	Method	Value	Duration	Species	cies Test desi		Fresh/salt water		
EC50	OECD 201: Alga, Growth inhibition Test	19000mg/l	96h	Pseudokirchnere subcapitata	ella STATIC SYSTEM		FRESH WATER	Experimental value	
EC50	OECD 201: Alga, Growth inhibition Test	19100 mg/l	96h	SKELETOEMA COSTATUM	STATIO SYSTE	TIC SALT WATER		Experimental value	
Long-tem to	oxicity to fish								
Parameter	Method	Value	Duration	Test design	Test design		lt water		
ChV	ECOSAR	2500mg/l	30days		<u> </u>		WATER	QSAR	
Long-term t	oxicity to aquatic inver	tibrates							
Parameter	Method	Value	Duration	Test design	Test design		lt water		
NOEC	EPA 600/4-89/001	13020mg/l	7days	Semi-static		FRESH WATER		Experimental value	
T	1			1					
Toxicity sec	International Method	Value	Duration	Test design	Test design		lt motor	Vormiirino	
Parameter					Test design Fres				
LC50	other	6983	10 days	STATIC SY	STATIC SYSTEM S		ATER	Experimental value	
Toxicity to	water micro-organisms								
Parameter	Method	Value	Duration	Test design	Test design		lt water	Species	
NOEC	other	20000 mg/l	18 days		F		WATER	PSEUDOMNIA SPUTIDA	
Conclusion Not harmfu Not harmfu Not harmfu	1 l to fishes (LC50(96h) > l to invertebrates(EC50 l to algae (EC50 (72h)> l to bacteria (EC50 >10	>1000 mg/l) (48h)>1000n 1000 mg/l) 00 mg/l)	ng/l)						
12.2 Persist	tence and degradabilit	y							
Biodegrada	tion in water	Value		Duration			1		
Method		value		Duration					
OECD 301F:manometric Respirometry Test 81.7% 28 days Experimental value									
Method	Phototransformation in air (D150 air)		Conc OH rad	Conc OH radicals Re		Reference		Remark	
AOPWIN vi9.2		0.83 days	1.5 x 10^6	1.5 x 10^6 OSAR					
Phototransfo	ormation in water (DT5	0 water)							
Method		Value	Conc OH rad	Conc OH radicals Re		Reference		Remark.	
other		2.3 years		C		Calculated value			
Conclusion Readily biod Photodegrad Biodegradat	degradable in water lation in water occurs si tion in the soil under an	lowly aerobic condi	tions						

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12.3 Bio accumulative potential										
BCF fishes										
species	Species duration			varue			Reference		Kelliark	
				0.09			Calculated	value		
Log Pow Method	Value			Remark		Reference				
Method		Tempera		value			Kemark			
Equivalent or simila	r to OECD 107	20.5°C		-1.07					Test data	
Conclusion Bioaccumulation: not applicable										
12.4 Mobility in soil										
Log Pow										
	Value		Referer	Reference		Method		Temperature		
	-1.07		Test da	ıta	Equivalent or similar to		r similar to	20.5°C		
Mobility in soil					OE	CD 107				J
internity in som	Parameter		Method		Val	Value		Reference		
Volatility (Henry'	Law Constan	tH)								
	Method		Value		Temperature			Reference		
EUSES calculation		1	0.00566		12°C		ESTIMAT		D VALUE	
Method	Fraction air	Fraction biota	a	Fraction sediment Fraction		Fraction	n soil Fraction wa		r Refe	rence
						10.10	5011	40.004		
Mackay level III 2.98%				0.07%		48.1% 48.8%		48.8%	Calcu	lated value
Conclusion Low potential for absorption in soil										
12.5 Results of	PBT and vPvB	assessment								
Substance does	not meet the scre	ening criter	ria for	persistency	nor bio	baccum	ulation so	is neither	PBT nor	vPvB.
12.6 Other adverse effects										
Global Warming Potential (GWP)										
Industrial designation or common name Lifetime Kadiative efficiency SAK# (100-yr) GwP 100-yr time nonzon GwP 500-yr time horizon										
Ozone-depleting potential (ODP)										
Ozone layer (Council Regulation (EC) no 1005/2009)								2009)		
Surface Water				Mild water pollutant (surface water)						
Ground Water				Ground water pollutant						
Air Contamination			I	Low potential for volatization from water surface						
Water Ecotoxicity reaction product										
13. DISPOSAL CONSIDERATIONS										
13.1 Waste treatment methods										
Provisions relating to waste										
Waste material	Waste material code (Directive 2008/98/EC, decision 2001/118/EC). Other organic solvents, washing liquids and									
mother liquors antifreeze fluids containing dangerous substances. Depending on branch of industry and production										

process, also other EURAL codes may be applicable. Hazardous waste according to Directive 2008/98/EC.

Disposal methods

Recycle by distillation. Remove to an authorized incinerator equipped with an afterburner and a flue gas scrubber. Remove waste in accordance with local and/ or national regulations. Obtain the consent of pollution control authorities before discharging to wastewater treatment plants. In appropriate low concentrations inhibition of the degradation of activated sludge is not anticipated. Do not discharge into surface water.

Packaging/Container

Waste material code packaging (Directive 2008/98/EC). Packaging containing residues of or contaminated by dangerous substances.

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14 TRANSPORT INFO	ORMATION						
14. IRANSFORT INFORMATION							
ADK/KID/IMDG/IATA							
14 2 Proper Shinning Name							
ADR/RID/IMDC/IATA							
Not considered dangerous goods							
14 3 Transport hazard class							
ADR/RID/IMDG/IATA							
Not considered dangerous goods							
14.4 Packing group							
ADR/RID/IMDG/IATA							
Not considered dangerous goods							
14.5 Environmental							
ADR/RID/IMDG/IATA							
Not considered dangerous goods							
14.6 Special precautions for users							
ADR/RID/IMDG/IATA							
Not considered dangerous goods							
14.7 Transport in bulk according	to Annex II of MAR	POL 73/78 and the IBC Code					
ADR/RID/IMDG/IATA							
Not considered dangerous goods							
15. REGULATORY I	NFORMATION						
15.1 Safety, health and environme	ntal regulations/leg	islation specific for the substance or mixture					
European legislation:							
REACH registration	REACH registration						
Substance is not classified as danger	ous, so no exposure	scenarios are available.					
National legislation							
-The Netherlands							
Waterbezwaarlijkheid (for NL)11							
Waste identification other lists of waste materials LWCA (the Netherlands) : KGA category 03							
-Germany	-Germany						
WGK	1	Classification water polluting in compliance					
		with Verwaltungsvorschrift					
		wassergefahrdender Stoffe (VwVwS) of 27					
		July 2005 (Anhang 2)					
TA-Luft	Propane-1,2-diol	TA-Luft Klasse 5.2.5					
15.2 Chemical safety assessment							
A chemical safety assessment has be	een performed.						
16. OTHER INFORMATION							
Label DSD							
Labels							
Not classified as dangerous according to the criteria of Regulation (EC) No 1272/2008							
Full text of R Phrases referred to under sections 2 and 3							
Full text of S Phrases referred to under sections 2 and 3							
Additional recommendations							
Remark							
Label CLP							
Pictograms							
Full text of H-Statements referred to under sections 2 and 3							
Source of key data used to compile the data sheet							
Supplier information							
Modifications from last revision							
The Safety Data Sheets have been revised throughout in accordance with EC Regulation 1907/2006 and amendments							
Date: 30/06/18							