

SAFETY DATA SHEET

CLARNATE® PC A1070, A1073, A1077, A1100, A1105, A1107, 1105FL, A1150, A1155, HL6157, A1220, A1225, A1227

Wanhua Chemical Group Co., LTD

Version No: 1.1 Safety Data Sheet Safety Data Sheet - Authored according to GB/T16483(2008) and GB/T17519(2013) Chemwatch Hazard Alert Code: 0

Issue Date: 05/06/2019 Print Date: 05/06/2019 L.GHS.CHN.EN

SECTION 1 IDENTIFICATION OF THE SUBSTANCE / MIXTURE AND OF THE COMPANY / UNDERTAKING

Product Identifier

Product name	CLARNATE® PC A1070, A1073, A1077, A1100, A1105, A1107, 1105FL, A1150, A1155, HL6157, A1220, A1225, A1227
Chemical Name	bisphenol A/ phosgene polymer
Chemical English Name	bisphenol A/ phosgene polymer
Synonyms	polycarbonate
Other means of identification	Not Available

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

Relevant identified uses	For blending, extrusion or injection moulding
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Details of the supplier of the safety data sheet

Registered company name	Wanhua Chemical Group Co., LTD
Address	No.17, Tianshan Rd, YEDA, Yantai China
Telephone	+86 535-8202188 +86 574-86756457
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Website	www.whchem.com
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Emergency telephone number

Association / Organisation	Wanhua Chemical Group Co., LTD
Emergency telephone numbers	+86 532-83889090
Other emergency telephone numbers	+86 535-8203123

SECTION 2 HAZARDS IDENTIFICATION

Classification of the substance or mixture

SUMMARY OF HAZARD IN AN EMERGENCY SITUATION

Solid.	
Does not mix with water.	
Sinks in water.Combustible.	

Classification ^[1] Not Applicable

Label elements

Hazard pictogram(s)	Not Applicable
SIGNAL WORD	NOT APPLICABLE

Hazard statement(s)

Not Applicable

Precautionary statement(s) Prevention

Not Applicable

Precautionary statement(s) Response

Not Applicable

CLARNATE® PC A1070, A1073, A1077, A1100, A1105, A1107, 1105FL, A1150, A1155, HL6157, A1220, A1225, A1227

Precautionary statement(s) Storage

Not Applicable

Precautionary statement(s) Disposal

Not Applicable

Physical and Chemical Hazard

Solid. Does not mix with water. Sinks in water.Combustible. Toxic smoke/fumes in a fire.

Health Hazards

Inhaled	The material is not thought to produce adverse health effects or irritation of the respiratory tract (as classified by EC Directives using animal models). Nevertheless, good hygiene practice requires that exposure be kept to a minimum and that suitable control measures be used in an occupational setting.
Ingestion	The material has NOT been classified by EC Directives or other classification systems as "harmful by ingestion". This is because of the lack of corroborating animal or human evidence. The material may still be damaging to the health of the individual, following ingestion, especially where pre-existing organ (e.g liver, kidney) damage is evident. Present definitions of harmful or toxic substances are generally based on doses producing mortality rather than those producing morbidity (disease, ill-health). Gastrointestinal tract discomfort may produce nausea and vomiting. In an occupational setting however, ingestion of insignificant quantities is not thought to be cause for concern.
Skin Contact	The material is not thought to produce adverse health effects or skin irritation following contact (as classified by EC Directives using animal models). Nevertheless, good hygiene practice requires that exposure be kept to a minimum and that suitable gloves be used in an occupational setting. 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.
Eye	Although the material is not thought to be an irritant (as classified by EC Directives), direct contact with the eye may cause transient discomfort characterised by tearing or conjunctival redness (as with windburn). Slight abrasive damage may also result. The material may produce foreign body irritation in certain individuals.
Chronic	Long-term exposure to the product is not thought to produce chronic effects adverse to health (as classified by EC Directives using animal models); nevertheless exposure by all routes should be minimised as a matter of course. On the basis, primarily, of animal experiments, concern has been expressed by at least one classification body 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.

Environmental Hazards

See Section 12

Other hazards

Limited evidence of a carcinogenic effect*. May possibly affect fertility*.

SECTION 3 COMPOSITION / INFORMATION ON INGREDIENTS

Substances

See section below for composition of Mixtures

Mixtures

CAS No	%[weight]	Name
25971-63-5	>99	bisphenol A/ phosgene polymer

SECTION 4 FIRST AID MEASURES

Description of first aid measures

Eye Contact	If this product comes in contact with eyes: • Wash out immediately with water. • If irritation continues, seek medical attention. • Removal of contact lenses after an eye injury should only be undertaken by skilled personnel.
Skin Contact	If skin or hair contact occurs: Flush skin and hair with running water (and soap if available). Seek medical attention in event of irritation.
Inhalation	 If dust is inhaled, remove from contaminated area. Encourage patient to blow nose to ensure clear passage of breathing. If irritation or discomfort persists seek medical attention.
Ingestion	 Immediately give a glass of water. First aid is not generally required. If in doubt, contact a Poisons Information Centre or a doctor.

Advise for rescue team (PPE requirement for rescue personnel)

Indication of any immediate medical attention and special treatment needed Treat symptomatically.

Page 3 of 10

CLARNATE® PC A1070, A1073, A1077, A1100, A1105, A1107, 1105FL, A1150, A1155, HL6157, A1220, A1225, A1227

Extinguishing media

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

Special hazards arising from the substrate or mixture

Fire Incompatibility	Avoid contamination with oxidising agents i.e. nitrates, oxidising acids, chlorine bleaches, pool chlorine etc. as ignition may result
Advice for firefighters	
Fire Fighting	 Alert Fire Brigade and tell them location and nature of hazard. Wear breathing apparatus plus protective gloves. Prevent, by any means available, spillage from entering drains or water courses. Use water delivered as a fine spray to control fire and cool adjacent area. DO NOT approach containers suspected to be hot. Cool fire exposed containers with water spray from a protected location. If safe to do so, remove containers from path of fire. Equipment should be thoroughly decontaminated after use.
Fire/Explosion Hazard	 Combustible solid which burns but propagates flame with difficulty; it is estimated that most organic dusts are combustible (circa 70%) - according to the dircumstances under which the combustion process occurs, such materials may cause fires and / or dust explosions. Organic powders when finely divided over a range of concentrations regardless of particulate size or shape and suspended in air or some other oxidizing medium may form explosive dust-air mixtures and result in a fire or dust explosion (including secondary explosions). Avoid generating dust, particularly doubd of dust in a comfined or unventilated space as dusts may form an explosive imixture with air, and any source of ignition, i.e. flame or spark, will cause fire or explosion. Dust clouds generated by the fine grinding of the solid are a particular hazard; accumulations of fine dust (420 micron or less) may burn rapidy and fiercely if ignited - particles exceeding this limit will generally not form flammable dust clouds; concertation itini (LEL) and upper explosive limit (LEL) are applicable to dust clouds but only the LEL is of practical use; - this is because of the inherent difficulty of achieving homogeneous dust clouds at the inhimmum amyound of energy required to ignite dust clouds. ME(2) Will be lower than the pure dust in air mixtures will be lower than the pure dust in air mixture. The Lower Explosive Limit (LEL) of the vapour/dust mixture may be formed with or mixture source or explosion (may release of large quantities of gaseous products; this in turn creates a subsequent pressure frise of thus wapors/mists or dusts. A dust explosion may release of large quantities of gaseous products; this in turn creates a subsequent pressure frise of dust clouds, and orten initiate a much larger secondary explosion enters the surrounding area, it will disturb any settled dust layers, forming a second dust cloud, and orten initiate a much larger secondary explosion netweres and layer requires and

SECTION 6 ACCIDENTAL RELEASE MEASURES

Personal precautions, protective equipment and emergency procedures

See section 8

Measures for Preventing Secondary Contamination

Refer to section above

Environmental precautions

See section 12

Methods and material for containment and cleaning up

Minor Spills	 Clean up all spills immediately. Avoid contact with skin and eyes. Wear impervious gloves and safety glasses. Use dry clean up procedures and avoid generating dust. Vacuum up (consider explosion-proof machines designed to be grounded during storage and use). Do NOT use air hoses for cleaning Place spilled material in clean, dry, sealable, labelled container.

CLARNATE® PC A1070, A1073, A1077, A1100, A1105, A1107, 1105FL, A1150, A1155, HL6157, A1220, A1225, A1227

Major Spills	 Clear area of personnel and move upwind. Alert Fire Brigade and tell them location and nature of hazard. Control personal contact with the substance, by using protective equipment and dust respirator. Prevent spillage from entering drains, sewers or water courses. Avoid generating dust. Sweep, shovel up. Recover product wherever possible. Put residues in labelled plastic bags or other containers for disposal. If contamination of drains or waterways occurs, advise emergency services. 	
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Personal Protective Equipment advice is contained in Section 8 of the SDS.

SECTION 7 HANDLING AND STORAGE

Safe handling	 Limit all unnecessary personal contact. Wear protective clothing when risk of exposure occurs. Use in a well-ventilated area. Avoid contact with incompatible materials. When handling, DD NOT eat, drink or smoke. Keep containers securely sealed when not in use. Avoid physical damage to containers. Always wash hands with scap and water after handling. Work clothes should be laundered separately. Use god occupational work practice. Observe manufacturer's storage and handling recommendations contained within this SDS. Atmosphere should be regularly checked against established exposure standards to ensure safe working conditions are maintained. Organic powders when finely divided over a range of concentrations regardless of particulate size or shape and suspended in air or some other oxidizing medium may form explosive dust-air mixtures and result in a fire or dust explosion (including secondary explosions) Minimise airborne dust and eliminate all ignition sources. Keep away from heat, hot surfaces, sparks, and flame. Establish good housekeeping practices. Remove dust accumulations on a regular basis by vacuuming or gentle sweeping to avoid creating dust clouds. Use continuous suction at points of dust generation to capture and minimise the accumulation of dusts. Particular attention should be given to overhead and hidden horizontal surfaces to minimise the probability of a "secondary" explosion. According to NFPA Standard 654, dust layers 1/32 in (0.8 mm) thick can be sufficient to warrant timmediate cleaning of the area. Do not use air hoses for cleaning. Minimise drives should be used. Control sources of static electricity. Dusts or their packages may accumulate static charges, and static discharge can be a source of ignition. Solids handling systems must be designed in accordance with applicable standards (e.g., NFPA including
	 cannot be grounded, and antistatic bags do not completely protect against development of static charges. Empty containers may contain residual dust which has the potential to accumulate following settling. Such dusts may explode in the presence of an appropriate ignition source. Do NOT cut, drill, grind or weld such containers. In addition ensure such activity is not performed near full, partially empty or empty containers without appropriate workplace safety authorisation or permit.
Other information	 Store in original containers. Keep containers securely sealed. Store in a cool, dry area protected from environmental extremes. Store away from incompatible materials and foodstuff containers. Protect containers against physical damage and check regularly for leaks. Observe manufacturer's storage and handling recommendations contained within this SDS. For major quantities: Consider storage in bunded areas - ensure storage areas are isolated from sources of community water (including stormwater, ground water, lakes and streams). Ensure that accidental discharge to air or water is the subject of a contingency disaster management plan; this may require consultation with local authorities.

Conditions for safe storage, including any incompatibilities

Suitable container	 Lined metal can, lined metal pail/ can. Plastic pail. Polyliner drum. Packing as recommended by manufacturer. Check all containers are clearly labelled and free from leaks.
Storage incompatibility	Avoid contamination of water, foodstuffs, feed or seed. ► Avoid reaction with oxidising agents

SECTION 8 EXPOSURE CONTROLS / PERSONAL PROTECTION

Control parameters

OCCUPATIONAL EXPOSURE LIMITS (OEL)

INGREDIENT DATA

Not Available

EMERGENCY LIMITS

Ingredient

TEEL-1

TEEL-2

TEEL-3

CLARNATE® PC A1070, A1073, A1077, A1100, A1105, A1107, 1105FL, A1150, A1155, HL6157, A1220, A1225, A1227

A1077, A1100, A1105, A1107, 1105FL, A1150, A1155, HL6157, A1220, A1225, A1227	ot Available	Not Available	Not Available	Not Available
	iginal IDLH ot Available		Revised IDLH Not Available	

MATERIAL DATA

These "dusts" have little adverse effect on the lungs and do not produce toxic effects or organic disease. Although there is no dust which does not evoke some cellular response at sufficiently high concentrations, the cellular response caused by P.N.O.C.s has the following characteristics:

· the architecture of the air spaces remain intact,

scar tissue (collagen) is not synthesised to any degree,

tissue reaction is potentially reversible.

Extensive concentrations of P.N.O.C.s may:

seriously reduce visibility

cause unpleasant deposits in the eyes, ears and nasal passages,

contribute to skin or mucous membrane injury by chemical or mechanical action, per se, or by the rigorous skin cleansing procedures necessary for their removal. [ACGIH] This limit does not apply:

to brief exposures to higher concentrations

nor does it apply to those substances that may cause physiological impairment at lower concentrations but for which a TLV has as yet to be determined.

This exposure standard applies to particles which

- are insoluble or poorly soluble* in water or, preferably, in aqueous lung fluid (if data is available) and
- have a low toxicity (i.e.. are not cytotoxic, genotoxic, or otherwise chemically reactive with lung tissue, and do not emit ionizing radiation, cause immune sensitization, or cause toxic effects other than by inflammation or by a mechanism of lung overload)

Exposure controls

Appropriate engineering	 Engineering controls are used to remove a hazard or place a barrier between the worker and the highly effective in protecting workers and will typically be independent of worker interactions to prote basic types of engineering controls are: Process controls which involve changing the way a job activity or process is done to reduce the ri Enclosure and/or isolation of emission source which keeps a selected hazard "physically" away free "removes" air in the work environment. Ventilation can remove or dilute an air contaminant if design the particular process and chemical or contaminant in use. Employers may need to use multiple types of controls to prevent employee overexposure. Local exhaust ventilation is required where solids are handled as powders or crystals; even will be powdered by mutual friction. Exhaust ventilation should be designed to prevent accumulation and recirculation of particula. If in spite of local exhaust an adverse concentration of the substance in air could occur, respinging the consist of: (a): particle dust respirators, if necessary, combined with an absorption cartridge; (b): filter respirators with absorption cartridge or canister of the right type; (c): fresh-air hoods or masks Build-up of electrostatic charge on the dust particle, may be prevented by bonding and grour Powder handling equipment such as dust collectors, dryers and mills may require additional Air contaminants generated in the workplace possess varying "escape" velocities which, in turn, required to efficiently remove the contaminant. 	ovide this high level of protectionsk. om the worker and ventilation gned properly. The design of a when particulates are relatively tes in the workplace. ratory protection should be co inding. protection measures such as	on. that strategically "adds" and ventilation system must y large, a certain proportion nsidered. Such protection explosion venting.
controls	Type of Contaminant:		Air Speed:
	direct spray, spray painting in shallow booths, drum filling, conveyer loading, crusher dusts, ga generation into zone of rapid air motion)	1-2.5 m/s (200-500 f/min.)	
	grinding, abrasive blasting, tumbling, high speed wheel generated dusts (released at high initia high rapid air motion).	I velocity into zone of very	2.5-10 m/s (500-2000 f/min.)
	Within each range the appropriate value depends on:		
	Lower end of the range	Upper end of the range	
	1: Room air currents minimal or favourable to capture	1: Disturbing room air curre	ents
	2: Contaminants of low toxicity or of nuisance value only	2: Contaminants of high tox	icity
	3: Intermittent, low production.	3: High production, heavy us	se
	4: Large hood or large air mass in motion	4: Small hood-local control	only
	Simple theory shows that air velocity falls rapidly with distance away from the opening of a simple square of distance from the extraction point (in simple cases). Therefore the air speed at the extra reference to distance from the contaminating source. The air velocity at the extraction fan, for exar for extraction of crusher dusts generated 2 metres distant from the extraction point. Other mecha within the extraction apparatus, make it essential that theoretical air velocities are multiplied by fac or used.	action point should be adjuste mple, should be a minimum of nical considerations, producin	d, accordingly, after 4-10 m/s (800-2000 f/min) g performance deficits
Personal protection			
	 Safety glasses with side shields Chemical goggles. Control longer may append appendix to the soft control longer may show the and appendix to the soft control longer may show the soft control	irritanta Aurrittan paliau dagu	ment describing the uppering

Contact

Eye and face protection

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

Page 6 of 10

CLARNATE® PC A1070, A1073, A1077, A1100, A1105, A1107, 1105FL, A1150, A1155, HL6157, A1220, A1225, A1227

	 should be removed at the first signs of eye redness or irritation - lens should be removed in a clean environment only after workers have washed hands thoroughly. [CDC NIOSH Current Intelligence Bulletin 59], [AS/NZS 1336 or national equivalent]
Skin protection	See Hand protection below
Hands/feet protection	The selection of suitable gloves does not only depend on the material, but also on further marks of quality which vary from manufacturer to manufacturer. Where the chemical is a proparation of several substances, the resistance of the glove material can not be calculated in advance and has therefore to be checked prior to the application. The exact 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 died throrughit, Application of a non-perfumed motisturies is recommended. Suitability and durability of glove type is dependent on usage. Important factors in the selection of gloves include: the requercy and duration of contact, duration of outation of contact, duration of contact, duration of contact and stance of glove material, a glove thickness and divertity. Select gloves tested to a relevant standard (eq. Europe EN 374, LSK F739, ASAZS 2161.1 or national equivalent). When notivinged of frequently repeated contact may occur, a glove with a protection class of 5 or higher (breakthrough time greater than 240 minutes according to EN 374, ASNZS 2161.10.1 or national equivalent) is recommended. Some glove polymer types are less affected by movement and this should be taken into account when considering gloves for long-term use. Contaminated gloves should be replaced. As defined in ASTM F739-66 in any application, gloves and the sale was affected by movement and this should be taken into account when considering gloves for long-term use. The context sequence than 9400 min the source and have a set of a specific chemical, as the permeation efficiency of the glove with a thickness typically greater than 0.35 mm, are recommended. Itshould be taken into account when considering on a substance of the most. Note: Contaminated degrades is not necessarily a good predictor of glove r
Body protection	See Other protection below
Other protection	No special equipment needed when handling small quantities. OTHERWISE: > Overalls. > Barrier cream. > Eyewash unit.

Respiratory protection

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

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

* - Negative pressure demand ** - Continuous flow

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

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

- The decision to use respiratory protection should be based on professional judgment that takes into account toxicity information, exposure measurement data, and frequency and likelihood of the worker's exposure ensure users are not subject to high thermal loads which may result in heat stress or distress due to personal protective equipment (powered, positive flow, full face apparatus may be an option).
- Published occupational exposure limits, where they exist, will assist in determining the adequacy of the selected respiratory protection. These may be government mandated or vendor recommended.
- Certified respirators will be useful for protecting workers from inhalation of particulates when properly selected and fit tested as part of a complete respiratory protection program.
- Use approved positive flow mask if significant quantities of dust becomes airborne.

Try to avoid creating dust conditions.

CLARNATE® PC A1070, A1073, A1077, A1100, A1105, A1107, 1105FL, A1150, A1155, HL6157, A1220, A1225, A1227

Information on basic physical and chemical properties

Appearance	Not Available		
Physical state	Solid	Relative density (Water = 1)	1.18-1.2
Odour	Not Available	Partition coefficient n-octanol / water	Not Available
Odour threshold	Not Available	Auto-ignition temperature (°C)	Not Available
pH (as supplied)	Not Applicable	Decomposition temperature	>380
Melting point / freezing point (°C)	Not Available	Viscosity (cSt)	Not Available
Initial boiling point and boiling range (°C)	Not Applicable	Molecular weight (g/mol)	Not Available
Flash point (°C)	Not Available	Taste	Not Available
Evaporation rate	Not Available	Explosive properties	Not Available
Flammability	Not Available	Oxidising properties	Not Available
Upper Explosive Limit (%)	Not Available	Surface Tension (dyn/cm or mN/m)	Not Applicable
Lower Explosive Limit (%)	Not Available	Volatile Component (%vol)	Not Available
Vapour pressure (kPa)	Not Applicable	Gas group	Not Available
Solubility in water	Immiscible	pH as a solution (1%)	Not Available
Vapour density (Air = 1)	Not Applicable	VOC g/L	Not Available

SECTION 10 STABILITY AND REACTIVITY

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

SECTION 11 TOXICOLOGICAL INFORMATION

POLYMER

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bisphenol A/ phosgene polymer	TOXICITY	IRRITATION
bisplienoi A/ phosgene polymer	Not Available	Not Available
	I	
Legend:	 Value obtained from Europe ECHA Registered Substances - Acute toxicity data extracted from RTECS - Register of Toxic Effect of chemical Substance 	
	v	
BISPHENOL A/ PHOSGENE POLYMER	No significant acute toxicological data identified in literature search.	
	The chemical structure of hydroxylated diphenylalkanes or bisphenols consi of endocrine disruptors that mimic oestrogens is widely used in industry, part	

CLARNATE® PC A1070, A1073, A1077, A1100, A1105, A1107, 1105FL, A1150, A1155, HL6157, A1220, A1225, A1227 & BISPHENOL A/ PHOSGENE

Bisphenols promoted cell proliferation and increased the synthesis and secretion of cell type-specific proteins. When ranked by proliferative potency, the longer the alkyl substituent at the bridging carbon, the lower the concentration needed for maximal cell yield; the most active compound contained two propyl chains at the bridging carbon. Bisphenols with two hydroxyl groups in the para position and an angular configuration are suitable for appropriate hydrogen bonding to the acceptor site of the oestrogen receptor.

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

Legend: 🗙 –

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

Page 8 of 10

CLARNATE® PC A1070, A1073, A1077, A1100, A1105, A1107, 1105FL, A1150, A1155, HL6157, A1220, A1225, A1227

SECTION 12 ECOLOGICAL INFORMATION

CLARNATE® DC 44070 44073					
CLARNATE® PC A1070, A1073,	ENDPOINT	TEST DURATION (HR)	SPECIES	VALUE	SOURCE
A1077, A1100, A1105, A1107, 1105FL, A1150, A1155, HL6157,	Not Available	Not Available	Not Available	Not Available	Not Available
A1220, A1225, A1227					
	ENDPOINT	TEST DURATION (HR)	SPECIES	VALUE	SOURCE
bisphenol A/ phosgene polymer	Not Available	Not Available	Not Available	Not Available	Not Available
		Not Available	Not Available	Not Available	Not Available
Legend:	(QSAR) - Aquatic To	CLID Toxicity Data 2. Europe ECHA Registe. xicity Data (Estimated) 4. US EPA, Ecotox c ration Data 7. METI (Japan) - Bioconcentrat	atabase - Aquatic Toxicity Dai	· ·	-
Substance does not meet the criteria As an environmental contaminant, bit the soil of only 1-10 days, its ubiquity time. Studies also indicate that it can of during treatment at municipal water tr Ecotoxicity: Fish LC50 (96 h): 4.6 mg/l (freshwate Fresh water invertebrates EC50 (96 Freshwater algae (96 h): 2.73 mg/l Marine water invertebrate EC50 (96 Freshwater algae (96 h): 1.1 mg/l Marine water plant EC50 (7 d): 20 mg In general, studies have shown that b Among freshwater organisms, fish ap	biodegradable g/l ogues, can be released fr for PBT or vPvB accordi sphenol A interferes with makes it an important po currently be found in mun reatment plants. er fish); 11 mg/l (saltwate h): 10.2 mg/l: NOEC 0.07 h): 1.1 mg/l; NOEC 0.17 yl: NOEC 7.8 mg/l bisphenol A can affect gro opear to be the most sen	o ,	XIII blants associated with the bac nitial assessment shows that a ted in the United States found 44 d); 0.064 mg/l (saltwater fi ic organisms. effects in fish, aquatic invertel	at low levels, bisphenol A c that 91-98% of bisphenol A sh 164 d) brates, amphibians and rep	an harm fish and organisms ov may be removed from water billes has been reported at
A 2009 review of the biological impact amphibians concluded that bisphenol A large 2010 study of two rivers in Ca while females made up only 55 per ce Although abundant data are available mutagenicity, and oestrogenic activity D. magna (48-h EC50 was 10 mg/l) natural estrogen 17 beta-oestradiol in bestrogenic activity as well as BPA. S nydroxydiphenyl/sulfone) and bis(4-h) Biodegradation is a major mechanism BPA-degrading bacteria have been is methyl group or the quaternary carbo piodegrade all bisphenols that have ai which has no substituent at the bridgi conditions. From this evidence, it was	I A has been shown to af inada found that areas co ent in uncontaminated are or the toxicity of bispher v using the Daphtoxkit (Cr according to the current the yeast screen, while i Some of the BPs showed ydroxyphenyl)sulfide) sho n for eliminating various e solated from enrichments in in the BPA molecule. J t least one methyl or meth ing carbon, is unlikely to l s clear that a specific method	nol-A (2,2-bis (4-hydroxydiphenyl)propane;(B reasel Ltd.), the umu test system, and the yea U.S. EPA acute toxicity evaluation standard, no mutagenicity was observed. All seven BP- l considerably higher oestrogenic activity thar wed oestrogenic activity. environmental pollutants. Studies on the biod of sludge from wastewater treatment plants ludging from these features of the biodegrada hylene group bonded at the carbon atom betw be metabolised by such a mechanism. New chanism for biodegradation of BPF does exist	, to impair development in crus uding bisphenol A showed fer PA) A variety of BPs were exa ist two-hybrid system, respecti and it was weakly oestrogenic is tested here showed moderal BPA, and others exhibited mu- egradation of bisphenols have The first step in the biodegra- ation mechanisms, it is possibl ween the two phenol groups. H ertheless BPF is readily degra- it in the natural ecosystem,	staceans and amphibians a nales made up 85 per cent with 5 orders of magnitude to slight acute toxicit, no ich lower activity. Bisphenc mainly focused on bisphe dation of BPA is the hydrox e that the same mechanisi owever, bisphenol F ([bis(4 ded by river water microorg	and to induce genetic aberration of the population of a certain fis y against Daphnia magna, PA. BPA was moderately toxic to lower activity than that of the mutagenicity, and weak of S (bis(4- nol A. A number of ylation of the carbon atom of a m used for BPA is used to -hydroxyphenyl)methane; BPF janisms under aerobic
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Other adverse effects

No data available

SECTION 13 DISPOSAL CONSIDERATIONS

Version No: 1.1	CLARNATE® PC A1070, A1073, A1077, A1100, A1105, A1107, 1105FL, A1150, A1155, HL6157, A1220, A1225, A1227	Print

Waste chemicals:	 DO NOT allow wash water from 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 sewer may be subject to local laws and regulations and these should be considered first. Where in doubt contact the responsible authority.
Contaminated packing materials:	Refer to section above
Precautions for Transport:	Refer to section above

SECTION 14 TRANSPORT INFORMATION

Labels Required

Marine Pollutant

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

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

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

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

NO

Not Applicable

Precautions for Transport

Suitable Containers

See section 7

SECTION 15 REGULATORY INFORMATION

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

BISPHENOL A/ PHOSGENE POLYMER(25971-63-5) IS FOUND ON THE FOLLOWING REGULATORY LISTS

China Inventory of Existing Chemical Substances

National Inventory Status

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

SECTION 16 OTHER INFORMATION

Revision Date	05/06/2019
Initial Date	05/06/2019

Other information

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

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

Definitions and abbreviations

PC – TWA: Permissible Concentration-Time Weighted Average PC – STEL: Permissible Concentration-Short Term Exposure Limit

IARC: International Agency for Research on Cancer

CLARNATE® PC A1070, A1073, A1077, A1100, A1105, A1107, 1105FL, A1150, A1155, HL6157, A1220, A1225, A1227

ACGIH: American Conference of Governmental Industrial Hygienists STEL: Short Term Exposure Limit TEEL: Temporary Emergency Exposure ${\sf Limit}_{\circ}$ IDLH: Immediately Dangerous to Life or Health Concentrations OSF: Odour Safety Factor NOAEL :No Observed Adverse Effect Level LOAEL: Lowest Observed Adverse Effect Level TLV: Threshold Limit Value LOD: Limit Of Detection OTV: Odour Threshold Value BCF: BioConcentration Factors

BEI: Biological Exposure Index

Disclaimer

The information in the SDS applies only for the specified product and does not include mixtures of this product with other substances and mixtures. The SDS provides product safety information for personnel trainned to use this product only.

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