# **CLH** report

# **Proposal for Harmonised Classification and Labelling**

Based on Regulation (EC) No 1272/2008 (CLP Regulation), Annex VI, Part 2

# International Chemical Identification: Tetramethylene dimethylacrylate

**EC Number: 218-218-1** 

**CAS Number: 2082-81-7** 

**Index Number:** Not available

Contact details for dossier submitter: Finnish Competent Authority

**Finnish Safety and Chemicals** 

Agency (Tukes)

**Finland** 

Version number: 1.0 Date: 13<sup>th</sup> July 2020

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# 1 IDENTITY OF THE SUBSTANCE

# 1.1 Name and other identifiers of the substance

Table 1: Substance identity and information related to molecular and structural formula of the substance

in the least to home the test of the least to the least t	ylprop-2-enoyloxy)butyl 2-methylprop-2-enoate nylprop-2-enoyl)oxy]butyl 2-methylprop-2-
4-[(2-incu	nylprop-2-enoyl)oxy]butyl 2-methylprop-2-
Butadiene	dimethacrylate
Butane-1,	4-diyl bis(2-methylacrylate)
Tetrameth	ylene dimethacrylate
Other names (usual name, trade name, abbreviation) 1,4-butane	ediol dimethacrylate
BDMA; 1	,4-BDDMA
ISO common name (if available and appropriate)	
EC number (if available and appropriate) 218-218-1	
EC name (if available and appropriate)  Tetrameth	ylene dimethacrylate
CAS number (if available) 2082-81-7	
Other identity code (if available)	
Molecular formula C12H18O	4
Structural formula	000000000000000000000000000000000000000
SMILES notation (if available) CC(=C)C(	(=0)OCCCCOC(=0)C(C)=C
Molecular weight or molecular weight range 226.27 g/r	nol
initial indication of optical activity and typical ratio of	table (the structure of the substance does not te stereo-isomerism)
Description of the manufacturing process and identity of the source (for UVCB substances only)  Not application of the manufacturing process and identity of the source (for UVCB substances only)	cable (the substance is not an UVCB)
Degree of purity (%) (if relevant for the entry in Annex VI)	% (w/w)

# 1.2 Composition of the substance

# **Table 2: Constituents (non-confidential information)**

Constituent	Concentration range (% w/w minimum and maximum in multiconstituent substances)	Current CLH in	Current self-
(Name and numerical		Annex VI Table 3.1	classification and
identifier)		(CLP)	labelling (CLP)
Tetramethylene dimethacrylate (CAS 2082-81-7)	95-99.63 % (w/w)	No entry in Annex VI	Eye Irrit. 2; H319 Skin Irrit. 2; H315 STOT SE 3; H335 Skin Sens. 1B; H317 Skin Sens. 1; H317

# Table 3: Impurities (non-confidential information) if relevant for the classification of the substance

No impurities relevant for classification.

# Table 4: Additives (non-confidential information) if relevant for the classification of the substance

No additives relevant for classification.

# 2 PROPOSED HARMONISED CLASSIFICATION AND LABELLING

# 2.1 Proposed harmonised classification and labelling according to the CLP criteria

Table 5:

					Classif	ication		Labelling			
	Index No	International Chemical Identification	EC No	CAS No	Hazard Class and Category Code(s)	Hazard statement Code(s)	Pictogram, Signal Word Code(s)	Hazard statement Code(s)	Suppl. Hazard statement Code(s)	Specific Conc. Limits, M-factors	Notes
Current Annex VI entry					No curren	t entry in Annex	VI				
Dossier submitters proposal	-	Tetramethylene dimethacrylate	218-218-1	2082-81-7	Skin Sens. 1B	H317	GHS07 Wng	H317	-	-	-
Resulting Annex VI entry if agreed by RAC and COM	-	Tetramethylene dimethacrylate	218-218-1	2082-81-7	Skin Sens. 1B	Н317	GHS07 Wng	Н317	-	-	-

Table 6: Reason for not proposing harmonised classification and status under public consultation

Hazard class	Reason for no classification	Within the scope of public consultation
Explosives	Hazard class not assessed in this dossier	No
Flammable gases (including chemically unstable gases)	Hazard class not assessed in this dossier	No
Oxidising gases	Hazard class not assessed in this dossier	No
Gases under pressure	Hazard class not assessed in this dossier	No
Flammable liquids	Hazard class not assessed in this dossier	No
Flammable solids	Hazard class not assessed in this dossier	No
Self-reactive substances	Hazard class not assessed in this dossier	No
Pyrophoric liquids	Hazard class not assessed in this dossier	No
Pyrophoric solids	Hazard class not assessed in this dossier	No
Self-heating substances	Hazard class not assessed in this dossier	No
Substances which in contact with water emit flammable gases	Hazard class not assessed in this dossier	No
Oxidising liquids	Hazard class not assessed in this dossier	No
Oxidising solids	Hazard class not assessed in this dossier	No
Organic peroxides	Hazard class not assessed in this dossier	No
Corrosive to metals	Hazard class not assessed in this dossier	No
Acute toxicity via oral route	Hazard class not assessed in this dossier	No
Acute toxicity via dermal route	Hazard class not assessed in this dossier	No
Acute toxicity via inhalation route	Hazard class not assessed in this dossier	No
Skin corrosion/irritation	Hazard class not assessed in this dossier	No
Serious eye damage/eye irritation	Hazard class not assessed in this dossier	No
Respiratory sensitisation	Hazard class not assessed in this dossier	No
Skin sensitisation	Harmonised classification proposed	Yes
Germ cell mutagenicity	Hazard class not assessed in this dossier	No
Carcinogenicity	Hazard class not assessed in this dossier	No
Reproductive toxicity	Hazard class not assessed in this dossier	No
Specific target organ toxicity- single exposure	Hazard class not assessed in this dossier	No
Specific target organ toxicity- repeated exposure	Hazard class not assessed in this dossier	No
Aspiration hazard	Hazard class not assessed in this dossier	No
Hazardous to the aquatic environment	Hazard class not assessed in this dossier	No
Hazardous to the ozone layer	Hazard class not assessed in this dossier	No

# 3 HISTORY OF THE PREVIOUS CLASSIFICATION AND LABELLING

For tetramethylene dimethacrylate there is no harmonized classification available, as the substance is not listed in Annex VI to the Regulation (EC) No 1272/2008 (CLP Regulation).

# 4 JUSTIFICATION THAT ACTION IS NEEDED AT COMMUNITY LEVEL

Justification that action is needed at Community level is required.

Reason for a need for action at Community level:

Differences in self-classification in the C&L Inventory

Disagreement by DS with current self-classification

# Further detail on need of action at Community level

According to Article 36(3) of the CLP Regulation, for a substance that fulfills the criteria for other hazard classes or differentiations than those of CMR or respiratory sensitisation (Category 1) and the substance is not an active substance under the Plant Protection Product Directive (PPPD) and Biocidal Product Directive (BPD), a harmonized classification and labelling proposal can be submitted if a justification is provided demonstrating the need for such action at community level. There is no entry in Annex VI to the CLP Regulation for tetramethylene dimethacrylate and there have been no previous classification and labelling discussions of the substance.

As of June 2020, the C&L Inventory contains in total 159 notifications for tetramethylene dimethacrylate with respect to skin sensitisation:

- Skin Sens. 1 (64 notifications)
- Skin Sens. 1B (95 notifications)

Furthermore, 110 notifiers did not classify the substance for skin sensitisation at all. None of the notifiers has classified the substance as Skin Sens. 1A.

Differences in self-classification between different notifiers in the C&L Inventory have been discovered, and the dossier submitter (DS) disagrees with the self-classifications Skin Sens. 1 and no classification proposed by the notifiers. Tetramethylene dimethacrylate is registered under REACH, and it is manufactured and/or imported in the European Economic Area in 1 000-10 000 tonnes per year. The widespread use of the substance supports action at community level: exposure to tetramethylene dimethacrylate is anticipated under circumstances of professional, industrial and consumer use, mainly via dermal route. Workers may be in direct contact with formulated products containing the substance during mixing (including by hand) or blending, and the products may be used with rollers or brushes or via dipping or pouring. Tetramethylene dimethacrylate is one of the most commonly patch tested (meth)acrylates that quite often induces positive reactions in clinical patients. There are over 100 published cases with a positive patch test reaction to the substance, which exceeds the limit for high frequency of occurrence of skin sensitisation.

# 5 IDENTIFIED USES

Tetramethylene dimethacrylate is used in different coating products, fillers, putties, plasters, modelling clay, paints, adhesives and sealants. It is used by consumers, by professional workers (widespread uses), in formulation or re-packing, at industrial sites and in manufacturing.

### 6 DATA SOURCES

The REACH registration dossier of tetramethylene dimethacrylate was used as the main data source for this CLH report. The unpublished full study reports were made available to the DS by the lead registrant. In addition, open literature publications and patient exposure data from the Finnish Institute of Occupational Health were used.

# 7 PHYSICOCHEMICAL PROPERTIES

**Table 7: Summary of physicochemical properties** 

Property	Value	Reference	Comment (e.g. measured or estimated)
Physical state at 20°C and 101,3 kPa	Liquid	REACH registration dossier	Observed
Melting/freezing point	-23°C at 1025 hPa	Anonymous 2006a	Measured OECD TG 102/EU Method A.1; differential scanning calorimetry
Boiling point	Not determined	Anonymous 2006a	No boiling point was detected prior to polymerisation at ca. 211°C at 1025 hPa
			OECD TG 103/EU Method A.2; differential scanning calorimetry
Relative density	1.024 at 20°C	Anonymous 2007	Measured OECD TG 109/DIN 51757; oscillating densitimeter method
Vapour pressure	0.1 Pa at 20°C	Anonymous 2006b	Measured OECD TG 104/EU Method A.4; dynamic method
Surface tension	Not assessed	REACH registration dossier	Based on chemical structure, no surface activity is to be expected. The test substance is not used as detergent.
Water solubility	243 mg/L at 20°C	Anonymous 2001	Measured OECD TG 105; flask method
Partition coefficient n- octanol/water	Log Pow 3.10 at 20°C	Anonymous 2010	Measured OECD TG 117/EU Method A.8; HPLC method
Flash point	139°C at 1013.25 hPa	Anonymous 2008a	Measured EU Method A.9; Pensky-Martens closed-cup method
Flammability	Not flammable	REACH registration dossier	Study technically not feasible (the substance is a liquid).
Explosive properties	Not explosive	REACH registration dossier	There are no chemical groups associated with explosive properties present in the molecule.
Self-ignition temperature	290°C at 101 325 Pa	Anonymous 2009	Measured EU Method A.15/DIN 51794
Oxidising properties	Not oxidising	REACH registration dossier	Oxidising properties are not expected on the basis of chemical structure.
Granulometry	Not applicable	REACH registration dossier	The substance is a liquid and is marketed or used in a non solid or granular form.

Property	Value	Reference	Comment (e.g. measured or estimated)
Stability in organic solvents and identity of relevant degradation products	Not assessed	REACH registration dossier	Stability of the substance is not considered critical; the substance has no particular reactivity towards typical solvents and is not used in solution.
Dissociation constant	Not assessed	REACH registration dossier	The substance does not contain any ionic, dissociable structures.
Viscosity	5.29 mm <sup>2</sup> /s at 20°C and 3.13 mm <sup>2</sup> /s at 40°C	Anonymous 2008b	Measured OECD TG 114/DIN 51562; Micro-Ubbelohde viscometer

# 8 EVALUATION OF PHYSICAL HAZARDS

Not assessed in this dossier.

# 9 TOXICOKINETICS (ABSORPTION, METABOLISM, DISTRIBUTION AND ELIMINATION)

**Table 8: Summary table of toxicokinetic studies** 

Method	Remarks	Results	Reference
Basic toxicokinetics in vitro Non-guideline GLP Key study Reliability: 2  Test material: 1,4- butanediol dimethacrylate Purity: not specified	Duration: phase I 120 min (samples collected at 0, 2, 5, 15, 30, 60 and 120 min), phase II 5 min (samples collected at 0 and 5 min) Concentrations: 0.25 mM (phase I), 0.05, 0.1 and 5.0 mM (phase II)  Negative controls in the rat liver microsome experiments included incubations with heat-inactivated microsomes, no microsomes and no NADPH. Positive control: methyl methacrylate	Test substance was rapidly converted to methacrylic acid (MAA) in whole rat blood (phase I) and rat liver microsomes (phase II) with hydrolysis half-lives of 4.10 min (blood) and 4.46 min (liver microsomes).  Absence of NADPH made little or no difference in hydrolysis rates. Heat inactivation significantly reduced the rates, and absence of microsomes resulted in no hydrolysis. $V_{max} \text{ (in vitro)} = 129 \text{ nmol/min/mg}$ $V_{max} \text{ (in vivo)} = 160 \text{ mg/h/g liver}$ $K_m \text{ (in vitro)} = 83  \mu \text{m}$ $K_m \text{ (in vivo)} = 19 \text{ mg/L}$	Anonymous (2013a)
Dermal absorption study (in silico modelling) Non-guideline Non-GLP Key study Reliability: 2  Test material: 1,4-butanediol dimethacrylate Purity: not specified	The physicochemical parameters of M <sub>w</sub> , log P and saturated aqueous solubility were used in the evaluation of 56 methacrylate compounds using a human skin model. An output of predicted steady-state flux was calculated using the principles defined in the Potts and Guy prediction model (1992).	The predicted steady-state flux of 1,4-butanediol dimethacrylate is 2.895 $\mu g/cm^2/h$ , indicating low relative dermal absorption.	Anonymous (2013b)
Basic toxicokinetics in vitro and in vivo	A series of in vitro and in vivo studies were used to develop PBPK models that predict the	Hydrolysis of MMA by rat liver microsomes: V <sub>max</sub> = 445.8 nmol/min/mg	Anonymous (2002)

Method	Remarks	Results	Reference
(read-across) Non-guideline GLP: not specified Key study Reliability: 1  Test material: methyl methacrylate (MMA) Lot: 98/15 Purity: > 99%	metabolism and fate of a series of methacrylates  Administration: i.v. injection  Liver microsome studies: human, rat  Dermal absorption studies: rat skin (epidermal membrane:  Wistar rat, whole skin: Fischer 344 rat), human abdominal skin	$K_m$ = 164.3 μm Clearance = 98.8% removed from blood liver flow T50% (body elimination time for 50% parent ester) = 4.4 min $C_{max}$ = 14.7 mg/L of methacrylic acid (MAA) in blood $T_{max}$ = 1.7 min to peak MAA concentration in blood Hydrolysis of MMA by human liver microsomes: $V_{max}$ = 1721 nmol/min/mg $K_m$ = 4103 mM Clearance = 419 μL/min/mg The studies confirmed that alkylmethacrylate esters are rapidly hydrolysed to MAA by ubiquitous carboxylesterases. First pass (local) hydrolysis of the parent ester has been shown to be significant for all routes of exposure. In vivo measurements of rat liver indicated this organ has the greatest esterase activity. Similar measurements for skin microsomes indicated approximately 20-fold lower activity than for liver. However, this activity was substantial and capable of almost complete first-pass metabolism of the alkyl-methacrylates.	
Basic toxicokinetics in vitro (read-across) Non-guideline GLP Supporting study Reliability: 2  Test material: methyl methacrylate (MMA) Purity: not specified	Duration: phase I 120 min (samples collected at 0, 2, 5, 15, 30, 60 and 120 min), phase II 5 min (samples collected at 0 and 5 min)  Negative controls in the rat liver microsome experiments included incubations with heat-inactivated microsomes, no microsomes and no NADPH.	The test substance was rapidly converted to methacrylic acid (MAA) in whole rat blood (phase I) and rat liver microsomes (phase II) with hydrolysis half-lives of 63 min (blood) and 0.29 min (liver microsomes).   Absence of NADPH made little or no difference in hydrolysis rates. Heat inactivation significantly reduced the rates, and absence of microsomes resulted in no hydrolysis. $V_{max} \text{ (in vitro)} = 475 \text{ nmol/min/mg} $ $V_{max} \text{ (in vivo)} = 128 \text{ mg/h/g liver} $ $K_m \text{ (in vitro)} = 289  \mu \text{m} $ $K_m \text{ (in vivo)} = 29 \text{ mg/L} $	Anonymous (2013c)

# 9.1 Short summary and overall relevance of the provided toxicokinetic information on the proposed classification(s)

A few toxicokinetic studies are available for tetramethylene dimethacrylate and structurally similar methyl methacrylate (MMA) (Table 8Table 8). Tetramethylene dimethacrylate has a molecular weight of 226.27 g/mol and it is in liquid form at  $20^{\circ}$ C. Water solubility of the substance is 243 mg/L at  $20^{\circ}$ C, and the octanol-water partition coefficient (log  $P_{OW}$ ) is 3.10.

#### **Absorption**

In general, the physico-chemical properties (molecular weight, physical state, water solubility, lipophilicity) of tetramethylene dimethacrylate favour absorption from the gastrointestinal tract.

The vapour pressure of the substance is 0.1 Pa at 20°C. This falls well below the general cut-off value of 0.5 kPa, indicating very low volatility and hence poor availability for inhalation as a vapour (ECHA 2017a). Solid particles, however, may be available for absorption after inhalation of an aerosolized substance, although this does not seem likely considering the size of the molecule. There are no studies regarding absorption of tetramethylene dimethacrylate from the respiratory tract.

On the basis of the molecular weight, tetramethylene dimethacrylate has a relatively low ability to be absorbed through the skin. The water solubility of the substance is moderate (between 100 and 10 000 mg/L) for partitioning from the stratum corneum into the epidermis (ECHA 2017a). Furthermore, the log  $P_{OW}$  (3.10) favours penetration into the stratum corneum and hence absorption across the skin. The predicted steady-state flux is 2.895  $\mu$ g/cm²/h (Anonymous 2013b). The ester bonds of tetramethylene dimethacrylate may be hydrolysed in the skin, although to a much lesser extent than in the gastrointestinal tract due to the lower level of enzymes. The breakdown products may then be absorbed and enter the bloodstream. Proof of sensitisation after dermal contact indicates that a sufficient amount of the substance is taken up via the dermal route to induce a positive reaction in the skin (Anonymous 2014; see Section 10.7 for details).

In the absence of more specific data, absorption can be assumed to occur via oral and dermal routes. Tetramethylene dimethacrylate is unlikely to be absorbed via inhalation.

# **Distribution**

Since tetramethylene dimethacrylate is expected to undergo enzymatic hydrolysis especially in the gastrointestinal tract, the breakdown products (acid and alcohol moieties) are likely to be widely distributed due to their small size and solubility in aqueous media. The parent compound has a high permeability across lipid membranes (log  $P_{\rm OW}$  3.10), but the degradation products do not contain any lipophilic groups. The available data do not show accumulation in any organ or tissue, either. No target organs have been identified for tetramethylene dimethacrylate.

#### Metabolism

Ester hydrolysis is the primary step in the metabolism of methacrylate esters. In the case of diol dimethacrylate esters (such as tetramethylene dimethacrylate), one of the ester bonds is first hydrolyzed to produce the corresponding mono-ester. The second ester bond is then hydrolyzed by carboxylesterases to produce methacrylic acid (MAA) and the corresponding alcohol, 1,4-butanediol. Tetramethylene dimethacrylate was rapidly converted to methacrylic acid in a basic toxicokinetics study conducted to investigate the in vitro hydrolysis rates (Anonymous 2013a). The hydrolysis half-lives were 4.46 minutes in rat liver microsomes and 4.10 minutes in whole rat blood. Similar metabolic pattern has been identified for a structurally similar substance, methyl methacrylate; it was hydrolyzed at a high rate to methacrylic acid, with a half-life of 4.4 minutes based on a PBPK estimation (Anonymous 2002). In the same study, the metabolism rates for alkyl-methacrylates were approximately 20 times lower in skin microsomes than in liver microsomes.

The primary methacrylic metabolite of tetramethylene dimethacrylate, methacrylic acid, will predominantly be metabolized in the liver through the valine pathway and the citric acid cycle (Cosmetic Ingredient Review 2005). 1,4-butanediol is, in turn, known to be rapidly transformed to gamma-hydroxybutyric acid and subsequently to succinic semialdehyde (NTP 1996). The aldehyde is then converted to succinic acid, which is degraded via the citric acid cycle.

Methacrylates are likely to have low reactivity with glutathione in vitro compared to the corresponding acrylates (Tanii & Hashimoto 1982, McCarthy et al. 1994). This is presumably due to steric hindrance of a nucleophilic addition at the double bond by the alpha-methyl side group. Therefore, glutathione conjugation may only play a minor role in the metabolism of alkyl and multifunctional methacrylate esters, such as tetramethylene dimethacrylate.

#### Excretion

The parent compound tetramethylene dimethacrylate is not likely to be excreted as such due to the rapid hydrolysis of the ester bonds. The metabolites of the substance will be cleared from blood circulation by physiological pathways, and the majority of the received dose will eventually be exhaled as CO<sub>2</sub>.

# 10 EVALUATION OF HEALTH HAZARDS

# **Acute toxicity**

# 10.1 Acute toxicity - oral route

Not assessed in this dossier.

# 10.2 Acute toxicity - dermal route

Not assessed in this dossier.

# 10.3 Acute toxicity - inhalation route

Not assessed in this dossier.

# 10.4 Skin corrosion/irritation

Not assessed in this dossier.

# 10.5 Serious eye damage/eye irritation

Not assessed in this dossier.

# 10.6 Respiratory sensitisation

Not assessed in this dossier.

#### 10.7 Skin sensitisation

Table 9: Summary table of animal studies on skin sensitisation

Method, guideline, deviations if any	Species, strain, sex, no/group	Test substance (including purity), vehicle, positive control	Dose levels, duration of exposure	Results	Reference
LLNA OECD TG 429 (2010) GLP Key study Reliability: 1 A pre-test was performed in 2 mice with concentration s of 50 and 100% to	CBA/CaOlaHsd female mice 5 per each treatment group, 5 in control group (vehicle only)	Tetramethylene dimethacrylate, purity 99.63%  Vehicle: acetone:olive oil (4+1 v/v), purity of the acetone 99.6%  Positive control: α-hexylcinnamaldehyde (CAS 101-86-0) in acetone:olive oil (4:1)	25, 50 and 100%  Induction: topical application to the dorsal surface of each ear lobe on days 1, 2 and 3 (volume: 25 µl).  I.v. injection of <sup>3</sup> H-methyl thymidine via a	Sensitising The SI values at 25, 50 and 100% were 2.74, 3.76, and 5.72, respectively.  EC3 value: 31.4% (w/v) Observations: no mortality occurred during the study. On day 3, all treated animals showed reduced spontaneous activity, ruffled fur and hunched posture. The animals treated with concentrations of 50	Anonymous (2014)

Method, guideline,	Species, strain, sex, no/group	Test substance (including purity),	Dose levels, duration of	Results	Reference
deviations if any		vehicle, positive control	exposure		
determine the highest non-irritant test concentration.			tail vein (20.0 µCi <sup>3</sup> HTdR per mouse, volume: 250 µl) on day 6.  Necropsy on day 6	and 100% showed eyelid closure and abnormal walk on day 3, and ruffled fur on day 4. 2/5 of the animals treated with a concentration 100% showed reduced spontaneous activity on day 4. All treated animals developed an erythema of the ear skin during the observation period (25%: score 1 on days 3 and 4; 50%: score 1 on days 3-5; 100%: score 1 on days 2 and 6, score 2 on days 3-5). Body weight was within normal range.	
GPMT OECD TG 406 (1981) GLP: not specified Reliability: 3 Supporting study A pre-test was performed to determine the highest non-irritant test concentration.	SSc:Al outbred female guinea pigs No. of animals not specified (with other chemicals in the same paper, 10-20 animals per test group had been used)	1,4-butanediol dimethacrylate (purity not specified, but obtained from commercial sources, hence, commercial grade assumed)  Vehicle: soybean oil or sbomek (soybean oil:2-butanone, 1:2) (intradermal induction), petrolatum (topical induction and challenge)  10% sodium lauryl sulphate (SLS) was used on day 7 as an adjuvant prior to topical induction.  Positive control: not specified	Induction (day 0): 1% intradermal injection Induction (day 8): 5% topical application Challenge (day 21): 25% topical application	Not sensitising  No further information available	Anonymous (1984a)
Freund´s complete adjuvant test Non-guideline GLP: not specified Reliability: 3 Weight of evidence A pre-test was performed to determine the	Dunkin-Hartley female guinea pigs 8 animals in the treatment group, 4-6 animals in the control group	1,4-butanediol dimethacrylate, purity 97%  Vehicle (topical induction and challenge): Aramek (methyl ethyl ketone:arachis oil 2:1)  Positive control: not specified	0.5 M (13%) for intradermal induction exposure (days 1-9) and 3 M (78%) for challenge and rechallenge exposures (days 21 and 35)	Sensitising After challenge on day 21 8/8 animals were sensitised, after rechallenge on day 35 5/8 animals were positive.	Anonymous (1983a)

Method, guideline, deviations if any	Species, strain, sex, no/group	Test substance (including purity), vehicle, positive control	Dose levels, duration of exposure	Results	Reference
highest non- irritant test concentration.  GPMT  Non-guideline  GLP: not specified  Reliability: 2  Weight of evidence  A pre-test was performed to determine the highest non- irritant test concentration.	Himalayan white spotted female guinea pigs 10 animals in the treatment group, 6 animals in the control group	1,4-butanediol dimethacrylate, purity 97%  Vehicle (topical induction): petrolatum or 80% ethanol  Vehicle (challenge): Aramek (methyl ethyl ketone:arachis oil 2:1)  Positive control: not specified	0.5 M (13%) for intradermal induction exposure (day 0), 100% for topical induction exposure (day 7), 1 M (26%) for challenge and rechallenge exposures (days 21 and 35)	Ambiguous  After challenge on day 21 0/10 animals were sensitised, after rechallenge on day 35 2/10 animals were positive.  According to the authors, a third challenge has been perfomed on day 49 which confirmed the results of the rechallenge, but the data are not shown in the publication.	Anonymous (1983b)
GPMT Non-guideline GLP: not specified Reliability: 2 Weight of evidence A pre-test was performed on three animals to determine the highest non-irritant test concentration.	Dunkin-Hartley female guinea pigs 10 animals in the treatment group, 10 animals in the control group	1,4-butanediol dimethacrylate, purity min. 95%  Vehicle (intradermal induction): olive oil:acetone 9:1  Vehicle (topical induction, challenge and rechallenge): petrolatum  Before topical induction, a pretreatment with 10% SLS (w/w) in petrolatum was used.  Positive control: not specified	2% (w/w) for intradermal induction exposure, 50% for topical induction and 1% (w/w) for challenge and rechallenge exposures (amount of test item ca. 0.015 g)  48 hours after the first challenge the animals received a booster dose intradermally (2%, without Freund's complete adjuvant).	Not sensitising 0/10 animals were sensitised in this test; however, it is not documented whether the scores were obtained after the first or second challenge.	Anonymous (1984b)

# Animal data

The sensitising potential of tetramethylene dimethylacrylate has been investigated in one murine local lymph node assay and in four guinea pig studies (Table 9).

# **LLNA**

The LLNA was conducted in accordance with OECD test guideline 429 (2010) and principles of GLP (Anonymous 2014). A pre-test was performed in two animals with concentrations of 50 and 100% to

determine the highest non-irritant test concentration. The mouse treated with the undiluted test substance showed slightly reduced spontaneous activity, and an erythema of the ear skin was observed in both animals (score 1 in the mouse treated with 50% concentration, score 1-2 in the mouse treated with 100% concentration). Furthermore, scabby ears were observed on day 5 in the animal treated with the undiluted test substance.

In the main study, three treated groups of five CBA/CaOlaHsd female mice aged 8-9 weeks and weighing 17.8-22.3 g (mean 20.3 g  $\pm$  1.2 g) were used. The animals were treated by topical application to the dorsal surface of left and right ears with test concentrations of 25, 50 and 100% in acetone/olive oil (4+1, v/v). The application volume, 25  $\mu$ l, was spread over the entire dorsal surface (diameter ~ 8 mm) of left and right ears once daily for three consecutive days. The control group of five mice received vehicle only. Five days after the topical application, all mice were given 250  $\mu$ l of 19.5  $\mu$ Ci  $^3$ H-methyl thymidine (corresponds to 78  $\mu$ Ci/ml  $^3$ H-methyl thymidine) by intravenous injection via the tail vein. The body weight of the animals recorded prior to the injection was within the normal range for the strain and age. All animals were euthanized approximately five hours after the injection. The left and right draining auricular lymph nodes were then excised and pooled per group. Single cell suspensions of lymph node cells were prepared from the pooled lymph nodes. The proliferative capacity of the cells was determined by the incorporation of  $^3$ H-methyl thymidine measured on a  $^3$ H-scintillation counter.

No mortality was observed during the study period. All treated animals showed unspecific clinical signs on day 3, including reduced spontaneous activity, ruffled fur and hunched posture. All tried to burrow themselves in the bedding one hour after the third application. Eyelid closure and abnormal walk were also observed in the mice treated with concentrations of 50 and 100%. On day 4, all animals treated with concentrations of 50 and 100% showed ruffled fur and two animals treated with the undiluted test substance showed reduced spontaneous activity. According to the authors, it cannot be confirmed whether these symptoms were signs of systemic toxicity or mere reactions to the irritant nature of the test substance. The body weight of the animals remained within the normal range.

A substance is regarded as a sensitiser in the LLNA if the exposure to one or more test concentration results in a three-fold or greater increase in incorporation of  ${}^{3}$ H-methyl thymidine compared with vehicle-treated controls (the ratio is termed as the Stimulation Index, SI). The estimated test substance concentration required to produce an SI is referred to as the EC3 value. In this study, Stimulation Indices of 2.74, 3.76, and 5.72 were determined at concentrations of 25, 50 and 100%, respectively (Table 10). The EC3 value was 31.4% (w/v).

Table 10: Calculation of Stimulation Indices per dose group

	Group calculation		
Test item concentration (%)	Mean DPM per animal (2 lymph nodes) <sup>a)</sup>	SD	SI
0 (control group)	999.4	398.8	1.00
25	2740.8	353.5	2.74
50	3757.0	1373.9	3.76
100	5714.8	1986.8	5.72

DPM = disintegrations per minute, SD = standard deviation, SI = Stimulation index

#### Guinea pig studies

In a GPMT study (mainly in accordance with the OECD test guideline 406, 1981), female guinea pigs (no. of animals not specified) were induced on day 0 with 1% intradermal injections of tetramethylene

<sup>&</sup>lt;sup>a)</sup> = Mean DPM/animal was determined by dividing the sum of the measured values from lymph nodes of all animals within a group by the number of animals in that group (5 animals)

dimethacrylate in the nape of the neck (Anonymous 1984a). The animals weighed 300-350 g at the initiation of the study and were approximately one month old. Purity of the test substance is not specified in the study report. On day 7, approximately 250 mg of 10% sodium lauryl sulphate (instead of Freund's adjuvant, as described in the test guideline) in petrolatum was gently massaged into the neck and left uncovered for 24 hours. Epicutaneous application of 5% tetramethylene dimethacrylate followed on day 8, and the dressing containing the test solution was left in place for 48 hours. The vehicle controls received the same treatment, but with an equivalent amount of petrolatum. Challenge exposure was performed on day 21 using an occlusive epicutaneous application with a 25% concentration, and readings were made on days 23 and 24 (after 48 and 72 hours, respectively). The vehicle controls received identical treatment. The test substance was not found to be sensitising in the study. There is no further information available on clinical signs or findings.

A Freund's complete adjuvant test (FCAT) and a GPMT were conducted together on groups of eight and ten albino female guinea pigs, respectively (Anonymous 1983a, 1983b). The purity of tetramethylene dimethacrylate was 97% in both studies. According to the authors, sensitisation to impurities cannot be completely excluded. The animals weighed 350-450 g at study initiation, but their ages are not specified in the study report. There were four to six animals in the control group in the FCAT and six animals in the control group in the GPMT. A pre-test with FCA-treated animals preceded both studies. In both the FCAT and the GPMT, the animals were induced with intradermal injections of 0.5 M tetramethylene dimethacrylate which, according to the authors, corresponds to a 13% concentration. In the FCAT, 3 M (78%) concentration was then used for challenge and rechallenge exposures. In the GPMT, a 100% concentration was used for the topical induction exposure and 1 M (26%) concentration was used for challenge and rechallenge exposures. For the closed patch induction on day 7 of the GPMT, petrolatum or 80% ethanol was used as a vehicle. Aramek (methyl ethyl ketone:arachis oil 2:1) was used as a vehicle in all challenges. Tetramethylene dimethacrylate was found to be sensitising in the FCAT, but the results in the GPMT were ambiguous. There is no information on mortality or clinical signs or findings in either of the studies.

In a GPMT study (in accordance with the method described by Magnusson and Kligman), 10 female albino guinea pigs weighing 300-400 g (ages not specified) were treated with tetramethylene dimethacrylate intradermal injection (2%) in olive oil/acetone to induce sensitisation (Anonymous 1984b). There were 10 guinea pigs in the control group. The animals were further sensitised by topical application of 50% tetramethylene dimethacrylate. Prior to topical induction, the animals were treated with 10% sodium lauryl sulphate in petrolatum. For challenge and rechallenge exposures, a concentration of 1% tetramethylene dimethacrylate in petrolatum was used. A booster dose was applied intradermally on the neck using the same concentration and vehicle 48 hours after the first challenge. The rechallenge occurred one week after the first challenge. None of the animals were sensitised in this test, but it is not documented whether the scores were obtained after the first or the second challenge. No clinical observations or macroscopical findings are described in the study report.

#### Human data

A total of 26 clinical studies have been identified for tetramethylene dimethacrylate (Table 11). The studies comprised a total of 128 patients who tested positive to the substance. In all studies, the diagnostic method was patch testing. Data on skin exposure to tetramethylene dimethacrylate is scarce.

Table 11: Summary table of human data on skin sensitisation

Type of data/report	Test substance	Relevant information about the study (as applicable)	Observations	Reference
CASE REPOR	CASE REPORTS ON SINGLE CASES			
Case report	Tetramethylene dimethacrylate (2%,	A 38-year-old female was sensitised to a glue used in the attachment of car rear-view	13 acrylic compounds provoked mild to extreme allergic reactions in a patch	Kanerva et al. (1995)

Type of data/report	Test substance	Relevant information about the study (as applicable)	Observations	Reference
	Chemotechnique's test substance i.e. in pet.)	mirrors to the windscreen (with 6 years of work history). She developed a dry and fissured dermatitis on fingers and palms of both hands. The dermatitis spread within a couple of weeks to lower arms, chest, neck and face, and she developed rhinitis, paresthesia of fingertips and gastrointestinal complaints.	test.  Positive reaction to test substance (++ on day 2, ++ on day 3, ++ on day 4).  Tetramethylene dimethacrylate was not mentioned in the safety data sheet of the glue or detected in chemical analysis.	
Case report	Tetramethylene dimethacrylate (2%, vehicle not specified)	A 47-year-old atopic female cosmetician developed dermatitis on her thumb within some weeks after starting to work with photobonded nails. The dermatitis spread to both hands, and after stronger exposure to UV-gel 3 months later, she developed a severe hand and face dermatitis.	Allergic reactions to 15 (meth)acrylates, a total of 31 were tested  Allergic reactions to the test substance (+ was the strongest reading on days 2, 3 and 4)  Tetramethylene dimethacrylate was not detected in chemical analyses of the nail products.	Kanerva et al. (1996)
Case report	Tetramethylene dimethacrylate (concentration and vehicle not defined)	47-year-old woman had used acrylic nails for 10 years. She presented with periungual dermatitis of all the fingers. Symptoms had begun 6 months earlier.	She tested positive to 11 acrylic compounds including the test substance.  Tetramethylene dimethacrylate reaction was + at 96 hours.	Paley et al. (2006)
PATIENT SEI	RIES			
Patient series	Tetramethylene dimethacrylate (2% in pet.)	7 patients occupationally sensitized to methacrylate- based dental composite products	1 patient reacted positively to the test substance out of 5 patients tested (20%). The test substance was not mentioned in safety data sheets of the products.	Kanerva et al. (1989)
Patient series	Tetramethylene dimethacrylate (2% in pet.), purity 97%	126 dental technicians were tested with (meth)acrylates in 1995-1999 in Department of Dermatology, Städtische Kliniken (Dortmund, DE)	Positive reaction to the test substance in 6 of 126 patients (4.8%), all the reactions were assessed clinically relevant i.e. the sensitised persons had handled tetramethylene dimethacrylate-containing products. Authors considered that the test substance was a weak sensitiser in comparison to methyl methacrylate due to low number of positive reactions despite common exposure.	Peiler et al. (2000)
Patient series	Tetramethylene dimethacrylate (2% in pet.)	A retrospective study of 13 833 patients tested for contact allergy at the Department of Dermatology, Catholic	Positive reaction to the test substance in 5 of 72 patients (6.9%) who were positive to some (meth)acrylate.	Geukens & Goossens (2001)

Type of data/report	Test substance	Relevant information about the study (as applicable)	Observations	Reference
		University (Leuven, BE) in 1978-1999 It is unclear how many patients were tested with (meth)acrylates.		
Patient series	Tetramethylene dimethacrylate (2% in pet.)	The incidence of allergic contact dermatitis was studied in 79 dentists and 46 dental nurses who were referred to the Institute of Occupational Medicine (Lodz, PL) in 1990-2000. All were tested with the European standard set, dental screening test and additional allergens.	In dentists sensitised to acrylic resins, 8 of 20 patients (40%) reacted positively to the test substance. There were no positive reactions to the test substance in dental nurses.	Kiec- Swierczynska & Krecisz (2002)
Patient series	Tetramethylene dimethacrylate (2% in pet.)	90 patients suspected of having dermatitis caused by (meth)acrylates were patch tested at the Department of Occupational and Environmental Dermatology (Malmö, SE) in 1995-2004	24 patients reacted to some (meth)acrylate. 16 of these patients were tested with the test substance, and 3 of them tested positive (18.8%).  It is unclear how many patients in total were tested with tetramethylene dimethacrylate.	Goon et al. (2007)
Patient series	Tetramethylene dimethacrylate (2% in pet.)	473 patients were tested with a (meth)acrylate series at Finnish Institute of Occupational Health (Helsinki, FI) in 1994-2006.  32 patients with allergic reaction to some (meth) acrylate and working in dental professions (dentist, dental nurse, dental technician) were identified.	Positive reactions to the test substance in 3 cases: 1 dentist (++ reaction), 1 dental nurse (++ reaction) and 1 dental technician (+ reaction).  Tetramethylene dimethacrylate was not mentioned in safety data sheets of the products used by these 3 patients.	Aalto-Korte et al. (2007)
Patient series	Tetramethylene dimethacrylate (2% in pet.)	473 patients were tested with a (meth)acrylate series at Finnish Institute of Occupational Health (Helsinki, FI) in 1994-2006.  Among 61 patients with allergic reaction to some (meth)acrylate, 10 patients with present occupational exposure to acrylic glues were identified.	Positive reaction to the test substance in 4 (40%) of 10 patients (++ in three patients, +++ in one patient). All 4 patients had handled methacrylate-based glues but tetramethylene dimethacrylate was not mentioned in the safety data sheets of the glues.	Aalto-Korte et al. (2008)
Patient series	Tetramethylene dimethacrylate (0.1% in pet.)	A retrospective study on 43 patients diagnosed with allergic contact dermatitis caused by (meth)acrylates in long-lasting nail polish at dermatology departments of 4 Spanish hospitals in 2013-2016	Positive reaction to the test substance in 1 patient out of 7 (20%) tested with the substance within the group of 43 patients.	Gatica-Ortega (2017)
Patient series	Tetramethylene dimethacrylate (2%	A retrospective study on 16 nail technicians with methacrylate	Positive reaction to the test substance in 2 of 16 patients	Fisch et al. (2019)

Type of data/report	Test substance	Relevant information about the study (as applicable)	Observations	Reference
	in pet.)	allergy who had been patch tested at the Department of Dermatology (Gävle and Malmö, SE) in 2007-2016	(12.5%).	
Patient series	Tetramethylene dimethacrylate (2% in pet.)	A retrospective study on patients suspected of nail manicure-related sensitisation to (meth)acrylates at dermatology departments of 3 Spanish hospitals in 2008-2017 A total of 208 patients were tested with (meth)acrylates.	66 patients reacted positively to at least one (meth)acrylate and the sensitisation was due to nail products.  In this group, positive reaction to the test substance in 6 of 26 patients (23.1%) tested with the substance.	Marrero- Alemán et al. (2019)
CROSS-SECT	TONAL STUDIES OF	N RISK OCCUPATIONS		
Cross- sectional study	Tetramethylene dimethacrylate (2% in pet.)	A questionnaire was sent to 1132 dental technicians and 173 answered. 55 cases were patch tested.	Tetramethylene dimethacrylate was positive in 1 (2%) case of those tested (N=55).	Rustemeyer & Frosch (1996)
Cross- sectional study	Tetramethylene dimethacrylate (Chemotechnique's test substance i.e. 2% in pet.)	49 out of 1038 dental technicians voluntarily participated in a study on patch testing at the Department of Dermatology in the Catholic University of Korea (Seoul, KR)	Positive reaction to the test substance in 1 case, 2.1% of those tested.  7 patients were positive to some acrylic substance. The test substance-positive case constituted 14% of this group.	Lee et al. (2001)
CLINICAL PA		ON SELECTED PATIENTS (AIM	ED TESTING WITH ACRYLIC	C
Patch test data, selected patients	Tetramethylene dimethacrylate (2% in pet.)	A retrospective study on 23 patients patch tested with (meth)acrylate series at the Nofer Institute of Occupational Medicine, Lodz (PL) in 1990- 1994	Positive reactions to the test substance in 2 (9.5%) dentists out of 21 patients tested with the substance.	Kiec- Swierczynska (1996)
Patch test data, selected patients	Tetramethylene dimethacrylate (2% in pet.)	The incidence of allergic reactions to certain methacrylates by the Information Network of Departments of Dermatology (Göttingen, DE) in 1992-1995	Positive reaction to the test substance in 13 of 2971 patients (0.4%).	Schnuch (1996)
Patch test data, selected patients	Tetramethylene dimethacrylate (2%; Chemotechnique's test substance i.e. in pet.)	A retrospective study on patients tested with (meth)acrylate patch test series at the Section of Dermatology in the Finnish Institute of Occupational Heath in 1885- 1995	Positive reaction to the test substance in 10 of 274 (3.6%) patients tested with the substance.  48 patients reacted positively to some (meth)acrylate. The test substance-positive cases constituted 20.8% of these.	Kanerva et al. (1997)
Patch test data, selected patients	Tetramethylene dimethacrylate (2%, Chemotechnique's test substance i.e.	A retrospective study of patch test records at the Section of Dermatology, University of Manchester (Salford, UK) in	Positive reaction to the test substance in 7 of 255 patients (2.7%) tested with the substance.	Tucker & Beck (1999)

Type of data/report	Test substance	Relevant information about the study (as applicable)	Observations	Reference
	in pet.)	1983-1998 440 patients with a history of exposure to (meth)acrylates were identified and patch tested with (meth)acrylates		
Patch test data, selected patients	Tetramethylene dimethacrylate (concentration or vehicle not stated)	A retrospective study on patients patch tested with dental screening series in 7 dermatology clinics in Finland in 1994-1998	There were 13 (0.5%) allergic reactions to the test substance in the 2408 patients tested. The frequency of allergic reactions varied between 0.1% and 2.2% in different clinics.	Kanerva et al. (2001)
Patch test data, selected patients	Tetramethylene dimethacrylate (2% in pet.)	109 patients (all dental personnel) were tested with a dental screening series at the Department of Occupational and Environmental Dermatology (Stockholm, SE) in 1995-1998	Positive reaction to the test substance in 6 (5.5%) of 109 patients tested with (meth)acrylates.  24 patients had allergic reactions to some (meth)acrylate. The 6 test substance-positive cases constituted 25% of these.	Wrangsjö et al. (2001)
Patch test data, selected patients	Tetramethylene dimethacrylate (2% in pet.)	A retrospective study of patch test records of 1632 patients tested with dental patient and/or dental personnel series at the Department of Occupational and Environmental Dermatology in Malmö University Central Hospital (SE) in 1995-2004	Positive reaction to the test substance in 9 (0.5%) out of 1642 patients tested.  48 patients reacted positively to at least one (meth)acrylate. The test substance-positive cases constituted 18.8% of these patients.	Goon et al. (2006)
Patch test data, selected patients	Tetramethylene dimethacrylate (2% in pet.)	A retrospective study on 451 patients suspected of having occupational contact dermatitis and tested with a (meth)acrylate series at Finnish Institute of Occupational Health (Helsinki, FI) in 1994-2009	Positive reaction to the test substance in 9 patients (2.0%) 66 patients reacted positively to at least one (meth)acrylate. The test substance-positive cases constituted 13.6% of this group.	Aalto-Korte et al. (2010) Includes the patients in Aalto-Korte et al. (2008) and Aalto-Korte et al. (2007)
Patch test data, selected patients	Tetramethylene dimethacrylate (2%; Chemotechnique's test substance i.e. in pet.)	A retrospective study on patients tested with (meth)acrylate series at the Department of Dermatology, University Medical Centre in Groningen (NL) in 1993-2012	Positive reactions in 6 of 151 (4.0%) patients tested with the substance.  24 patients reacted positively to some (meth)acrylate. The positive reactions to tetramethylene dimethacrylate constituted 25% of these.	Christoffers et al. (2013)
Patch test data, selected patients	Tetramethylene dimethacrylate (2% in pet.)	122 patients were tested with an extended series of (meth)acrylates at the Department of Dermatology (Coimbra, PT) in 2006-2013	Positive reaction to the test substance in 5 (4.1%) patients.  37 patients reacted positively to (meth)acrylates. The tetramethylene dimethacrylate-positive cases	Ramos et al. (2014)

Type of data/report	Test substance	Relevant information about the study (as applicable)	Observations	Reference
			constituted 13.5% of these.	
Patch test data, selected patients	Tetramethylene dimethacrylate (2% in pet.)	475 patients were tested with a (meth)acrylate series at the Cutaneous Allergy Unit (Birmingham, UK) in 2002-2015	Positive reactions to the test substance in 10 (2.1%) patients tested with the substance.  52 patients reacted positively to (meth)acrylates. The positive reactions to tetramethylene dimethacrylate constituted 19% of these.	Spencer et al. (2016)

Diagnostic patch testing is conducted in order to diagnose contact allergy to a substance and performed according to international standards by dermatologists (Johansen et al. 2015). The results of such tests are usually reported as number of patients/subjects with positive reactions in relation to the total number of tested (frequency of positive patch tests). An important factor of assessing prevalence of positive reactions in diagnostic patch test is how the group of patients is defined, i.e. if they are selected in some way or not. Selected patients can be, for instance, patients with dermatitis suspected of having contact with acrylic compounds or special occupational groups (aimed testing). Consecutive or unselected patients are groups of patients for whom allergic contact dermatitis is generally suspected.

There are no studies on diagnostic patch tests with tetramethylene dimethacrylate in general population or unselected clinical patients.

Tetramethylene dimethacrylate has been commonly tested as part of the (meth)acrylate series since the 1980s. Its established test concentration is 2% in petrolatum. A total of 11 diagnostic patch test studies on selected patients could be identified for the substance. The frequency of positive reactions varied between 0.4% and 9.5% (median 2.7%).

No strict workplace studies could be identified for tetramethylene dimethacrylate. However, two cross-sectional studies on dental technicians, who are at risk of developing a contact allergy due to exposure to acrylic compounds at work, share a similar design. Only the workers with skin symptoms were patch tested in these studies. The frequency of positive patch test reactions to the substance was 2% in both studies (1/55 and 1/49 of the tested patients; Rustemeyer & Frosch 1996, and Lee et al. 2001, respectively).

The rest of the identified studies were either case reports of single cases (n=4) or reports describing patient series (n=10) without clearly stating the frequency of reaction to tetramethylene dimethacrylate in all patients tested during the same time period. Specific exposure to the substance was described by Peiler et al. (2000) in all six dental technicians who tested positive to it. In the 1990s in Germany, tetramethylene dimethacrylate was commonly found in the products used by dental technicians and virtually all workers were exposed to the substance. The authors considered that tetramethylene dimethacrylate was a weak sensitiser compared to methyl methacrylate because the frequency of contact allergy was low (4.8%), despite common exposure. Dental technicians' skin exposure to tetramethylene dimethacrylate may also vary within countries, as for instance in Finland only two dental technicians out of eight had used products containing the substance (Aalto-Korte et al. 2007).

# Table 12: Summary table of other studies relevant for skin sensitisation

No other data is available.

# 10.7.1 Short summary and overall relevance of the provided information on skin sensitisation

Animal data

In the key LLNA (OECD TG 429, 2010), three treated groups of five mice were administered tetramethylene dimethacrylate topically at concentrations of 25, 50 and 100% in acetone/olive oil (4+1, v/v) (Anonymous 2014). The control group of five mice received vehicle only. No mortality was observed during the study period. All treated animals showed unspecific clinical signs on day 3, including reduced spontaneous activity, ruffled fur and hunched posture. All tried to burrow themselves in the bedding one hour after the third application. Eyelid closure and abnormal walk were also observed in the mice treated with concentrations of 50 and 100%. On day 4, all animals treated with concentrations of 50 and 100% showed ruffled fur and two animals treated with the undiluted test substance showed reduced spontaneous activity. According to the authors, it cannot be confirmed whether these symptoms were signs of systemic toxicity or mere reactions to the irritant nature of the test substance. A dose-related increase in the stimulation index (SI) values was observed and the threshold positive value of 3 was exceeded at concentrations of 50% and 100%. The EC3 value was 31.4% (w/v).

Four guinea pig studies conducted in the 1980s are also available for evaluation of skin sensitisation potential of tetramethylene dimethacrylate. Only one of them, Anonymous 1984a, complies with the OECD test guideline (TG 406, 1981), although with clear deviations (number of animals and positive control not specified, purity of the substance not known) and therefore not suitable for classification purposes. The remaining three studies, a non-guideline Freund's complete adjuvant test (FCAT) and two non-guideline GPMTs, are of better methodological quality, apart from their unspecified positive controls and a rather low number of animals (Anonymous 1983a, 1983b and 1984b). Tetramethylene dimethacrylate was found to be sensitising in the FCAT (8/8 sensitised animals after the first challenge, 5/8 sensitised animals after rechallenge) (Anonymous 1983a). The substance was not a skin sensitiser in one GPMT, whereas the results were ambiguous in the other (Anonymous 1984b and Anonymous 1983b, respectively).

#### Human data

A total of 26 clinical patch test studies were identified for tetramethylene dimethacrylate. There are no studies in general population or unselected clinical patients. Tetramethylene dimethacrylate is usually tested as part of the (meth)acrylate patch test series, and a total of 11 diagnostic patch test studies on selected patients could be identified for the substance. The frequency of positive reactions varied between 0.4% and 9.5% (median 2.7%) in the studies.

There are no strict workplace studies for tetramethylene dimethacrylate. However, in two cross-sectional studies dental technician was identified as a risk occupation for contact allergy following exposure to acrylic compounds, such as tetramethylene dimethacrylate. The rest of the identified studies were either case reports of single cases (n=4) or reports describing patient series (n=10) without clearly stating the frequency of reaction to the substance in all patients tested during the same time period.

# 10.7.2 Comparison with the CLP criteria

Substances are classified as Category 1 skin sensitisers where data are not sufficient for sub-categorisation, if there is evidence in humans that the substance can lead to sensitisation by skin contact in a substantial number of persons, or if there are positive results from an appropriate animal test (Annex I, Table 3.4.2 of the CLP Regulation).

Substances are classified as Sub-category 1A skin sensitisers where there is evidence of a high frequency of occurrence in humans and/or a high potency in animals. Such evidence includes

Human evidence: diagnostic patch test data where there is a relatively high and substantial incidence of reactions in a defined population in relation to relatively low exposure.

GPMT:  $\geq$ 30% responding at  $\leq$ 0.1% intradermal induction dose or  $\geq$ 60% responding at >0.1% to  $\leq$ 1% intradermal induction dose.

LLNA: EC3 value ≤2%.

Substances are classified as Sub-category 1B skin sensitisers where there is evidence of a low to moderate frequency of occurrence in humans and/or a low to moderate potency in animals. Such evidence includes:

Human evidence: diagnostic patch test data where there is a relatively low but substantial incidence of reactions in a defined population in relation to relatively high exposure.

GPMT:  $\geq$ 30% to <60% responding at >0.1% to  $\leq$ 1% intradermal induction dose or  $\geq$ 30% responding at >1% intradermal induction dose.

LLNA: EC3 value >2%.

In the key LLNA (conducted in compliance with OECD TG 429 and GLP), tetramethylene dimethacrylate showed an EC3 value of 31.4% (w/v), indicating a low to moderate skin sensitisation potency. Sub-category 1A can therefore be excluded. According to the Guidance on the Application of the CLP Criteria (ECHA 2017b, Table 3.4.4), the results allow classification in Skin Sens. 1B. Four guinea pig tests from the 1980s are also available for assessment; however, due to their methodological limitations, they mainly serve as supporting information to be used as part of a weight-of-evidence evaluation.

### Human data

According to the classification criteria human evidence for Sub-categories 1A and 1B, respectively, can include the following type of data (ECHA 2017b, Section 3.4.2.2.3.1.):

	Human data
Sub-category 1A	(a) positive responses at ≤ 500 μg/cm2 (HRIPT, HMT – induction threshold);
	(b) diagnostic patch test data where there is a relatively high and substantial incidence of reactions in a defined population in relation to relatively low exposure;
	(c) other epidemiological evidence where there is a relatively high and substantial incidence of allergic contact dermatitis in relation to relatively low exposure.
Sub-category 1B	(a) positive responses at > 500 μg/cm2 (HRIPT, HMT – induction threshold);
	(b) diagnostic patch test data where there is a relatively low but substantial incidence of reactions in a defined population in relation to relatively high exposure;
	(c) other epidemiological evidence where there is a relatively low but substantial incidence of allergic contact dermatitis in relation to relatively high exposure.

HRIPT: Human Repeat Insult Patch Test; HMT: Human Maximisation Test

The Guidance on the Application of the CLP Criteria further outlines how high or low frequency of occurrence of skin sensitisation shall be assessed (ECHA 2017b, Section 3.4.2.2.3.1., Table 3.2):

Human diagnostic patch test data	High frequency	Low/moderate frequency	Tetramethylene dimethacrylate
General population studies	≥ 0.2 %	< 0.2 %	No studies
Dermatitis patients (unselected, consecutive)	≥ 1.0 %	< 1.0 %	No studies
Selected dermatitis patients (aimed testing, usually special test series)	≥ 2.0 %	< 2.0 %	11 studies 0.4%-9.5% (median 2.8%)
Workplace studies:			
1: all or randomly selected workers	≥ 0.4 %	< 0.4 %	No studies

2: selected workers with known exposure or dermatitis	≥ 1.0 %	< 1.0 %	2 studies: 2%
Number of published cases	≥ 100 cases	< 100 cases	128 patch-test- positive cases

There are no studies on general population or on unselected consecutive dermatitis patients.

Frequencies of positive patch tests in 11 selected dermatitis patient materials (aimed testing) have varied between 0.4% and 9.5% (median 2.7%), but they are mostly above the limit of high frequency.

There are no workplace studies on all or randomly selected workers. In two cross-sectional studies on dental technicians mimicking workplace studies (on selected workers) the frequency of positive patch tests was 2%, i.e. above the cut-off value of 1.0% for high frequency.

The number of published patch-test-positive cases, 128, also exceeds the cut-off value for high frequency (≥ 100).

Positive patch test reactions to tetramethylene dimethacrylate are relatively common in patients sensitised to methacrylates, but specific exposure to the substance in sensitised or tested patients has rarely been described in the literature. Both the exposure and the lack of exposure to tetramethylene dimethacrylate are typically difficult to assess in clinical work due to the unavailability of chemical analyses. Positive test reactions may also arise from cross-reactivity to other methacrylates, yet true exposure to tetramethylene dimethacrylate in clinical patients cannot be excluded. Of the identified literature, only Peiler et al. (2000) confirmed exposure to the substance in all six dental technicians who gave a positive reaction to it.

To conclude, the frequency of positive reactions to tetramethylene dimethacrylate in diagnostic patch tests can be considered high. However, there is no adequate information enabling the assessment of true exposure to the substance. Animal data is sufficient for sub-categorization, and human data supports the classification of tetramethylene dimethacrylate as a skin sensitiser. Based on the key LLNA, Sub-category 1A can be excluded and Sub-category 1B is justified.

# 10.7.3 Conclusion on classification and labelling for skin sensitisation

Based on the available data, the proposed classification and labelling for skin sensitisation is **Sub-category 1B**. The corresponding hazard statement is **H317: May cause an allergic skin reaction**.

# 10.8 Germ cell mutagenicity

Not assessed in this dossier.

#### 10.9 Carcinogenicity

Not assessed in this dossier.

#### 10.10 Reproductive toxicity

Not assessed in this dossier.

# 10.11 Specific target organ toxicity-single exposure

Not assessed in this dossier.

# 10.12 Specific target organ toxicity-repeated exposure

Not assessed in this dossier.

#### 10.13 Aspiration hazard

Not assessed in this dossier.

# 11 EVALUATION OF ENVIRONMENTAL HAZARDS

#### 11.1 Rapid degradability of organic substances

Not assessed in this dossier.

#### 11.2 Environmental transformation of metals or inorganic metals compounds

Not assessed in this dossier.

#### 11.3 Bioaccumulation

Not assessed in this dossier.

# 11.4 Acute aquatic hazard

Not assessed in this dossier.

### 11.5 Long-term aquatic hazard

Not assessed in this dossier.

#### 12 EVALUATION OF ADDITIONAL HAZARDS

#### 12.1 Hazardous to the ozone layer

Not assessed in this dossier.

#### 13 ADDITIONAL LABELLING

The label on the packaging of mixtures not classified as sensitising but containing tetramethylene dimethacrylate, classified as Skin Sens. 1B; H317, in a concentration of  $\geq 0.1\%$  shall bare the statement EUH208 (CLP Annex II, Section 2.8).

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#### 15 ANNEXES

Confidential Annex on toxicokinetic studies