



Safety Information Sheet for Medical Devices

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Revision date: 23/10/2019 **Supersedes date:** Initial issue.
Transportation version number: 1.00 (23/10/2019)

A safety data sheet is not required for this Product. This Safety Information Sheet has been created on a voluntary basis.

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

1.1. Product identifier

3M FILTEK Z250 UNIVERSAL RESTORATIVE (ALL SHADES EXCEPT B0.5 and B1)

Product Identification Numbers

LE-F100-0078-3	LE-F100-2472-6	70-2010-2225-1	70-2010-2226-9	70-2010-2227-7
70-2010-2228-5	70-2010-2229-3	70-2010-2232-7	70-2010-2233-5	70-2010-2234-3
70-2010-2237-6	70-2010-2238-4	70-2010-2241-8	70-2010-2242-6	70-2010-2243-4
70-2010-2244-2	70-2010-2245-9	70-2010-2248-3	70-2010-2249-1	70-2010-2250-9
70-2010-2253-3	70-2010-2254-1	70-2010-2259-0	70-2010-2260-8	70-2010-2564-3
70-2010-2565-0	70-2010-2566-8	70-2010-2567-6	70-2010-2568-4	70-2010-2571-8
70-2010-2572-6	70-2010-2573-4	70-2010-2576-7	70-2010-2577-5	70-2010-2578-3
70-2010-2585-8	70-2010-2586-6	70-2010-2587-4	70-2010-2588-2	70-2010-2589-0
70-2010-2592-4	70-2010-2593-2	70-2010-2594-0	70-2010-2597-3	70-2010-2598-1
70-2010-2599-9	70-2010-2712-8	70-2010-2723-5	70-2010-2724-3	70-2010-2725-0
70-2010-2726-8	70-2010-3200-3	70-2010-3201-1	70-2010-3202-9	70-2010-3203-7
70-2010-5064-1	70-2010-5200-1	70-2010-5201-9	70-2010-5202-7	70-2010-5203-5
70-2010-5204-3	70-2010-5207-6	70-2010-5208-4	70-2010-5209-2	70-2010-5212-6
70-2010-5213-4	70-2010-5214-2	70-2010-5216-7	70-2010-5217-5	70-2010-5218-3
70-2010-5219-1	70-2010-5565-7	70-2010-9546-3	70-2010-9547-1	70-2010-9548-9
70-2010-9549-7	70-2010-9550-5	70-2010-9552-1	70-2010-9553-9	70-2010-9554-7
70-2010-9555-4	70-2010-9556-2	70-2010-9557-0	70-2010-9558-8	70-2010-9559-6
70-2010-9560-4	70-2010-9561-2	70-2010-9563-8	70-2010-9564-6	70-2010-9565-3
70-2010-9566-1	70-2010-9567-9	70-2010-9568-7	70-2010-9605-7	70-2014-1122-3
70-2014-1123-1	70-2014-1124-9	70-2014-1125-6	70-2014-1126-4	70-2014-1127-2
70-2014-1128-0	70-2014-1129-8	70-2014-1130-6	70-2014-1131-4	70-2014-1132-2
70-2014-1133-0	AH-0105-6585-3	UU-0091-3103-6	UU-0091-3104-4	UU-0091-3105-1
UU-0091-3106-9	UU-0091-3107-7	UU-0091-3109-3	UU-0091-3110-1	UU-0091-3111-9
UU-0091-3112-7	UU-0091-3113-5	UU-0091-3114-3		

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7000054199	7000054200	7000054201	7000054202	7000054203
7000054205	7000054206	7000054207	7000054208	7000054209
7000003072	7000003073	7000003074	7000003075	7000003076
7000003078	7000003079	7000054210	7000054211	7000003084
7000054212	7000003085	7000030424	7000054236	7000030425
7000054237	7000030426	7000030428	7000030429	7000054239
7000054240	7000054241	7000054242	7000030433	7000030434
7000030435	7000030436	7000030437	7000030440	7000030441
7000030442	7000030445	7000030446	7000030447	7100111620
7100111621	7100111787	7100111812	7100111789	7100111781
7100111851	7000003194	7000003195	7000003196	7000003197
7000003198	7000003200	7000003201	7100009086	7000030530
7000003202	7000003203	7100111854	7010387258	7100111821
7100111796	7000054359	7010302362	7010387631	7010342916
7010387632	7010317413	7010342917	7010290952	7010387634
7010342918	7100111723	7100111703	7010387652	7010387626
7100111730	7010387627	7010342913	7010296472	7010342914
7010387629	7010317412	7100111726	7010342915	7100140987
7100140984	7100140981	7100140980	7100140989	7100141017
7100140988	7100140983	7100140986	7100140979	7100140990
7100141009	7100149332	7100149331	7100149330	7100149329
7100149328	7100149339	7100149338	7100149337	7100149336
7100149335	7100149334			

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

Identified uses

Medical device; refer to Instructions for Use

Restrictions on Use

For use only by dental professionals

1.3 Details of the supplier of the safety information sheet for medical devices

Address: 3M United Kingdom PLC, 3M Centre, Cain Road, Bracknell, Berkshire, RG12 8HT.
Telephone: +44 (0)1344 858 000
E Mail: tox.uk@mmm.com
Website: www.3M.com/uk

1.4. Emergency telephone number

+44 (0)1344 858 000

SECTION 2: Hazard identification

2.1. Classification of the substance or mixture

CLP REGULATION (EC) No 1272/2008

This product is a medical device as defined in Directive 93/42/EEC (MDD), which is invasive or used in direct physical contact with the human body, and therefore is exempt from the requirements of classification and labelling according to Regulation (EC) No. 1272/2008 (CLP; Article 1, paragraph 5). Although not required, the classification and label information, as applicable, is provided below.

CLASSIFICATION:

Skin Sensitization, Category 1 - Skin Sens. 1; H317

For full text of H phrases, see Section 16.

2.2. Label elements

CLP REGULATION (EC) No 1272/2008

SIGNAL WORD

WARNING.

Symbols:

GHS07 (Exclamation mark) |

Pictograms



Ingredients:

Ingredient	CAS Nbr	EC No.	% by Wt
Urethane dimethacrylate (UDMA)	72869-86-4	276-957-5	1 - 10
Bis-GMA	1565-94-2	216-367-7	1 - 10
Triethylene glycol dimethacrylate	109-16-0	203-652-6	< 5

HAZARD STATEMENTS:

H317 May cause an allergic skin reaction.

PRECAUTIONARY STATEMENTS

Prevention:

P280E Wear protective gloves.

Response:

P333 + P313 If skin irritation or rash occurs: Get medical advice/attention.

2.3. Other hazards

For information on hazards and safe use, please consider the corresponding sections of this document.

SECTION 3: Composition/information on ingredients

Ingredient	CAS Nbr	EC No.	% by Wt	Classification
Silane treated ceramic	444758-98-9		75 - 85	Substance not classified as hazardous
Dimethacrylate (BIS-MEPP)	41637-38-1	609-946-4	1 - 10	Aquatic Chronic 4, H413
Urethane dimethacrylate (UDMA)	72869-86-4	276-957-5	1 - 10	Aquatic Chronic 3, H412 Skin Sens. 1B, H317
Bis-GMA	1565-94-2	216-367-7	1 - 10	Skin Sens. 1B, H317
Triethylene glycol dimethacrylate (REACH Reg. No.:01-2119969287-21)	109-16-0	203-652-6	< 5	Skin Sens. 1, H317

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Aluminum oxide (REACH Reg. No.:01-2119529248-35)	1344-28-1	215-691-6	<= 1	Substance with a Community level exposure limit in the workplace
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Note: Any entry in the EC# column that begins with the numbers 6, 7, 8, or 9 are a Provisional List Number provided by ECHA pending publication of the official EC Inventory Number for the substance. Please see section 16 for the full text of any H statements referred to in this section

For information on ingredient occupational exposure limits or PBT or vPvB status, see sections 8 and 12 of this SIS

SECTION 4: First aid measures

4.1. Description of first aid measures

Inhalation

Remove person to fresh air. If you feel unwell, get medical attention.

Skin contact

Immediately wash with soap and water. Remove contaminated clothing and wash before reuse. If signs/symptoms develop, get medical attention.

Eye contact

Flush with large amounts of water. Remove contact lenses if easy to do. Continue rinsing. If signs/symptoms persist, get medical attention.

If swallowed

Rinse mouth. If you feel unwell, get medical attention.

SECTION 5: Fire-fighting measures

5.1. Extinguishing media

In case of fire: Use a fire fighting agent suitable for ordinary combustible material such as water or foam to extinguish.

5.2. Special hazards arising from the substance or mixture

None inherent in this product.

Hazardous Decomposition or By-Products

Substance

Carbon monoxide.
Carbon dioxide.

Condition

During combustion.
During combustion.

5.3. Advice for fire-fighters

Wear full protective clothing, including helmet, self-contained, positive pressure or pressure demand breathing apparatus, bunker coat and pants, bands around arms, waist and legs, face mask, and protective covering for exposed areas of the head.

SECTION 6: Accidental release measures

6.1. Personal precautions, protective equipment and emergency procedures

Evacuate area. Ventilate the area with fresh air. For large spill, or spills in confined spaces, provide mechanical ventilation to disperse or exhaust vapours, in accordance with good industrial hygiene practice. Refer to other sections of this SIS for information regarding physical and health hazards, respiratory protection, ventilation, and personal protective equipment.

6.2. Environmental precautions

Avoid release to the environment.

6.3. Methods and material for containment and cleaning up

Collect as much of the spilled material as possible. Place in a closed container approved for transportation by appropriate authorities. Clean up residue. Seal the container. Dispose of collected material as soon as possible.

SECTION 7: Handling and storage

Refer to Instructions for Use (IFU) for more information.

SECTION 8: Exposure controls/personal protection

8.1 Control parameters

Occupational exposure limits

If a component is disclosed in section 3 but does not appear in the table below, an occupational exposure limit is not available for the component.

Ingredient	CAS Nbr	Agency	Limit type	Additional comments
Aluminum oxide	1344-28-1	UK HSC	TWA(as inhalable dust):10 mg/m ³ ;TWA(as respirable dust):4 mg/m ³	

UK HSC : UK Health and Safety Commission

TWA: Time-Weighted-Average

STEL: Short Term Exposure Limit

CEIL: Ceiling

Biological limit values

No biological limit values exist for any of the components listed in Section 3 of this safety information sheet.

8.2. Exposure controls

8.2.1. Engineering controls

Use in a well-ventilated area.

8.2.2. Personal protective equipment (PPE)

Eye/face protection

Select and use eye/face protection to prevent contact based on the results of an exposure assessment. The following eye/face protection(s) are recommended:
Safety glasses with side shields.

Applicable Norms/Standards

Use eye protection conforming to EN 166

Skin/hand protection

See Section 7.1 for additional information on skin protection.

Respiratory protection

None required.

SECTION 9: Physical and chemical properties

9.1. Information on basic physical and chemical properties

Appearance	
Physical state	Solid.
Colour	Multicolor
Specific Physical Form:	Paste
Odor	Slight Acrylate
pH	<i>Not applicable.</i>
Boiling point/boiling range	<i>Not applicable.</i>
Melting point	<i>No data available.</i>
Flammability (solid, gas)	Not classified
Explosive properties	Not classified
Oxidising properties	Not classified
Flash point	No flash point
Autoignition temperature	<i>No data available.</i>
Flammable Limits(LEL)	<i>Not applicable.</i>
Flammable Limits(UEL)	<i>Not applicable.</i>
Relative density	2.1 [Ref Std:WATER=1]
Water solubility	Negligible
Viscosity	approximately 300,000 mPa-s
Density	2.1 g/cm ³

9.2. Other information

EU Volatile Organic Compounds	<i>No data available.</i>
Molecular weight	<i>No data available.</i>
Percent volatile	<i>No data available.</i>

SECTION 10: Stability and reactivity

10.1 Reactivity

This material is considered to be non reactive under normal use conditions

10.2 Chemical stability

Stable.

10.3 Possibility of hazardous reactions

Hazardous polymerisation will not occur.

10.4 Conditions to avoid

None known.

10.5 Incompatible materials

None known.

10.6 Hazardous decomposition products

<u>Substance</u>	<u>Condition</u>
None known.	

Refer to section 5.2 for hazardous decomposition products during combustion.

SECTION 11: Toxicological information

The information below may not agree with the EU material classification in Section 2 and/or the ingredient

classifications in Section 3 if specific ingredient classifications are mandated by a competent authority. In addition, statements and data presented in Section 11 are based on UN GHS calculation rules and classifications derived from 3M assessments.

11.1 Information on Toxicological effects

Signs and Symptoms of Exposure

Based on test data and/or information on the components, this material may produce the following health effects:

Inhalation

This product may have a characteristic odour; however, no adverse health effects are anticipated.

Skin contact

Contact with the skin during product use is not expected to result in significant irritation. Allergic skin reaction (non-photo induced): Signs/symptoms may include redness, swelling, blistering, and itching.

Eye contact

Contact with the eyes during product use is not expected to result in significant irritation.

Ingestion

May be harmful if swallowed.

Gastrointestinal irritation: Signs/symptoms may include abdominal pain, stomach upset, nausea, vomiting and diarrhoea.

Toxicological Data

If a component is disclosed in section 3 but does not appear in a table below, either no data are available for that endpoint or the data are not sufficient for classification.

Acute Toxicity

Name	Route	Species	Value
Overall product	Dermal		No data available; calculated ATE >5,000 mg/kg
Overall product	Ingestion		No data available; calculated ATE 2,000 - 5,000 mg/kg
Silane treated ceramic	Dermal		LD50 estimated to be > 5,000 mg/kg
Silane treated ceramic	Ingestion		LD50 estimated to be 2,000 - 5,000 mg/kg
Dimethacrylate (BIS-MEPP)	Dermal	Professional judgement	LD50 estimated to be > 5,000 mg/kg
Urethane dimethacrylate (UDMA)	Dermal	Professional judgement	LD50 estimated to be > 5,000 mg/kg
Dimethacrylate (BIS-MEPP)	Ingestion	Rat	LD50 > 2,000 mg/kg
Urethane dimethacrylate (UDMA)	Ingestion	Rat	LD50 > 5,000 mg/kg
Bis-GMA	Ingestion		LD50 estimated to be 2,000 - 5,000 mg/kg
Bis-GMA	Dermal	Professional judgement	LD50 estimated to be 2,000 - 5,000 mg/kg
Triethylene glycol dimethacrylate	Dermal	Professional judgement	LD50 estimated to be > 5,000 mg/kg
Triethylene glycol dimethacrylate	Ingestion	Rat	LD50 10,837 mg/kg
Aluminum oxide	Dermal		LD50 estimated to be > 5,000 mg/kg
Aluminum oxide	Inhalation-Dust/Mist (4 hours)	Rat	LC50 > 2.3 mg/l
Aluminum oxide	Ingestion	Rat	LD50 > 5,000 mg/kg

ATE = acute toxicity estimate

Skin Corrosion/Irritation

Name	Species	Value
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Silane treated ceramic	similar compounds	No significant irritation
Bis-GMA	Not available	Minimal irritation
Triethylene glycol dimethacrylate	Guinea pig	Mild irritant
Aluminum oxide	Rabbit	No significant irritation

Serious Eye Damage/Irritation

Name	Species	Value
Silane treated ceramic	similar compounds	Mild irritant
Bis-GMA	Not available	Moderate irritant
Triethylene glycol dimethacrylate	Professional judgement	Moderate irritant
Aluminum oxide	Rabbit	No significant irritation

Skin Sensitisation

Name	Species	Value
Silane treated ceramic	similar compounds	Not classified
Dimethacrylate (BIS-MEPP)	Guinea pig	Not classified
Urethane dimethacrylate (UDMA)	Guinea pig	Sensitising
Bis-GMA	Guinea pig	Sensitising
Triethylene glycol dimethacrylate	Human and animal	Sensitising

Respiratory Sensitisation

For the component/components, either no data is currently available or the data is not sufficient for classification.

Germ Cell Mutagenicity

Name	Route	Value
Dimethacrylate (BIS-MEPP)	In Vitro	Not mutagenic
Bis-GMA	In Vitro	Some positive data exist, but the data are not sufficient for classification
Triethylene glycol dimethacrylate	In Vitro	Some positive data exist, but the data are not sufficient for classification
Aluminum oxide	In Vitro	Not mutagenic

Carcinogenicity

Name	Route	Species	Value
Silane treated ceramic	Inhalation	similar compounds	Some positive data exist, but the data are not sufficient for classification
Triethylene glycol dimethacrylate	Dermal	Mouse	Not carcinogenic
Aluminum oxide	Inhalation	Rat	Not carcinogenic

Reproductive Toxicity

Reproductive and/or Developmental Effects

Name	Route	Value	Species	Test result	Exposure Duration
Bis-GMA	Ingestion	Not classified for female reproduction	Mouse	NOAEL 0.8 mg/kg/day	prematuring & during gestation
Bis-GMA	Ingestion	Not classified for male reproduction	Mouse	NOAEL 0.8 mg/kg/day	prematuring & during gestation
Bis-GMA	Ingestion	Not classified for development	Mouse	NOAEL 0.8 mg/kg/day	prematuring & during gestation
Triethylene glycol dimethacrylate	Ingestion	Not classified for female reproduction	Mouse	NOAEL 1 mg/kg/day	1 generation
Triethylene glycol dimethacrylate	Ingestion	Not classified for male reproduction	Mouse	NOAEL 1 mg/kg/day	1 generation
Triethylene glycol dimethacrylate	Ingestion	Not classified for development	Mouse	NOAEL 1 mg/kg/day	1 generation

Target Organ(s)

Specific Target Organ Toxicity - single exposure

For the component/components, either no data is currently available or the data is not sufficient for classification.

Specific Target Organ Toxicity - repeated exposure

Name	Route	Target Organ(s)	Value	Species	Test result	Exposure Duration
Silane treated ceramic	Inhalation	pulmonary fibrosis	Not classified	similar compounds	NOAEL Not available	
Bis-GMA	Ingestion	endocrine system liver nervous system kidney and/or bladder	Not classified	Mouse	NOAEL 0.8 mg/kg/day	prematuring & during gestation
Triethylene glycol dimethacrylate	Dermal	kidney and/or bladder blood	Not classified	Mouse	NOAEL 833 mg/kg/day	78 weeks
Aluminum oxide	Inhalation	pneumoconiosis	Some positive data exist, but the data are not sufficient for classification	Human	NOAEL Not available	occupational exposure
Aluminum oxide	Inhalation	pulmonary fibrosis	Not classified	Human	NOAEL Not available	occupational exposure

Aspiration Hazard

For the component/components, either no data is currently available or the data is not sufficient for classification.

Please contact the address or phone number listed on the first page of the SIS for additional toxicological information on this material and/or its components.

The product was evaluated by a toxicologist to be safe for its intended use.

SECTION 12: Ecological information

The information below may not agree with the EU material classification in Section 2 and/or the ingredient classifications in Section 3 if specific ingredient classifications are mandated by a competent authority. In addition, statements and data presented in Section 12 are based on UN GHS calculation rules and classifications derived from 3M assessments.

12.1. Toxicity

No product test data available.

Material	CAS #	Organism	Type	Exposure	Test endpoint	Test result
Silane treated ceramic	444758-98-9		Data not available or insufficient for classification			
Dimethacrylate (BIS-MEPP)	41637-38-1	Green algae	Endpoint not reached	72 hours	EC50	>100 mg/l
Dimethacrylate (BIS-MEPP)	41637-38-1	Green algae	Experimental	72 hours	NOEC	0.05 mg/l
Urethane dimethacrylate (UDMA)	72869-86-4	Green algae	Endpoint not reached	72 hours	Effect Growth Rate Conc 50%	>100 mg/l
Urethane dimethacrylate (UDMA)	72869-86-4	Water flea	Experimental	48 hours	EC50	>100 mg/l
Urethane dimethacrylate (UDMA)	72869-86-4	Zebra Fish	Experimental	96 hours	LC50	10.1 mg/l
Urethane dimethacrylate (UDMA)	72869-86-4	Green algae	Endpoint not reached	72 hours	Effect Conc. 10% - Growth Rate	>100 mg/l

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Bis-GMA	1565-94-2		Data not available or insufficient for classification			
Triethylene glycol dimethacrylate	109-16-0	Green Algae	Experimental	72 hours	EC50	>100 mg/l
Triethylene glycol dimethacrylate	109-16-0	Zebra Fish	Experimental	96 hours	LC50	16.4 mg/l
Triethylene glycol dimethacrylate	109-16-0	Green algae	Experimental	72 hours	NOEC	18.6 mg/l
Triethylene glycol dimethacrylate	109-16-0	Water flea	Experimental	21 days	NOEC	32 mg/l
Aluminum oxide	1344-28-1	Fish	Experimental	96 hours	LC50	>100 mg/l
Aluminum oxide	1344-28-1	Green Algae	Experimental	72 hours	EC50	>100 mg/l
Aluminum oxide	1344-28-1	Water flea	Experimental	48 hours	LC50	>100 mg/l
Aluminum oxide	1344-28-1	Green Algae	Experimental	72 hours	NOEC	>100 mg/l

12.2. Persistence and degradability

Material	CAS Nbr	Test type	Duration	Study Type	Test result	Protocol
Silane treated ceramic	444758-98-9	Data not available or insufficient			N/A	
Dimethacrylate (BIS-MEPP)	41637-38-1	Estimated Biodegradation	28 days	CO2 evolution	7-12 % weight	OECD 301B - Modified sturm or CO2
Urethane dimethacrylate (UDMA)	72869-86-4	Experimental Biodegradation	28 days	CO2 evolution	22 %CO2 evolution/THC O2 evolution (does not pass 10-day window)	OECD 301B - Modified sturm or CO2
Bis-GMA	1565-94-2	Estimated Biodegradation	28 days	BOD	32 % weight	OECD 301C - MITI test (I)
Triethylene glycol dimethacrylate	109-16-0	Experimental Biodegradation	28 days	CO2 evolution	85 % weight	OECD 301B - Modified sturm or CO2
Aluminum oxide	1344-28-1	Data not available or insufficient			N/A	

12.3 : Bioaccumulative potential

Material	Cas No.	Test type	Duration	Study Type	Test result	Protocol
Silane treated ceramic	444758-98-9	Data not available or insufficient for classification	N/A	N/A	N/A	N/A
Dimethacrylate (BIS-MEPP)	41637-38-1	Estimated Bioconcentration		Bioaccumulation factor	6.6	Estimated: Bioconcentration factor
Urethane dimethacrylate (UDMA)	72869-86-4	Experimental Bioconcentration		Log Kow	3.39	Other methods
Bis-GMA	1565-94-2	Estimated Bioconcentration		Bioaccumulation factor	5.8	Estimated: Bioconcentration factor
Triethylene glycol dimethacrylate	109-16-0	Experimental Bioconcentration		Log Kow	2.3	Other methods
Aluminum oxide	1344-28-1	Data not available or insufficient for classification	N/A	N/A	N/A	N/A

12.4. Mobility in soil

Please contact manufacturer for more details

12.5. Results of the PBT and vPvB assessment

This material does not contain any substances that are assessed to be a PBT or vPvB

12.6. Other adverse effects

No information available.

SECTION 13: Disposal considerations

13.1 Waste treatment methods

Dispose of contents/ container in accordance with the local/regional/national/international regulations.

Refer to Instructions for Use (IFU) for more information.

EU waste code (product as sold)

180106* Chemicals consisting of or containing dangerous substances.

SECTION 14: Transportation information

70-2010-2225-1,	70-2010-2226-9,	70-2010-2227-7,	70-2010-2228-5,
70-2010-2229-3,	70-2010-2232-7,	70-2010-2233-5,	70-2010-2234-3,
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70-2010-2243-4,	70-2010-2244-2,	70-2010-2245-9,	70-2010-2248-3,
70-2010-2249-1,	70-2010-2250-9,	70-2010-2253-3,	70-2010-2254-1,
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70-2010-5208-4,	70-2010-5209-2,	70-2010-5212-6,	70-2010-5213-4,
70-2010-5214-2,	70-2010-5565-7,	70-2014-1122-3,	70-2014-1123-1,
70-2014-1124-9,	70-2014-1125-6,	70-2014-1126-4,	70-2014-1127-2,
70-2014-1128-0,	70-2014-1129-8,	70-2014-1130-6,	70-2014-1131-4,
70-2014-1132-2,	70-2014-1133-0		

Not hazardous for transportation

70-2010-9550-5

70-2010-9552-1

70-2010-9553-9

70-2010-9554-7

70-2010-9555-4

70-2010-9556-2

70-2010-9557-0

70-2010-9558-8

70-2010-9559-6

70-2010-9560-4

70-2010-5216-7

70-2010-9561-2

70-2010-9563-8

70-2010-9564-6

70-2010-9565-3

70-2010-9566-1

70-2010-9567-9

70-2010-9568-7

70-2010-9605-7

AH-0105-6585-3

UU-0091-3103-6

70-2010-5217-5

UU-0091-3104-4

UU-0091-3105-1

UU-0091-3106-9

UU-0091-3107-7

UU-0091-3109-3

UU-0091-3110-1

UU-0091-3111-9

UU-0091-3112-7

UU-0091-3113-5

UU-0091-3114-3

70-2010-5218-3

70-2010-5219-1

70-2010-9546-3

70-2010-9547-1

70-2010-9548-9

70-2010-9549-7

ADR/IATA/IMDG: Not hazardous for transport.

SECTION 15: Regulatory information

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

Global inventory status

Contact the manufacturer for more information

SECTION 16: Other information

List of relevant H statements

H317	May cause an allergic skin reaction.
H412	Harmful to aquatic life with long lasting effects.
H413	May cause long lasting harmful effects to aquatic life.

Revision information:

Revision information not available

The product to which this Safety Information Sheet applies is classified as a medical device according to the EU Medical Device Regulation EU 2017/745. _x000D_
Medical devices which are invasive or used in direct physical contact with the human body are exempt from the requirements of classification and labelling according to Regulation (EC) No. 1272/2008 (CLP; Article 1, paragraph 5). _x000D_
The EU Medical Device Regulation does not foresee the use of Safety Data sheets for medical devices which are invasive or used in direct physical contact with the human body, as the safe use of the product is described through the Instructions for Use and /or the labelling for the product. Nevertheless, the 3M Safety Information Sheet is provided as a further service to customers to provide additional toxicology and chemical information on the product. In case of further questions, please contact your 3M representative listed on the Safety Information Sheet.

3M United Kingdom Safety Information Sheets are available at www.3M.com/uk