

15/10/2019 -Issue 1

Technical Data Sheet- TDS08 (Lachrymal Cannulae)

Please see Appendix 1 for product specific information including ordering references & materials

**1. GENERAL INFORMATION****1.1 Intended Use**

Surgically invasive device for probing and irrigating the lachrymal ducts. They are intended for transient use.

**1.2 Certification**

All Sterimedix Ltd products are manufactured at the site below

STERIMEDIX FACILITY	NOTIFIED BODY
Thornhill Road North Moons Moat Redditch Worcestershire B98 9ND United Kingdom	NB Number 0123 TUV SUD Product Service GmbH, Ridlerstr. 65, D-80339 Munich

Sterimedix has appointed a European Representative (EC Rep) as per the below

***Bausch & Lomb GmbH, Brunsbutteler Damm, 165-173, 13581, Berlin, Germany***

**1.3 Packaging Materials****Primary packaging****Either -**

Blister - Pentamed® Film (PET'G'/'A'PET/PET'G')

Blister lid - Tyvek® with all over lacquer

**or –**

In Protector without blister – Protector – Polyethylene

**Secondary packaging –**

Sterile: Box - White Board

Sterile or Non-Sterile: Bulk - Double PE sealed bags labelled on the inner bag

**1.4 Materials of Concern**

MATERIAL	DECLARATION
DEHP/Phthalates	Devices do not contain referenced material
Latex	Devices do not contain referenced material
PVC	Devices do not contain referenced material
Bisphenol A (BPA)	Some devices may contain very low levels of BPA CAS 80-05-7 as a residue from the epoxy synthesis process which is used as a bonding agent. However, levels are below the 0.1% SVHC threshold

15/10/2019 -Issue 1

Technical Data Sheet- TDS08 (Lachrymal Cannulae)

Substances of animal origin	Devices do not contain referenced material
Conflict Minerals	Devices are free from any Conflict Minerals referring to gold, tin, tantalum and tungsten, the derivatives of cassiterite, columbite-tantalite and wolframite, extracted in the Democratic Republic of the Congo (DRC) and surrounding countries.

**1.5 REACH**

Based on current available information Sterimedix has not been able to identify any substances above 0.1% in any of its final devices which have been included on the Candidate List as a SVHC.

**1.6 Instructions for Use (IFU)**

As per Annex I, 13.1, Paragraph 4 by way of exception, no such instructions for use are needed for devices in Class I or IIa if they can be used safely without any such instructions. Therefore, unless otherwise stated, Sterimedix does not provide IFU's.

**1.7 Shelf Life**

Shelf life for all Sterile devices is 5 years

Non-Sterile N/A

There are no specific storage or transport conditions. Generic instructions such as Store in a cool dry place, keep out of direct sunlight, do not use if box is damaged are displayed on the box packaging with the appropriate symbology.

**1.8 Sterilisation, Resterilisation/Reprocessing**

All Sterimedix devices are validated to be sterilised via EtO Cycle 25 only at the Sterigenics Plant in Derbyshire\*. After initial sterilisation, Sterimedix devices may be resterilised using the same EtO cycle once more. Sterimedix devices can be sterilised a maximum of two times\*\*.

\*Those using another EtO sterilisation cycle for Sterimedix devices are responsible for validating their own cycle to ensure sterility has been achieved and can be maintained.

\*\*Some devices may have already undergone two sterilisation cycles before leaving Sterimedix. This will be represented by a "Z" at the end of the Lot number of the batch.

Further information be found in declaration "ST41 Sterimedix Sterilisation reprocessing instructions".

**1.9 Sterile & Non-Sterile products**

Non-Sterile devices are identical to Sterile devices in materials, manufacturing and control, with the exception of the packaging- and sterilisation-process

**1.10 Classification**

Class IIa in accordance with Annex IX Rule 6

# Sterimedix Ltd

Quality System Document QD 2125 Issue B  
Technical Data Sheet Template

15/10/2019 -Issue 1

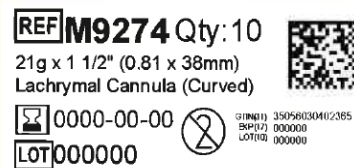
## Technical Data Sheet- TDS08 (Lachrymal Cannulae)

### 1.11 GMDN Code

GMDN code is 10575

## 2. LABELLING

### 2.1 Example Label(s)



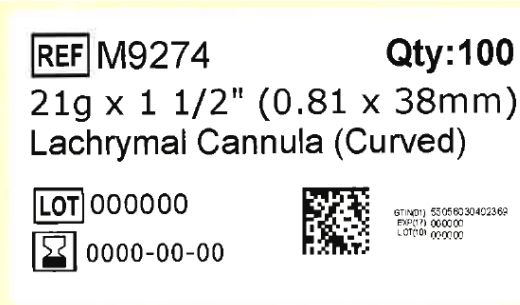
LBL0008 – Sterile boxes of 10



LBL0020 – Sterile bulk



LBL0012 – Non-Sterile bulk



LBL0024 – Sterile boxes of 100

15/10/2019 -Issue 1

Technical Data Sheet- TDS08 (Lachrymal Cannulae)

**3. DISCLAIMER**

All information has been collated to the best of our knowledge and based on information currently available to us. This document can be updated without further notification.

15/10/2019 -Issue 1

### Technical Data Sheet- TDS08 (Lachrymal Cannulae)

#### Appendix 1- Product specific information

PRODUCT SUB-GROUP	PRODUCT CODE	LABEL DESCRIPTION	HUB COLOUR	ORDERING REFERENCE	MATERIALS
Lachrymal Cannulae	M9272	23g x 1/2" (0.64 x 12.5mm) 0.3mm Port Lachrymal Cannula (Bullet tip)	Blue	M9272-02/10	Polypropylene, Stainless Steel Grade 304, Single Part Epoxy Adhesive
	M9274	21g x 1 1/2" (0.81 x 38mm) Lachrymal Cannula (Curved)	Green	M9274, M9274-02/10, M9274-02/100, 9274-02/Y, 9274NS-02/Y	
	M9275	26g x 3/4" (0.45 x 19mm) Short Lachrymal Cannula (Straight)	Brown	M9275-02/10, M9275-02/100, 9275-03NSP/Y	
	M9276	26g x 1 1/4" (0.45 x 32mm) Lachrymal Cannula (Curved)	Brown	M9276-02/10, M9276-02/100, 9276NS-02/Y	
	M9277	25g x 1 1/8" (0.5 x 29mm) Lachrymal Cannula (Curved)	Orange	M9277, M9277-02/10, M9277-02/100, 9277NS-02/Y	
	M9278	25g x 7/8" (0.5 x 22mm) 0.2mm Port Lachrymal Cannula (Straight)	Orange	M9278-02/10, M9278-02/100, 9278-03NSP/Y	
	M9279	19g/23g Reducing x 1 1/2" (1.1/0.64 x 38mm) Lachrymal Cannula	Cream	M9279-02/10, M9279-02/100	
	M9282	26g (0.45 x 30mm) Lachrymal Cannula (Curved)	Brown	9282-02/Y	

KEY	
REFERENCE	DEFINITION
-02/X	Sterile, Blister Packed, Box of X
-03NSP/Y	Non-Sterile, in a Printed Protector, in Bulk
NS-02/Y	Non-Sterile, Blister Packed, in Bulk
-02/Y	Sterile, in Bulk