

DEMO — Fictional product. This page demonstrates the eIFU.one platform; it does not belong to a real medical device and is not for clinical use.

Carotid Shunt Set — Electronic Instructions for Use

Sample Medical Ltd. · REF KM-CS-2210 · MDR Class IIb · Version 1

GTIN / UDI-DI	09506000134000
Catalogue no. (REF)	KM-CS-2210
Regulation	MDR 2017/745 · EU 2021/2226
Intended user	Healthcare professional

1. Intended purpose

The manufacturer's approved intended-purpose text appears here.

eIFU.one takes this section from the manufacturer's approved PDF; human approval is required before publishing.

2. Device description

Device components and materials appear here.

Each revision is archived as a separate version; the address (this page) never changes.

3. Contraindications, warnings and precautions

Approved warning texts appear here.

Healthcare professionals can ask the AI assistant about this section; answers draw only on this IFU.

4. Directions for use

Step-by-step directions appear here.

Scanning the GS1 DataMatrix on the label via /01/{GTIN}/10/{lot} opens the version for that lot.

5. Storage and disposal

Storage conditions and disposal information appear here.

The IFU stays available at this address for the expected device lifetime; the period is tracked.

6. Paper copy request

The paper IFU request channel appears here.

Users can request a paper copy free of charge (EU 2021/2226). The channel is defined on the device record.

7. Symbols

Explanation of the ISO 15223-1 symbols used on the label appears here.