Version:

2

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#### Important Information 1



Read this Instruction for Use carefully before every application and keep it easily accessible for all users or the respective specialist staff.



Carefully read the warnings marked with this symbol. Improper use of the products may result in serious injuries to the patient, the users or third parties.

#### Scope 2

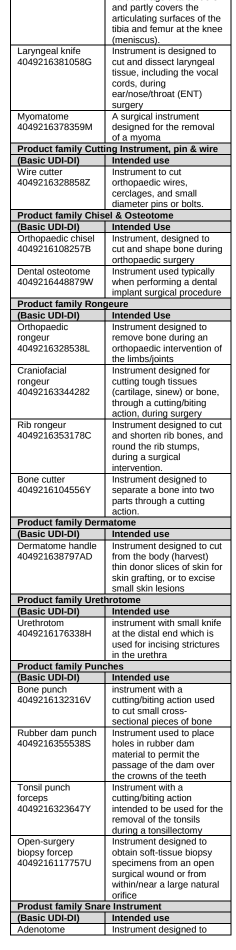
The instruments must be used according to their intended use in the medical fields and by respectively trained and qualified staff only. The treating physician and/or user is responsible for choosing the equipment for specific applications and/or operative use, for the appropriate training and information, and for the sufficient experience regarding the handling of the equipment.

#### 3 Products / Intended use

The cutting instruments are intended for surgically invasive and partly also for non-surgically invasive treatments in various fields of medicine (of less than 60 min.). They correspond to risk class I/Ir.

Product family Scissors		
(Basic UDI-DI)	Intended use	
General-purpose	Instrument designed to cut	
surgical scissors,	a variety of tissues during	
4049216387279Q	open surgery	
Nail scissors	A hand-held device	
4049216134957Z	designed specifically for	
	cutting the nails of a	
	patient.	
Nail clippers,	A device with an integral	
reusable	or exchangeable knife-like	
40492161269584	blade used to remove	
	facial or body hair from a	
	patient.	
Bandage scissors,	A hand-held manual	
reusable	instrument intended to be	
4049216134817N	used to cut bandages and	
later e sul s	cloth.	
Intraocular	Ophthalmic surgical	
scissors 40492161348884	instrument intended to be used to cut intraocular	
40492101348884	tissue of the anterior	
	segment (e.g., iris)	
Ear scissors,	Instrument designed to cut	
reusable	tissue during ear surgery	
4049216334147V	lioodo dalling our ourgory	
Suture scissors	A surgical instrument used	
40492161350275	during surgery to cut	
	suture or ligature material	
Tonsil scissors	Surgical instrument used	
4049216165207U	to cut tonsil tissue during	
	ear/nose/throat (ENT)	
	surgery.	
Nasal scissors	A surgical instrument used	
4049216134957Z	to cut tissue during	
	ear/nose/throat (ENT) or	
	plastic surgery in or on the	
	nose and its associated	
Line hills at a start	structures.	
Umbilical cord	Instrument designed specifically to cut the	
scissors, reusable 4049216326027R	umbilical cord after birth	
Cast cutting	Surgical instrument with	
scissors	specially strong, curved	
4049216463148Q	blades that is used to cut	
10102104001400	small plaster cast	
	dressings and layers of	
	thick material, e.g.,	
	bandages.	
	. 0	

Rectal scissors	A surgical instrument
40492161350173	designed to cut tissue
	during rectal surgery.
Enucleation	An ophthalmic surgical
scissors 40492161348782	instrument used to cut tissue during eye surgery
40492101340702	involving enucleation of
	the eye and/or its related
	structures (i.e., the
	removal of the eyeball to,
	e.g., remove a malignant tumour or to relieve
	intolerable pain in a blind
	eye).
Rigid endoscopy	Rigid instrument used in
scissors	endoscopic procedures for
4049216464218S	cutting tissue or suture material during an
	endoscopic procedure.
Product family Micr	o scissors
(Basic UDI-DI)	Intended use
Micro scissors	A hand-held, manual
4049216443548N	surgical instrument
	designed to cut a blood vessel (i.e., to cut over a
	blood vessel or cut
	longitudinally to split it
	open) during open
	vascular surgery
Product family Cutt	
(Basic UDI-DI) Scalpel handle,	A device that is an
reusable	interchangeable
4049216122356W	component of a scalpel
	and that functions as a
	handle designed to mount
Scalpel, reusable,	a compatible blade Instrument constructed as
4049216351417X	a one-piece handle and
10102100011111	scalpel blade (not an
	exchangeable component)
	used by the operator to
	manually cut or dissect tissue.
Amputation knife	Instrument which is used
4049216351317U	in surgery for the
	amputation of a limb
Periodontal knife	Dental instrument used to
40492164154488	excise the gums and other
	oral soft tissue during a periodontal intervention
Tonsil knife	A surgical instrument
40492161225574	intended for the removal of
	the tonsils during a
Nee al lucif	surgical intervention
Nasal knife 4049216381038C	Instrument designed to cut and dissect internal nasal
-0-32 1030 10300	tissue, including the nasal
	septum, during
	ear/nose/throat (ENT)
For Init-	surgery
Ear knife 4049216122456Z	Instrument intended to cut and dissect tissues of the
	ear during a surgical
	intervention
Ophthalmic knife	Ophthalmic surgical
4049216327648L	instrument designed to
	make precise incisions in the eye and surrounding
	tissues during ophthalmic
	surgery.
Cartilage knife	Instrument designed for
4049216378409E	cutting, shaving or shaping
	cartilage during a surgical intervention.
Cut-throat razor	A device with an integral
4049216450788U	or exchangeable knife-like
	blade used to remove
	facial or body hair from a
	patient.
Cast/plastor knife	Instrument which is used
Cast/plaster knife 4049216351407V	Instrument which is used to cut or trim the plaster of
Cast/plaster knife 4049216351407V	to cut or trim the plaster of a cast.
4049216351407V Meniscus knife	to cut or trim the plaster of a cast. A surgical instrument
4049216351407V	to cut or trim the plaster of a cast. A surgical instrument designed to apply
4049216351407V Meniscus knife	to cut or trim the plaster of a cast. A surgical instrument



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fibrocartilage that borders

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09.02.2023

Cutting, Removing Instruments



40492161002563	excise hypertrophic
	lymphoid tissue in the
	nasopharynx
Lens loop	Instrument designed for
40492161231975	eye lens extraction and
	gentle manipulation and/or
	irrigation of eye tissues
	during an ophthalmic
	surgical procedure.
Nasal snare	Instrument intended to be
4049216156768M	inserted into the naris for
	the removal of tissue,
	typically polyps, tumours,
	and other abnormal tissue
	from the nasal cavity
	during ear/nose/throat
Ear snare	(ENT) surgery
4049216310206N	the ear to remove tissue,
40492103102001	typically tumorous or
	damaged tissue, from the
	ear during ear/nose/throat
	(ENT) surgery
Tonsil snare	Instrument inserted into
4049216136327K	the oral cavity to remove
10102101000271	the tonsils during
	ear/nose/throat (ENT)
	surgery
L	ourgory

## 4 Contraindication

Valid from:

The instruments may only be used for their intended purpose by appropriately trained and qualified personnel. The products are not intended for use on the heart and the central circulatory and nervous system.

The products are not intended for connection to active medical devices. There is a risk of injury to patients and users when using RF, RF or laser devices simultaneously.

## Product specific contraindications

## Manual dermatome:

- Bacterially contaminated wound bed
- Non-perfused wound bed (tendons, bones, joint capsule)
- Heavy mechanical stress at the recipient site
- Exposed vessels or nerves
- Exposed implantsAesthetic defects on the face
- Relative: defects on the flexion side of joints (secondary shrinkage of grafts)

### Urethrotome:

- Urinary tract infection
- Coagulation disorders

## Snare Instruments:

- Agranulocytosis
- Leukemia
- Coagulation disordersCardiovascular insufficiency

# 5 Complications / Side effect

### ⚠ General:

After contact with the instrument, hypersensitivity reactions can be triggered in a patient with material intolerances to stainless steel. In the event of such a reaction, the procedure must be discontinued immediately and the necessary steps taken.

In the course of market monitoring, further potential complications / side effects were identified:

# $\underline{\Lambda}$ Treatment-related complications / side effects / risks

### General:

- Injury to surrounding vessels and tissues
- Injury to nerves

2.2\_IFU\_Cutting\_Removing\_v02.doc

## Manual Dermatome:

 Removal of transplantat that are too deep because the dermatome was not set correctly: In this case, the transplantat can either be immediately refixed at the removal site like a graft or this lifting site can be covered with another correctly obtained graft.

2

## Urethrotome:

- latrogenic lesion of the urethra due to inadequate control of the incision.
- Bleeding from the urethraPenile or scrotal hematoma

Version:

- Penne or scrotal nematoma
  Urinary tract infection, urethritis, prostatitis,
- epididymitisUrethral perforation with formation of a via
- falsaPenile deviation
- Fernie deviation

## Snare Instruments:

- Risks associated with tonsillectomy
- RebleedingRevision surgery due to secondary bleeding
- tooth damage
- nerve damage
- Airway obstruction (edema)
- Emphysema
- Tasting disorders

# $\underline{\Lambda}$ Product-related complications / side effects / risks

In the course of market monitoring, further potential complications / side effects could be identified: **Scissors:** 

## Fracture

- Possible unwanted perforation
- Cutting instruments:
- Incorrect information in the IFU regarding disassembly of the instrument.
- Breakage of the blades
  - Breakage of working ends due to lever
- movementsSwallowing of components after breakage

#### Cutting instruments, pins & wires:

Breakage of the cutting edge, componentsCutting edges defect

## Chisels, Osteotomes:

- Breakage of the cutting edge, components
- Cutting edges defect
- Rust on blade
- Blades not compatible with handle
- Wrong labeling
- Residues (reprocessing)
  Blunt blades

## Rongeure

- Breakage of jaws
- 6 Precautions and Warnings

#### Attention!

The instruments are designed for surgical use only and must not be used for any other purpose. Improper handling and care as well as improper use can lead to premature wear of the instruments.

#### A Material intolerance

Under no circumstances must the instruments be used if the user or specialist staff become aware of the patient being intolerant to the material.

#### ⚠ Functional Impairment

Surgical instruments corrode and become impaired in their functionality if they come into contact with aggressive substances. It is therefore necessary to observe the storage and Sterilization instructions.

#### 🗥 Operating Conditions

The aforementioned products require correct maintenance and care in order to guarantee that the products operate safely. In addition to this, functionality testing and a visual check should be performed prior to each application. For this reason, please pay attention to the respective chapters in this Instruction for Use.

#### 

Should the products be reassembled after disassembly, individual parts must not be replaced v02 with parts from other manufacturers! If the intended purpose of the product entails certain parts being exchanged (e.g. different attachments), no parts from different manufacturers must be used! We recommend to also purchase other accessories (e.g. detergents) at CM Instrumente GmbH.

#### A Storage

There are no specific storage requirements concerning the products. Nevertheless, we recommend storing medical products in a clean and dry environment.

#### ▲ Creutzfeldt Jakob Disease

With regard to the reprocessing of medical devices that have been used on patients or suspected patients suffering from or suspected of suffering from Creutzfeldt-Jacob disease (CJD) or its variant (vCJD), the requirements specified in the corresponding appendix of the guidelines for hospital hygiene and infection prevention and the requirements specified by publications in the Federal Health Gazette must be adhered to. The medical devices that were used on this group of patients must be disposed of by incineration (European Waste Catalogue EAK 18 01 03) without risk. Dry heat, ethanol, formaldehyde and glutaraldehyde have a fixing but no inactivating effect on TSE pathogens. Of the sterilization methods available, only steam sterilization (especially 134°C, 18 minutes) has been shown to have a limited effect.

#### ⚠ Pointed / sharp instruments

Care must be taken when handling instruments with sharp points or edges.

#### 7 Combination products & accessories

The products are not applied with other products except scalpels and offered without accessories. Scalpels with interchangeable blades Scalpels can be combined with blades according to DIN EN 27740. The scalpels are designed to be compatible with figures 3, 4 according to DIN 58849-2.

#### 8 Liability and Warranty

As a manufacturer, CM Instrumente GmbH is not liable for consequential damage resulting from improper use or handling. This particularly applies to use which is not compliant with the defined intended use, or non-compliance with the instructions on preparation and Sterilization. This also applies to repairs or changes to the product which are not carried out by authorized staff of the manufacturer. These disclaimers also apply to warranty services.

#### 9 Sterility

#### ▲ State upon Delivery

Medical products are delivered in a non-sterile condition and need to be prepared and sterilised by the user prior to the first application and any subsequent application according to the following instructions.

#### 10 Reprocessing

#### ⚠ Warnings

- Frequent reprocessing impairs the quality of the products.
- City water to be used must comply with COUNCIL DIRECTIVE 98/83/EC of 3 November 1998 on the quality of water intended for human consumption.
- This treatment instruction specifies the detergents and disinfectants used for validation. If an alternative detergent and disinfectant (RKI or VAH listed) is used, the responsibility rests with the reprocessor.
- Reassemble disassembled products before sterilization.
- Reprocessing may only be performed by qualified medical personnel. Machine reprocessing must be qualified and validated by the user. The washer-disinfectors must

2/4

Instruction for Use Cutting, Removing Instruments				
Valid from:	09.02.2023	Version:	2	

fully comply with the requirements of DIN EN ISO 15883.

#### **∆Use Site**

The first steps of a proper reprocessing take place in the operating theatre. Coarse contaminations must be removed prior to storing the instruments if possible. For this purpose, the instruments should be rinsed under cold tap water (<40°C). If this procedure is not sufficient to remove the obvious soiling, a soft plastic brush can be used to remove soiling.

Whenever possible, dry removal (moistened, closed system) should be the method of choice. A drying of any residues should be avoided! Wherever possible, dry disposal is to be preferred, since with wet disposal the prolonged lying of the medical devices in solutions can lead to material damage (e.g. corrosion). Long periods of waiting until the reprocessing, for instance overnight or over the weekend, must be avoided with both types of removal (<60 minutes).

#### ⚠Transport

The products must be disposed of in a dry state immediately (<60 min) after use, if possible. This means that the products have to be transported in a closed container from the place of application to the purification, so that the products do not dry up.

#### Preparing the Decontamination

The products must be disassembled prior to the following reprocessing steps and/or must be exposed to the following reprocessing steps in an open condition, where possible. Rinse residue must be avoided. The products must be reprocessed in appropriate screen baskets or rinsing shields (choose size according to product). The products must be positioned in the cleaning basket at a minimum clearance from one another. Avoid overlapping so that the damaging of the products during the cleaning process can be excluded.

#### Pre-cleaning

1. Pre-clean products completely under cold water (city water drinking water quality <40°C) with a soft brush.

2. Flush cavities and hard-to-reach areas, gaps and slots on the instrument with cold water (city water drinking water quality <40°C) for 60 sec using a water pressure gun.

3. Soak products in an alkaline cleaner (0.5 % Neodisher Mediclean forte) in an ultrasonic bath at 35 kHz for 5 min.

4. Rinse products under cold water (city water

drinking water quality <40°C) for 15 sec. 5. Flush cavities and hard-to-reach areas, gaps and slots on the instrument with cold water (city water drinking water quality <40°C) for 30 sec using a water pressure gun.

#### **Cleaning/disinfection**

Automated cleaning and/or disinfection process

(Miele Disinfector G7835 CD as per ISO 15883):

- 1 Pre-clean for 1 minute
- Drain water
- Pre-clean for 4 minutes
- Drain water
- Clean for 6 minutes at 58°C +/- 1°C using 0.5 % alkaline detergent (0,5 % Neodisher Mediclean forte)
- Drain water
- Neutralize for 3 minutes (0.1% 0,1 % NeodisherZ) with cold tap water suitable for use as drinking water <40°C Drain water
- Clean for 2 minutes with FD water <40°C

## Automated Disinfection

Automated thermal disinfection in a cleaning and disinfection device taking into consideration the national requirements for the A0 value; for instance, A0 value 3000:

>5 minutes at >95°C

2.2\_IFU\_Cutting\_Removing\_v02.doc

#### Automated Drying

Automated drying in accordance with the drying operation of the cleaning and disinfection device for at least 30 minutes at 92°C +/- 2°C.

#### 11 Sterilization

(Typ B Autoclave by Tuttmauer as per DIN EN 13060)

Sterilization of products with a fractionated prevacuum method (according to DIN EN ISO 17665-1) taking into consideration the respective national requirements. The Sterilization of the products must be conducted in suitable Sterilization packaging according to DIN EN ISO 11607-1 and EN 868

The Sterilization must be completed using a fractionated pre-vacuum method with the following parameters:

- 134°C.
- 5 minutes hold time
- 3 pre-vacuum cycles

Drying in vacuum for at least 20 minutes

The Instruction for Use of the manufacturer of the autoclave and the recommended directions for maximum loading with goods to be sterilised must be observed. The autoclave must be installed. maintained, validated and calibrated in accordance with requirements.

#### Additional Information

The reprocessor is responsible for ensuring that the actual reprocessing, including the used equipment, materials and the staff involved in the reprocessing facility, achieves the desired results. This typically requires the validation and routine monitoring of the method and the equipment used.

#### 12 Maintenance-Control-Inspection

Cool down the instruments to room temperature!

#### Visual inspection (before assembly)

Check the surface of the instruments or the individual components before assembly. Pay particular attention to checking joints (final part), profiles, grooves and other structures that are difficult to access:

Is there any residual soiling or residue? If so, manual re-cleaning and renewed complete mechanical cleaning and disinfection

- Are traces of corrosion (rust, pitting) visible? Is the surface damaged by cracks (including
- hairline cracks) or other signs of wear? Is the instrument labeling no longer legible?

If so, the instrument in question must be marked and immediately sorted out and replaced.

#### Assembly and maintenance

Assemble the disassembled instruments in a functionally correct manner.

- Treat moving parts, such as joints, threads and sliding surfaces, manually with suitable, medically approved instrument oil (steam sterilizable care product based on paraffin/white oil, biocompatible according to EU standard). EU standard).
- Distribute the oil in the joint by opening and closing several times, remove excess care product with a clean, lint-free cloth.

Do not use mineral oil or silicone lubricant! Do not immerse instruments completely in the care product!

#### Function test

During the functional check, pay particular attention to the following aspects and possible malfunctions:

- No damage, such as broken tips, bent or loose parts (screws)
- Proper closure of jaws
- Correct and safe function of detents and locks
- Easy and even movement of handles, as backlash-free as possible v02

- Proper cutting function of shears
- Re- and spring pressure in order (punches, gouge pliers etc.)
- Continuity of lumens
- No other signs of wear, e.g. on seals, insulation or coatings If defects are found during the functional test, the

instruments must be marked and excluded from further use without fail

#### 13 Lifespan of the Products

The service life of the products results from their function, gentle reprocessing in accordance with these instructions and careful handling when handling the instruments. Therefore, a limit to the number of reprocessing cycles cannot be set across the board. Nevertheless, 100 reprocessing cycles were simulated, which showed no impairment of functionality, biocompatibility and identification of the products. The user recognizes the end of the usage cycle by the possible defects and limiting properties of the products indicated under maintenance, inspection and testing.

#### 14 Service and Repair

#### $\triangle$ Service and Repair

Do not carry out any repairs or changes to the product yourself. Authorized staff of the manufacturer are solely responsible for such work. Should you wish to make complaints or queries, or offer us any advice regarding our products, please feel free to contact us.

#### 🗥 Returns

Defective or non-compliant products must go through the entire reprocessing process before being sent back for repairs/service

#### 15 Packaging, Storage and Disposal

Standard packaging of the products for Sterilization according to ISO 11607 and EN 868.

Store sterile products in a dry, clean, and dust-free environment, secured against damage, at moderate temperatures.

The medical products of the manufacturer should be stored and kept in single packaging, boxes or protective containers. Please handle the instruments with care during transportation, storage and reprocessing. The user and/or specialist staff intended for this is responsible for ensuring that the sterile state is maintained after the Sterilization.

The disposal of the products, packaging as well as the accessories must be performed in accordance with current rules and laws. No specific instruction regarding this matter is provided by the manufacturer.

#### 16 Reporting obligations

Product defects which have occurred during proper use of our products should be reported directly to us as the manufacturer or to your supervising specialist dealer.

Defects in which patients, users or third parties have been harmed by the products (so-called reportable incidents) must be reported immediately to the manufacturer and, if necessary, to your competent, responsible authority. This reporting of incidents must take place immediately after they occur so that important reporting deadlines can be met.

The affected products must be discarded, reprocessed and sent to the manufacturer for examination. Your servicing dealer will be pleased to help you with this. After receipt of your notification, we will inform you

within a reasonable time frame about the further measures required.

#### 17 Additional information

If the chemicals and machines described here are not available, and if the reprocessing process cannot be carried out as described, it is the user's responsibility to validate his process accordingly.



	Instruction for Use Cutting, Removing Instruments			
Valid from:	09.02.2023	Version:	2	

Further information on the reprocessing of medical devices:

- Internet: http://www.rki.de
- Internet: http://www.a-k-i.org
- Hygiene requirements for the reprocessing of medical devices Recommendation of the Commission for Hospital Hygiene and Infection Prevention at the Robert Koch Institute (RKI) and the Federal Institute for Drugs and Medical Devices (BfArM) on the "Hygiene requirements for the reprocessing of medical devices"
- DIN 96298-4 Functional control in the reprocessing process

#### 18 Other applicable documents

Instructions for the proper disassembly of the listed products can be found on our homepage: <a href="http://www.cm-instrumente.de/ifu">www.cm-instrumente.de/ifu</a>

Disassembly instructions for instruments

## 19 Description of Symbols Used

$\triangle$	Attention!	
Ĩ	Observe the Instruction fo Use	
REF	Item number	
LOT	Lot designation	
CExxxx	CE labeling, if necessary m identification number of the notified body.	
mile	Indication of a non-sterile product	
	Name and address of the manufacturer	
	Manufacturing date	
MD	Medical device	
UDI	Unique Device Identification, code for identifying a product	
SRN	Registration number of the manufacturer in the EUDAMED database	