



# **EU Quality Management Certificate**



This is to certify that the company

#### **TONTARRA Medizintechnik GmbH**

Daimlerstraße 15 78573 Wurmlingen Germany

SRN: DE-MF-000005520

has established, implemented and maintains a Quality Management System in accordance with

#### Annex IX, Chapter I and III of the Regulation (EU) 2017/745

Conformity Assessment based on a Quality Management System and on Assessment of Technical Documentation

for the device categories and products listed in the Annex of this certificate.

The conformity of the Quality Management System has been verified in an audit and is subject to regular surveillance in accordance with Annex IX, Chapter 1, Section 3. Limitations to this certificate are listed in the Annex.

Devices listed in the Annex may bear the CE marking with the identification number of the Notified Body (0297).

For placing of devices of class III and devices class IIb implantable according to Article 52(4) subparagraph 2 listed in the Annex on the market, an additional certificate according to Annex IX, Chapter II is required.

Certificate registration no. 448891 MDR2017Q

Certificate ID 1000220878
Effective date 2025-03-06
Expiry date 2027-12-14
Frankfurt am Main, 2025-03-06



#### **DQS Medizinprodukte GmbH**

Heinrich von Mettenheim Managing Director





**Certificate ID: 1000220878** 

#### Device categories and variants covered by this certificate:

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Antrum punching

Risk classification: Ir

Basic-UDI-DI: 405018300000369XW

Intended purpose: Antrum punches are used to punch holes in hard fabric.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Biopsy curettes

Risk classification: Ir

Basic-UDI-DI: 405018300000227X9

Intended purpose: Biopsy curettes are used to scrape endometrial secretions and / or

tissue from the uterus.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Gallstone scoops

Risk classification: Ir

Basic-UDI-DI: 405018300000207X3

Intended purpose: Gallstone scoops are used to remove gallstones.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Dissectors

Risk classification: Ir

Basic-UDI-DI: 405018300000206WZ

Intended purpose: Dissectors are used to separate soft tissue or body structures from

each other.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Surgical spoon

Risk classification: Ir

Basic-UDI-DI: 405018300000394XV

405018300000289XX 405018300000394XV 405018300000408XF 405018300000422X9

Intended purpose: A surgical spoon is used to scrape out or remove tissue.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Chisel Risk classification: Ir

Basic-UDI-DI: 405018300000135X3

405018300000136X5 405018300000386XW 405018300000510X7

Intended purpose: A surgical instrument used to cut or contour bone.



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Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Gouge Risk classification: Ir

Basic-UDI-DI: 405018300000292XL

Intended purpose: Gouges are used to separate or to work hard tissue or bones.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Punching

Risk classification: Ir

Basic-UDI-DI: 405018300000395XX

Intended purpose: An ear punch is designed to punch holes in bone and hard tissue.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Osteotome blades

Risk classification: Ir

Basic-UDI-DI: 405018300000293XN

Intended purpose: A device that is an interchangeable component of the orthopaedic

osteotome and acts as a cutting blade.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Elevators

Risk classification: Ir

Basic-UDI-DI: 405018300000353XF

Intended purpose: Elevators of the product group Elevators, others are used to lift and

position tissue and organs.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Elevators

Risk classification: Ir

Basic-UDI-DI: 405018300000389Y4

Intended purpose: Elevators for the ENT area are used to lift and position tissue, suture

material and nerves.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Elevators

Risk classification: Ir

Basic-UDI-DI: 405018300000290XG

Intended purpose: Elevators for orthopaedics are used to lift and dissect bones, tissue

and nerves. They are also used to clean and scrape bones and to

expose fractures.



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Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Elevators

Risk classification: Ir

Basic-UDI-DI: 405018300000349XO

Intended purpose: Elevators are used to lift and position tissue and nerves.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Enucleators

Risk classification: Ir

Basic-UDI-DI: 405018300000339XM

Intended purpose: Enucleators are used to evacuate intact tissue.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Thread catcher

Risk classification: Ir

Basic-UDI-DI: 405018300000146X8

Intended purpose: Thread catchers are used to catch / hold the sewing thread and push

the knot.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Fixation instrument

Risk classification: Ir

Basic-UDI-DI: 405018300000557XZ

Intended purpose: Fixation instruments are used to fix tissue.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Flat drill Risk classification: Ir

Basic-UDI-DI: 405018300000509XN

Intended purpose: Flat drills are used for drilling shallow cavities and for drilling into

cartilage

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Milling cutter

Risk classification: Ir

Basic-UDI-DI: 405018300000327XE

Intended purpose: Milling cutters are used for further processing of drilled holes.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Implatation fork

Risk classification: Ir

Basic-UDI-DI: 405018300000201WP

Intended purpose: Forks are used to hold and lift fabric.



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MDN 1208 - Non-active non-implantable instruments Device category:

Product name: Tissue claw

Risk classification:

405018300000164XA Basic-UDI-DI:

Intended purpose: Fabric claws are used to hold fabric in place.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Tick Risk classification:

405018300000187XN Basic-UDI-DI:

Hooks are used to separate or expose tissue from other anatomical Intended purpose:

parts.

MDN 1208 - Non-active non-implantable instruments Device category:

Product name: Hand drill

Risk classification: Ιr

Basic-UDI-DI: 405018300000325XA

Intended purpose: Hand drills are used to drill holes in bones using a drill.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Chisel handle

Risk classification:

Basic-UDI-DI: 405018300000507XJ

Intended purpose: Handpieces are attached to the proximal end of a surgical instrument.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Sphenoidal punch

Risk classification:

Basic-UDI-DI: 405018300000361XE

Intended purpose: Cuneiform punches are designed to punch holes in bones.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Bone files

Risk classification:

Basic-UDI-DI: 405018300000303WY

Intended purpose: Bone files are used to smooth, grind or shorten bones.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Surgical punch

Risk classification:

Basic-UDI-DI: 405018300000295XS

Intended purpose: A surgical punch for bones is intended to punch holes in bones and

hard tissue and remove tissue.





# Annex to EU Quality Management Certificate SRN of Manufacturer: DE-MF-000005520 Certificate ID: 1000220878

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Cartilage punch

Risk classification: Ir

Basic-UDI-DI: 405018300000304X2

Intended purpose: Cartilage files are used for smoothing, grinding and shortening

cartilage tissue.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Curettes
Risk classification: Ir

Basic-UDI-DI: 405018300000137X7

405018300000340X6 405018300000342XA 405018300000396XZ 405018300000420X5

Intended purpose: Curettes are designed for scraping, shaping and cleaning skin, tissue

or bone

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Micro-instruments

Risk classification: Ir

Basic-UDI-DI: 405018300000088XK

Intended purpose: Knives are used for cutting various fabrics.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Myoma drill

Risk classification: Ir

Basic-UDI-DI: 405018300000250X4

Intended purpose: Myoma drills are used to stabilise and manipulate fibroids.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Needles Risk classification: Ir

Basic-UDI-DI: 405018300000338XK

Intended purpose: Needles are used to puncture tissue.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Nail drill Risk classification: Ir

Basic-UDI-DI: 405018300000324X8

Intended purpose: Nail Drills are used for puncturing and drilling of finger and toenails.



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Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Obturators

Risk classification: Ir

Basic-UDI-DI: 405018300000504XC

Intended purpose: Obturators are used for the atraumatic insertion of sockets at the

surgical site.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Ear curettes

Risk classification: Ir

Basic-UDI-DI: 405018300000388Y2

Intended purpose: Ear curettes are intended to scrape, form and clean skin, tissue and

bones.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Ear loops

Risk classification: Ir

Basic-UDI-DI: 405018300000384XS

Intended purpose: Ear loops are used to cut off tissue.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Osteotomes

Risk classification: Ir

Basic-UDI-DI: 405018300000291XJ

Intended purpose: Orthopaedic osteotomes are used to shape and model bone material.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Preperation electrode

Risk classification:

Basic-UDI-DI: 405018300000248XH

405018300000482XT

Intended purpose: Preparation electrodes (cold electrodes) are designed to separate the

myoma from the surrounding myometrium and to dilate stenotic

tissue.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Raspatorium

Risk classification: Ir

Basic-UDI-DI: 405018300000377XV

Intended purpose: Rasps for the middle ear are used to smooth, shape and clean bony

structures.





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Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Rasps Risk classification: Ir

Basic-UDI-DI: 405018300000368XU

Intended purpose: Rasps for the nose are used to smooth, form and clean bony

structures.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Rasps Risk classification: Ir

Basic-UDI-DI: 405018300000301WU

Intended purpose: Graters for bones are used to flatten, shape and clean bones.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Revolution punching

Risk classification: Ir

Basic-UDI-DI: 405018300000296XU

Intended purpose: Revolution punches are designed to punch holes in bone and hard

tissue and remove the tissue.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Rongeurs

Risk classification: Ir

Basic-UDI-DI: 405018300000317XB

Intended purpose: Rongeurs are designed to punch holes in cartilage and remove the

tissue.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Scissor punching

Risk classification: Ir

Basic-UDI-DI: 405018300000413X8

Intended purpose: Scissor punches are used to punch holes in soft fabric.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Twist drill

Risk classification: Ir

Basic-UDI-DI: 405018300000326XC

Intended purpose: Twist drills are used to drill holes in bones.





# Annex to EU Quality Management Certificate SRN of Manufacturer: DE-MF-000005520 Certificate ID: 1000220878

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Sternum chisel

Risk classification: Ir

Basic-UDI-DI: 405018300000508XL

Intended purpose: Sternum chisels are used to split the breastbone.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Plunger Risk classification: Ir

Basic-UDI-DI: 405018300000302WW

Intended purpose: Plungers are used to compact or wedge structures.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Trephines

Risk classification: Ir

Basic-UDI-DI: 405018300000421X7

Intended purpose: Trephines are used to take tissue samples.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Trocars
Risk classification: Ir

Basic-UDI-DI: 405018300000158XF

Intended purpose: Trocars are used to gain access to the surgical site.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Uterine curettes

Risk classification: Ir

Basic-UDI-DI: 405018300000229XD

Intended purpose: Uterine curettes are used to scrap tissue.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Fistula probes

Risk classification: Ir

Basic-UDI-DI: 405018300000156XB

Intended purpose: Fistula probes are used in abnormal body passages or connection

(fistulas).

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Gall duct probes

Risk classification: Ir

Basic-UDI-DI: 405018300000222WX

Intended purpose: Gall duct probes are used to explore the bile duct.



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Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Goitre probes/ hollow probes

Risk classification: Ir

Basic-UDI-DI: 405018300000027WX

405018300000145X6

Intended purpose: Hollow probes / goiter probes are used to form a guide

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Probes
Risk classification: Ir

Basic-UDI-DI: 405018300000417XG

Intended purpose: Meniscus probes are used to examine the inside of the joint and for

therapeutic interventions in the joint.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Probes Risk classification: Ir

Basic-UDI-DI: 405018300000157XD

Intended purpose: Probes are intended to reach difficult accessible places or remote

locations to examine the place.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Uterine depressors

Risk classification: Ir

Basic-UDI-DI: 405018300000240WZ

Intended purpose: Uterine depressors are used to move or retract the uterus from its

position to visualise other structures.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Uterine probes

Risk classification: Ir

Basic-UDI-DI: 405018300000241X3

Intended purpose: Uterine probes are used to examine and explore the uterus.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Applicators for clips

Risk classification: Ir

Basic-UDI-DI: 405018300000143X2

405018300000272XE

Intended purpose: Applikatoren für Clips werden dazu verwendet Clips zu halten, sie

anzubringen bzw. zu applizieren und wieder zu entfernen.



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Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Applicator

Risk classification: Ir

Basic-UDI-DI: 405018300000139XB

Intended purpose: Applicators are used to place and secure cerclage wire

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Contact pliers

Risk classification: Ir

Basic-UDI-DI: 405018300000139XB

Intended purpose: Applicators are used to place and secure haemostatic clips.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Extractors

Risk classification: Ir

Basic-UDI-DI: 405018300000423XB

Intended purpose: Extractors are used to catch varices.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Ring applicators

Risk classification: I

Basic-UDI-DI: 405018300000479Y6

405018300000480XP

Intended purpose: Ring applicators are used to position the silicone rings to clamp the

woman's fallopian tubes.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Screwdriver / extraction bolt

Risk classification: Ir

Basic-UDI-DI: 405018300000332X7

405018300000333X9

Intended purpose: Screwdriver for bones are used to introduce or remove surgical

screws.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Underbanding needles/ palatal needles

Risk classification: Ir

Basic-UDI-DI: 405018300000140WU

405018300000142WY

Intended purpose: The ligature needle holds the surgical thread over which a ligature is

performed.



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Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Guide needles

Risk classification: Ir

Basic-UDI-DI: 405018300000029X3

Intended purpose: The Redon guide needle is used to guide the plastic drain from the

inside of the body through the skin.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Thread catcher

Risk classification: Ir

Basic-UDI-DI: 405018300000149XE

405018300000513XD 405018300000528XS

Intended purpose: Thread catchers are used to catch the suture material.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Thread guiding pliers

Risk classification: Ir

Basic-UDI-DI: 405018300000514XF

Intended purpose: Suture guide forceps are used to close wounds after laparoscopic

procedures.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Thread fork

Risk classification:

Basic-UDI-DI: 405018300000512XB

Intended purpose: Thread forks are used to knot the thread.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Suture holding forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000568Y6

Intended purpose: Suture holding forceps are used for wound closure after surgery.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Thread holding pliers

Risk classification: Ir

Basic-UDI-DI: 405018300000148XC

Intended purpose: Thread holding pliers are used to close wounds after laparoscopic

procedures.





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Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Management tool

Risk classification: Ir

Basic-UDI-DI: 405018300000535XP

Intended purpose: Guides for drill bits are used to direct or guide drill bits in order to

position them.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Knot guide / knot slider

Risk classification: Ir

Basic-UDI-DI: 405018300000080X3

405018300000526XN 405018300000529XU

Intended purpose: Knot guides / knot pushers are used to push seam knots to the point

of suitable seam tension.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Needle Risk classification: Ir

Basic-UDI-DI: 405018300000511X9

Intended purpose: Needles for suturing the fascia are used to close wounds after

laparoscopic procedures

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Paracentesis needles

Risk classification: Ir

Basic-UDI-DI: 405018300000378XX

Intended purpose: Paracentesis needles are used to pierce the eardrum.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Reverdin needles

Risk classification: I

Basic-UDI-DI: 405018300000141WW

Intended purpose: Reverdin needles are used to pierce thick skin or firm tissue and sew

over an inserted thread

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Abortion forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000225X5

Intended purpose: Abortion forceps are used to grasp, hold and secure or remove the

ovaries and placenta pieces.



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Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Baby Duckbill punches

Risk classification: Ir

Basic-UDI-DI: 405018300000351XB

Intended purpose: Foreign body grasping forceps / foreign body forceps are used to

grasp, hold and remove foreign bodies.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Chalazion tweezers

Risk classification: Ir

Basic-UDI-DI: 405018300000536XR

Intended purpose: Chalazion forceps are used to reverse the eyelid and expose chalazion

for drainage and removal.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Cholangiography forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000548XY

Intended purpose: Cholangiography forceps are used to insert the cholangiography

catheter into the bile duct in order to inject the contrast solution.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Cilia tweezers

Risk classification: Ir

Basic-UDI-DI: 405018300000403X5

Intended purpose: Cilia tweezers are used to grip or manipulate the eyelashes.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Intestinal/tissue forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000210WQ

Intended purpose: Intestinal/Tissue grasping forceps are used to grasp and / or compress

internal intestinal structures or tissue.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Gripper tongs

Risk classification: Ir

Basic-UDI-DI: 405018300000003WH

Intended purpose: The grasping tongs are used for endoscopic diagnosis and treatment

in urology.



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MDN 1208 - Non-active non-implantable instruments Device category:

Product name: Wire cutters

Risk classification:

405018300000299Y2 Basic-UDI-DI:

Wire pliers / wire crimping pliers / wire clamping pliers / locking pliers Intended purpose:

are used to hold, tighten, cut and / or twist surgical wire.

MDN 1208 - Non-active non-implantable instruments Device category:

Product name: Pliers Risk classification: Ir

Basic-UDI-DI: 405018300000003WH

> 405018300000538XV 405018300000547XW

Intended purpose: Gripping tongs are used to grip and hold fabric.

MDN 1208 - Non-active non-implantable instruments Device category:

Product name: Flat nose pliers

Risk classification: Ir

Basic-UDI-DI: 405018300000298XY

Intended purpose: Flat nose pliers / parallel nose pliers are used to bend and cut wires.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Foreign body forceps

Risk classification:

Basic-UDI-DI: 405018300000171X7

405018300000254XC

Foreign body grasping forceps / foreign body forceps are used to Intended purpose:

grasp, hold and remove foreign bodies.



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Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Grasping forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000171X7

405018300000254XC 405018300000298XY 405018300000351XB 405018300000371XH 405018300000374XP 405018300000549Y2 405018300000550XK 405018300000551XM 405018300000552XP 405018300000569Y8

405018300000570XR 405018300000571XT 405018300000572XV

Intended purpose: Foreign body grasping forceps / foreign body tongs are used to grasp,

hold and remove foreign bodies.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Foreign body lever

Risk classification: Ir

Basic-UDI-DI: 405018300000390XM

Intended purpose: Foreign body levers are used to loosen foreign bodies in the ear.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Vascular forceps

Risk classification:

Basic-UDI-DI: 405018300000166XE

Intended purpose: Vascular forceps are used to grip and hold delicate tissue

atraumatically.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Iris tweezers

Risk classification: Ir

Basic-UDI-DI: 405018300000404X7

Intended purpose: Iris forceps are used to grasp, hold or manipulate tissues of the eye or

surrounding tissues.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Capsule forceps





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Risk classification:

Basic-UDI-DI: 405018300000266XK

Intended purpose: Capsule graspers are used to grip, hold and manipulate soft tissue.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Capsule tweezers

Risk classification: Ir

Basic-UDI-DI: 405018300000543XN

Intended purpose: Capsule forceps are used to hold lens capsules during surgery.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Catheter insertion instruments

Risk classification: Ir

Basic-UDI-DI: 405018300000268XP

Intended purpose: Catheter insertion instruments are used to insert and position

catheters in the urinary tract.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Catheter insertion forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000375XR

Intended purpose: Catheter insertion forceps are used to grip and hold the endotracheal

tube during intubation or to remove foreign bodies from the airways.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Laryngeal polyp forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000373XM

Intended purpose: Laryngeal polyp forceps are used to remove polyps from the vocal

cords.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Clamp holder

Risk classification: Ir

Basic-UDI-DI: 405018300000214WY

Intended purpose: Clamp holders are used to hold the clamp in position.



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Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Clamp closing pliers

Risk classification: Ir

Basic-UDI-DI: 405018300000211WS

405018300000539XX

Intended purpose: Clamp closing pliers are used to close clamps.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Bone holding forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000306X6

Intended purpose: Bone holding forceps are used to grasp and hold bones.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Bone splinter forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000297XW

Intended purpose: Bone splitting forceps are used to cut bones.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Cartilage grasping forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000316X9

Intended purpose: Cartilage grasping forceps are used to grasp and hold cartilage

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Cartilage crushing

Risk classification: Ir

Basic-UDI-DI: 405018300000334XB

Intended purpose: Cartilage crushers are used to crush pieces of cartilage that have been

removed.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Laryngoscope spoon forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000372XK

Intended purpose: Laryngoscope spoon forceps are used to grip, hold and manipulate

soft tissue / anatomical structures.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Spoon tongs / ear tongs



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Risk classification:

Basic-UDI-DI: 405018300000383XQ

Intended purpose: Spoon tongs / ear tongs are used to grip, manipulate, compress or pull

fabric, equipment or accessories.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Marker Risk classification: Ir

405018300000425XF Basic-UDI-DI:

Markers enable precise marking of (skin) flaps for excision Intended purpose:

MDN 1208 - Non-active non-implantable instruments Device category:

Product name: Meniscus grasping forceps

Risk classification:

Basic-UDI-DI: 405018300000415XC

Meniscus grasping forceps are used to grip, hold and compress the Intended purpose:

meniscus.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Needle holder

Risk classification:

405018300000144X4 Basic-UDI-DI:

> 405018300000337XH 405018300000352XD

Needle holders are designed to hold the sewing needles firmly during Intended purpose:

sewing.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Nail extrusion forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000540XG

Intended purpose: Nail extraction forceps are used to split and remove ingrown nails.

MDN 1208 - Non-active non-implantable instruments Device category:

Product name: Micro-tying tweezers / suture tweezers

Risk classification: Ir

Basic-UDI-DI: 405018300000147XA

405018300000406XB

Intended purpose: Suture tweezers / micro-tying tweezers are used to grip and hold

suture material.

MDN 1208 - Non-active non-implantable instruments Device category:

Product name: Ear and nose tweezers

Risk classification:

Basic-UDI-DI: 405018300000380XI



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Intended purpose: Ear and nose forceps are used to grasp or hold tissue on the nose or

ear.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Ear polyp forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000381XL

Intended purpose: Ear polyp forceps are used to grasp, hold and remove soft tissue or

polyps in the ear.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Organ grasping forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000167XG

Intended purpose: Organ grasping forceps are used to grasp organs atraumatically.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Patella forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000541XJ

Intended purpose: Patella forceps are used to grip and hold the patella.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: PE pliers Risk classification: Ir

Basic-UDI-DI: 405018300000160X2

Intended purpose: PE forceps are used for the endoscopic removal of tissue samples.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Tweezers / Tissue grasping tweezers / Hook tweezers / Fixation

tweezers

Risk classification: Ir

Basic-UDI-DI: 405018300000165XC

405018300000183XE 405018300000186XL 405018300000336XF 405018300000405X9 405018300000407XD

Intended purpose: Tweezers / tissue grasping tweezers / hook tweezers / fixing tweezers

are used to grasp, manipulate, squeeze, pull or connect tissue,

materials or accessories.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Forceps Risk classification: Ir

Basic-UDI-DI: 405018300000546XU



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Intended purpose: Forceps for changing the valves are used to change the valves.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Haemorrhoidal forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000237XC

Intended purpose: Polyp forceps / polyp forceps / haemorrhoidal forceps are used to

grasp, hold and remove polyps in the uterus

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Polyp forceps / polyp forceps / nasal forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000237XC

405018300000365XN 405018300000382XN 405018300000537XT

Intended purpose: Polyp forceps / polyp forceps / nasal forceps are used to grasp, hold

and remove polyps.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Pyloric spreader

Risk classification: Ir

Basic-UDI-DI: 405018300000545XS

Intended purpose: Pyloric spreaders are used to separate the pylorus or other

gastrointestinal structures from adjacent structures.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Reduction forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000305X4

405018300000307X8

Intended purpose: Reduction forceps are used to hold and reposition bones.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Sponge forceps / grain forceps / swab forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000161X4

405018300000162X6 405018300000163X8

Intended purpose: Sponge forceps / grain forceps / swab forceps are used to grip surgical

sponges.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Tendon pulling forceps / tendon grasping forceps / instruments

Risk classification: Ir

Basic-UDI-DI: 405018300000313X3



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405018300000314X5

Intended purpose: Tendon tunneling forceps / Tendon grasping forceps / instruments are

used to grasp, hold, pull and guide tendons.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Side cutters / bolt cutters / head cutters

Risk classification: Ir

Basic-UDI-DI: 405018300000300WS

Intended purpose: Side cutting pliers / bolt cutting pliers / head cutting pliers are used to

cut orthopedic wires or pins.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Septum straightening forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000363XJ

Intended purpose: Septum straightening forceps / septum crimping forceps are used to

grip and straighten the septum.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Sinus clamps

Risk classification: Ir

Basic-UDI-DI: 405018300000163X8

Intended purpose: Sinus clamps are used to grip surgical sponges.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Sinuscopy forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000364XL

Intended purpose: Sinuscopy forceps are used to grasp, hold and remove polyps and to

open the maxillary, frontal and sphenoid sinuses.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Splinter tweezers

Risk classification: Ir

Basic-UDI-DI: 405018300000174XD

Intended purpose: Splinter tweezers are used to grasp, hold and remove small foreign

bodies.



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Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Biopsy forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000159XH

405018300000212WU 405018300000232X2 405018300000252X8 405018300000256XG 405018300000346XJ 405018300000347XL 405018300000542XL

Intended purpose: Rigid biopsy forceps are used for the endoscopic removal of tissue

samples.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Stone grasping forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000257XJ

Intended purpose: Stone grasping forceps are used for endoscopic diagnosis and

treatment in urology.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Tampon forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000362XG

Intended purpose: Tampon forceps are used to apply gauze bandages.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Tenaculum grasping forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000234X6

Intended purpose: Tenaculum forceps are used for grasping and holding cervical tissue.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Tonsil forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000359XT

Intended purpose: Tonsil forceps are intended to hold the tonsil.



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Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Tumour grasping forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000175XF

Intended purpose: Tumour grasping forceps are used to grasp, hold and remove the

tumour.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Tunnelling pliers

Risk classification: Ir

Basic-UDI-DI: 405018300000318XD

Intended purpose: Tunnelling pliers are used to grip and hold vessels.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Uterine grasping forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000235X8

405018300000236XA 405018300000238XE

Intended purpose: Uterine grasping forceps are used to lift the vaginal wall and to grasp,

hold and remove vaginal overgrowths, polyps and malignant diseases.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Uterine clamp forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000226X7

Intended purpose: Uterine clamp forceps are used to grip and clamp tissue in the uterus.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Cotton swabs

Risk classification: Ir

Basic-UDI-DI: 405018300000176XH

Intended purpose: Cotton swabs are used to hold a plaster at their distal end to clean or

apply a substance to superficial wounds or body orifices.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Dental forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000544XQ

Intended purpose: Dental forceps are used to grip a dressing that is applied orally.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Gallstone forceps

Risk classification: Ir



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Basic-UDI-DI: 405018300000208X5

Intended purpose: Forceps for the gallbladder are designed to grasp or manipulate the

gallstones.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Birth forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000223WZ

Intended purpose: Obstetric forceps are used to grasp the newborn's head and facilitate

the birth process.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Lung grasping forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000168XJ

Intended purpose: Forceps for the lung are designed to grasp, manipulate or support the

lung atraumatically.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Kidney stone forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000261X9

Intended purpose: Kidney forceps are designed to grasp and remove kidney stones.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Tonsil forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000359XT

Intended purpose: Tonsil forceps are intended to hold the tonsil.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Pliers for bones

Risk classification: Ir

Basic-UDI-DI: 405018300000308XA

405018300000322X4 405018300000370XF 405018300000418XJ

Intended purpose: Pliers for bones are used for gripping, cutting or breaking bones.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Pliers



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Risk classification:

Basic-UDI-DI: 405018300000239XG

Intended purpose: IUD removal forceps / IUD removal instruments are used to remove

IUDs.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Anastomosis clamps

Risk classification: Ir

Basic-UDI-DI: 405018300000215X2

405018300000279XU

Intended purpose: Anastomosis clamps are used to grip and temporarily close blood

vessels.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Aneurysm clips

Risk classification: Ir

Basic-UDI-DI: 405018300000275XL

Intended purpose: Aneurysm clips are used to grip blood vessels and completely block

the blood flow.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Aortic clamps

Risk classification: Ir

Basic-UDI-DI: 405018300000277XQ

405018300000280XD

Intended purpose: Aortic clamps are used to grip and compress large vessels with thick

and dense walls.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Artery clamps

Risk classification: Ir

Basic-UDI-DI: 405018300000178XM

Intended purpose: Haemostatic forceps are used to grip and compress arteries.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Auricular clamps

Risk classification: Ir

Basic-UDI-DI: 405018300000285XP

Intended purpose: Auricular clamps are used to grip the atrium while a suture is placed or

bleeding is stopped.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Bronchus clamps

Risk classification: Ir



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Basic-UDI-DI: 405018300000185XJ

Intended purpose: Bronchus clamps are used to grip the bronchi and compress them

atraumatically.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Bulldog clamps

Risk classification: Ir

Basic-UDI-DI: 405018300000271XC

Intended purpose: Bulldog clamps are used to temporarily compress vessels

atraumatically.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Coarctation clamps

Risk classification: Ir

Basic-UDI-DI: 405018300000180X8

405018300000281XF 405018300000283XK

Intended purpose: Coarctation clamps are used to grip and compress the aorta.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Pressure clamps

Risk classification: Ir

Basic-UDI-DI: 405018300000179XP

Intended purpose: Pressure clamps are used to grip blood vessels and control blood flow.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Bowel clamps

Risk classification: Ir

Basic-UDI-DI: 405018300000213WW

Intended purpose: Bowel clamps are used to grip the bowel and temporarily close the

end of the bowel.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Drum clamps

Risk classification: Ir

Basic-UDI-DI: 405018300000169XL

405018300000518XP

Intended purpose: Seizing clamps / thoracic clamps are used to grasp and compress

tissues.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Bile duct clamps

Risk classification: Ir

Basic-UDI-DI: 405018300000209X7



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Intended purpose: Bile duct clamps are used to grasp and clamp bile ducts.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Clamps Risk classification: Ir

Basic-UDI-DI: 405018300000177XK

405018300000274XJ

Intended purpose: Vascular clamps / paediatric vascular clamps / peripheral vascular

clamps / dissecting clamps / ligature clamps / aorta ligature clamps / atrauma clamps are used to grip vessels and compress them for

temporary haemostasis

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Haemorrhoid clamps

Risk classification: Ir

Basic-UDI-DI: 405018300000516XK

Intended purpose: Haemorrhoid clamps are used to grip and hold the haemorrhoids in

place.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Hysterectomy clamps

Risk classification: Ir

Basic-UDI-DI: 405018300000233X4

Intended purpose: Hysterectomy clamps are used to grasp and compress tissue and

structures of the uterus.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Iliac clamps

Risk classification: Ir

Basic-UDI-DI: 405018300000523XG

Intended purpose: Iliac clamps are used to grip and compress the iliac arteries.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Circumcision device

Risk classification: Ir

Basic-UDI-DI: 405018300000263XD

Intended purpose: Circumcision clamps are used to compress the foreskin of the penis

during circumcision and to control bleeding.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Bone holding clamps

Risk classification: Ir



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405018300000309XC Basic-UDI-DI:

Intended purpose: Bone holding clamps are used to grasp bone and hold them in

position.

MDN 1208 - Non-active non-implantable instruments Device category:

Product name: Scalp clamps

Risk classification: Ir

Basic-UDI-DI: 405018300000348XN

Intended purpose: Scalp clamps are used to grip the scalp of children to facilitate delivery.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Ligature clamps

Risk classification: Ir

Basic-UDI-DI: 405018300000184XG

Intended purpose: Ligature clamps are used to grip and fix tissue.

MDN 1208 - Non-active non-implantable instruments Device category:

Product name: Mylohyoid muscle clamps

Risk classification: Ir

Basic-UDI-DI: 405018300000276XN

Intended purpose: Mylohyoid muscle clamps are used to grip and compress the

mylohyoid muscle.

MDN 1208 - Non-active non-implantable instruments Device category:

Product name: Renal pedicle clamps

Risk classification:

Basic-UDI-DI: 405018300000262XB

Intended purpose: Renal pedicle clamps are used to grip and compress the renal pedicle.

MDN 1208 - Non-active non-implantable instruments Device category:

Product name: Peritoneal clamps

Risk classification:

405018300000217X6 Basic-UDI-DI:

Peritoneal clamps are used to grip and compress the peritoneum. Intended purpose:

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Profunda clamps

Risk classification:

Basic-UDI-DI: 405018300000278XS

Intended purpose: Profunda clamps are used to grip and compress vessels.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Ramus clamps



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Risk classification:

Basic-UDI-DI: 405018300000191XD

Intended purpose: Ramus clamps are used to grip and compress the mandibular ramus.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Anastomosis clamps

Risk classification: Ir

Basic-UDI-DI: 405018300000216X4

Intended purpose: Rectal anastomosis clamps are used to grip tissue atraumatically in

order to stop bleeding.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Anastomosis clamps

Risk classification: Ir

Basic-UDI-DI: 405018300000216X4

Intended purpose: Rectal clamps are used to atraumatically grasp, connect, compress or

support the rectum, the rectal valves or the anal canal.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Tendon interlacing clamps

Risk classification: Ir

Basic-UDI-DI: 405018300000315X7

Intended purpose: Tendon interlacing clamps are used to hold and pull on injured

tendons.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Tobacco pouch seam clamps

Risk classification: Ir

Basic-UDI-DI: 405018300000519XR

Intended purpose: Tobacco pouch seam clamps are used to grip and fix the tobacco

pouch seam.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Tangential clamps

Risk classification: Ir

Basic-UDI-DI: 405018300000284XM

Intended purpose: Tangential clamps are used to regulate the blood flow through the

main vessels.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: URO-TANGENTIAL clamps

Risk classification: Ir

Basic-UDI-DI: 405018300000267XM



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Intended purpose: URO-TANGENTIAL clamps are used to grip and compress urine-

forming and urine-draining tissues and structures.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Vasectomy clamps

Risk classification: In

Basic-UDI-DI: 405018300000265XH

Intended purpose: Vasectomy clamps are used to grip and fix the vas deferens during

vasectomy.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Vena cava clamps

Risk classification:

Basic-UDI-DI: 405018300000265XH

Intended purpose: Vena cava clamps are used to grip the vena cava and regulate the

blood flow.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Vein strippers

Risk classification: Ir

Basic-UDI-DI: 405018300000287XT

405018300000428XM

Intended purpose: Vein strippers are used to pull veins or remove them for

transplantation.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Tendon strippers

Risk classification: Ir

Basic-UDI-DI: 405018300000330X3

Intended purpose: Tendon strippers are used to pull tendons or remove them for

transplantation.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Bone rongeur forc

Risk classification: Ir

Basic-UDI-DI: 405018300000310WV

Intended purpose: Bone rongeur forceps are used to cut through small bones and

remove larger bones and bone fragments

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Amniotome

Risk classification: Ir



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Basic-UDI-DI: 405018300000224X3

Intended purpose: An amniotome is used to cut or tear the membrane surrounding the

foetus.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Zygomatic bone reamers

Risk classification: In

Basic-UDI-DI: 405018300000331X5

Intended purpose: Zygomatic reamers are used to pierce the zygomatic arch and sew it

over a wire.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Orthopaedic reamers / bone reamers

Risk classification:

Basic-UDI-DI: 405018300000294XQ

Intended purpose: Orthopaedic reamers / bone reamers are designed to form and resect

holes in bones.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Puncture needles

Risk classification: Ir

Basic-UDI-DI: 405018300000356XM

Intended purpose: Puncture needles are used to puncture tissue in order to take tissue

samples.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Adenotomes

Risk classification: In

Basic-UDI-DI: 405018300000358XR

Intended purpose: Adenotomes are used to remove severely enlarged palatine tonsils

(tonsils).

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Amputation saws

Risk classification: Ir

Basic-UDI-DI: 405018300000129X8

Intended purpose: Amputation saws are used to saw through bones.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Bayonet knives

Risk classification: Ir

Basic-UDI-DI: 405018300000092XA

Intended purpose: Bayonet knives are used to cut through fabric.



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Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Banana knives

Risk classification: Ir

Basic-UDI-DI: 405018300000094XE

Intended purpose: Banana knives are used to cut through fabric.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Cartilage resection knives

Risk classification: Ir

Basic-UDI-DI: 405018300000416XE

Intended purpose: Cartilage resection knives are used to cut through cartilage.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Episiotomy scissors

Risk classification: Ir

Basic-UDI-DI: 405018300000122WS

Intended purpose: Episiotomy scissors are used for cutting the perineum and posterior

vaginal wall.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Knife
Risk classification: Ir

Basic-UDI-DI: 405018300000385XU

Intended purpose: The instrument is used for incisions in the ENT area (throat/nose/ear).

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Tonsil knife

Risk classification: Ir

Basic-UDI-DI: 405018300000360XC

Intended purpose: The tonsil knife is used to surgically remove the palatine tonsils.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Dermatomes

Risk classification: Ir

Basic-UDI-DI: 405018300000100WG

Intended purpose: Dermatomes are used to obtain uniform skin flaps that serve as grafts.

Dermatomes are also used to cut out small skin lesions.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Saw blades

Risk classification: Ir

Basic-UDI-DI: 405018300000134WZ

Intended purpose: The blade provides the cutting edge for cutting materials or tissues.



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Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Blade Risk classification: Ir

Basic-UDI-DI: 405018300000101WJ

Intended purpose: The blade is used as cutting edge for the extraction of even skin flaps

or to remove smaller skin lesions.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Cutting pliers

Risk classification: Ir

Basic-UDI-DI: 405018300000311WX

Intended purpose: The cutting pliers for ribs are used to cut through bones and ribs.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Wire saws

Risk classification: Ir

Basic-UDI-DI: 405018300000131WT

Intended purpose: Wire saws are used to cut smoothly through boiling.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Wire cutters

Risk classification: I

Basic-UDI-DI: 405018300000127X4

Intended purpose: Wire cutters are used for cutting wire.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Suture cutter

Risk classification: Ir

Basic-UDI-DI: 405018300000107WW

Intended purpose: Suture cutter are used to endoscopically cut the sewing thread.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Fistula knives

Risk classification: Ir

Basic-UDI-DI: 405018300000090X6

Intended purpose: Fistula knives are used for tissue dissection and making incision.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Fistula scissors

Risk classification: Ir

Basic-UDI-DI: 405018300000124WW

Intended purpose: Fistula scissors are used for cutting fistulas.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Vascular scissors



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Risk classification:

Basic-UDI-DI: 405018300000108WY 405018300000117WZ

Intended purpose: Vascular scissors are used for cutting vessels.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Hook knives

Risk classification: Ir

Basic-UDI-DI: 405018300000095XG

Intended purpose: Hook knives are used to cut through fabric.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Hook scissors

Risk classification: Ir

Basic-UDI-DI: 405018300000350X9

Intended purpose: Hook scissors are used to cut tissue endoscopically.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Micro scissors

Risk classification: Ir

Basic-UDI-DI: 405018300000335XD

Intended purpose: Capsule shears are used to cut capsule structures.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Cartilage scissors

Risk classification: Ir

Basic-UDI-DI: 405018300000109X2

Intended purpose: Cartilage scissors are used to cut cartilage tissue.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Ligature scissors

Risk classification: Ir

Basic-UDI-DI: 405018300000104WQ

Intended purpose: Ligature scissors are used to cut suture or ligature material.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Lung flap and rectal scissors

Risk classification: Ir

Basic-UDI-DI: 405018300000118X3

Intended purpose: Lung flap and rectal scissors are used to cut or dissect tissue.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Meniscus knives



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Risk classification:

Basic-UDI-DI: 405018300000091X8

405018300000093XC

Intended purpose: Meniscus knives are used to cut meniscus fragments.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Amputation knife

Risk classification: Ir

Basic-UDI-DI: 405018300000089XM

Intended purpose: Knives for amputation are intended for the amputation of limbs.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Micro laryngeal scissors

Risk classification: Ir

Basic-UDI-DI: 405018300000125WY

Intended purpose: Micro laryngeal scissors are used to make fine incisions in laryngeal

surgery.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Micro tumour scissors

Risk classification: Ir

Basic-UDI-DI: 405018300000106WU

Intended purpose: Micro tumour scissors are used to separate tumours from the

surrounding tissue.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Myoma knives

Risk classification: Ir

Basic-UDI-DI: 405018300000242X5

Intended purpose: Myoma knives are used to cut cone-shaped tissue from the cervix.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Nose saws

Risk classification: Ir

Basic-UDI-DI: 405018300000130WR

Intended purpose: Nose saws are used to cut bone tissue.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Nasals scissors

Risk classification: Ir

Basic-UDI-DI: 405018300000112WP

Intended purpose: Nasals scissors are used for cutting or dissecting tissue of the nose.





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Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Ear scissors

Risk classification: Ir

Basic-UDI-DI: 405018300000113WR

Intended purpose: Ear scissors are used to cut or dissect tissue in the ear.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Rhinoplasty knives

Risk classification: Ir

Basic-UDI-DI: 405018300000085XD

Intended purpose: Rhinoplasty knives are used to make incisions.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Scissors for the eye

Risk classification: Ir

Basic-UDI-DI: 405018300000110WK

Intended purpose: Scissors for the eye are used to cut structures / tissues on the eye and

/ or the iris.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Scissors for the gynaecology

Risk classification: Ir

Basic-UDI-DI: 405018300000123WU

Intended purpose: Scissors for the gynaecology are used to cut tissues and structures in

the female reproductive system.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Scissors for plastic surgery

Risk classification: Ir

Basic-UDI-DI: 405018300000424XD

Intended purpose: Plastic surgery scissors are used for cutting in shape-altering or

reconstructive procedures

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Tonsil scissors

Risk classification: Ir

Basic-UDI-DI: 405018300000114WT

Intended purpose: Tonsil scissors are used to remove the tonsils through cutting.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Scissor attachments

Risk classification: Ir

Basic-UDI-DI: 405018300000126X2

Intended purpose: Scissor attachments are used to cut tissue endoscopically.



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Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Scissors
Risk classification: Ir

Basic-UDI-DI: 405018300000105WS

405018300000119X5

Intended purpose: Scissors are used for cutting or preparing different tissues or

materials.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Tendon scissors

Risk classification: Ir

Basic-UDI-DI: 405018300000111WM

Intended purpose: Tendon scissors are used for cutting tendons.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: BALLENGER septum knife

Risk classification: Ir

Basic-UDI-DI: 405018300000086XF

Intended purpose: Septum knives are used to cut cartilage and soft tissue.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Sickle knives

Risk classification: Ir

Basic-UDI-DI: 405018300000096XJ

Intended purpose: Sickle knives are used to cut through fabric.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Scalpel knives

Risk classification: Ir

Basic-UDI-DI: 405018300000084XB

Intended purpose: Scalpel knives are used to sharply cut through tissue.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Stricture knives

Risk classification: Ir

Basic-UDI-DI: 405018300000098XN

Intended purpose: Stricture knives are used to widen the urethra by making incisions.



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Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Trigeminal scissors

Risk classification: Ir

Basic-UDI-DI: 405018300000116WX

Intended purpose: Trigeminal scissors are used to cut the trigeminal nerve.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Conversion instrument

Risk classification: Ir

Basic-UDI-DI: 405018300000419XL

Intended purpose: Conversion instrument are used to guide flexible wire saws around

bones.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Ureteral slitting scissors

Risk classification: Ir

Basic-UDI-DI: 405018300000253XA

Intended purpose: Ureteral slitting scissors are used to slit open urethral strictures.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Urethrotomes

Risk classification:

Basic-UDI-DI: 405018300000428XM

405018300000499YC

Intended purpose: Urethrotomes are used to slit open urethral strictures.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Dental scissors

Risk classification: Ir

Basic-UDI-DI: 405018300000115WV

Intended purpose: Dental scissors are used to cut or prepare gums.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Aortic hooks

Risk classification: Ir

Basic-UDI-DI: 405018300000270XA

Intended purpose: Aortic hooks are used to exert traction on the aorta.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Atrial hooks

Risk classification: Ir

Basic-UDI-DI: 405018300000269XR

Intended purpose: Atrial hooks are used to grip and retract the atria.



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Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Abdominal spatulas

Risk classification:

405018300000203WT Basic-UDI-DI:

Intended purpose: Abdominal spatulas are used to hold back soft tissue organs.

MDN 1208 - Non-active non-implantable instruments Device category:

Product name: Bladder spatula

Risk classification:

405018300000204WV Basic-UDI-DI:

Bladder spatulas are used to grasp and withdraw the bladder. Intended purpose:

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Alar Retractor

Risk classification:

Basic-UDI-DI: 405018300000527XQ

Intended purpose: The LATERA® Alar retractor is used with LATERA® implants in nasal

surgery.

MDN 1208 - Non-active non-implantable instruments Device category:

Product name: **Dilators** Risk classification: Ιr

Basic-UDI-DI: 405018300000259XN

405018300000428XM

Intended purpose: Dilators for the choledochal duct are designed to examine or dilate the

bile duct.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: **Dilators** Risk classification: Ir

Basic-UDI-DI: 405018300000357XP

405018300000401WZ

Intended purpose: Dilators are used to dilate the nephrostomy tract.

MDN 1208 - Non-active non-implantable instruments Device category:

Product name: Retractors

Risk classification: Ιr

Basic-UDI-DI: 405018300000354XH

Intended purpose: Endo retractors are used to hold disturbing structures out of the

surgical site.

MDN 1208 - Non-active non-implantable instruments Device category:

Product name: **Exploratory hooks** 

Risk classification:

Basic-UDI-DI: 405018300000343XC



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Intended purpose: Exploratory hooks are used to explore nerve tissue and the vascular

system.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Fistula hooks

Risk classification: Ir

Basic-UDI-DI: 405018300000197XR

Intended purpose: Fistula hooks are used to grip, stabilise and retract sensitive soft

tissue.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Retractor

Risk classification: Ir

Basic-UDI-DI: 405018300000195XM

405018300000200WM

Intended purpose: Flexible retractors are used to separate tissue or other anatomical

parts.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Nerve and vascular hooks

Risk classification: Ir

Basic-UDI-DI: 405018300000345XG

Intended purpose: Vessel hooks are used to grip and retract vessels.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Skin hooks

Risk classification: Ir

Basic-UDI-DI: 405018300000199XV

Intended purpose: Skin hooks are used to fix tissue or to apply traction to the skin.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Knee hooks

Risk classification: Ir

Basic-UDI-DI: 405018300000323X6

Intended purpose: Knee hooks are used to grip fabric and pull it back at a right angle.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Bone lever

Risk classification: Ir

Basic-UDI-DI: 405018300000312WZ

Intended purpose: Bone lever are used to lift and position bones.



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Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Lid retractors

Risk classification: Ir

Basic-UDI-DI: 405018300000517XM

Intended purpose: Lid retractors are used to grip and retract the upper eyelid and

eyelashes.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Lung spatula

Risk classification: Ir

Basic-UDI-DI: 405018300000288XV

Intended purpose: Lung spatulas are used to manipulate lung tissue, surfaces or vessels.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Dislocation lever

Risk classification: Ir

Basic-UDI-DI: 405018300000290XG

Intended purpose: Dislocation levers are used to remove the humeral head.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Nose wing hook

Risk classification: Ir

Basic-UDI-DI: 405018300000366XQ

Intended purpose: Nose wing hook are used to grasp and retract nostrils.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Nerve root hook

Risk classification: Ir

Basic-UDI-DI: 405018300000219XA

405018300000344XE 405018300000521XC

Intended purpose: Nerve root hooks are used to grip and retract nerve roots / nerves.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Progenitors / chin holders

Risk classification: Ir

Basic-UDI-DI: 405018300000192XF

Intended purpose: Progenitors / chin holders are used to grip and retract soft tissues of

the lower jaw.



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Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Prostate hook

Risk classification: Ir

Basic-UDI-DI: 405018300000260X7

Intended purpose: Prostate hooks are used to grip and retract the prostate.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Ramus hooks

Risk classification: Ir

Basic-UDI-DI: 405018300000193XH

Intended purpose: Ramus hooks are used to grasp and retract in front of the descending

ramus of the lower jaw.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Mucosal hook

Risk classification: Ir

Basic-UDI-DI: 405018300000190XB

Intended purpose: Mucosal hooks are used to enlarge mucosal incisions.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Spatulas Risk classification: Ir

Basic-UDI-DI: 405018300000205WX

405018300000329XJ 405018300000534XM

Intended purpose: Spatulas are used to introduce material into bone cavities, to

manipulate tissue, or to remove material from a surface or vessel.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Spina nasalis hook

Risk classification: Ir

Basic-UDI-DI: 405018300000367XS

Intended purpose: Spina nasalis hooks are used to grasp and retain periosteal tissue on

the spina nasalis.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Pointed hooks

Risk classification: Ir

Basic-UDI-DI: 405018300000376XT

Intended purpose: Pointed hooks for the middle ear are used to remove foreign bodies

from the ear and to fix structures in the ear or exert traction on the

tissue.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Split separators



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Risk classification:

Basic-UDI-DI: 405018300000341X8

405018300000524XJ

Intended purpose: Split separators are used to lengthen bone cuts.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: S-retractors

Risk classification: Ir

Basic-UDI-DI: 405018300000221WV

Intended purpose: S-retractors are used to widen the opening to the surgical site in order

to remove the gallbladder from the body.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Strabismus hooks

Risk classification: Ir

Basic-UDI-DI: 405018300000194XK

Intended purpose: Strabismus hooks are used to hold the eye muscle or exert traction on

it.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Dilator Risk classification: Ir

Basic-UDI-DI: 405018300000532XH

Intended purpose: Tracheal dilators are used to widen tracheal structures and passages.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Surgical tracheal hooks are used to fix tracheal tissue or to apply

traction to the tissue.

Risk classification: Ir

Basic-UDI-DI: 405018300000202WR

Intended purpose: Surgical tracheal hooks are used to fix tracheal tissue or to apply

traction to the tissue.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Vascular dilator

Risk classification: Ir

Basic-UDI-DI: 405018300000286XR

Intended purpose: Vascular dilators are used to widen a vessel.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Vein hooks

Risk classification: Ir

Basic-UDI-DI: 405018300000525XL

Intended purpose: Vein hooks are used to grasp and retract veins.



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Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Visual dilator

Risk classification: Ir

Basic-UDI-DI: 405018300000530XD

Intended purpose: Dilators are used to dilate lumens in the body.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Soft tissue hook

Risk classification: Ir

Basic-UDI-DI: 405018300000196XP

Intended purpose: Soft tissue hooks are used to grip, stabilise and retract soft tissue.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Wisdom tooth hook

Risk classification: Ir

Basic-UDI-DI: 405018300000533XK

Intended purpose: Wisdom tooth hooks are used to remove impacted wisdom teeth from

their alveolar processes.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Retractor Risk classification: Ir

Basic-UDI-DI: 405018300000188XQ

Intended purpose: Retractors are used to separate tissue or other anatomical parts to

expose or access organs or tissues.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Biopsy forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000218X8

Intended purpose: Biopsy forceps for flexible endoscopy are used to remove tissue

endoscopically.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Scissors forceps

Risk classification: Ir

Basic-UDI-DI: 405018300000120WN

Intended purpose: Scissors forceps are used for the endoscopic cutting of sutures.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Handles of polypectomy snares

Risk classification: Ir

Basic-UDI-DI: 405018300000618XU

Intended purpose: The HF polypectomy snares are intended for use in endoscopies of the

gastrointestinal tract for the removal (ectomy) of colorectal polyps.



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(ectomy) of colorectal polyps for therapeutic or diagnostic purposes

for histological examination in

in conjunction with an endoscope using high-frequency current.

Device category: MDA 0312 - Other active non-implantable surgical devices

Product name: Polypectomy snares

Risk classification: IIb

Basic-UDI-DI: 405018300000617XS

Intended purpose: The HF polypectomy snares are intended for use in endoscopies of the

gastrointestinal tract to remove (ectomize) (ectomy) of colorectal polyps for therapeutic or diagnostic purposes for histological examination in used in conjunction with an endoscope by means of

highfreuency current

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: ADSON suction tubes

Risk classification: IIa

Basic-UDI-DI: 405018300000021WK

Intended purpose: ADSON tube is used to aspirate deposits, residues, blood, and other

fluids from the surgical site to enable a better view of the surgical field.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: ANDREWS-PYNCHON Suction tube

Risk classification: IIa

Basic-UDI-DI: 405018300000015WQ

Intended purpose: ANDREWS-PYNCHON is used to remove fluid collections, blood and

debris from the operating site, in order to view critical structures, such

as blood vessels.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: BARON Suction tube

Risk classification: IIa

Basic-UDI-DI: 405018300000023WP

Intended purpose: BARON suction tube is used for atraumatic cleaning of the operating

field, through which the view of blood vessels and other structures is

improved.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: BELLUCCI Suction tube

Risk classification: IIa

Basic-UDI-DI: 405018300000593Y5



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Intended purpose: The BELLUCCI suction tube serves as a channel for the aspiration of

blood, fluids and other tissue residues during a surgical procedure.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: COOLEY Suction tube

Risk classification: IIa

Basic-UDI-DI: 405018300000591XZ

Intended purpose: The COOLEY suction tube is used to improve visibility of the surgical

field by aspirating liquids, debris and particles.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: DE BAKEY Suction tube

Risk classification: IIa

Basic-UDI-DI: 405018300000016WS

Intended purpose: DE BAKEY suction tube severs as a conductor for the suction of blood,

fluids and other tissue residues during a surgical procedure.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: FERGUSSON (POPPEN) Suction tube

Risk classification: IIa

Basic-UDI-DI: 405018300000019WY

Intended purpose: FERGUSSON (POPPEN) suction tube is used for the atraumatic suction

of liquids from cavities, especially during otological surgery.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: FRAZIER Suction tube

Risk classification: IIa

Basic-UDI-DI: 405018300000017WU

Intended purpose: FRAZIER suction tube is used to atraumatically clean the surgical site

and to expose vital structures.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: FUKUSHIMA Suction tube

Risk classification: IIa

Basic-UDI-DI: 405018300000022WM

Intended purpose: FUKUSHIMA suction tube is a flexible suction tube that is used to

aspirate liquids from areas that are difficult to access.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: GUILLEN suction raspatorium



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Risk classification:

Basic-UDI-DI: 405018300000594Y7

IIa

Intended purpose: The GUILLEN suction raspatory is used to scrape tissue and aspirate

fluids during a rhinological procedure.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: MAGILL Suction tube

Risk classification: IIa

Basic-UDI-DI: 405018300000592Y3

Intended purpose: The MAGILL suction tube is used to aspirate blood and other fluids, as

well as small tissue fragments from the surgical area.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: PLESTER Suction tube

Risk classification: IIa

Basic-UDI-DI: 405018300000018WW

Intended purpose: PLESTER suction tube is used to remove liquids, debris of foreign

bodies from the nasal cavity, maxillary antrum, or other small

anatomical cavities.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: POOLE Suction tube

Risk classification: IIa

Basic-UDI-DI: 405018300000590XX

Intended purpose: The POOLE suction tube is used to suck out large accumulations of

liquid from cavities.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: SCHUKNECHT Suction tube

Risk classification: IIa

Basic-UDI-DI: 405018300000024WR

Intended purpose: SCHUKNECHT suction tube is used for atraumatic cleaning of the

operating field, through which the view of blood vessels and other

strictures is improved.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Van Eicken sinus irrigation tube

Risk classification: IIa

Basic-UDI-DI: 405018300000154X7



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Intended purpose: VAN EICKEN frontal sinus irrigation tube is used to irrigate the surgical

field, clean infected tissue, and prevent it from spreading.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: WALTON-YANKAUER Suction tube

Risk classification: IIa

Basic-UDI-DI: 405018300000588YC

Intended purpose: The WALTON-YANKAUER suction tube is used to reject fluids and

debris to clear the view of the surgical site.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: YANKAUER Suction tube

Risk classification: IIa

Basic-UDI-DI: 405018300000008WT

Intended purpose: YANKAUER suction tube is used to remove fluids and residues, thus

clearing the surgical site.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: YASARGIL Suction tube

Risk classification: IIa

Basic-UDI-DI: 405018300000020WH

Intended purpose: YASARGIL suction tube is used to aspirate debris and liquids and thus

enable an improved view of the operating field.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Heparin suction cannula with button

Risk classification: IIa

Basic-UDI-DI: 405018300000152X3

Intended purpose: The heparin irrigation cannulas are used to inject solution into vessels.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Infusion cannula

Risk classification: IIa

Basic-UDI-DI: 405018300000153X5



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Intended purpose: The infusion cannula is used to create an atraumatic access to the

venous system and to inject and remove fluids from there.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: MOCK heparin needle with button

Risk classification: IIa

Basic-UDI-DI: 405018300000151WZ

Intended purpose: The MOCK Heparin needles are used to probe small blood vessels or

inject solutions via an atraumatic access.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: NOVAK suction curette

Risk classification: IIa

Basic-UDI-DI: 405018300000230WW

Intended purpose: The NOVAK suction curette is used to scrape tissue from the uterus

and to aspirate fluids..

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: RANDALL suction curette

Risk classification: IIa

Basic-UDI-DI: 405018300000230WW

Intended purpose: The RANDALL suction curette is used to grasp the uterine wall and

scrape away some of the tissue of the uterine cavity.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: MiniFESS Straight Suction

Risk classification: IIa

Basic-UDI-DI: 405018300000155X9

Intended purpose: The suction / irrigation cannulas are intended for aspiration of liquids

and fragments. They are also intended for irrigation with liquids. The suction / irrigation cannulas can be used together with navigation systems to better navigate the instrument in the human body during

application.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Suction/irrigation systems



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Risk classification:

Basic-UDI-DI: 405018300000355XK

IIa

Intended purpose: The suction/ irrigation systems for laparoscopy are used to rinse the

operating area and/ or to suck off the liquids and tissue fragments

that have accumulated during the procedure.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: SCHMID cannula

Risk classification: IIa

Basic-UDI-DI: 405018300000150WX

Intended purpose: The SCHMID irrigation cannulas are used to inject solution into

vessels.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: VERES insufflation cannula

Risk classification: IIa

Basic-UDI-DI: 405018300000076XC

Intended purpose: A VERES Insufflation Cannula is used to insufflate carbon dioxide

through an atraumatic access to the peritoneal cavity and thus to

generate a pneumoperitoneum.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: LANDAU Trocar

Risk classification: IIa

Basic-UDI-DI: 405018300000595Y9

Intended purpose: LANDAU trocars are used to provide access to a body cavity and to

aspirate fluids via the tube or to irrigate the surgical site via an

irrigation tube.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Puncture cannula Needle

Risk classification: IIa

Basic-UDI-DI: 405018300000356XM

Intended purpose: Puncture needles are used to puncture tissues and organs, to inject

fluids or a contrast medium, or to aspirate fluids.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: OLSEN SWAN cholangiography forceps

Risk classification: IIa



**Certificate ID: 1000220878** 

Basic-UDI-DI: 405018300000573XX

Intended purpose: Cholangiography forceps are intended to be used to grasp the bile

ducts and to place a contrast catheter for intraoperative visualization

of the gallbladder or bile ducts.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Cystoscopes

Risk classification: IIa

Basic-UDI-DI: 405018300000580XU

Intended purpose: Cystoscopes are intended for endoscopic visualization of inner

anatomical structures in the area of the urethra and urinary bladder.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Hysteroscopes

Risk classification: IIa

Basic-UDI-DI: 405018300000581XW

Intended purpose: Hysteroscopes are designed for endoscopic visualization of inner

anatomical structures in the uterine region.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Nephroscopes

Risk classification: IIa

Basic-UDI-DI: 405018300000582XY

Intended purpose: Nephroscopes are designed for endoscopic visualization of inner

anatomical structures in the area of the kidney.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Ureterorenoscopes

Risk classification: IIa

Basic-UDI-DI: 405018300000583Y2

Intended purpose: Ureterorenoscopes are intended for endoscopic visualization of inner

anatomical structures in the area of the ureters and renal pelvises.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Arthroscopes

Risk classification: IIa

Basic-UDI-DI: 405018300000584Y4

Intended purpose: Rigid arthroscopes are designed for endoscopic visualization of inner

anatomical structures in joints and skeletal structures.

Device category: MDN 1208 - Non-active non-implantable instruments



**Certificate ID: 1000220878** 

Product name: Laparoscopes

Risk classification: IIa

Basic-UDI-DI: 405018300000585Y6

Intended purpose: Laparoscopes are intended for endoscopic visualization of inner

anatomical structures in the area of the abdominal cavity and the

organs located therein.

Device category: MDA 0312 - Other active non-implantable surgical devices

Product name: HF- electrodes

Risk classification: IIb

Basic-UDI-DI: 405018300000586Y8

Intended purpose: HF electrodes are intended for endoscopically controlled tissue

resection, vaporization, enucleation, incision or coagulation in urology

and gynecology.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Electrode holder

Risk classification: IIa

Basic-UDI-DI: 405018300000587YA

Intended purpose: Electrode holders, HF surgery are used to hold and fix the endoscope

as well as for endoscopically controlled insertion of the electrode (cold electrode, HF electrode, stricture gauge) or the laser probe into the

surgical site.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Sheats Risk classification: IIa

Basic-UDI-DI: 405018300000601XB

Intended purpose: Sheaths are used to create a working or irrigation channel during

surgical procedures for diagnosis and therapy in urology and

gynecology

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: inserts Risk classification: IIa

Basic-UDI-DI: 405018300000602XD

Intended purpose: Working inserts enable the introduction of working instruments into

the surgical field via the integrated working channel. They are used in conjunction with sheaths and endoscopes during surgical procedures

for diagnosis and therapy in urology and gynecol.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: optical bridges

Risk classification: IIa



**Certificate ID: 1000220878** 

Basic-UDI-DI: 405018300000603XF

Intended purpose: Optical bridges enable visualization as well as the insertion of

instruments into the surgical field. They are used in conjunction with sheaths and endoscopes in surgical procedures for diagnosis and

therapy in urology and gynecology.

Device category: MDA 0312 - Other active non-implantable surgical devices

Product name: Endoor Cut scissor inserts

Risk classification: IIb

Basic-UDI-DI: 405018300000574XZ

Intended purpose: The Endoor Cut disposable scissor inserts are used in conjunction with

a reusable handle and shaft for cutting while simultaneously

coagulating soft tissue sections, organs or foreign bodies via natural

or surgically created accesses.

Device category: MDA 0312 - Other active non-implantable surgical devices

Product name: Endoor C coagulation forceps

Risk classification: IIb

Basic-UDI-DI: 405018300000575Y3, 405018300000576Y5, 405018300000578Y9,

405018300000579YB, 405018300000577Y7

Intended purpose: The dismountable, monopolar Endoor C coagulation forceps are used

for endoscopically controlled grasping, manipulation and cutting during simultaneous coagulation, as well as for dissection and

sampling of soft tissue sections, organs or foreign bodies via naturally

or surgically created accesses.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Rib spreader

Risk classification: IIa

Basic-UDI-DI: 405018300000596YB

Intended purpose: The Rib spreader is used to spread the ribs during a thoracotomy to

provide better access to the surgical field.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Ring retractor system

Risk classification: IIa

Basic-UDI-DI: 405018300000597YD

Intended purpose: The Ring Retractor System is used to fix the abdominal cavity, organs,

skin as well as internal tissues to provide better access to the surgical

field.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: GELPI Retractor



**Certificate ID: 1000220878** 

Risk classification:

IIa

Basic-UDI-DI:

405018300000598YF

Intended purpose:

The GELPI retractor is used to retract thick and hard tissue to provide

better access to the surgical field.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: Abdominal Retractor

Risk classification: IIa

Basic-UDI-DI: 405018300000600X9, 405018300000606XM,405018300000607XP,

405018300000608XR

Intended purpose: The Abdominal Retractor is used to hold tissues and organs in place to

provide better access to the surgical field.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: CHARNLEY retractor

Risk classification: IIa

Basic-UDI-DI: 405018300000604XH

Intended purpose: The CHARNLEY retractor is used to fix tendons and skin for better

access to the surgical field. The weight system is used to pull apart

hard fibrous tissue.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: retractors

Risk classification: IIa

Basic-UDI-DI: 405018300000609XT, 405018300000611XE, 405018300000612XG,

405018300000613XJ, 405018300000614XL, 405018300000619XW, 405018300000620XF, 405018300000621XH, 405018300000622XK,

405018300000623XM, 405018300000624XP

Intended purpose: The retractor is used to retract incision surfaces and wounds to

provide better access to the surgical field.

Device category: MDN 1208 - Non-active non-implantable instruments

Product name: LEYLA retractor

Risk classification: IIa

Basic-UDI-DI: 405018300000616XQ

Intended purpose: Retractors / holding systems with adjustable angle enable hands-free

use of additional instruments during an operation. They can be designed rigidly or individually adaptable, such as spreader with rail

(rigid), Leyla retractor system (flexible).



**Certificate ID: 1000220878** 

#### **Examinations and tests performed:**

448891\_A209802MED\_02 dated 10.07.2022 448891\_A212926MED\_03 polypectomy snares dated 2024-12-09 448891\_A212926MED\_04 Suction/irrigation instrument dated 2025-01-27

#### Further conditions for or limitations to the validity of the certificate:

In the case of reusable surgical instruments, the involvement of the Notified Body in the conformity assessment procedure is limited to the aspects related to reuse, in particular cleaning, disinfection, sterilization, maintenance and functional testing, as well as the related instructions for use.

#### Reference to previous certificates:

Revision	Date of Issue	Certificate-ID	Description of change
01	2022-12-15	170779053	Addition of Product "polypectomy
			snares TF16"
02	2025-01-26	1000209540	Addition of Product "Suction/irrigation
			instruments TF12"
03	2025-01-31	1000214224	Addition of product files using the "TF-
			02, TF-04, TF-07, TF-10 and TF11"
			sampling procedure