





Instruction for Use - Bi-Track® - Guiding Catheter SU

Attention:

This medical device may only be purchased by specialists, doctors and medical personnel and may only be used in accordance with these instructions for use and the defined area of application.

Description / Versions

This guiding catheter has two lumens for simultaneous placement of two .035" guide wires. The tube of the catheter is Fr. 8 with a total length of 195 cm. Distally, a metal ring was integrated into the tube in order to be able to monitor the advance radiologically.

Products

This user manual is valid for the products listed below:

E200-BTR

Important Note

Please ensure you read this user manual carefully before each use and keep it somewhere that is easily accessible for users and/or appropriate specialist staff.

Ensure you carefully read the warnings. Patients, users or third parties may suffer serious injury as a result of the improper use of the products.

Content and Packaging

- · 1 outer box
- 1 Bi-Track® Guiding Catheters SU (Single Use) individually sterile packed
- · 1 Instruction for use

Patient Population

The patient population or target patient group is derived from the indication determined by the responsible doctor who diagnostically or therapeutically treats the patient as part of an endoscopic procedure (the leading procedure per se) according to the intended use of the medical device.

There are no known restrictions to the patient population or the target patient group.

Use of the product on minors:

The user can use the product on minors in the case that the physiological and anatomical conditions of the patient permit the use of the product.

Use of the product on women who are pregnant or breastfeeding:

The indication for the use of the product on women who are pregnant or breastfeeding must be narrowed down by the user based on the respective individual physiological and anatomical conditions of the patient.

1. Scope of Application

The products listed above must only be used by physicians with appropriate specialist training in gastroenterology. The products are intended solely for the medical sector as illustrated below and must therefore be used within an operating environment suitable for this purpose. It is essential that both the user as well as the appropriate specialist staff familiarize themselves with the instruments before the user makes use of them.

2. Application Period

These instruments are intended to be used uninterrupted for a period of less than 60 minutes under normal conditions (MDD 93/42 EEC).

3. Intended Use

Bi-Track® Guiding Catheter are designed for transpapillary placement to place simultaneously two Guide Wires as preparation for a double stent-implantation in both ducts of Hepatic Fork (Ductus hepaticus communis and Ductus cysticus).

The application is carried out via the working channel of a duodenoscope. Indications

· Endoscopic implantation of biliary polymeric stents

Contraindications

- · Not fasting patients
- Diabetic patients
- · Fragility of the intestinal wall: e.g. highly florid inflammation of the colon (e.g. ulcerative colitis, diverticulitis, ulcerative colitis, toxic megacolon)
- · Peritonitis, acute abdomen, e.g. intestinal perforation, ileus
- Co-morbidity, e.g. severe cardiopulmonary diseases and decompensation
- Uncontrollable haemorrhagic diatheses
- Recently created gastrointestinal anastomosis.
- Stent migration
- Stent occlusion

4. Complications / Side Effects / Cross-reactions

Sedation during endoscopic examination increases the risk of hypoxemia, hypercapnia, hypotension, arrhythmias and aspiration due to reduced protective reflexes. Hypoxemia also occurs without sedation during endoscopic examinations due to the feed of the endoscope. Possible injuries in connection with endoscopic examinations can be:

- perforations,
- bleeding and Post-ERCP bleeding,
- · infections such as acute pancreatitis,
- Post ERCP cholangitis and cholecystitis
- Long term consequences of ERCP

Warnings / Precautions

These instructions must be followed, as well as instructions from compatible components and hospital regulations for infection prevention, safe use, cleaning and sterilization. Failure to do so may result in serious injury to the patient and/or the

The following applies to the product:

- Sterile only if packaging is undamaged or unopened!
- For single use only! Do not reuse, reprocess or sterilize several times. Reuse, reprocessing or repeated sterilization of the instrument may affect its structural integrity and cause malfunction, resulting in contamination, infection and serious
- If the instrument accidentally becomes dirty before treatment, it must be disposed of immediately! No cleaning agents may be applied.
- Do not use after the expiration date!
- All components should be carefully checked for compatibility and integrity before use. Do not use defective instruments! If defects occur, dispose of the instrument and replace it with a new one.
- Never use the product outside the recommended technical specifications (intended use)
- Never tamper with the structural conditions of the instrument, avoid kinks and other damage, and immediately discontinue use in the event of a malfunction!
- Wear protective clothing (gloves, goggles, gown, etc.) is absolutely necessary!
- Comparison of the technical data of the product with those of the endoscope used. The working channel diameter must be at least 0.2 mm larger than the outer diameter of the instrument.
- Never force instruments into the working channel!

Appropriate preparations for complications must be made before starting the application!

6. Compatibility

- **Guide Wires**
- Duodenoscopes

Function Test

The medical devices must be checked with regard to the following aspects prior to use:

- Expiration date
- Undamaged packaging
- · Damage on the product (cracks, bending, deformation)

Inspect products for immaculate surfaces, correct assembly and functionality.

Products that have failed the function test must not be used as their sterility and product safety cannot be ensured. Dispose of these products accordingly, or please return them to the manufacturer.

Preparation / Application

- Check the instrument for reliable function and irregularities before the procedure.
- If you notice any irregularities, replace the instrument with a new one!

Note: To facilitate the application, an endoscopic sphincterotomy prior to guide wire placement is recommended. (Please observe HF-generator and sphincterotome manufacturer's instructions).

- Place the duodenoscope close to the papilla.
- Insert (if possible, under fluoroscopic control) the Guiding Catheter through the working channel into the biliary duct and through the stenosis into the Ductus Cysticus.
- Insert successively both Guide Wires through the corresponding Luer-Lock-Connector into the catheter.
- Check the correct egression of the Guide Wires (Guide Wire 1 distal, Guide Wire 2 lateral) into both ducts of Hepatic Fork.
- Remove carefully the Guiding Catheter over the placed Guide Wires to use this wires for the planned stent placement.
- 6. Dispose the catheter after application.

Reprocessing and Sterilisation

These instruments are delivered in sterile condition and CANNOT EFFECTIVELY be cleaned, disinfected and sterilised after single use on account of the design which can no longer be removed and must be disposed of after single use.

The products are only sterile if the packaging is undamaged and unopened and if the shelf life is not exceeded. Products whose packaging is damaged or whose shelf life has expired must be discarded









Instruction for Use - Bi-Track® - Guiding Catheter SU

10. Sterility

Delivery condition

Disposable medical devices are supplied in an ETO gas-sterilized manner. Renewed preparation and sterilization following use is no longer possible and is prohibited! The product may only be used once.

Resterilisation should not be performed following expiry of the shelf life, i.e. the product must be disposed of according to clinical provisions.

11. Shelf Life of Products

The shelf life of the product is typically 3 years after the date of manufacture under

12. Liability and Warranty

As the manufacturer here, ENDO-FLEX shall not be liable for any damage or consequential damage arising from improper use or handling. This shall in particular apply to any use not in line with the defined purpose or failure to observe the preparation and sterilisation instructions and warnings. This shall also apply to repairs or modifications to the product made by individuals who are not authorised to do so by the manufacturer.

13. Service/Repairs

Do not carry out any repairs or modifications to the product. Only staff authorized by the manufacturer shall be responsible and intended to carry out such work. If you have any concerns, complaints or comments regarding our products, please get in touch with us. No liability whatsoever shall be assumed in the event of repairs carried out by individuals not authorized by the manufacturer.

14. Transport and Storage Conditions

- Products may only be transported and stored in the packaging provided for this
- Products must be stored dry and protected from sunlight at room temperature.
- Do not place any objects on the storage packaging and the sterile barrier system!
- Ensure the instruments are not kept near to chemicals, disinfectants or radioactive radiation!

15. Disposal

The products, packaging materials and accessories must all be disposed of in accordance with the nationally applicable regulations and laws. No specific instructions regarding this are given by the manufacturer.

16. Symbols used

REF

Symbol for "Item Number"

LOT

Symbol for "Batch code"

Symbol for "Manufacturer"

Symbol for "Date of manufacture"

Symbol for "Observe instructions for use"

STERILE EO Symbol for "Sterilized with ethylene oxide"



Symbol for "non-reuse"



Symbol for "Do not re-sterilize"



Symbol for "Do not use if packaging is damaged"



Symbol for "Expiry Date"



Symbol for "keep dry"



Symbol for "protect from sunlight"



Symbol for "Caution"

