

# EC Certificate

**mdc medical device certification GmbH**

Notified Body 0483  
herewith certifies that

**ENDO-FLEX GmbH**  
**Alte Hünxer Straße 115**  
**46562 Voerde**  
**Germany**

for the scope

**Endoscopic instruments, HF-instruments and accessories,  
Needle systems and Drainage systems  
(see attachment)**

has introduced and applies a

**Quality System**

for the design, manufacture and final inspection.

The mdc audit has proven that this quality system  
meets all requirements according to

**Annex II – excluding Section 4  
of the Council Directive 93/42/EEC**

of 14 June 1993 concerning medical devices.

The surveillance will be held as specified in Annex II, Section 5.

Valid from	2019-01-04
Valid until	2023-01-23
Registration no.	D1033500036
Report no.	P18-01361-131197
Stuttgart	2019-01-04



Head of Certification Body



**Attachment of the certificate**

**No. D1033500036**

Date 2019-01-04

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Product category	Product	Class
Drainage systems	Nasal Biliary Drainage Probes SU	Ila
Endoscopic instruments	Stone extraction Balloons SU	Ila
	Scissors RU	Ila
	Cytology Brushes SU	Ila
	Spray Catheters SU/RU	Ila
	Suture Punches RU	Ila
	Foreign Body Retrievers / Polyp Retrievers SU/RU	Ila
	Biopsy Forceps SU/RU	Ila
	Multi Band Ligation Device SU	Ila
Needle systems	Fibrin Application Needles SU/RU	Ila
	FNA Systems for ultrasound endoscopy SU	Ila
	Transbronchial Aspiration Needles SU	Ila
	Injection Needles SU/RU	Ila
Drainage systems	Biliary Stents SU	Ilb
	Pancreatic Stents SU	Ilb
	Self-expanding Stents SU (Biliary, Bronchial/Tracheal, Colonic, Duodenal, Esophageal)	Ilb
HF-instruments and accessories	Handles incl. HF connector RU	Ilb
	Cysto Gastro Sets SU	Ilb
	Sphincterotomes SU/RU	Ilb
	Polypectomy Snares, Mukosectomy Snares SU/RU	Ilb
	HOT Biopsy Forceps SU/RU	Ilb



  
 Head of Certification Body

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Notified Body 0483  
herewith certifies that

**ENDO-FLEX GmbH**  
**Alte Hünxer Straße 115**  
**46562 Voerde**  
**Germany**

for the scope

**endoscopic instruments**  
**(see attachment)**

has introduced and applies a

**Quality System**

for the aspects of manufacture concerned with securing and  
maintaining sterile conditions as specified in Annex V, Section 3.

The mdc audit has proven that this quality system  
meets all requirements according to

**Annex V – Section 3**  
**of the Council Directive 93/42/EEC**

of 14 June 1993 concerning medical devices.

The surveillance will be held as specified in Annex V, Section 4.

Valid from	2019-01-04
Valid until	2023-01-23
Registration no.	D1033500037
Report no.	P18-01361-131199
Stuttgart	2019-01-04



Head of Certification Body



**Attachment of the certificate**

**No. D1033500037**

Date 2019-01-04

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Product category	Product	Class
endoscopic instruments	E.R.C.P. Catheters SU/RU Suction / Flushing Catheters SU Stone Extraction Baskets SU/RU Lithotripsy Baskets / Lithotripsy Spirals SU/RU Guiding Catheters SU/RU Pushers SU/RU Stent Placement Sets SU/RU Biliary Dilation Catheters SU Polyp & Foreign Body Retriever "EasyCollect" SU Guide Wires SU/RU Dilation Balloons SU	I (steril)



Head of Certification Body

# EC Certificate

## Full Quality Assurance System

Certificate No.:  
**13422-2018-CE-CZS-NA-PS Rev. 2.0**

Project No.:  
**PRJC-575486-2017-PRC-CZE**

Valid Until:  
**01 November 2023**

This is to certify that the quality system of:

**Biosintex S.R.L.**

4 Vladiceasca Str.  
077168 Snagov  
Romania

For design, production and final product inspection/testing of:

**Sterile surgical sutures**

Has been assessed with respect to:

**The conformity assessment procedure described in Annex II of  
Council Directive 93/42/EEC on Medical Devices, as amended**

and found to comply.

Further details of the product(s) and conditions for certification are given overleaf.

Place and Date:  
**Høvik, 17 October 2019**



PROD 021  
Notified Body No.: 2460

**For: DNV GL Presafe AS**



**Palani Damodharan**

The Certificate has been digitally signed.  
See [www.presafe.com/digital\\_signatures](http://www.presafe.com/digital_signatures) for more info

Notice: The Certificate is subject to terms and conditions as set out in the Certification Agreement. Failure to comply may render this Certificate invalid.

# EC Certificate

## Full Quality Assurance System

Certificate No.:  
**13422-2018-CE-CZS-NA-PS Rev 2.0**

Project No.:  
**PRJC-575486-2017-PRC-CZE**

Valid Until:  
**01 November 2023**

### Jurisdiction

Application of Council Directive 93/42/EEC of 14 June 1993, adopted as "Forskrift om Medisinsk Utstyr" by the Norwegian Ministry of Health and Care Services.

### Certificate history:

Revision	Description	Issue Date
	Original Certificate	2018-11-01
1.0	Change of product name	2019-09-11
2.0	<b>Editorial change</b> <b>BICRIL changed to DACRIL</b> <b>BICRIL RAPID changed to DACRIL RAPID</b> <b>BICRIL 910 changed to DACRIL 910</b>	<b>2019-10-17</b>

### Products covered by this Certificate:

Product Description	Product Name	Class
<b>Surgical suture with /without needle</b>	<b>DACRIL-</b> Polyglycolic acid multifilament coated absorbable <b>DACRIL RAPID-</b> Polyglycolic acid multifilament coated fast absorbable <b>DACRIL 910</b> - Poly(glycolide-co-Lactide) (90/10) multifilament coated absorbable <b>PDO-x</b> - Polydioxanone monofilament absorbable <b>MONO-x</b> - Poly(glycolide-co-caprolactone) (75/25) monofilament absorbable <b>BIOPRO-</b> Polypropylene monofilament non-absorbable	III*

\* Design assessment is covered by a separate EC-Design Examination Certificate No.:  
13464-2018-CE-CZS-NA-PS

# EC Certificate

## Full Quality Assurance System

Certificate No.:  
**13422-2018-CE-CZS-NA-PS Rev 2.0**

Project No.:  
**PRJC-575486-2017-PRC-CZE**

Valid Until:  
**01 November 2023**

### Sites covered by this certificate

Site Name	Address
BIOSINTEX S.R.L.	4 Vladiceasca Str., RO 077168, Snagov, Romania

# EC Certificate

## Full Quality Assurance System

Certificate No.:  
**13422-2018-CE-CZS-NA-PS Rev 2.0**

Project No.:  
**PRJC-575486-2017-PRC-CZE**

Valid Until:  
**01 November 2023**

### Terms and conditions

The certificate is subject to the following terms and conditions:

- Any producer (see 2001/95/EC for a precise definition) is liable for damage caused by a defect in his product(s), in accordance with directive 85/374/EEC, as amended, concerning liability of defective products.
- The certificate is only valid for the products and/or manufacturing premises listed above.
- The Manufacturer shall fulfil the obligations arising out of the quality system as approved and uphold it so that it remains adequate and efficient.
- The Manufacturer shall inform Presafe of any intended updating of the quality system and Presafe will assess the changes and decide if the certificate remains valid.
- Periodical audits will be held, in order to verify that the Manufacturer maintains and applies the quality system. Presafe reserves the right, on a spot basis or based on suspicion, to pay unannounced visits.

The following may render this Certificate invalid:

- Changes in the quality system affecting production.
- Periodical audits not held within the allowed time window.

### Conformity declaration and marking of product

When meeting with the terms and conditions above, the producer may draw up an EC declaration of conformity and legally affix the CE mark followed by the Notified Body identification number of Presafe.

End of Certificate



# MANAGEMENT SYSTEM CERTIFICATE

Certificate No.:  
257642-2018-AQ-CZE-NA-PS rev. 2.0

Project No.:  
PRJC-575485-2017-MSC-CZE

Initial Certification Date:  
11 April 2019

Valid Until:  
11 April 2022

This is to certify that the management system of:

**BIOSINTEX S.R.L.**

4 Vladiceasca Str. 077168, Snagov, Ilfov County, Romania

Complies with the requirements of:

**ISO 13485:2016/NS-EN ISO 13485:2016**

The Certificate is valid for the following scope:

**DESIGN, DEVELOPMENT, MANUFACTURING AND TRADE OF  
STERILE SURGICAL SUTURES, WITH/ WITHOUT NEEDLES.**

Place and date:  
Hovik, 01 February 2021

For:  
DNV GL PRESAFE AS



*Tone Kolpus*

Tone Elise Kolpus

The certificate is digitally verified by Blockchain technology. For more info, see  
[www.dnvgl.com/resources/our/verifierinthe-blockchain.htm](http://www.dnvgl.com/resources/our/verifierinthe-blockchain.htm)



Notice: The Certificate is subject to terms and conditions as set out in the Certification Agreement. Failure to comply may render this Certificate invalid.  
ACCREDITED UNIT: DNV GL PRESAFE AS, Vertasveien 3, N-1363 Hovik, Norway • Registered Enterprise No: NO 097 067 401 MVA



The management system of

## Teleflex Medical

2917 Weck Drive, Research Triangle Park, NC, 27709, United States

has been assessed and certified as meeting the requirements of

### Directive 93/42/EEC

on medical devices, Annex II (excluding Section 4)

For the following products

The scope of registration appears on page 2 of this certificate.

This certificate is valid from 11 September 2018 until 14 July 2023  
and remains valid subject to satisfactory surveillance audits.

Re certification audit due before 27 May 2021


Issue 29. Certified since 26 September 2000

Certification is based on reports numbered WWW/MC/06866

Multiple certificates have been issued for this scope

The main certificate is numbered US97/10879.00

Authorised by



SGS United Kingdom Ltd, Notified Body 0120

2026 Worle Parkway, Weston-super-Mare, BS22 6WA UK  
t +44 (0)1934 522917 f +44 (0)1934 522137 www.sgs.com

SGS CE 02 0315 M2

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# Teleflex Medical

## Directive 93/42/EEC

on medical devices, Annex II (excluding section 4)

Issue 29

Detailed scope

Sterile Hem-o-lok Ligation Clips.  
Sterile Deknatel® PTFE pledgets.  
Sterile Polyester Nonabsorbable Surgical Sutures (POLYLENE/ "cottony"™ II, "silky" II POLYDEK®, TEVDEK® II, NextSitch®, Capio™, Fx®, NiceLoop™, TEVDEK®).  
Sterile DEKLENE® II, DEKLENE® MAXXTM, CAPIOTM and FIXT® polypropylene non-absorbable surgical sutures.  
Sterile BONDEK® and BONDEK® Plus Polyglycolic Acid Synthetic Absorbable Surgical Sutures.  
Sterile Polyglytone 6211™ Monofilament Absorbable Surgical Sutures.  
Sterile MONODEK® Polydioxanone Absorbable Surgical Sutures.  
Sterile Hem-o-lok Automatic Clip Applicators.  
Metal Ligation System.

Sterile External stapling system (including stainless steel staples, staplers and removers), Sterile, Efx endo fascial closuresystem (abdominal access), Sterile, Efx shield fascial closure system (abdominal access), Sterile, Efx classic fascial closuresystem (abdominal access)

Sterile stainless steel surgical Sutures  
Sterile FORCE FIBER® surgical sutures.  
Sterile Chest drainage and autotransfusion systems,  
Sterile Thoracic Catheters,  
Sterile and Non-sterile Aortic Punch,  
Non-sterile Self Retaining Tissue retractor/blades

Non-sterile Anaesthesia and respiratory Circuits including breathing bags and water traps,  
Non-sterile Heated Humidifiers, Non-sterile Non-Pre-filled Humidifiers and Nebulizers, Non-sterile Small Volume Nebulizers, Sterile Pre-filled Humidifiers and Nebulizers (saline or water) with adaptors, Sterile Pre-filled unit dose vial /solution for nebulisation, Non-sterile Respiratory therapy Adaptors and connectors, Sterile Column and Reservoirs including adaptors, Non-sterile Nasal cannula (including gas sampling), Non-sterile Cannula and Supply Tubing, Nonsterile CPAP Cannula System, Non-sterile Manual resuscitators and PEEP valves, Non-sterile Respiratory and anaesthesia masks, Non-sterile Gas scavenging mask, Sterile Endotracheal tubes, Sterile Endobronchial tubes, Non-sterile Suction and Aspirating Tubes, Sterile Vented Thoracic Chest Seal, Sterile Operative Cholangiogram Catheters, Sterile Abdominal Access and Insufflation devices, Sterile Capillary drains, Sterile Percutaneous Surgical System (MiniLap and Grip graspers), Sterile Percutaneous Surgical System (Mini Polar electrosurgical probe and MiniGrip Bipolar Graspers), Percutaneous surgical System (Interchangeable electrosurgical tool tips) for laparoscopic surgery, Non-sterile Heat and Moisture Exchangers

Where the above scope includes class III medical device(s), a valid EC Design Examination Certificate according to Annex II (Section 4) is a mandatory requirement for each device in addition to this certificate to place that device on the market

