

# **Disposable Endoscopic Scissors**

# Instructions for Use



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#### About this Manual

This manual provides information on how to operate the HERizon Disposable Endoscopic Scissors. It is essential to read and understand all the information in this manual before using the device. Pay attention to all warnings, contraindications, precautions and adverse events in this manual and other related materials. Failure to thoroughly understand and follow all instructions may result in harm to the patient or user of the device.

#### Reuse Warning

The HERizon Disposable Endoscopic Scissors are supplied STERILE using Ethylene Oxide. Do not use if sterile barrier is damaged. If damage is found, call your Minerva Surgical representative.

The HERizon Disposable Endoscopic Scissors are for single use only. Do not reuse, reprocess or resterilize. Reuse, reprocessing or resterilization may compromise the structural integrity of the device and/or lead to device failure which, in turn, may result in patient injury, illness or death.

Reuse, reprocessing or resterilization may also create a risk of contamination of the device and/or cause patient infection or cross-infection, including, but not limited to, the transmission of infectious disease(s) from one patient to another. Contamination of the device may result in patient injury, illness or death.

After use, dispose of product and packaging in accordance with hospital, administrative and/or local government policy.

#### **Device Description**

The HERizon Disposable Endoscopic Scissors (Catalog # JY-JD-17-350-B) are supplied sterile, sterilized by ethylene oxide, and single use only.

The HERizon Disposable Endoscopic Scissors are composed of a handle, knob, scissors bar, and scissors head (Figure 1).

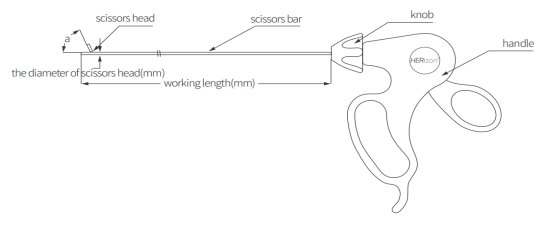


FIGURE 1: DISPOSABLE ENDOSCOPIC SCISSORS

## Intended Use/Indications for Use

The HERizon Disposable Endoscopic Scissors are designed to cut tissue during endoscopic procedures.

#### Contraindications

Hysteroscopy may be contraindicated by the following conditions, depending on their severity or extent:

- inability to distend the uterus
- cervical stenosis
- cervical/vaginal infection
- uterine bleeding or menses

- known pregnancy
- invasive carcinoma of the cervix
- recent uterine perforation
- medical contraindication or intolerance to anesthesia

#### Warnings/Precautions

- Do not attempt to disassemble the Scissors.
- Handle the Scissors with care.
- If the Scissors do not function properly, replace with another properly functioning Scissors.
- Do not place the Disposable Scissors in environments with high temperatures, humidity, dust, direct sunlight or corrosive gases.
- Only those personnel who have been trained or whose knowledge and practical experience are sufficient to ensure correct operation should operate and use this device.
- Only those who have read the appropriate IFUs and know the correct use of the system and accessories can operate
  it.
- If this system is to be used with other equipment, the user should consult applicable IFUs and receive appropriate training on the use of the equipment and use it strictly in accordance with its IFU.
- This product is for prescription only.
- The disposable endoscopic Scissors are ethylene oxide (EO) sterilized and for single use only.

#### **Environmental Protection**

Follow local governing ordinances and hospital practice regarding the disposal of the HERizon Disposable Endoscopic Scissors.

The shelf life of HERizon Disposable Endoscopic Scissors is 3 years.

### **How Supplied**

DO NOT USE IF PACKAGE IS OPENED OR DAMAGED. DO NOT USE IF LABELING IS INCOMPLETE OR ILLEGIBLE.

The HERizon Disposable Endoscopic Scissors are supplied sterile and are intended for single use. The shelf box contains:

- One (1) Disposable Endoscopic Scissors
- One (1) Disposable Endoscopic Scissors eIFU Insert

#### Set Up and Operation

IMPORTANT: In addition to these instructions, follow the instruction manuals or IFUs of the products used in conjunction with this product. See <a href="https://www.minervasurgical.com/IFU\_Symbol-Glossary">www.minervasurgical.com/IFU\_Symbol-Glossary</a> for additional information. Refer to the HERizon Medical Image Processor System Operator's Manual for additional details and instructions on system set up and operation. Refer to Figure 1 for device image.

- 1. Using sterile technique, remove the scissors from the packaging. Inspect the scissors and tip for damages prior to use.
- 2. Prior to inserting into the endoscope working channel, grip the front handle slot with your fingers and back handle slot with your thumb. Test squeeze the handle 1-2 times to verify and confirm proper movement of the Head
- 3. Introduce the Scissors into the working channel of the Endoscope. The Head must be in the closed position by squeezing the handle.
- 4. Once inserted into the working channel of the endoscope, advance the Scissors Head towards the target area.
- 5. Adjust the knob to align the Head with the target tissue.
- 6. Squeeze and release the handle to close and open the Head to conduct tissue dissection.
- 7. Squeeze the handle such that the Scissors are in the closed position and slowly withdraw the entire device from the endoscope.

# Storage and Handling

The HERizon Disposable Endoscopic Scissors must be kept in a cool and dry environment.

#### Maintenance

Do not use after expiration date as specified on the device label. Single Use Items must not be repaired or reused.

#### **Limited Warranty**

Minerva Surgical warrants that the device(s) supplied is free from defects in materials or workmanship. The Disposable Endoscopic Scissors has a 3 year shelf life and is warranted throughout the intended shelf life, so long as the product has been maintained in accordance with procedures documented in the IFU.

The following circumstances are not covered by the warranty and will cause all liability statements to be invalid: faults caused by non-standard or improper use, incorrect treatment, and modification or repair of device. The packaging materials of devices and accessories sold by Minerva Surgical are all designed to protect them from damage.

## Customer Service/Technical Support

Contact Minerva Surgical Customer Service for customer support.

Call +1 (855) 646-7874

# Symbol Glossary

Symbol Glo Symbol	Symbol Title	Standard	Description
	Contents	ISO 15223-1	Package content
UDI	Unique device Identifier	ISO 15223-1	Indicates a carrier that contains unique device identifier information
	Date of manufacture	ISO 15223-1	Indicates the date when the medical device was manufactured
***	Manufacturer	ISO 15223-1	Indicates the medical device manufacturer
53	Use-by date	ISO 15223-1	Indicates the date after which the medical device is not to be used
LOT	Batch code	ISO 15223-1	Indicates the manufacturer's batch code so that the batch or lot can be identified
REF	Catalog number	ISO 15223-1	Indicates the manufacturer's catalog number so that the medical device can be identified
STERNIZE	Do not resterilize	ISO 15223-1	Indicates a medical device that is not to be resterilized
<b>(a)</b>	Do not use if package is damaged and consult instructions for use	ISO 15223-1	Indicates that a medical device should not be used if the package has been damaged or opened and that the user should consult the instructions for use for additional information
	Do not re-use	ISO 15223-1	Indicates a medical device that is intended for single use only
	Consult instructions for use	ISO 15223-1	Indicates the need for the user to consult the instructions for use
*	Keep away from sunlight	ISO 15223-1	Indicates a medical device that needs protection from light sources

<del>*</del>	Keep dry	ISO 15223-1	Indicates a medical device that needs to be protected from moisture
	Temperature limit	ISO 15223-1	Indicates the temperature limits to which the medical device can be safely exposed
	Fragile, handle with care	ISO 15223-1	Indicates a medicaldevice that can bebroken or damaged if nothandled carefully.
	Stacking limit by six	ISO 7000 : 2018	To indicate that the items shall not be vertically stacked beyond the specified number
75%	Humidity limitation	ISO 15223-1	Indicates the range of humidity to which the medical device can be safely exposed
STERILEEO	Single sterile barrier system	ISO 15223-1	Indicates a single sterile barrier system
Rx only	U.S. Federal Law restricts this device to sale by or on the order of a licensed practitioner.	NA	Indicates this is a restricted device.

For Additional Product Information, Scan the QR Code



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