



November 9, 2022

Shanghai AnQing Medical Instrument Co., Ltd  
Shuwen Fan  
RA Manager  
3 & 4 Floor, No.2 Building, 366 Huiqing Road,  
East Zhangjiang High-Tech Park  
Shanghai, 201201  
China

Re: K222162  
Trade/Device Name: Flexible Cystoscope (Model: CY50H-20EU, CY50H-20US, CY55H-24EU, CY55H-24US)  
Regulation Number: 21 CFR§ 876.1500  
Regulation Name: Endoscope and Accessories  
Regulatory Class: II  
Product Code: FAJ  
Dated: October 11, 2022  
Received: October 11, 2022

Dear Shuwen Fan:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. Although this letter refers to your product as a device, please be aware that some cleared products may instead be combination products. The 510(k) Premarket Notification Database located at <https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfpmn/pmn.cfm> identifies combination product submissions. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration. Please note: CDRH does not evaluate information related to contract liability warranties. We remind you, however, that device labeling must be truthful and not misleading.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the Federal Register.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies.

You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part 801); medical device reporting (reporting of medical device-related adverse events) (21 CFR 803) for devices or postmarketing safety reporting (21 CFR 4, Subpart B) for combination products (see <https://www.fda.gov/combination-products/guidance-regulatory-information/postmarketing-safety-reporting-combination-products>); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820) for devices or current good manufacturing practices (21 CFR 4, Subpart A) for combination products; and, if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21 CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to <https://www.fda.gov/medical-devices/medical-device-safety/medical-device-reporting-mdr-how-report-medical-device-problems>.

For comprehensive regulatory information about medical devices and radiation-emitting products, including information about labeling regulations, please see Device Advice (<https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance>) and CDRH Learn (<https://www.fda.gov/training-and-continuing-education/cdrh-learn>). Additionally, you may contact the Division of Industry and Consumer Education (DICE) to ask a question about a specific regulatory topic. See the DICE website (<https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/contact-us-division-industry-and-consumer-education-dice>) for more information or contact DICE by email ([DICE@fda.hhs.gov](mailto:DICE@fda.hhs.gov)) or phone (1-800-638-2041 or 301-796-7100).

Sincerely,

**Mark J. Antonino -S**

Mark J. Antonino, M.S.  
Assistant Director  
DHT3B: Division of Reproductive,  
Gynecology and Urology Devices  
OHT3: Office of GastroRenal, ObGyn,  
General Hospital and Urology Devices  
Office of Product Evaluation and Quality  
Center for Devices and Radiological Health

Enclosure

## Indications for Use

510(k) Number (if known)  
K222162

Device Name

Flexible Cystoscope (Model: CY50H-20EU, CY50H-20US, CY55H-24EU, CY55H-24US)

Indications for Use (Describe)

AnQing Medical Single Use Flexible Cystoscope is used to provide visualization and operative access during diagnostic and therapeutic endoscopic procedures of the lower urinary tract, including the bladder and urethra.

The Flexible Cystoscope is intended for use in a hospital environment or medical office environment.  
The Flexible Cystoscope is designed for use in adults.

Type of Use (Select one or both, as applicable)

Prescription Use (Part 21 CFR 801 Subpart D)

Over-The-Counter Use (21 CFR 801 Subpart C)

### CONTINUE ON A SEPARATE PAGE IF NEEDED.

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510(k) Premarket Notification Submission  
Flexible Cystoscope

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510(k) Summary

Date Prepared: July 8, 2022  
Manufacturer: Shanghai AnQing Medical Instrument Co., Ltd.  
3 & 4 Floor, No.2 Building, 366 Huiqing Rd, East  
Zhangjiang High-Tech Park, 201201  
Shanghai, China

Contact Person: Shuwen Fan  
RA Manager  
Shanghai AnQing Medical Instrument Co., Ltd.  
Tel: +86-21-61117375  
Fax: +86-21-61117374  
[ra\\_dept@anqing-sh.com](mailto:ra_dept@anqing-sh.com)

**Identification of the Device:**

Proprietary/Trade Name: Flexible Cystoscope  
Model: CY50H-20EU, CY50H-20US, CY55H-24EU,  
CY55H-24US  
Common name: Flexible Cystoscope  
Classification Name: Endoscope and Accessories  
Regulatory Number: 21 CFR Part 876.1500  
Product Code: FAJ  
Device Class: Class II  
Review Panel: Gastroenterology/Urology

**Identification of the Legally Marketed Predicate Device:**

Trade Name: Ambu aScope 4 Cysto  
Common name: Flexible Cystoscope  
Classification Name: Cystoscope And Accessories, Flexible/Rigid  
Regulatory Number: 21 CFR Part 876.1500  
Product Code: FAJ  
Device Class: Class II  
Review Panel: Gastroenterology/Urology  
Submitter/510(k) Holder: Ambu A/S.  
Clearance: K193095 (cleared April 2, 2020)  
This predicate has not been subject to a design-related recall.



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**Device Description:**

The Flexible Cystoscope (Model: CY50H-20EU, CY50H-20US, CY55H-24EU, CY55H-24US) is a sterile single-use endoscope which is used with the video processor (Model: EOS-H-01, FDA cleared #K211169) produced by AnQing for providing endoscopic imaging of the lower urinary tract, including the bladder and urethra for the purpose of diagnosis and treatment.

The CY50H-20 and CY55H-24 difference is the size of Distal End Outer Diameter and Working Channel Inner Diameter. The US and EU model are identical except the deflection versions, which is opposite from each other (EU version or US version).

The Flexible Cystoscope is a single-use endoscope, which consists of Handle, Insertion Section, Distal Tip, and Endoscope Connector.

The handle includes a deflection lever, a lever lock, a push button for picture taking/video recording and a Luer port for insertion of accessory devices and irrigation to the working channel. The insertion section contains one working channel and wiring to transmit the image signals to the Video Processor. The distal bending section of the insertion section is controlled by the user via the deflection lever on the handle. The distal end of the insertion section contains a CMOS sensor for capturing image and transmitting it to the Video Processor, LEDs for illumination, and the distal opening of the working channel. The endoscope connector connects the endoscope handle to the video processor, which provides power and processes video signals from the endoscope.

**Mechanism of action:**

The light emitted by the LED cold light source at the distal tip of the disposable Flexible Cystoscope is irradiated into the body cavity, and the light reflected from the cavity enters the optical system and is captured by the CMOS image sensor. The CMOS acquisition image is controlled by the CMOS drive circuit, and the RGB video signal is output to the Video Processor via the VI circuit. The Video Processor receives video signals from the endoscope, processes the video signals, and outputs the processed video signal to the attached monitor. The video processor also controls the brightness of the LEDs on the endoscope.

Flexible Cystoscope has the following physical and performance characteristics:

- Maneuverable tip controlled by the user
- Flexible insertion cord
- Camera and LED light source at the distal tip
- Sterilized by Ethylene Oxide
- For single use



**Indications for Use:**

AnQing Medical Single Use Flexible Cystoscope is used to provide visualization and operative access during diagnostic and therapeutic endoscopic procedures of the lower urinary tract, including the bladder and urethra.

The Flexible Cystoscope is intended for use in a hospital environment or medical office environment.

The Flexible Cystoscope is designed for use in adults.

**Comparison with Predicate Device:**

The Flexible Cystoscope and its predicate device, the Ambu aScope 4 Cysto (K193095), have the same intended use, sterilization Methods and similar physical characteristics, optical characteristics.

**Substantial Equivalence:**

The Flexible Cystoscope employs the same fundamental scientific technology as its predicate device, as below table:

	Subject Device (CY50H-20EU, CY50H-20US, CY55H-24EU, CY55H-24US)	Predicate Device, Ambu aScope 4 Cysto (K193095)	Comparison
Indications For Use			
Indications For Use:	<p>AnQing Medical Single Use Flexible Cystoscope is used to provide visualization and operative access during diagnostic and therapeutic endoscopic procedures of the lower urinary tract, including the bladder and urethra.</p> <p>The Flexible Cystoscope is intended for use in a hospital environment or medical office environment.</p>	<p>Ambu® aScope™ 4 Cysto is a sterile, single-use, flexible cystoscope intended to be used for endoscopic access to and examination of the lower urinary tract. The Ambu® aScope™ 4 Cysto is intended to provide visualization via Ambu® displaying unit and can be used with endoscopic</p>	<p>Equivalent  See Note1.</p>



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Flexible Cystoscope

	Subject Device (CY50H-20EU, CY50H-20US, CY55H-24EU, CY55H-24US)	Predicate Device, Ambu aScope 4 Cysto (K193095)	Comparison
	The Flexible Cystoscope is designed for use in adults.	accessories. Ambu® aScope™ 4 Cysto is intended for use in a hospital environment or medical office environment. Ambu® aScope™ 4 Cysto is designed for use in adults.	
Physical Characteristics			
Type of Scope	Flexible	Flexible	Same
Outer diameter (mm)	Max. 5.0 mm (CY50H-20EU, CY50H-20US) Max. 5.5 mm (CY55H-24EU, CY55H-24US)	Max. 6.0 mm	Similar See Note2.
Inner diameter (mm)	Min. 2.0 mm (CY50H-20EU, CY50H-20US) Min. 2.4 mm (CY55H-24EU, CY55H-24US)	Min. 2.2 mm	Similar See Note2.
Working length	380 mm	390 mm (15.4")	Similar See Note2.
Deflection	210°up, 180°down	210° up, 120° down	Similar See Note2.
Optical Characteristics			
Type of Imager	CMOS	CMOS	Same
Field of View	110°	120°	Similar See Note2.
Direction of	0°	0°	Same



510(k) Premarket Notification Submission  
Flexible Cystoscope

	Subject Device (CY50H-20EU, CY50H-20US, CY55H-24EU, CY55H-24US)	Predicate Device, Ambu aScope 4 Cysto (K193095)	Comparison
View			
Depth of Field	5mm~100mm	3mm~100mm	Similar See Note2.
Light Source	Internal LED	Internal LED	Same
Patient Contacting Materials			
General material type of main patient-contact part	Compliance with ISO10993-1	Compliance with ISO10993-1	Similar See Note2.
Duration and type of contact	“Surface –Mucosal Membrane” with a contact duration of “Limited (< 24 hours)”	“Surface –Mucosal Membrane” with a contact duration of “Limited (< 24 hours)”	Same
Sterilization Methods			
Number of Users	Single-Use	Single-Use	Same
Sterilization	EO Sterilized, SAL 10 <sup>-6</sup>	EO Sterilized, SAL 10 <sup>-6</sup>	Same
Technological Characteristics			
Environment of use	Healthcare facility/hospital	Healthcare facility/hospital	Same
Energy source	Electricity	Electricity	Same
<p>Note:</p> <p>1. The indications for use statement for the subject Cystoscope is very similar to that of the predicate device. A slightly different is wording description. The differences do not alter the intended use of the device nor do they raise different questions of safety and effectiveness of the device relative to the predicate.</p> <p>2. The subject and predicate device have the same fundamental technology, type of scope, type of imager, light source, Sterilization method, number of users and environment. The subject Cystoscope differs from the predicate in Outer diameter, Inner diameter,</p>			





510(k) Premarket Notification Submission  
Flexible Cystoscope

	Subject Device (CY50H-20EU, CY50H-20US, CY55H-24EU, CY55H-24US)	Predicate Device, Ambu aScope 4 Cysto (K193095)	Comparison
Working length, Deflection, Field of View, Depth of Field, and patient-contacting materials. These differences do not raise different questions of safety and effectiveness as compared to the predicate, and can be evaluated through performance testing.			

**Summary of Testing:**

**Summary of Non-Clinical Tests:**

Electrical Safety and Electromagnetic Compatibility Summary

The electrical safety and EMC data included in the submission is in compliance with the following FDA recognized standards:

- ANSI/AAMI ES:60601-1:2005/A2:2010
- IEC 60601-2-18:2009: Part 2-18
- IEC 60601-1-2:2014

**Bench Testing Summary**

Photobiological safety

The LEDs in submitted Cystoscope were tested according to the following FDA recognized standards:

- IEC 62471:2006 Medical electrical equipment, Photobiological safety of lamps and lamp systems

**Mechanical and Optical Performance**

The Flexible Cystoscope was designed to comply with applicable parts of ISO 8600. Optical measurements were performed according to applicable part of ISO 8600 standard.

Mechanical characteristics were tested and include leakage tightness, bending, deflection endurance, withstand of channel.

In addition, comparative testing related to image quality parameters was performed for submitted Flexible Cystoscope and the predicate device to support substantial equivalence.



### **Biocompatibility Summary**

The biocompatibility evaluation for the patient contacting components of the Flexible Cystoscope was performed according to ISO 10993-1 and FDA Guidance. The following tests were conducted based contact category of “Surface – Mucosal Membrane” with a contact duration of “Limited (< 24 hours):

- Cytotoxicity: ISO 10993-5:2009/(R) 2014
- Sensitization, Intracutaneous reactivity/irritation: ISO 10993-10:2010
- Material-mediated pyrogenicity: ISO 10993-11:2017
- Acute systemic toxicity: ISO 10993-11:2017

### **Sterilization and shelf life testing**

The sterilization method has been validated to ISO 11135:2014, which has thereby determined the routine control and monitoring parameters.

EO/ECH residual test was performed according to ISO 10993-7:2008.

The shelf life of the Flexible Cystoscope is determined based on stability study which includes ageing test according to ASTM F1980-16, Standard Guide for Accelerated Aging of Sterile Barrier.

### **Package Validation**

Package validation was conducted according to ISO 11607-1:2019 and ISO 11607-2:2019, and F88/F88M-15, ASTM F 1929-15

Transport and shipping testing as per ASTM D4169-16.

### **Summary of Clinical Tests:**

The subject of this premarket submission, did not require clinical studies to support substantial equivalence.

### **Conclusion:**

The conclusions drawn from the nonclinical tests demonstrate that the subject device, the Flexible Cystoscope is substantially equivalent to the predicate device.