

June 17, 2022

Yangzhou Fartley Medical Instrument Technology Co., Ltd. % Ethan Liu RA Specialist Shanghai Thinkwell Consulting Co., Ltd Room 211/6F, Xinling Road, Minhang District Shanghai, Shanghai 201100 China

Re: K220210

Trade/Device Name: Disposable Endoscopy Adapter Set

Regulation Number: 21 CFR§ 876.1500

Regulation Name: Endoscope and Accessories

Regulatory Class: II Product Code: ODC Dated: May 9, 2022 Received: May 16, 2022

Dear Ethan Liu:

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 <u>Federal Register</u>.

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.

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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-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/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) or phone (1-800-638-2041 or 301-796-7100).

Sincerely,

Shanil P. Haugen, Ph.D.
Assistant Director
DHT3A: Division of Renal, Gastrointestinal,
Obesity and Transplant Devices
OHT3: Office of GastroRenal, ObGyn,
General Hospital and Urology Devices
Office of Product Evaluation and Quality
Center for Devices and Radiological Health

Enclosure

DEPARTMENT OF HEALTH AND HUMAN SERVICES Food and Drug Administration

Indications for Use

Form Approved: OMB No. 0910-0120

Expiration Date: 06/30/2023 See PRA Statement below.

510(k) Number <i>(if known)</i>	
K220210	
Device Name	
Disposable Endoscopy Adapter Set	
Indications for Use (Describe)	

The Disposable Endoscopy Adapter Set is a collection of several sterile units. It is intended to be fitted to multiple endoscope working channels/ports to control the flow of fluids, gases and other materials.

- Disposable Air/Water Valve: This unit is intended to be fitted to an endoscope air/water channel to control the inflow of medical gases and water, whilst preventing back-flow.
- Disposable Suction Valve: This unit is intended to be fitted to an endoscope suction channel to control the operations of suction, whilst preventing inflow of air.
- Disposable Biopsy Valve: This unit is intended to be fitted to an endoscope biopsy port to prevent leakage of gases and body fluids during an endoscopic procedure.
- Disposable Water Jet Adapter: This unit is intended to provide irrigation via sterile water supply during GI endoscopic procedures when used in conjunction with an irrigation pump.

Type of Use (Select one or both, as applicable)	
X Prescription Use (Part 21 CFR 801 Subpart D)	Over-The-Counter Use (21 CFR 801 Subpart C)

CONTINUE ON A SEPARATE PAGE IF NEEDED.

This section applies only to requirements of the Paperwork Reduction Act of 1995.

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510(k) Summary

This summary of 510(k) safety and effectiveness information is being submitted in accordance with requirements of 21 CFR Part 807.92.

5.1 Submitter

Submitted by:	Yangzhou Fartley Medical Instrument Technology Co., Ltd.	
	Address:Beizhou Road, Lidian Town, Guangling District, Yangzhou	
	225106 Jiangsu, China	
Contact Person:	Ethan Liu	
	RA Specialist	
	Shanghai Thinkwell Consulting Co., Ltd	
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	China.	
	Phone: 0086-15216699240	
	Email: xtdeepwater@126.com	
Date Prepared:	Jan 27, 2022	

5.2 Device

Device Name:	Disposable Endoscopy Adapter Set
Classification Name:	Endoscope Channel Accessory
Regulatory Class:	II
Regulation Number:	21 CFR 876.1500
Regulation Name:	Endoscope and Accessories
Product Code:	ODC

5.3 Predicate Device

Device Name:	Disposable Endoscope Valves Set, Disposable Endoscope
	Valves Set B, K200769
Manufacturer:	Wilson Instruments (SHA) Co., LTD
Classification Name:	Endoscope Channel Accessory
Regulatory Class:	II
Regulation Number:	21 CFR 876.1500
Regulation Name:	Endoscope and Accessories
Product Code:	ODC

5.4 Device Description

The Disposable Endoscopy Adapter Set is a collection of several sterile units. It is intended to be fitted to multiple endoscope working channels/ports to control the flow of fluids, gases and other materials. The sterile units may consist of Disposable Air/Water Valve, Disposable Suction Valve, Disposable Biopsy Valve and Disposable Water Jet Adapter. Disposable Endoscopy Adapter Set



may be configured as single valve or multiple valves in any combination up to a maximum of 4 valves.

5.5 Indication for Use:

The Disposable Endoscopy Adapter Set is a collection of several sterile units. It is intended to be fitted to multiple endoscope working channels/ports to control the flow of fluids, gases and other materials.

- Disposable Air/Water Valve: This unit is intended to be fitted to an endoscope air/water channel to control the inflow of medical gases and water, whilst preventing back-flow.
- Disposable Suction Valve: This unit is intended to be fitted to an endoscope suction channel to control the operations of suction, whilst preventing inflow of air.
- Disposable Biopsy Valve: This unit is intended to be fitted to an endoscope biopsy port to prevent leakage of gases and body fluids during an endoscopic procedure.
- Disposable Water Jet Adapter: This unit is intended to provide irrigation via sterile water supply during GI endoscopic procedures when used in conjunction with an irrigation pump.

5.6 Comparison of Technological Characteristics

The Disposable Endoscopy Adapter Set has substantially equivalent device design, configuration, packaging fundamental technology, sterilization process and intended use as those featured in the predicate device Wilson Instruments (SHA) Co., LTD's Disposable Endoscope Valves Set, Disposable Endoscope Valves Set B, K200769. The differences between the proposed device and the predicate devices do not raise any questions regarding its safety and effectiveness. The differences are listed in the table below.

Item	Disposable Endoscopy	Disposable Endoscope Valves	Discussion
	Adapter Set(Proposed	Set, Disposable Endoscope	
	Device)	Valves Set B, K200769	
Indication	The Disposable Endoscopy	The Disposable Endoscope	Substantially
for Use	Adapter Set is a collection	Valves Set is a collection of	equivalent
	of several sterile units. It is	several sterile units. It is	
	intended to be fitted to	intended to be fitted to	
	multiple endoscope	multiple endoscope working	
	working channels/ports to	channels/ports to control the	
	control the flow of fluids,	flow of fluids, gases and other	
	gases and other materials.	materials. It includes an	
		air/water valve, a suction	
	- Disposable Air/Water	valve, a biopsy valve and a	
	Valve: This unit is intended	water connector.	
	to be fitted to an endoscope	- Disposable Air/Water Valves:	
	air/water channel to control	This unit is intended to be	
	the inflow of medical gases	fitted to an endoscope	
	and water, whilst	air/water channel to control the	
	preventing back-flow.	inflow of medical gases and	
	- Disposable Suction Valve:	water, whilst preventing	



Item	Disposable Endoscopy	Disposable Endoscope Valves	Discussion
Item	Adapter Set(Proposed	Set, Disposable Endoscope	Discussion
	Device)	Valves Set B, K200769	
	,	back-flow.	
	This unit is intended to be		
	fitted to an endoscope	- Disposable Suction Valves:	
	suction channel to control	This unit is intended to be	
	the operations of suction,	fitted to an endoscope suction	
	whilst preventing inflow of	channel to control the	
	air.	operations of suction, whilst	
	- Disposable Biopsy Valve:	preventing inflow of air.	
	This unit is intended to be	- Disposable Biopsy Valves:	
	fitted to an endoscope	This unit is intended to be	
	biopsy port to prevent	fitted to an endoscope biopsy	
	leakage of gases and body	port to prevent leakage of	
	fluids during an endoscopic	gases and body fluids during	
	procedure.	an endoscopic procedure.	
	- Disposable Water Jet	- Disposable Endoscope Water	
	Adapter: This unit is	Connectors: This unit is	
	intended to provide	intended to provide irrigation	
	irrigation via sterile water	via sterile water supply during	
	supply during GI	GI endoscopic procedures	
	endoscopic procedures	when used in conjunction with	
	when used in conjunction	an irrigation pump.	
	with an irrigation pump.	The Discoult Endougn	
		The Disposable Endoscope	
		Valves Set B is a collection of sterile device intended to be	
		fitted to multiple endoscope working channels/ports to	
		enable an endoscope operator	
		control the function of the	
		working channels/ports and	
		prevent retrograde flow of	
		fluids, gases, and other	
		materials. It includes a suction	
		valve, an air/water valve. This	
		is a single-use device.	
		Air/Water Valves: This device	
		is intended to be fitted to an	
		endoscope air/water channel to	
		enable the operator to control	
		inflow of medical gases and	
		water, whilst preventing	
		backflow Suction Valves:	
		vackiiow Suction valves:	



Item	Disposable Endoscopy	Disposable Endoscope Valves	Discussion
	Adapter Set(Proposed	Set, Disposable Endoscope	
	Device)	Valves Set B, K200769	
		The device is intended to be	
		fitted to an endoscope suction	
		channel to enable the operator	
		to control suction whilst	
		preventing inflow of air.	
Product	ODC	ODC	Same
Code			
Regulation	21 CFR 876.1500	21 CFR 876.1500	Same
Number			
Classification	II	II	Same
Air/Water Val	ve		
Compatible	Olympus	WS-VO-01:Olympus	Substantially
Endoscopes	140/160/180/190/240/260/2	140/160/180/190/240/260/2 90	equivalent
	90 series endoscopes;	series endoscopes	supported by
	PENTAX GI Video	WS-VP-01: PENTAX GI	bench testing.
	Endoscope 90K/90i Series,	Video Endoscope 90K/90i	
	K10/i10 Series;	Series, K10/i10 Series	
	Fujifilm® 500/600/		
	700 series endoscopes		
Sterile	EO Sterilization	EO Sterilization	Same
Material	Silicone Rubber, ABS,	WS-VO-01: SIR, ABS, TPE,	Substantially
	SUS304	Stainless steel 304	equivalent
		WS-VP-01: SIR, ABS, TPE,	supported by
		Stainless steel 304	biocompatibility
			testing.
Environment	Hospital/clinics	Hospital/clinics	Same
Use			
Suction Valve			
Compatible	Olympus	WS-VO-02: Olympus	Substantially
Endoscopes	140/160/180/190/240/260/2	140/160/180/190/240/260/2 90	equivalent
	90 series endoscopes;	series endoscopes	supported by
	PENTAX GI Video	WS-VP-02: PENTAX GI	bench testing.
	Endoscope 90K/90i Series;	Video Endoscope 90K/90i	
	K10/i10 Series;	Series; K10/i10 Series	
	Fujifilm® 500/600/		
	700 series endoscopes		
Sterile	EO Sterilization	EO Sterilization	Same
Material	ABS, Silicone Rubber,	We Vo 02, cip Ape TDE	Substantially.
wiateriai		WS-VO-02: SIR, ABS, TPE,	Substantially
	SUS304	Stainless steel 304, Stainless	equivalent
		steel 17-4PH	supported by



Item	Disposable Endoscopy	Disposable Endoscope Valves	Discussion
Teem	Adapter Set(Proposed	Set, Disposable Endoscope	Discussion
	Device)	Valves Set B, K200769	
	Device)	WS-VP-02: SIR, ABS, TPE,	biocompatibility
		Stainless steel 304	testing.
Environment	Hospital/clinics	Hospital/clinics	Same
Use	110spital/ennies	110spital/ellilles	Same
Biopsy Valve			
Compatible	Olympus	Olympus	Substantially
Endoscopes	140/160/180/190/240/260/2	140/160/180/190/240/260/290	equivalent
Linuoscopes	90 series endoscopes;	series endoscopes	supported by
	PENTAX GI Video	series endoscopes	bench testing.
	Endoscope 90K/90i Series;		benen testing.
	K10/i10 Series;		
	Fujifilm® 500/600/		
	700 series endoscopes		
Sterile	EO Sterilization	EO Sterilization	Same
Material	Silicone Rubber	Silicone Rubber	Same
Environment	Hospital/clinics	Hospital/clinics	Same
Use			
Water Jet Adaj	pter		
Compatible	Olympus	Olympus	Substantially
Endoscopes	140/160/180/190/240/260/2	140/160/180/190/240/260/290	equivalent
	90 series endoscopes;	series endoscopes	supported by
	PENTAX GI Video		bench testing.
	Endoscope 90K/90i Series;		
	K10/i10 Series;		
	Fujifilm® 500/600/		
	700 series endoscopes		
Sterile	EO Sterilization	EO Sterilization	Same
Material	Metal Type:PC,	NBR, SIR, ABS	Substantially
	SUS303,Silicone Rubber		equivalent
	D1 4' T DC C'1'		supported by
	Plastic Type: PC, Silicone		FF
	Rubber PC, Silicone		biocompatibility
	* *		
Environment	* *	Hospitals/clinics	biocompatibility

5.7 Non-clinical Performance Data

The proposed device meets the requirements of ISO 10993 "Biological Evaluation of Medical Devices – Part 1: Evaluation and Testing", ISO 11135-1 "Sterilization of Health Care products Ethylene Oxide - Part 1: Requirements for Development, Validation, and Routine Control of Sterilization processes for Medical Devices", and ISO 10993-7 "Biological evaluation of medical



devices - Part 7: ethylene oxide sterilization residuals".

The following bench tests were performed on Disposable Endoscopy Adapter Set: Appearance, Physical properties. The results of all testing were passing.

5.8 Clinical Test Data

No Clinical Study is included in this submission.

5.9 Conclusion

In accordance with the Federal Food, Drug and Cosmetic Act, 21 CFR Part 807, Based on the information provided in this premarket notification, Yangzhou Fartley Medical Instrument Technology Co., Ltd. has demonstrated that proposed device Disposable Endoscopy Adapter Set is substantially equivalent to Wilson Instruments (SHA) Co., LTD's currently marketed Disposable Endoscope Valves Set, Disposable Endoscope Valves Set B, K200769.