

August 19, 2022

Stryker Marlene Fraga Sr. Staff, Regulatory Affairs Specialist, Software Interoperability 5900 Optical Court San Jose, California 95138

Re: K222130

Trade/Device Name: 1688 4K Camera System with Advanced Imaging Modality

Regulation Number: 21 CFR 876.1500

Regulation Name: Endoscope And Accessories

Regulatory Class: Class II Product Code: GCJ, GWG Dated: July 13, 2022 Received: July 18, 2022

#### Dear Marlene Fraga:

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 <a href="https://www.accessdata.fda.gov/scripts/cdrh/efdocs/efpmn/pmn.cfm">https://www.accessdata.fda.gov/scripts/cdrh/efdocs/efpmn/pmn.cfm</a> 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. 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 <a href="https://www.fda.gov/combination-products/guidance-regulatory-information/postmarketing-safety-reporting-combination-products">https://www.fda.gov/combination-products/guidance-regulatory-information/postmarketing-safety-reporting-combination-products</a>); 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 <a href="https://www.fda.gov/medical-devices/medical-device-safety/medical-device-reporting-mdr-how-report-medical-device-problems">https://www.fda.gov/medical-device-problems</a>.

For comprehensive regulatory information about medical devices and radiation-emitting products, including information about labeling regulations, please see Device Advice (<a href="https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance">https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance</a>) and CDRH Learn (<a href="https://www.fda.gov/training-and-continuing-education/cdrh-learn">https://www.fda.gov/training-and-continuing-education/cdrh-learn</a>). 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 (<a href="https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/contact-us-division-industry-and-consumer-education-dice">https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/contact-us-division-industry-and-consumer-education-dice">https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/contact-us-division-industry-and-consumer-education-dice</a>) for more information or contact DICE by email (DICE@fda.hhs.gov) or phone (1-800-638-2041 or 301-796-7100).

Sincerely,

for Long Chen, Ph.D.
Assistant Director
DHT4A: Division of General Surgery Devices
OHT4: Office of Surgical
and Infection Control Devices
Office of Product Evaluation and Quality
Center for Devices and Radiological Health

Enclosure

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# DEPARTMENT OF HEALTH AND HUMAN SERVICES Food and Drug Administration

#### **Indications for Use**

Form Approved: OMB No. 0910-0120 Expiration Date: 06/30/2020

Expiration Date: 06/30/2020 See PRA Statement below.

510(k) Number (if known)	
K222130	
Device Name 1688 4K Camera System with Advanced Imaging Modality (AIM)	
Indications for Use (Describe)	

The 1688 Video Camera is indicated for use in general laparoscopy, nasopharyngoscopy, ear endoscopy, sinuscopy, neurosurgery and plastic surgery whenever a laparoscope/ endoscope/ arthroscope/ sinuscope is indicated for use. The 1688 Video Camera is indicated for adults and pediatric patients.

A few examples of the more common endoscope surgeries are Laparoscopic cholecystectomy, Laparoscopic hernia repair, Laparoscopic appendectomy, Laparoscopic pelvic lymph node detection, Laparoscopically assisted hysterectomy, Laparoscopic and thorascopic anterior spinal fusion, Anterior cruciate ligament reconstruction, Knee arthroscopy, Small joint arthroscopy, Decompression fixation, Wedge resection, Lung biopsy, Pleural biopsy, Dorsal sympathectomy, Pleurodesis, Internal mammary artery dissection for coronary artery bypass, Coronary artery bypass grafting where endoscopic visualization is indicated and Examination of the evacuated cardiac chamber during performance of valve replacement.

The users of the 1688 Video Camera are general and pediatric surgeons, gynecologists, cardiac surgeons, thoracic surgeons, plastic surgeons, orthopedic surgeons, ENT surgeons, neurosurgeons and urologists.

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)		
Type of Use (Select one or both, as applicable)	Prescription Use (Part 21 CFR 801 Subpart [	O) 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.

#### \*DO NOT SEND YOUR COMPLETED FORM TO THE PRA STAFF EMAIL ADDRESS BELOW.\*

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# **stryker**

## 510(k) Summary

#### **Submitter:**

Applicant:	Stryker Endoscopy
	5900 Optical Court
	San Jose, CA 95138
Contact Person:	Marlene Fraga
	Sr. Staff Regulatory Affairs Specialist, Software Interoperability
	Email: marlene.fraga@stryker.com
Date Prepared:	July 13, 2022

#### **Subject Device:**

Name of Device:	1688 4K Camera System with Advanced Imaging Modality
Common or	3-chip Video Camera
Usual Name	
Classification	Laparoscope, General and Plastic Surgery (21 C.F.R. §876.1500)
Name:	Endoscope, Neurological (21 C.F.R. §882.1480)
Regulatory Class:	II
Product Code:	GCJ
	GWG
510(k) Review	General & Plastic Surgery
Panel:	Neurology

#### **Predicate Device(s):**

1688 4K Camera System with Advanced Imaging Modality	K211202
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#### **Reference Device(s):**

1688 4K Camera System with Advanced Imaging Modality	K220895, K212511
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#### **Device Description:**

The 1688 4K Camera System with Advanced Imaging Modality is an endoscopic camera system that produces live video in the surgical field during surgical endoscopic procedures. The system is sensitive in the visible and infrared spectrum. The optical image is transferred from the surgical site to the camera head by a variety of rigid and flexible endoscopes, which are attached to the camera head. The 1688 4K Camera System consists of three main components: (1) a camera control unit (CCU); (2) a camera head with an integral cable that connects to the CCU; and (3) a coupler for attaching an endoscope to the camera head.



# **Indications for Use:**

This Submission  System (K211202)  Intended Use  Endoscopic white light and near- infrared illumination and imaging during endoscopic procedures.  System (K211202)  Same as subject device device  Same as subject device	688 4K Camera System (K212511) me as subject
This Submission System (K211202) Same as subject device Use Same as subject device Same as subject device device Same as subject device device	(K212511) ume as subject vice
Intended UseEndoscopic white light and near- infrared illumination and imaging during endoscopic procedures.Same as subject deviceSame as subject deviceSame as subject device	me as subject vice
Use infrared illumination and imaging device device during endoscopic procedures.	vice
during endoscopic procedures.	
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Indications The 1600 Video Communicated Comm	me as subject
Indications   The 1688 Video Camera is indicated   Same as subject device   Same as subject   Same	
for Use for use in general laparoscopy, device device	vice
nasopharyngoscopy, ear endoscopy,	
sinuscopy, neurosurgery and plastic	
surgery whenever a laparoscope/	
endoscope/ arthroscope/ sinuscope is	
indicated for use. The 1688 Video	
Camera is indicated for adults and	
pediatric patients.	
A C	
A few examples of the more common	
endoscope surgeries are Laparoscopic cholecystectomy, Laparoscopic hernia	
repair, Laparoscopic appendectomy,	
Laparoscopic pelvic lymph node	
detection, Laparoscopically assisted	
hysterectomy, Laparoscopic and	
thorascopic anterior spinal fusion,	
Anterior cruciate ligament	
reconstruction, Knee arthroscopy,	
Small joint arthroscopy,	
Decompression fixation, Wedge	
resection, Lung biopsy, Pleural	
biopsy, Dorsal sympathectomy,	
Pleurodesis, Internal mammary artery	
dissection for coronary artery bypass,	
Coronary artery bypass grafting where	
endoscopic visualization is indicated	
and Examination of the evacuated	
cardiac chamber during performance	
of valve replacement.	
The users of the 1688 Video Camera	
are general and pediatric surgeons,	
gynecologists, cardiac surgeons,	
thoracic surgeons, plastic surgeons,	
orthopedic surgeons, ENT surgeons,	
neurosurgeons and urologists.	



## **Comparison of Technological Characteristics with the Predicate Device:**

K222130

	Subject Device	Predicate Device	Reference Devices		
Item	1688 4K Camera System (This Submission)	1688 4K Camera System (K211202)	1688 4K Camera System (K220895)	1688 4K Camera System (K212511)	
Manufacturer	Stryker	Same as subject device	Same as subject device	Same as subject device	
Imaging Modes	White Light Near-infrared fluorescence Near-infrared transillumination	Same as subject device	Same as subject device	Same as subject device	
Camera System Components	Camera Control Unit	Camera Control Unit	Same as subject device	Same as subject device	
	Camera Head(s) – Standard, Integrated, Inline, Pendulum, Autoclave	Camera Head(s) – Standard, Integrated, Inline, Pendulum			
	Coupler(s) – AIM 4K, AIM 4K Autoclave	Coupler(s) – AIM 4K			
Principles of Operations  Safety Standards	Via an optical light guide, endoscope and coupler, light is projected from a light source and either reflected or absorbed and fluoresced onto one or more complementary metal oxide semiconductor image sensors which acquire a continuous stream of image data. The image data is processed to provide a video stream that is then sent to a display for viewing.  IEC 60601-1 IEC 60601-1-6	Same as subject device  Same as subject device	Same as subject device  Same as subject device	Same as subject device  Same as subject device	
Standards	IEC 60601-1-6 IEC 60601-2-18 IEC 60601-1-2				
Modes of Operation	Alternate Frame processing Simultaneous Frame processing	Same as subject device	Same as subject device	Same as subject device	
Image Sensor	CMOS image sensor	Same as subject device	Same as subject device	Same as subject device	
Image Processing/ Video Output	Digital	Same as subject device	Same as subject device	Same as subject device	
Resolution	4K (up to 3840 x 2160)	Same as subject device	Same as subject device	Same as subject device	
Frame Rate	60 frames per second	Same as subject device	Same as subject device	Same as subject device	
Camera Head Cable Proximal PCBA Capacitor	Soft-termination capacitor	Conventional termination capacitor	Conventional termination capacitor	Conventional termination capacitor	



#### **Performance Data:**

The following performance data were provided in support of the substantial equivalence determination:

Test	Method		Result
Performance Testing	•	Environmental	Pass
	•	RF Interference	
Electrical Safety and	•	Electrical Safety (ANSI AAMI ES 60601-1, IEC 60601-1-6,	Pass
EMC		IEC 60601-2-18)	
	•	Electromagnetic Compatibility (IEC 60601-1-2)	

### **Conclusions:**

The 1688 4K Camera System is substantially equivalent in design, intended use, principles of operation, technological characteristics and safety features to the predicate devices. In summary, the 1688 4K Camera System is substantially equivalent with respect to safety and effectiveness to the legally marketed predicate device.