

December 28, 2021

K1 Medical LLC % Joseph Azary Consultant Joseph Azary 543 Long Hill Avenue Shelton, Connecticut 06484

Re: K213209

Trade/Device Name: EZ-TRAX Zimmer G7 Acetabular System / Taperloc Complete Hip System Containment Device
Regulation Number: 21 CFR 880.6850
Regulation Name: Sterilization wrap
Regulatory Class: Class II
Product Code: KCT
Dated: September 27, 2021
Received: September 29, 2021

Dear Joseph Azary:

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 <u>https://www.fda.gov/medical-devices/medical-device-safety/medical-device-reporting-mdr-how-report-medical-device-problems</u>.

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

Sincerely,

For Clarence Murray III, Ph.D. Assistant Director DHT4B: Division of Infection Control and Plastic Surgery Devices OHT4: Office of Surgical and Infection Control Devices Office of Product Evaluation and Quality Center for Devices and Radiological Health

Enclosure

Indications for Use

510(k) Number (if known)

K213209

Device Name

EZ-TRAX[™] Zimmer G7 Acetabular System / Taperloc Complete Hip System Containment Device

The **EZ-TRAX**[™] Zimmer G7 Acetabular System / Taperloc Complete Hip System Containment Device is intended for use as an accessory in healthcare facilities to organize, enclose, reprocess, transport, and store Zimmer G7 Acetabular System / Taperloc Complete Hip System devices between surgical uses. The **EZ-TRAX**[™] Zimmer G7 Acetabular System / Taperloc Complete Hip System Containment Device is not intended on its own to maintain sterility; it is intended to be used in conjunction with a legally marketed, validated, FDA-cleared sterile barrier system.

Cycle	Temperature	Exposure Time	Drying Time
Dynamic Air Removal	270 F/132 C	4 minutes	10 minutes

Validations included the worst case load configurations of the EZ-TRAX™ Zimmer G7 Acetabular System / Taperloc Complete Hip System Containment Devices.

- Contents in the validated configuration include reusable surgical instruments (impactors, trials, acetabular reamers, broaches, etc)
- No lumened devices were validated within the tray system as part of the product load. The EZ-TRAX™ Zimmer G7 Acetabular System / Taperloc Complete Hip System Containment Device does not have any lumen claims.
- Healthcare facilities should not exceed 25 pounds (EZ-TRAX[™] Containment Device + Sterile Barrier System + Zimmer G7 Acetabular System / Taperloc Complete Hip System Devices).

The EZ- TRAX[™] Zimmer G7 Acetabular System / Taperloc Complete Hip System Containment Device is offered in the following size:

Brand Name	Model	Dimensions	Dry Weight
EZ-TRAX™	BASE.ASSY.AL.24.12.2.4	L) 22.97" (W) 11.18" (H) 2.44"	4.25 lbs

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) SUMMARY K1 Medical LLC's EZ-TRAX Containment Device K213209

Submitter

K1 Medical LLC 56 Newton Road

Woodbridge, CT 06525

Contact Person: Joseph Azary

Phone: 203-242-6670

Date Prepared: December 23, 2021

Name of Device: EZ-TRAXTM Zimmer G7 Acetabular System / Taperloc Complete Hip System Containment Device

Common or Usual Name: Sterilization Cassette

Classification Name: Sterilization Wrap

Regulatory Class: Class II, 21 CFR 880.6850

Product Code: KCT

II. PREDICATE DEVICE:

EZ-TRAX™Persona Knee Containment Device K211007

III.DEVICE DESCRIPTION

The EZ-TRAX[™]Zimmer G7 Acetabular System / Taperloc Complete Hip System Containment Device is intended for use as an accessory in healthcare facilities to organize, enclose, reprocess, transport and store G7 Acetabular System / Taperloc Complete Hip System Devices between surgical uses. The EZ-TRAX[™]Zimmer G7 Acetabular System / Taperloc Complete Hip System Containment Device is not intended on its own to maintain sterility; it is intended to be used in conjunction with a legally marketed, validated, FDA cleared sterile barrier system. The subject device protects the inteterior components during transportation, sterilization, and storage.

The EZ-TRAX[™]Zimmer G7 Acetabular System / Taperloc Complete Hip System Containment Device is composed of intrinsically stable metals and thermoplastic polymers. The trays and lids are composed of anodized aluminum with stainless steel handles. The dividers are composed of Aluminum and the posts are composed of medical grade thermoplastic polymers.

The lids and bottom of trays are fully perforated with an evenly distributed hole pattern. The sides of the trays are partially perforated. The trays are used with locking lids.

The trays were designed to be used for sterilization via steam sterilization and used in standard autoclaves found in hospitals and healthcare facilities. The trays were designed in such a way to withstand repeated steam sterilization cycles.

IV. INTENDED USE / INDICATIONS FOR USE

The **EZ-TRAX™** Zimmer G7 Acetabular System / Taperloc Complete Hip System Containment Device is intended for use as an accessory in healthcare facilities to organize, enclose, reprocess, transport, and store Zimmer G7 Acetabular System / Taperloc Complete Hip System devices between surgical uses. The **EZ-TRAX™** Zimmer G7 Acetabular System / Taperloc Complete Hip System Containment Device is not intended on its own to maintain sterility; it is intended to be used in conjunction with a legally marketed, validated, FDA-cleared sterile barrier system.

Cycle	Temperature	Exposure Time	Drying Time
Dynamic Air Removal	270 F/132 C	4 minutes	10 minutes

Validations included the worst case load configurations of the EZ-TRAXTM Zimmer G7 Acetabular System / Taperloc Complete Hip System Devices.

- Contents in the validation configuration include: reusable surgical instruments (impactors, trials, acetabular reamers, broaches, etc).
- No lumended devices were validated within the tray system as part of the product load.

The EZ-TRAX[™] Zimmer G7 Acetabular System / Taperloc Complete Hip System Containment Device does not have any lumen claims.

Healthcare facilities should not exceed 25 pounds (EZ-TRAX[™] Containment Device + sterile barrier system + Zimmer G7 Acetabular System / Taperloc Complete Hip System containment devices).

The EZ-TRAX[™] Zimmer G7 Acetabular System / Taperloc Complete Hip System Containment Device is offered in the following sizes:

Brand Name	Model	Dimensions	Dry Weight
EZ-TRAX™	Base.Assy.AL.24.12.2.4	(L) 22.97" (W) 11.18" (H) 2.44"	4.25 lbs

VII. TECHNOLOGICAL CHARACTERISTICS COMPARISON TABLE

PROVIDED BELOW IS A TECHNOLOGICAL COMPARISON OF THE SUBJECT DEVICE TO THE PREDICATE DEVICE.

TRADENAME	K1 Medical LLC EZ-TRAX [™] Zimmer G7 Acetabular System / Taperloc Complete Hip System Containment Device K213209	K1 Medical LLC EZ-TRAX [™] Persona Knee Containment Device K211007	Comparison
Fundamental Scientific Technology	Sterilization Cassette	Sterilization Cassette	Identical
Product Code	KCT	КСТ	Identical
Material Composition	Thermoplastic polymers, Aluminum, and stainless steel	Thermoplastic polymers, Aluminum, and stainless steel	Identical
Design	Base, lid with locking latch, and individual inserts	Base, lid with locking latch and individual inserts	Identical
Dimensions	Worst Case 24 x 12 x 2.4"	Worst case 24 x 12 x 2.4"	Identical
Configuration	Perforated bases, lids, and inserts	Perforated bases, lids and inserts	Identical
Air Permeance	Yes	Yes	Identical
Percent Perforation	Evenly distributed hole pattern	Evenly distributed hole pattern	Identical
Sterilization Method	Dynamic Air Removal	Dynamic Air Removal	Identical
Sterilization Parameters	Dynamic Air Removal Temperature: 270F Exposure Time: 4 minutes Drying Time: 10 minutes	Dynamic Air Removal Temperature: 270F Exposure Time: 4 minutes Drying Time: 10 minutes	Identical
Reusable	Yes	Yes	Identical
Microbial Barrier Properties	Used with FDA cleared sterile barrier system	Used with FDA cleared sterile barrier system	Identical
Material Compatibility	Materials compatible with sterilization method	Materials compatible with sterilization method	Identical
Biocompatibility	Biocompatible	Biocompatible	Identical

Intended Use/	The EZ-TRAX™ Zimmer G7	The EZ-TRAX™ Persona Knee	Same
Indications for	Acetabular System / Taperloc	Containment Device is intended for	
Use	Complete Hip System Containment	use as an accessory in healthcare	
	Device is intended for use as an	facilities to organize, enclose,	
	accessory in healthcare facilities to	reprocess, transport, and store	
	organize, enclose, reprocess,	Persona Knee Devices between	
	transport, and store G7 Acetabular	surgical uses. The EZ- TRAX™	
	System / Taperloc Complete Hip	Persona Knee Containment Device is	
	System devices between surgical	not intended on its own to maintain	
	uses. The EZ-TRAX™ Zimmer G7	sterility; it is intended to be used in	
	Acetabular System / Taperloc	conjunction with a legally marketed,	
	Complete Hip System Containment	validated, FDA-cleared sterile barrier	
	Device is not intended on its own to	system.	
	maintain sterility; it is intended to be		
	used in conjunction with a legally	Cycle: Dynamic Air Removal	
	marketed, validated, FDA-cleared	Temperature: 270 F/132 C	
	sterile barrier system.	Exposure Time: 4 minutes	
		Drying Time: 10 minutes	
	Cycle: Dynamic Air Removal		
	Temperature: 270 F/132 C	Sterilization validations included the	
	Exposure Time: 4 minutes	worst case load configurations of	
	Drying Time: 10 minutes	Persona Knee Containment Device	
		utilized the Persona Knee Devices	
	Validations include the worst case	required to perform an arthroplasty	
	load configurations of the EZ-	procedure.	
	TRAX™ Zimmer G7 Acetabular		
	System / Taperloc Complete Hip	- Contents in the validated	
	System Devices.	configuration include: reusable	
		surgical instruments (impactors, trials,	
		MICS, sagital saws,.arrays, etc.)	
	- Contents in the validated		
	configuration include: reusable	- No lumened devices were validated	
	surgical instruments (impactors,	within the tray system as part of the	
	trials, acetabular reamers,	product load. The EZ- TRAX™	
	broaches, etc.).	Persona Knee Containment Device	
	No lumonod dovisoo woro	does not have any lumen claims.	
	- No lumened devices were		
	validated within the tray system as	-Healthcare facilities should not	
	part of the product load. The EZ- TRAX™ Zimmer G7 Acetabular	exceed 25 pounds (EZ-TRAX™	
		Containment Device + sterile barrier	
	System / Taperloc Complete Hip System Containment Device does	system + Persona Knee	
	not have any lumen claims.	Containment Devices).	
	-Healthcare facilities should not		
	exceed 25 pounds (EZ-TRAX TM		
	Containment Device + sterile barrier		
		I	

system + Zimmer G7 Acetabular	
System / Taperloc Complete Hip	
System Devices).	
System Devices).	
The EZ-TRAX [™] Zimmer G7	
Acetabular System / Taperloc	
Complete Hip System Containment	
Device is offered in the following size:	

EZ-TRAX™ BASE.ASSY.AL.24.12.2.4 (L) 22.97" (W) 11.18" (H) 2.44" Dry weight 4.25 lbs.	The EZ-TRAX [™] Persona Knee Containment Device is offered in the following sizes: Brand Name Model Dimensions EZ-TRAX [™] BASE.ASSY.AL.24. 12.2.4 (L) 22.97" (W) 11.18" (H) 2.44" (Dry
	Weight 4.25 lbs). EZ-TRAX™ BASE.ASSY.AL.21. 12.4 (L) 19.97" (W) 11.18" (H) 4.187"

Summary of Non-Clinical Testing

Provided below is the summary of the non-clinical testing that was performed per specification of the standard and test methodology listed below. The results of the performance testing demonstrated the subject device met the acceptance criteria of the standard and the test methodology.

VIII. NON-CLINICAL PERFORMANCE TESTING TABLE

Test Methodology	Purpose	Acceptance Criteria	Results
Material Compatibility	To verify the device did not degrade or lose functionality	Material compatibility pre-vacuum 132C for 4 minutes.	Passed
AAMI TIR12:2010 AAMI ST81:2004/R2016 ISO 17665-1:2006/R2013	after 25 reprocessing cycles at worst case conditions.	Mechanical Washing and Steam Sterilization	
		The testing subjected the device to repetitive cleaning and sterilization processing cycles at parameters that represented worst case conditions.	
		Chemical indicators were utilized to demonstrate steam penetration. The study found no degradation or lack of functionality after 25 cycles.	
Mechanical Cleaning Validation – Hemoglobin** AAMI TIR30:2011 ASTM F32018-18 ASTM F3293-18	To verify the manufacturer's cleaning instructions are effective for removing gross amounts of soil to a hemoglobin level less than 2.2 ug/cm2 per device.	The mechanical cleaning validation of the EZ-TRAX TM containment system concluded that the manufacturer's cleaning instructions are efficacious for removing gross amounts of soil from the EZ-TRAX TM containment system to a hemoglobin level less than 2.2 ug/cm2 per device.	Passed
Mechanical Cleaning Validation – Protein Analysis** AAMI TIR30:2011 ASTM F32018-18 ASTM F3293-18	To verify the manufacturer's cleaning instructions are effective for removing gross amounts of soil to a protein level of less than 6.4 ug/cm2 per device.	The mechanical cleaning validation of the EZ-TRAX TM containment system protein concluded that the manufacturer's cleaning instructions are efficacious for removing gross amounts of soil from the EZ-TRAX TM containment system to a protein level less than 6.4 ug/cm2 per device.	Passed
MEM Elution Cytotoxicity ISO 10993-5:2009 / R2014	To verify that the device meets requirements of ISO 10993-5 and is not considered cytotoxic.	The cytotoxicity testing was conducted per ISO 10993-5:2009 /R2014 and concluded that test articles met the requirements of the test and are NOT considered to have a cytotoxic potential.	Passed
Sterilization Validation AAMI ST77:2013 ISO 14937:2009 AAMI ST8:2013	To verify that the device could achieve a sterility assurance level of 10 ⁻⁶ after processing in pre-vacuum steam sterilization cycle of 132C (270F) for 4 minutes.	The sterilization validation of the EZ- TRAX TM Containment System included pre-vacuum steam 132C for 4 minutes. The conclusion is the device could achieve a Sterility Assurance Level (SAL) of 10 ⁻⁶ after processing in the following pre- vacuum steam sterilization cycle 132C (270F) for 4 minutes.	Passed
Thermal Profile Study AAMI ST77:2013	To verify that adequate sterilant penetration can be achieved when processed in pre-vacuum steam sterilization cycle of 132C (270F) for 4 minutes.	The thermal profile study of the EZ- TRAX [™] Containment Device included pre-vacuum steam 132C for 4 minutes. The study demonstrated that adequate sterilant penetration can be achieved. The EZ- TRAX [™] Containment Device can reach and maintain a steady state thermal conditions throughout the exposure phase when processed in the following pre- vacuum steam sterilization cycle 132C (270F) for 4 minutes.	Passed
Drying Time Test AAMI ST77:2013 ISO 17665-1:2006/R2013	To verify that the device is properly dried following processing in pre-vacuum steam sterilization cycle of 132C for 4.0 minutes and dry time of 10 minutes.	The results demonstrate EZ-TRAX TM meets or exceeds the minimum acceptance criteria for dry time. The EZ-TRAX TM is considered properly dried following processing in the steam prevacuum sterilization cycle of 132C / 270F, Exposure Time 4.0 minutes and Dry Time 10.0 minutes minimum.	Passed
Handle 100 lbs force test AAMI ST77:2013	To verify that tray handles did not break or show evidence of distortion, cracking or other	None of the tray handles broke loose, showed evidence of permanent distortion, cracking or other evidence of failure when tested with force of 50 lbs.	Passed

	failure following testing with force of 50 lbs.		
Sterilization Validation of the EZ-TRAX [™] Zimmer G7 Acetabular & Taperloc Containment Device (unorganized) steam pre- vacuum 132C (270F) for 4 minutes.	To verify that a Sterility Assurance Level of 10 ⁻⁶ can be achieved after processing in pre-vacuum steam.	The testing verified that a Sterility Assurance Level of 10 ⁻⁶ can be achieved after processing EZ-TRAX TM G7 Acetabular System / Taperloc Complete Hip System Containment Device in a steam pre-vacuum cycle at 132C (270F) and 4.0 minutes.	Passed
AAMI ST8:2013 TIR12:2010 ST77:2013/R2018 ST79:2017 ISO 14937:2009/R2013 ISO 17665-1:2006/R2013			

IX. CONCLUSIONS

The conclusions drawn from the nonclinical tests demonstrate that the EZ-TRAX[™]Zimmer G7 Acetabular System / Taperloc Complete Hip System Containment Device is as safe, as effective and performs as well as or better than the legally marketed device EZ-TRAX[™]Persona Knee Containment Device (K211007).