June 27, 2023



L&K BIOMED Co., Ltd. Katherine Kim RA #101, 201, 202 16-25 Dongbaekjungang-ro 16 beon-gil Giheung-gu Yongin-si, Gyeonggi-do, 17015 South Korea

Re: K231636

Trade/Device Name: LnK Spinal Fixation System /OpenLoc-L Spinal Fixation System, AccelFix Spinal Fixation System
Regulation Number: 21 CFR 888.3070
Regulation Name: Thoracolumbosacral Pedicle Screw System
Regulatory Class: Class II
Product Code: NKB
Dated: June 2, 2023
Received: June 5, 2023

Dear Katherine Kim:

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. 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,

Colin O'neill -S

Colin O'Neill, M.B.E. Assistant Director DHT6B: Division of Spinal Devices OHT6: Office of Orthopedic Devices Office of Product Evaluation and Quality Center for Devices and Radiological Health

Enclosure

Indications for Use

510(k) Number *(if known)* K231636

Device Name

LnK Spinal Fixation System /OpenLoc-L Spinal Fixation System

Indications for Use (Describe)

The LnK Spinal Fixation System /OpenLoc-L Spinal Fixation System is a non-cervical spinal fixation device intended for use as a posterior pedicle screw fixation system (T1-S2/ilium), or as an anterolateral fixation system (T8-L5). All components in the system are limited to skeletally mature patients. This device is indicated as an adjunct to fusion for all of the following indications regardless of the intended use: degenerative disc disease (defined as discogenic back pain with degeneration of the disc confirmed by history and radiographic studies); spondylolisthesis; trauma (i.e., fracture or dislocation); deformities or curvatures (i.e., scoliosis, kyphosis, and/or lordosis); tumor; stenosis, and failed previous fusion (pseudoarthrosis).

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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Indications for Use

510(k) Number *(if known)* K231636

Device Name AccelFix Spinal Fixation System

Indications for Use (Describe)

The AccelFix Spinal Fixation System is a non-cervical spinal fixation device intended for use as a posterior pedicle screw fixation system (T1-S2/ilium), or as an anterolateral fixation system (T8-L5). All components in the system are limited to skeletally mature patients. These devices are indicated as an adjunct to fusion for all of the following indications regardless of the intended use: degenerative disc disease (defined as discogenic back pain with degeneration of the disc confirmed by history and radiographic studies); spondylolisthesis; trauma (i.e., fracture or dislocation); deformities or curvatures (i.e., scoliosis, kyphosis, and/or lordosis); tumor; stenosis, and failed previous fusion (pseudoarthrosis).

 Prescription Use (Part 21 CFR 801 Subpart D)
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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Paperwork Reduction Act (PRA) Staff

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

PRAStaff@fda.hhs.gov

Form Approved: OMB No. 0910-0120

Expiration Date: 06/30/2023

See PRA Statement below.



510(k) SUMMARY

The following 510(k) summary is being submitted as required by 21 CFR 807.92(a):

1. SUBMITTER

Submitter's Name:	L&K BIOMED Co., Ltd.	
Submitter's Address:	#101, 201, 202 16-25, Dongbaekjungang-ro 16 beon-gil	
	Giheung-gu, Yongin-si, Gyeonggi-do, 17015, Korea	
Submitter's Telephone:	+82-2-6717-1983	
Contact Person:	Katherine Kim	
	khkim@lnkbiomed.com / ra@lnkbiomed.com	
Prepared Date	June 2, 2023	

2. DEVICE IDENTIFICATION

Trade or Proprietary Name	LnK Spinal Fixation System /OpenLoc-L Spinal Fixation System	
Common or Usual Name	Spinal interlaminal fixation orthosis Spinal intervertebral body fixation orthosis Thoracolumbosacral pedicle screw system	
Regulation class / Number	Class II, 21 CFR 888.3070	
Regulation Name	Thoracolumbosacral pedicle screw system	
Product Code	NKB, KWP, KWQ	
Classification Panel	Spinal Devices (DHT6B)	

Trade or Proprietary Name	AccelFix Spinal Fixation System	
Common or Usual Name	Spinal interlaminal fixation orthosis Spinal intervertebral body fixation orthosis	
	Thoracolumbosacral pedicle screw system	
Regulation class / Number	Class II, 21 CFR 888.3070	
Regulation Name	Thoracolumbosacral pedicle screw system	
Product Code	NKB, KWP, KWQ	
Classification Panel	Spinal Devices (DHT6B)	

3. PREDICATE OR LEGALLY MARKETED DEVICES WHICH ARE SUBSTANTIALLY EQUIVALENT.

The additional components of the LnK Spinal Fixation System, OpenLoc-L Spinal Fixation System 18and AccelFix Spinal Fixation System are considered substantially equivalent to the predicate devices. The systems have same design, materials, scientific technology, and indications for use.

LnK Spinal Fixation System /OpenLoc-L Spinal Fixation System

Primary Predicate Device: LnK Spinal Fixation System (K230245) Additional Predicate Devices: LnK Spinal Fixation System/OpenLoc-L Spinal Fixation System (K120270, K143363, K171813, K183168, K 200790, K223565) Olympic Posterior Spinal Fixation System(K181139)

AccelFix Spinal Fixation System

Primary Predicate Device: AccelFix Spinal Fixation System (K182544) Additional Predicate Devices: AccelFix Spinal Fixation System (K200794, K223565, K230245) Olympic Posterior Spinal Fixation System(K181139)



4. MATERIALS

MATERIALS					
LnK Spinal Fixation System/ OpenLoc-L	Ti-6Al-4V ELI titanium alloy (ASTM F136) and				
Spinal Fixation System	Cobalt-28Chromium-6Molybdenum-4Vanadium				
AccelFix Spinal Fixation System	ELI (ASTM F1537)				

The additional S-Rod is manufactured from Ti-6Al-4V ELI titanium alloy (ASTM F136) and Cobalt-28Chromium-6Molybdenum-4Vanadium ELI (ASTM F1537). This this is the same material used in the predicate devices.

5. DESCRIPTION OF THE DEVICE

LNK SPINAL FIXATION SYSTEM /OPENLOC-L SPINAL FIXATION SYSTEM

The LNK SPINAL FIXATION SYSTEM, OPENLOC-L SPINAL FIXATION SYSTEM are available in various sizes. This system is comprised of screws, set screws, rods, crosslinks, connectors and hooks. The screws are available from 4.0 mm to 10.5 mm diameters with lengths ranging from 20 mm to 150 mm. The rods are available from 5.0 mm, 5.5 mm, 6.0 mm and 6.35 mm diameter with lengths ranging from 40 mm to 600 mm. Both straight rods and curved rods have four types of design that consist of standard type, hex type, stopper type and double stopper.

ACCELFIX SPINAL FIXATION SYSTEM

The AccelFix Spinal Fixation System consists of screws, rods, crosslinks, set screws, cross-link connectors, and hooks. The screws are available from 5.0 mm, 5.5 mm, 6.0 mm, 7.0, 7.5, 8.0, 8.5, 9.0 and 9.5mm diameters with working lengths ranging from 20 mm to 150 mm. The rods are available from 5.5 mm, 6.0mm and 6.35mm diameter with lengths ranging from 40 mm to 600 mm. Both straight rods and curved rods have four types of design that consist of standard type, hex type, stopper type and double stopper.

6. INDICATION FOR USE

LNK SPINAL FIXATION SYSTEM /OPENLOC-L SPINAL FIXATION SYSTEM

The LnK Spinal Fixation System /OpenLoc-L Spinal Fixation System is a non-cervical spinal fixation device intended for use as a posterior pedicle screw fixation system (T1-S2/ilium), or as an anterolateral fixation system (T8-L5). All components in the system are limited to skeletally mature patients. This device is indicated as an adjunct to fusion for all of the following indications regardless of the intended use: degenerative disc disease (defined as discogenic back pain with degeneration of the disc confirmed by history and radiographic studies); spondylolisthesis; trauma (i.e., fracture or dislocation); deformities or curvatures (i.e., scoliosis, kyphosis, and/or lordosis); tumor; stenosis, and failed previous fusion (pseudoarthrosis).

ACCELFIX SPINAL FIXATION SYSTEM

The AccelFix Spinal Fixation System is a non-cervical spinal fixation device intended for use as a posterior pedicle screw fixation system (T1-S2/ilium), or as an anterolateral fixation system (T8-L5). All components in the system are limited to skeletally mature patients. These devices are indicated as an adjunct to fusion for all of the following indications regardless of the intended use: degenerative disc disease (defined as discogenic back pain with degeneration of the disc confirmed by history and radiographic studies); spondylolisthesis; trauma (i.e., fracture or dislocation); deformities or curvatures (i.e., scoliosis, kyphosis, and/or lordosis); tumor; stenosis, and failed previous fusion (pseudoarthrosis).



7. **PERFORMANCE DATA**

The additional components to be added through this submission do not require additional mechanical testing. None of the additional components is the worst case of the LnK Spinal Fixation System, OpenLoc-L Spinal Fixation System and AccelFix Spinal Fixation System. Therefore, we substitute mechanical test data of additional components of LnK Spinal Fixation System, OpenLoc-L Spinal Fixation System and AccelFix Spinal Fixation System with the predicate device (LnK Spinal Fixation System, OpenLoc-L Spinal Fixation System -K120270, K143363, K171813, K183168, K200790, K223565, K230245 / AccelFix Spinal Fixation System: K182544, K200794, K223565, K230245).

8. SUMMARY OF TECHNICAL CHARACTERISTICS WITH THE PREDICATE DEVICE

The subject and predicate devices have nearly identical technological characteristics and the minor differences do not raise any new issues of safety and effectiveness. Specifically, the following characteristics are identical between the subject and predicates:

- Instruction for use
- Material
- Design with components
- Dimension
- Sterilization Method

The following technological similarities and differences exist between the subject and predicate devices:

LNK SPINAL FIXATION SYSTEM, OPENLOC-L SPINAL FIXATION SYSTEM

INDICATION FOR USE		
Devices		Similarities
LnK Spinal Fixation System/OpenLoc-L Spinal Fixation System	Subject	
LnK Spinal Fixation System(K230245)	Primary Predicate	
LnK Spinal Fixation System/OpenLoc-L		Instruction for use including indication is
Spinal Fixation System (K120270, K143363,	A 111.1 1	similar.
K171813, K183168, K 200790, K223565)	Additional Predicate	
Olympic Posterior Spinal Fixation		
System(K181139)		

Materials		
Devices		Similarities
LnK Spinal Fixation System/OpenLoc-L Spinal Fixation System	Subject	
LnK Spinal Fixation System(K230245)	Primary Predicate	Similar materials • Ti-6A1-4V ELI titanium alloy (ASTM
LnK Spinal Fixation System/OpenLoc-L Spinal Fixation System (K120270, K143363, K171813, K183168, K 200790, K223565)	Additional Predicate	 F136) Cobalt-28Chromium-6Molybdenum- 4Vanadium ELI (ASTM F1537)
Olympic Posterior Spinal Fixation System(K181139)		



Design with Components			
Devices		Similarities	Dissimilarities
LnK Spinal Fixation System/OpenLoc-L Spinal Fixation System	Subject	Screws, Set Screws, Rods, Crosslinks, Connectors, Hooks Rods include Z-rod	
LnK Spinal Fixation System(K230245)	Primary Predicate	Screws, Set Screws, Rods, Crosslinks, Connectors, Hooks	Rods do not include S-rod
LnK Spinal Fixation System/OpenLoc-L Spinal Fixation System (K120270, K143363, K171813, K183168, K 200790, K223565)	Additional Predicate	Screws, Set Screws, Rods, Crosslinks, Connectors, Hooks	Rods do not include S-rod
Olympic Posterior Spinal Fixation System(K181139)	Additional Predicate	Screws, Set Screws, Rods, Crosslinks, Connectors, Hooks Rods include Z-rod	

Dimension			
Devices		Similarities	
LnK Spinal Fixation System/OpenLoc-L Spinal Fixation System	Subject	$\begin{tabular}{ c c c c c c c c c c c c c c c c c c c$	
LnK Spinal Fixation System(K230245)	Primary Predicate	$\begin{tabular}{ c c c c c c c c c c c c c c c c c c c$	
LnK Spinal Fixation System/OpenLoc-L Spinal Fixation System (K120270, K143363, K171813, K183168, K 200790, K223565)	Additional Predicate	$\begin{tabular}{ c c c c c c c c c c c c c c c c c c c$	
Olympic Posterior Spinal Fixation System(K181139)	Additional Predicate	$\begin{tabular}{ c c c c c c c c c c c c c c c c c c c$	



Sterilization Method		
Devices		Similarities
LnK Spinal Fixation System/OpenLoc-L Spinal Fixation System	Subject	 Sterile device: Gamma radiation Non-sterile device: recommended steam sterilization (autoclave sterilization)
LnK Spinal Fixation System(K230245)	Primary Predicate	 Sterile device: Gamma radiation Non-sterile device: recommended steam sterilization (autoclave sterilization)
LnK Spinal Fixation System/OpenLoc-L Spinal Fixation System (K120270, K143363, K171813, K183168, K 200790, K223565)	Additional Predicate	 Sterile device: Gamma radiation Non-sterile device: recommended steam sterilization (autoclave sterilization)
Olympic Posterior Spinal Fixation System(K181139)	Additional Predicate	• Non-sterile

ACCELFIX SPINAL FIXATION SYSTEM

INDICATION FOR USE		
Devices		Similarities
AccelFix Spinal Fixation System	Subject	
AccelFix Spinal Fixation System (K182544)	Primary Predicate	
AccelFix Spinal Fixation System (K200794, K223565, K230245)	Additional	Instruction for use including indication are similar.
Olympic Posterior Spinal Fixation System(K181139)	Predicate	

Materials				
Devices		Similarities		
AccelFix Spinal Fixation System	Subject			
AccelFix Spinal Fixation System (K182544)	Primary Predicate	Similar materials • Ti-6Al-4V ELI titanium alloy (ASTM F136)		
AccelFix Spinal Fixation System (K200794, K223565, K230245)	Additional	Cobalt-28Chromium-6Molybdenum- 4Vanadium ELI (ASTM F1537)		
Olympic Posterior Spinal Fixation System(K181139)	Predicate			

Design with Components			
Devices		Similarities	Dissimilarities
AccelFix Spinal Fixation System	Subject	Screws, Set Screws, Rods, Crosslinks, Connectors, Hooks Rods include Z-rod	
AccelFix Spinal Fixation System (K182544)	Primary Predicate	Screws, Set Screws, Rods, Crosslinks, Connectors, Hooks	Rods do not include S-rod
AccelFix Spinal Fixation System (K200794, K223565, K230245)	Additional Predicate	Screws, Set Screws, Rods, Crosslinks, Connectors, Hooks	Rods do not include S-rod
Olympic Posterior Spinal Fixation System(K181139)	Additional Predicate	Screws, Set Screws, Rods, Crosslinks, Connectors, Hooks Rods include Z-rod	



Dimension					
Devices		Similarities			
AccelFix Spinal Fixation System	Subject	_	Screws Rods Z-Rod	OD Ø Length OD Ø Length OD Ø Length	4.0~10.5 mm 20~150mm 5.5/6.0/6.35 mm 40~600 mm 5.5/6.0mm 320mm
		S-Rod	S-Rod	OD Ø Length	5.0/5.5/6.0mm 320mm
AccelFix Spinal Fixation System (K182544)	Primary Predicate	:	Screws Rods	OD Ø Length OD Ø Length	4.0~10.5 mm 20~ 150mm 5.5/6.0/6.35 mm 40~600 mm
AccelFix Spinal Fixation System (K200794, K223565, K230245)	Additional Predicate	_	Screws Rods S-Rod	OD Ø Length OD Ø Length OD Ø Length	4.0~10.5 mm 20~ 150mm 5.5/6.0/6.35 mm 40~600 mm 5.0/5.5/6.0mm 320mm
Olympic Posterior Spinal Fixation System(K181139)	Additional Predicate	_	Screws Rods Z-Rod	OD Ø Length OD Ø Length OD Ø Length	4.0~10 mm 20~ 120mm 5.5/6.0/6.35 mm - 5.5/6.0mm -

Sterilization Method				
Devices		Similarities		
AccelFix Spinal Fixation System	Subject	 Sterile device: Gamma radiation Non-sterile device: recommended steam sterilization (autoclave sterilization) 		
AccelFix Spinal Fixation System (K182544)	Primary Predicate	 Sterile device: Gamma radiation Non-sterile device: recommended steam sterilization (autoclave sterilization) 		
AccelFix Spinal Fixation System (K200794, K223565, K230245)	Additional Predicate	 Sterile device: Gamma radiation Non-sterile device: recommended steam sterilization (autoclave sterilization) 		
Olympic Posterior Spinal Fixation System(K181139)	Additional Predicate	• Non-sterile		

9. SUBSTANTIAL EQUIVALENCE AND CONCLUSION

The subject additional components of LnK Spinal Fixation System/ OpenLoc-L Spinal Fixation System and AccelFix Spinal Fixation System have been found to be substantially equivalent to the predicate devices with respect to technical characteristics and intended use.

The information provided within this premarket notification supports substantial equivalence of the subject device to the predicate devices. The overall data lead to the conclusion that the additional components of LnK Spinal Fixation System/ OpenLoc-L Spinal Fixation System and AccelFix Spinal Fixation System are substantially equivalent to the predicate devices (K120270, K143363, K171813, K183168, K200790, K182544, K200794, K223565, K230245).