



June 9, 2021

Onkos Surgical
Matthew Vernak
Vice President, Quality, Regulatory, and Product Development
77 East Halsey Rd
Parsippany, New Jersey 07054

Re: K203815

Trade/Device Name: ELEOS Limb Salvage System with BioGrip
Regulation Number: 21 CFR 888.3350
Regulation Name: Hip Joint Metal/Polymer Semi-Constrained Cemented Prosthesis
Regulatory Class: Class II
Product Code: JDI, KRO, JWH, LPH, LZO, KWY
Dated: May 12, 2021
Received: May 13, 2021

Dear Matthew Vernak:

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

Sincerely,

For:

William Jung, PhD
Director
DHT6A: Division of Joint
Arthroplasty Devices
OHT6: Office of Orthopedic Devices
Office of Product Evaluation and Quality
Center for Devices and Radiological Health

Enclosure

4. Indications for Use Statement (Form FDA 3881)

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 (if known) K203815	
Device Name ELEOS™ Limb Salvage System with BioGrip™	
Indications for Use (Describe) ELEOS™ Limb Salvage System Hip Components: Indicated for use in total hip arthroplasty for reduction or relief of pain and/or improved hip function in skeletally mature patients with the following conditions:	
1) Non-inflammatory degenerative joint disease such as osteoarthritis, avascular necrosis, ankylosis, protrusion acetabuli, and painful hip dysplasia; 2) Inflammatory degenerative joint disease such as rheumatoid arthritis; 3) Correction of functional deformity 4) Revision procedures where other treatments or devices have failed; and, 5) Treatment of fractures that are unmanageable using other techniques. Limb salvage system is also indicated for procedures where radical resection and replacement of the proximal, distal and/ or total femur is required with the following conditions:	
1) Patients suffering from severe arthropathy of the hip that does not respond to any conservative therapy or better alternative surgical treatment; 2) Surgical intervention for severe trauma, revision hip arthroplasties, and/or Oncology indications. 3) Metastatic diseases	
ELEOS™ Limb Salvage System Knee Components: Indicated for cemented use in knee arthroplasty for reduction or relief of pain and/or improved knee function in skeletally mature patients with the following conditions:	
1) Non-inflammatory degenerative joint disease including osteoarthritis, traumatic arthritis, or avascular necrosis; 2) Inflammatory degenerative joint disease including rheumatoid arthritis; 3) Correction of functional deformity; 4) Revision procedures where other treatments or devices have failed; and 5) Treatment of fractures that are unmanageable using other techniques.	
Limb salvage system is also indicated for procedures where radical resection and replacement of the proximal tibia is required with the following conditions:	
1) Patients suffering from severe arthropathy of the knee that does not respond to any conservative therapy or better alternative surgical treatment; 2) Surgical intervention for severe trauma, revision knee arthroplasties, and/or Oncology indications. 3) Metastatic diseases	
Type of Use (Select one or both, as applicable)	
<input checked="" type="checkbox"/> Prescription Use (Part 21 CFR 801 Subpart D) <input type="checkbox"/> Over-The-Counter Use (21 CFR 801 Subpart C)	
CONTINUE ON A SEPARATE PAGE IF NEEDED.	

Premarket Notification: ELEOS™ Limb Salvage System with BioGrip™

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.

The burden time for this collection of information is estimated to average 79 hours per response, including the time to review instructions, search existing data sources, gather and maintain the data needed and complete and review the collection of information. Send comments regarding this burden estimate or any other aspect of this information collection, including suggestions for reducing this burden, to:

Department of Health and Human Services
Food and Drug Administration
Office of Chief Information Officer
Paperwork Reduction Act (PRA) Staff
PRASStaff@fda.hhs.gov

"An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB number."

5. 510(k) Summary

5.1. Submitter

Onkos Surgical, Inc.
77 East Halsey Road
Parsippany, NJ 07054
Phone: (551) 579-1081
Contact Person: Matthew Vernak
Email: mvernak@onkossurgical.com

Date Prepared: 08-Jun-2021

5.2. Device

Name of Device: ELEOS™ Limb Salvage System with BioGrip™

Common Name: Limb Salvage System

Classification Name: 21 CFR 888.3350, Prosthesis, Hip, Semi-Constrained, Metal/Polymer, Cemented
21 CFR 888.3510, Prosthesis, Knee, Femorotibial, Constrained, Metal Polymer, Cemented
21 CFR 888.3560, Knee joint patellofemorotibial polymer/metal/polymer semi- constrained cemented prosthesis
21 CFR 888.3358 Prosthesis, Hip, Semi-Constrained, Metal/Polymer, Porous, Uncemented
21 CFR 888.3353, Prosthesis, Hip, Semi-Constrained, Metal/Ceramic/Polymer Cemented or Non-Porous, Uncemented
21 CFR 888.3390 Hip joint femoral (hemi-hip) metal/polymer cemented or uncemented prosthesis

Regulatory Class: II

Product Code(s): JDI KRO JWH LPH LZO KWY

5.3. Predicate and Reference Devices

ELEOS™ Limb Salvage System with BioGrip™, Onkos Surgical, Inc., K203090 (Predicate Device)
ELEOS™ Limb Salvage System, Onkos Surgical, Inc., K161520 (Predicate Device)
Modular Endoprosthesis Tumour System (METS), Stanmore, K121029 (Reference Device)
Modular Endoprosthesis Tumour System (METS), Stanmore, K121056 (Reference Device)

5.4. Device Description

The Onkos Surgical ELEOS™ Limb Salvage System consists of components that are used in the reconstruction of the lower limb. The reconstruction applications are proximal femur, distal femur, total femur, proximal tibia, and hinged knee. The ELEOS Limb Salvage components are femoral head, proximal femur, mid-section, stem, distal hinge femur, tibial hinge assembly, axial pin, tibial poly spacer, tibial sleeve, male-male mid-section, resurfacing hinge femur, and proximal tibia, optional components include modular collars, patella, stem extensions, tibial wedges and augments. Instrumentation is provided non-sterile in surgical trays which are to be re-processed per the validated instructions stated below.

Components	RECONSTRUCTION APPLICATIONS				
	Proximal Femur	Distal Femur	Total Femur	Proximal Tibia	Hinged Knee
Femoral head	✓		✓		
Proximal Femur	✓		✓		
Mid-Section	✓	✓	✓	✓	
Segmental Stem	✓	✓		✓	
Modular Collar^{1, 2}	✓	✓		✓	
Distal Femur		✓	✓		
Tibial Hinge Component		✓	✓	✓	✓
Axial Pin		✓	✓	✓	✓
Tibial Poly Spacer		✓	✓	✓	✓
Tibial Baseplate		✓	✓		✓
Male-Male Mid-Section			✓		
Resurfacing Hinge Femur				✓	✓
Proximal Tibia¹				✓	
Patella ²		✓	✓	✓	✓
Wedges and Augments ²		✓	✓		✓
Stem Extensions ²		✓	✓	✓	✓

1 - Bolded components are offered with BioGrip™ porous technology. Porous modular collars are offered with or without HA treatment.

2 - These implants are optional for each procedure. The surgeon shall use his/her medical judgement to determine if these implants are necessary based on factor such as patient bone quality, joint stability and pathology.

The implants are single use devices.

5.5. Indications for Use

ELEOS™ Limb Salvage System Hip Components:

Indicated for use in total hip arthroplasty for reduction or relief of pain and/or improved hip function in skeletally mature patients with the following conditions:

- 1) Non-inflammatory degenerative joint disease such as osteoarthritis, avascular necrosis, ankylosis, protrusion acetabuli, and painful hip dysplasia;
- 2) Inflammatory degenerative joint disease such as rheumatoid arthritis;
- 3) Correction of functional deformity
- 4) Revision procedures where other treatments or devices have failed; and,
- 5) Treatment of fractures that are unmanageable using other techniques.

Limb salvage system is also indicated for procedures where radical resection and replacement of the proximal, distal and/or total femur is required with the following conditions:

- 1) Patients suffering from severe arthropathy of the hip that does not respond to any conservative therapy or better alternative surgical treatment;
- 2) Surgical intervention for severe trauma, revision hip arthroplasties, and/or Oncology indications.
- 3) Metastatic diseases

ELEOS™ Limb Salvage System with Knee Components:

Indicated for cemented use in knee arthroplasty for reduction or relief of pain and/or improved knee function in skeletally mature patients with the following conditions:

- 1) Non-inflammatory degenerative joint disease including osteoarthritis, traumatic arthritis, or avascular necrosis;
- 2) Inflammatory degenerative joint disease including rheumatoid arthritis;
- 3) Correction of functional deformity;
- 4) Revision procedures where other treatments or devices have failed; and
- 5) Treatment of fractures that are unmanageable using other techniques.

Limb salvage system is also indicated for procedures where radical resection and replacement of the proximal tibia is required with the following conditions:

- 1) Patients suffering from severe arthropathy of the knee that does not respond to any conservative therapy or better alternative surgical treatment;
- 2) Surgical intervention for severe trauma, revision knee arthroplasties, and/or Oncology indications.
- 3) Metastatic diseases

5.6. Comparison of Technological Characteristics with the Predicate Device

The primary difference between the subject device, which is a line extension to the predicate device, is the provision of modular collars to match patient anatomy. Specifically, the stems have been modified to accept porous or solid modular collars. The porous collars are produced using an additive manufacturing process with the option to be treated with a nano HA surface treatment. The implant incorporates a hybrid stem and a modular collar with or without HA. Mechanical (both static and fatigue) and biocompatibility testing have demonstrated that the subjected device is substantially equivalent to the predicate system. Furthermore, the technological characteristic of a modular collar has been cleared with Stanmore's METS,

reference K121029 and K121056.

5.7. Performance Data

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

5.7.1 Biocompatibility

The biocompatibility evaluation for the ELEOS Limb Salvage System with BioGrip™ was conducted in accordance with FDA's Guidance for Industry and FDA Staff, "Use of International Standard ISO 10993, Biological Evaluation of Medical Devices Part 1: Evaluation and Testing within a Risk Management Process" issued June 16, 2016. The subject device is a permanent contact device manufactured from printed Ti6Al4V. The following biocompatibility tests were performed on each material group to ensure biocompatibility:

- Cytotoxicity (per ISO 10993-5, Biological evaluation of medical devices - Part 5: Tests for in vitro cytotoxicity)
- Sensitization (per ISO 10993-10, Biological evaluation of medical devices - Part 10: Tests for irritation and skin sensitization)
- Irritation (per ISO 10993-10, Biological evaluation of medical devices - Part 10: Tests for irritation and skin sensitization)
- Systemic Toxicity: Acute (per ISO 10993-11, Biological evaluation of medical devices - Part 11: Tests for systemic toxicity)
- Systemic Toxicity: Material Mediated Pyrogenicity (per ISO 10993-11, Biological evaluation of medical devices - Part 11: Tests for systemic toxicity, and ST72:2011/(R)2016, Bacterial endotoxins - Test methods routine monitoring and alternatives to batch testing)
- Genotoxicity (per ISO 10993-3, Biological evaluation of medical devices - Part 3: Tests for genotoxicity carcinogenicity and reproductive toxicity)
- Chemical Characterization (per ISO 10993-18, Biological evaluation of medical devices - Part 18: Chemical characterization of medical device materials within a risk management process)
- Toxicological Risk Assessment (per ISO 10993-17, Biological evaluation of medical devices - Part 17: Establishment of allowable limits for leachable substances)

The bacterial endotoxin test was performed to verify that the subject implants meet the 20 endotoxin units (EU)/device pyrogen limit specification. Testing was successfully performed, and it was confirmed that the subject devices meet the 20 EU/device testing limit for general medical devices that are implanted as outlined in ANSI/AAMI ST72: Bacterial endotoxins – Test methods, routine monitoring, and alternatives to batch testing and USP<161>, Medical Devices – Bacterial Endotoxin and Pyrogen Tests. Testing to monitor pyrogens will be performed periodically.

5.7.2. Mechanical Testing

Onkos Surgical has evaluated the subject devices to demonstrate substantial equivalence to the

predicate devices. The following FDA Guidance documents were consulted to select the bench tests:

- *FDA Guidance*: “Guidance Document for Testing Orthopedic Implants with Modified Metallic Surfaces Apposing Bone or Bone Cement”, Issued 28-Apr-1994

The following bench testing was completed and the results of support the subject devices are equivalent to the predicate device:

- ASTM F1044 - Standard Test Method for Shear Testing of Calcium Phosphate Coatings and Metallic Coatings
- ASTM F1147 - Standard Test Method for Tension Testing of Calcium Phosphate and Metallic Coatings
- ASTM F1160: Standard Test Method for Shear and Bending Fatigue Testing of Calcium Phosphate and Metallic Medical and Composite Calcium Phosphate/Metallic Coatings
- ASTM F1854 - Standard Test Method for Stereological Evaluation of Porous Coatings on Medical Implants
- ASTM F1978 - Standard Test Method for Measuring Abrasion Resistance of Metallic Thermal Spray Coatings by Using the Taber Abraser
- ISO 7206-4 - Implants for surgery — Partial and total hip joint prostheses — Part 4: Determination of endurance properties and performance of stemmed femoral components
- ISO 13179-1:2014: Implants for surgery — Plasma sprayed unalloyed titanium coatings on metallic surgical implants

5.8 Clinical Data

Clinical data was not deemed necessary for the subject device.

5.9 Design Validation

Design Validation was performed for the subject implant using a cadaver lab. Design Validation demonstrated the subject implant and existing instruments function as intended and user needs were met.

5.10 Conclusions

Based on the test results and supporting documentation provided in this premarket notification, the subject ELEOS Limb Salvage System with BioGrip™ is substantially equivalent to the predicate device, ELEOS Limb Salvage System with BioGrip™. The content in this premarket notification demonstrates that:

- any differences do not raise new questions of safety and effectiveness;
- and the proposed device is at least as safe and effective as the legally marketed predicate device.



Onkos Surgical
Matthew Vernak
Vice President, Quality, Regulatory and Product Development
77 East Halsey Rd
Parsippany, New Jersey 07054

February 5, 2021

Re: K203588

Trade/Device Name: ELEOS™ Limb Salvage System

Regulation Number: 21 CFR 888.3510

Regulation Name: Knee Joint Femorotibial Metal/Polymer Constrained Cemented Prosthesis

Regulatory Class: Class II

Product Code: KRO, JWH, JDI, LPH, LZO

Dated: December 8, 2020

Received: December 8, 2020

Dear Matthew Vernak:

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.

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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>.

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Sincerely,

Vesa Vuniqui
Assistant Director
DHT6A: Division of Joint Arthroplasty Devices
OHT6: Office of Orthopedic Devices
Office of Product Evaluation and Quality
Center for Devices and Radiological Health

Enclosure

DEPARTMENT OF HEALTH AND HUMAN SERVICES
Food and Drug Administration

Form Approved: OMB No. 0910-0120
Expiration Date: 06/30/2023
See PRA Statement below.

Indications for Use

510(k) Number (if known)
K203588

Device Name
ELEOST™ Limb Salvage System

Indications for Use (Describe)

ELEOST™ Limb Salvage System with Hip Components:

Indicated for use in total hip arthroplasty for reduction or relief of pain and/or improved hip function in skeletally mature patients with the following conditions:

- 1) Non-inflammatory degenerative joint disease such as osteoarthritis, avascular necrosis, ankylosis, protrusion acetabuli, and painful hip dysplasia;
- 2) Inflammatory degenerative joint disease such as rheumatoid arthritis;
- 3) Correction of functional deformity
- 4) Revision procedures where other treatments or devices have failed; and,
- 5) Treatment of fractures that are unmanageable using other techniques.

Limb salvage system is also indicated for procedures where radical resection and replacement of the proximal, distal and/or total femur is required with the following conditions:

- 1) Patients suffering from severe arthropathy of the hip that does not respond to any conservative therapy or better alternative surgical treatment;
- 2) Surgical intervention for severe trauma, revision hip arthroplasties, and/or Oncology indications.
- 3) Metastatic diseases

ELEOST™ Limb Salvage System with Knee Components:

Indicated for cemented use in knee arthroplasty for reduction or relief of pain and/or improved knee function in skeletally mature patients with the following conditions:

- 1) Non-inflammatory degenerative joint disease including osteoarthritis, traumatic arthritis, or avascular necrosis;
- 2) Inflammatory degenerative joint disease including rheumatoid arthritis;
- 3) Correction of functional deformity;
- 4) Revision procedures where other treatments or devices have failed; and
- 5) Treatment of fractures that are unmanageable using other techniques.

Limb salvage system is also indicated for procedures where radical resection and replacement of the proximal tibia is required with the following conditions:

- 1) Patients suffering from severe arthropathy of the knee that does not respond to any conservative therapy or better alternative surgical treatment;
- 2) Surgical intervention for severe trauma, revision knee arthroplasties, and/or Oncology indications.
- 3) Metastatic diseases

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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Department of Health and Human Services
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PRAStaff@fda.hhs.gov

“An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB number.”

Premarket Notification: ELEOS™ Limb Salvage System

5. 510(k) Summary**5.1. Submitter**

Onkos Surgical, Inc.
 77 East Halsey Road
 Parsippany, NJ 07054
 Phone: (551) 579-1081
 Contact Person: Matthew Vernak
 Email: mvernak@onkossurgical.com

Date Prepared: 04-Feb-2021

Name of Device: ELEOS™ Limb Salvage System

Common Name: Limb Salvage System

Classification Name: 21 CFR 888.3510, Prosthesis, Knee, Femorotibial, Constrained, Metal Polymer, Cemented
 21 CFR 888.3560, Knee joint patellofemorotibial polymer/metal/polymer semi- constrained cemented prosthesis
 21 CFR 888.3350, Prosthesis, Hip, Semi-Constrained, Metal/Polymer, Cemented
 21 CFR 888.3358 Prosthesis, Hip, Semi-Constrained, Metal/Polymer, Porous, Uncemented
 21 CFR 888.3353, Prosthesis, Hip, Semi-Constrained, Metal/Ceramic/Polymer Cemented or Non-Porous, Uncemented

Regulatory Class: II

Product Code(s): KRO JWH JDI LPH LZO

5.3. Predicate Device

ELEOS™ Limb Salvage System, Onkos Surgical, Inc., K161520 (Predicate)
 Orthopaedic Salvage System (OSS), Biomet, K052685 (Reference Device)

5.4. Device Description

The Onkos Surgical ELEOS™ Limb Salvage System consists of components that are used in the reconstruction of the lower limb. The reconstruction applications are proximal femur, distal femur, total femur, proximal tibia, and hinged knee. The ELEOS Limb Salvage components are femoral head, proximal femur, mid-section, stem, distal femur, tibial hinge component, axial pin, tibial poly spacer, tibial baseplate, male-male mid-section, resurfacing femur, proximal tibia,

Premarket Notification: ELEOS™ Limb Salvage System

tapered screws, patella, stem extension, tibial wedges and augments. Instrumentation is provided non-sterile in surgical trays which are to be re-processed per validated instructions.

Components	RECONSTRUCTION APPLICATIONS				
	Proximal Femur	Distal Femur	Total Femur	Proximal Tibia	Hinged Knee
Femoral head	✓		✓		
Proximal Femur	✓		✓		
Mid-Section	✓	✓	✓	✓	
Segmental Stem	✓	✓		✓	
Distal Femur		✓	✓		
Tibial Hinge Component		✓	✓	✓	✓
Axial Pin		✓	✓	✓	✓
Tibial Poly Spacer		✓	✓	✓	✓
Tibial Baseplate ¹		✓	✓		✓
Male-Male Mid-Section			✓		
Resurfacing Femur ¹				✓	✓
Proximal Tibia				✓	
Tapered Screws ¹		✓	✓	✓	✓
Patella ²		✓	✓	✓	✓
Wedges and Augments ²		✓	✓		✓
Stem Extensions ²		✓	✓	✓	✓

1 – The Resurfacing Femur and Tibial Baseplate have the option for addition of a tapered screw between the respective component and a stem extension. The surgeon shall use their medical judgement to determine if the additional fixation is necessary based on factors such as patient bone quality, joint stability, and pathology.

2 – These implants are optional for each procedure. The surgeon shall use their medical judgement to determine if these implants are necessary based on factor such as patient bone quality, joint stability, and pathology.

The implants are single use devices.

5.5. Indications for Use

ELEOS™ Limb Salvage System with Hip Components:

Indicated for use in total hip arthroplasty for reduction or relief of pain and/or improved hip function in skeletally mature patients with the following conditions:

- 1) Non-inflammatory degenerative joint disease such as osteoarthritis, avascular necrosis, ankylosis, protrusion acetabuli, and painful hip dysplasia;
- 2) Inflammatory degenerative joint disease such as rheumatoid arthritis;
- 3) Correction of functional deformity
- 4) Revision procedures where other treatments or devices have failed; and,

Premarket Notification: ELEOS™ Limb Salvage System

- 5) Treatment of fractures that are unmanageable using other techniques.

Limb salvage system is also indicated for procedures where radical resection and replacement of the proximal, distal and/or total femur is required with the following conditions:

- 1) Patients suffering from severe arthropathy of the hip that does not respond to any conservative therapy or better alternative surgical treatment;
- 2) Surgical intervention for severe trauma, revision hip arthroplasties, and/or Oncology indications.
- 3) Metastatic diseases

ELEOS™ Limb Salvage System Knee Components:

Indicated for cemented use in knee arthroplasty for reduction or relief of pain and/or improved knee function in skeletally mature patients with the following conditions:

- 1) Non-inflammatory degenerative joint disease including osteoarthritis, traumatic arthritis, or avascular necrosis;
- 2) Inflammatory degenerative joint disease including rheumatoid arthritis;
- 3) Correction of functional deformity;
- 4) Revision procedures where other treatments or devices have failed; and
- 5) Treatment of fractures that are unmanageable using other techniques.

Limb salvage system is also indicated for procedures where radical resection and replacement of the proximal tibia is required with the following conditions:

- 1) Patients suffering from severe arthropathy of the knee that does not respond to any conservative therapy or better alternative surgical treatment;
- 2) Surgical intervention for severe trauma, revision knee arthroplasties, and/or Oncology indications.
- 3) Metastatic diseases

5.6. Comparison of Technological Characteristics with the Predicate Device

The primary difference between the subject device and the predicate device is the option to use a tapered screw between the resurfacing femur and tibial baseplate components and corresponding stem extension. The tapered screw will be manufactured from Titanium-6Aluminum-4Vanadium (TAV) ELI per ASTM F136 and offered in two sizes (0.650" for use with the tibial baseplate and 1" for use with the resurfacing femur). To accommodate the screw, the resurfacing femur was modified to remove material between the condyles to allow insertion of a tapered screw between the resurfacing femur and stem extension after impaction of the tapered connections. This modification was incorporated as not to affect any of the bearing surfaces. Additionally, a tapered hole was incorporated into the sleeve of the tibial baseplate to allow insertion of a tapered screw between the tibial baseplate and stem extension after impaction of the tapered connections.

Premarket Notification: ELEOS™ Limb Salvage System

5.7. Performance Data

The following performance data are provided in support of substantial equivalence:

5.7.1 Mechanical Testing

Onkos Surgical has evaluated the subject devices to demonstrate substantial equivalence to the predicate devices. The following bench testing was completed and the results of support the subject devices are equivalent to the predicate device:

- Fatigue testing per ASTM F1800, Standard Practice for Cyclic Fatigue Testing of Metal Tibial Tray Components of Total Knee Joint Replacements and ASTM F2083, Standard Specification for Knee Replacement Prosthesis
- Evaluation of fretting corrosion for mating surfaces post fatigue testing
- Evaluation of screw loosening post fatigue testing

5.8 Clinical Data

Clinical data was not deemed necessary for the subject device.

5.9 Conclusions

Based on the test results and supporting documentation provided in this premarket notification, the subject ELEOS Limb Salvage System is substantially equivalent to the predicate, ELEOS Limb Salvage System. The content in this premarket notification demonstrates that:

- any differences do not raise new questions of safety and effectiveness;
- and the proposed device is at least as safe and effective as the legally marketed predicate device.



December 11, 2020

Onkos Surgical
Matthew (Matt) Vernak
Vice President, Quality, Regulatory, and Product Development
77 East Halsey Rd
Parsippany, New Jersey 07054

Re: K203090

Trade/Device Name: ELEOS™ Limb Salvage System featuring BIOGRIP™
Regulation Number: 21 CFR 888.3350
Regulation Name: Hip Joint Metal/Polymer Semi-Constrained Cemented Prosthesis
Regulatory Class: Class II
Product Code: JDI, KRO, JWH, LPH, LZO
Dated: October 13, 2020
Received: October 13, 2020

Dear Matthew (Matt) Vernak:

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

Sincerely,

Ting Song, Ph.D., R.A.C.
Assistant Director
DHT6A: Division of Joint
Arthroplasty 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)

K203090

Device Name

ELEOST™ Limb Salvage System with BIOGRIP™

Indications for Use (Describe)

ELEOST™ Limb Salvage System with Hip Components:

Indicated for use in total hip arthroplasty for reduction or relief of pain and/or improved hip function in skeletally mature patients with the following conditions:

- 1) Non-inflammatory degenerative joint disease such as osteoarthritis, avascular necrosis, ankylosis, protrusion acetabuli, and painful hip dysplasia;
- 2) Inflammatory degenerative joint disease such as rheumatoid arthritis;
- 3) Correction of functional deformity
- 4) Revision procedures where other treatments or devices have failed; and,
- 5) Treatment of fractures that are unmanageable using other techniques.

Limb salvage system is also indicated for procedures where radical resection and replacement of the proximal, distal and/or total femur is required with the following conditions:

- 1) Patients suffering from severe arthropathy of the hip that does not respond to any conservative therapy or better alternative surgical treatment;
- 2) Surgical intervention for severe trauma, revision hip arthroplasties, and/or Oncology indications.
- 3) Metastatic diseases

ELEOST™ Limb Salvage System with Knee Components:

Indicated for cemented use in knee arthroplasty for reduction or relief of pain and/or improved knee function in skeletally mature patients with the following conditions:

- 1) Non-inflammatory degenerative joint disease including osteoarthritis, traumatic arthritis, or avascular necrosis;
- 2) Inflammatory degenerative joint disease including rheumatoid arthritis;
- 3) Correction of functional deformity;
- 4) Revision procedures where other treatments or devices have failed; and
- 5) Treatment of fractures that are unmanageable using other techniques.

Limb salvage system is also indicated for procedures where radical resection and replacement of the proximal tibia is required with the following conditions:

- 1) Patients suffering from severe arthropathy of the knee that does not respond to any conservative therapy or better alternative surgical treatment;
- 2) Surgical intervention for severe trauma, revision knee arthroplasties, and/or Oncology indications.
- 3) Metastatic diseases

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)

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Department of Health and Human Services
Food and Drug Administration
Office of Chief Information Officer
Paperwork Reduction Act (PRA) Staff
PRAStaff@fda.hhs.gov

“An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB number.”

Premarket Notification: ELEOS™ Limb Salvage System with BIOGRIP™

5. 510(k) Summary**5.1. Submitter**

Onkos Surgical, Inc.
 77 East Halsey Road
 Parsippany, NJ 07054
 Phone: (551) 579-1081
 Contact Person: Matthew Vernak
 Email: mvernak@onkossurgical.com

Date Prepared: 11-Oct-2020

5.2. Device

Name of Device: ELEOS™ Limb Salvage System with BIOGRIP™

Common Name: Limb Salvage System

Classification Name: 21 CFR 888.3350, Prosthesis, Hip, Semi-Constrained, Metal/Polymer, Cemented
 21 CFR 888.3510, Prosthesis, Knee, Femorotibial, Constrained, Metal Polymer, Cemented
 21 CFR 888.3560, Knee joint patellofemorotibial polymer/metal/polymer semi- constrained cemented prosthesis
 21 CFR 888.3358 Prosthesis, Hip, Semi-Constrained, Metal/Polymer, Porous, Uncemented
 21 CFR 888.3353, Prosthesis, Hip, Semi-Constrained, Metal/Ceramic/Polymer Cemented or Non-Porous, Uncemented

Regulatory Class: II

Product Code(s): JDI KRO JWH LPH LZO

5.3. Predicate Device

ELEOS™ Limb Salvage System, Onkos Surgical, Inc., K161520

5.4. Device Description

The Onkos Surgical ELEOS™ Limb Salvage System consists of components that are used in the reconstruction of the lower limb. The reconstruction applications are proximal femur, distal femur, total femur, proximal tibia, and hinged knee. The ELEOS Limb Salvage components are femoral head, proximal femur, mid-section, stem, distal hinge femur, tibial hinge assembly, axial pin, tibial poly spacer, tibial sleeve, male-male mid-section, resurfacing hinge femur, and proximal tibia, patella, stem extension, modular collar, tibial wedges and augments.

Premarket Notification: ELEOS™ Limb Salvage System with BIOGRIP™

Instrumentation is provided non-sterile in surgical trays which are to be re-processed per the validated instructions stated below.

Components	RECONSTRUCTION APPLICATIONS				
	Proximal Femur	Distal Femur	Total Femur	Proximal Tibia	Hinged Knee
Femoral head	✓		✓		
Proximal Femur	✓		✓		
Mid-Section	✓	✓	✓	✓	
Segmental Stem	✓	✓		✓	
Modular Collar	✓	✓		✓	
Distal Femur		✓	✓		
Tibial Hinge Component		✓	✓	✓	✓
Axial Pin		✓	✓	✓	✓
Tibial Poly Spacer		✓	✓	✓	✓
Tibial Baseplate		✓	✓		✓
Male-Male Mid-Section			✓		
Resurfacing Hinge Femur				✓	✓
Proximal Tibia¹				✓	
Patella ²		✓	✓	✓	✓
Wedges and Augments ²		✓	✓		✓
Stem Extensions ²		✓	✓	✓	✓

1 – Bolded components are offered with BIOGRIP™ porous technology.

2 - These implants are optional for each procedure. The surgeon shall use his/her medical judgement to determine if these implants are necessary based on factor such as patient bone quality, joint stability and pathology.

The implants are single use devices.

5.5. Indications for Use

ELEOS™ Limb Salvage System with Hip Components:

Indicated for use in total hip arthroplasty for reduction or relief of pain and/or improved hip function in skeletally mature patients with the following conditions:

- 1) Non-inflammatory degenerative joint disease such as osteoarthritis, avascular necrosis, ankylosis, protrusion acetabuli, and painful hip dysplasia;
- 2) Inflammatory degenerative joint disease such as rheumatoid arthritis;
- 3) Correction of functional deformity
- 4) Revision procedures where other treatments or devices have failed; and,
- 5) Treatment of fractures that are unmanageable using other techniques.

Limb salvage system is also indicated for procedures where radical resection and replacement of the proximal, distal and/or total femur is required with the following conditions:

Premarket Notification: ELEOS™ Limb Salvage System with BIOGRIP™

- 1) Patients suffering from severe arthropathy of the hip that does not respond to any conservative therapy or better alternative surgical treatment;
- 2) Surgical intervention for severe trauma, revision hip arthroplasties, and/or Oncology indications.
- 3) Metastatic diseases

ELEOS™ Limb Salvage System with Knee Components:

Indicated for cemented use in knee arthroplasty for reduction or relief of pain and/or improved knee function in skeletally mature patients with the following conditions:

- 1) Non-inflammatory degenerative joint disease including osteoarthritis, traumatic arthritis, or avascular necrosis;
- 2) Inflammatory degenerative joint disease including rheumatoid arthritis;
- 3) Correction of functional deformity;
- 4) Revision procedures where other treatments or devices have failed; and
- 5) Treatment of fractures that are unmanageable using other techniques.

Limb salvage system is also indicated for procedures where radical resection and replacement of the proximal tibia is required with the following conditions:

- 1) Patients suffering from severe arthropathy of the knee that does not respond to any conservative therapy or better alternative surgical treatment;
- 2) Surgical intervention for severe trauma, revision knee arthroplasties, and/or Oncology indications.
- 3) Metastatic diseases

5.6. Comparison of Technological Characteristics with the Predicate Device

The primary difference between the subject implant, which is a line extension to the predicate device system, is the manufacturing process used. Specifically, an additive manufacturing processes is used to create the ELEOS BIOGRIP™ Proximal Tibia. The implant incorporates an additive porous surface (replacing titanium plasma) and additional suture holes are incorporated to better facilitate approximation of soft tissue and support bony apposition. Mechanical (both static and fatigue) and biocompatibility testing have demonstrated that the subjected device is substantially equivalent to the predicate system.

5.7. Performance Data

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

5.7.1 Biocompatibility

The biocompatibility evaluation for the ELEOS BIOGRIP™ Proximal Tibia was conducted in accordance with FDA's Guidance for Industry and FDA Staff, "Use of International Standard ISO 10993, Biological Evaluation of Medical Devices Part 1: Evaluation and Testing within a

Premarket Notification: ELEOS™ Limb Salvage System with BIOGRIP™

Risk Management Process” issued June 16, 2016. The subject device is a permanent contact device manufactured from printed Ti6Al4V. The following biocompatibility tests were performed on each material group to ensure biocompatibility:

- Cytotoxicity (per ISO 10993-5, Biological evaluation of medical devices - Part 5: Tests for in vitro cytotoxicity)
- Sensitization (per ISO 10993-10, Biological evaluation of medical devices - Part 10: Tests for irritation and skin sensitization)
- Irritation (per ISO 10993-10, Biological evaluation of medical devices - Part 10: Tests for irritation and skin sensitization)
- Systemic Toxicity: Acute (per ISO 10993-11, Biological evaluation of medical devices - Part 11: Tests for systemic toxicity)
- Systemic Toxicity: Material Mediated Pyrogenicity (per ISO 10993-11, Biological evaluation of medical devices - Part 11: Tests for systemic toxicity, and ST72:2011/(R)2016, Bacterial endotoxins - Test methods routine monitoring and alternatives to batch testing)
- Genotoxicity (per ISO 10993-3, Biological evaluation of medical devices - Part 3: Tests for genotoxicity carcinogenicity and reproductive toxicity)
- Chemical Characterization (per ISO 10993-18, Biological evaluation of medical devices - Part 18: Chemical characterization of medical device materials within a risk management process)
- Toxicological Risk Assessment (per ISO 10993-17, Biological evaluation of medical devices - Part 17: Establishment of allowable limits for leachable substances)

The bacterial endotoxin test was performed to verify that the subject implants meet the 20 endotoxin units (EU)/device pyrogen limit specification. Testing was successfully performed, and it was confirmed that the subject devices meet the 20 EU/device testing limit for general medical devices that are implanted as outlined in ANSI/AAMI ST72: Bacterial endotoxins – Test methods, routine monitoring, and alternatives to batch testing and USP<161>, Medical Devices – Bacterial Endotoxin and Pyrogen Tests. Testing to monitor pyrogens will be performed periodically.

5.7.2. Mechanical Testing

Onkos Surgical has evaluated the subject devices to demonstrate substantial equivalence to the predicate devices. The following FDA Guidance documents were consulted to select the bench tests:

- *FDA Guidance*: “Guidance Document for Testing Orthopedic Implants with Modified Metallic Surfaces Apposing Bone or Bone Cement”, Issued 28-Apr-1994

The following bench testing was completed and the results of support the subject devices are equivalent to the predicate device:

- ASTM F1044 - Standard Test Method for Shear Testing of Calcium Phosphate Coatings and Metallic Coatings

Premarket Notification: ELEOS™ Limb Salvage System with BIOGRIP™

- ASTM F1147 - Standard Test Method for Tension Testing of Calcium Phosphate and Metallic Coatings
- ASTM F1160: Standard Test Method for Shear and Bending Fatigue Testing of Calcium Phosphate and Metallic Medical and Composite Calcium Phosphate/Metallic Coatings
- ASTM F1854 - Standard Test Method for Stereological Evaluation of Porous Coatings on Medical Implants
- ASTM F1978 - Standard Test Method for Measuring Abrasion Resistance of Metallic Thermal Spray Coatings by Using the Taber Abraser
- ASTM F2083, Standard Specification for Knee Replacement Prosthesis
- ISO13179-1:2014: Implants for surgery — Plasma sprayed unalloyed titanium coatings on metallic surgical implants

5.8 Clinical Data

Clinical data was not deemed necessary for the subject device.

5.9 Design Validation

Design Validation was performed for the subject implant using a cadaver lab. Design Validation demonstrated the subject implant and existing instruments function as intended and user needs were met.

5.10 Conclusions

Based on the test results and supporting documentation provided in this premarket notification, the subject ELEOS Limb Salvage System with BIOGRIP™ is substantially equivalent to the predicate device, ELEOS Limb Salvage System. The content in this premarket notification demonstrates that:

- any differences do not raise new questions of safety and effectiveness;
- and the proposed device is at least as safe and effective as the legally marketed predicate device.



Food and Drug Administration
10903 New Hampshire Avenue
Document Control Center - WO66-G609
Silver Spring, MD 20993-0002

October 28, 2016

Onkos Surgical, Inc.
Jan Triani
Sr. Director Quality Assurance and Regulatory Affairs
77 East Halsey Road
Parsippany, New Jersey 07054

Re: K161520

Trade/Device Name: ELEOS Limb Salvage System
Regulation Number: 21 CFR 888.3510
Regulation Name: Knee joint femorotibial metal/polymer constrained cemented prosthesis
Regulatory Class: Class II
Product Code: KRO, JDI, JWH, LPH, LZO
Dated: September 28, 2016
Received: September 30, 2016

Dear Jan Triani:

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. 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); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820); and if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

If you desire specific advice for your device on our labeling regulation (21 CFR Part 801), please contact the Division of Industry and Consumer Education at its toll-free number (800) 638-2041 or (301) 796-7100 or at its Internet address

<http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm>. 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 <http://www.fda.gov/MedicalDevices/Safety/ReportaProblem/default.htm> for the CDRH's Office of Surveillance and Biometrics/Division of Postmarket Surveillance.

You may obtain other general information on your responsibilities under the Act from the Division of Industry and Consumer Education at its toll-free number (800) 638-2041 or (301) 796-7100 or at its Internet address

<http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm>.

Sincerely yours,

Mark N. Melkerson -S

Mark N. Melkerson
Director
Division of Orthopedic Devices
Office of Device Evaluation
Center for Devices and
Radiological Health

Enclosure

Indications for Use

510(k) Number (if known)

K161520

Device Name

ELEOS LIMB SALVAGE SYSTEM

Indications for Use (Describe)

ELEOS™ Limb Salvage System Hip Components:

Indicated for use in total hip arthroplasty for reduction or relief of pain and/or improved hip function in skeletally mature patients with the following conditions:

- 1) Non-inflammatory degenerative joint disease such as osteoarthritis, avascular necrosis, ankylosis, protrusion acetabuli, and painful hip dysplasia;
- 2) Inflammatory degenerative joint disease such as rheumatoid arthritis;
- 3) Correction of functional deformity
- 4) Revision procedures where other treatments or devices have failed; and,
- 5) Treatment of fractures that are unmanageable using other techniques.

Limb salvage system is also indicated for procedures where radical resection and replacement of the proximal, distal and/or total femur is required with the following conditions:

- 1) Patients suffering from severe arthropathy of the hip that does not respond to any conservative therapy or better alternative surgical treatment;
- 2) Surgical intervention for severe trauma, revision hip arthroplasties, and/or Oncology indications.
- 3) Metastatic diseases

ELEOS™ Limb Salvage System Knee Components:

Indicated for cemented use in knee arthroplasty for reduction or relief of pain and/or improved knee function in skeletally mature patients with the following conditions:

- 1) Non-inflammatory degenerative joint disease including osteoarthritis, traumatic arthritis, or avascular necrosis;
- 2) Inflammatory degenerative joint disease including rheumatoid arthritis;
- 3) Correction of functional deformity;
- 4) Revision procedures where other treatments or devices have failed; and
- 5) Treatment of fractures that are unmanageable using other techniques.

Limb salvage system is also indicated for procedures where radical resection and replacement of the proximal tibia is required with the following conditions:

- 1) Patients suffering from severe arthropathy of the knee that does not respond to any conservative therapy or better alternative surgical treatment;
- 2) Surgical intervention for severe trauma, revision knee arthroplasties, and/or Oncology indications.
- 3) Metastatic diseases

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)

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Office of Chief Information Officer
Paperwork Reduction Act (PRA) Staff
PRASStaff@fda.hhs.gov

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K161520

Onkos Surgical, Inc.
ELEOS™ Limb Salvage Systems

October 27, 2016

5. 510(k) Summary

I. SUBMITTER

Onkos Surgical, Inc.
77 East Halsey Road
Parsippany, NJ 07054

Phone: (201)543-9388

Contact Person: Jan Triani
Email: jtriani@onkossurgical.com

Date Prepared: October 27, 2016

II. DEVICE

Name of Device:	ELEOS™ Limb Salvage System
Common Name:	Limb Salvage System
Classification Name:	21 CFR 888.3350, Prosthesis, Hip, Semi-Constrained, Metal/Polymer, Cemented 21 CFR 888.3510, Prosthesis, Knee, Femorotibial, Constrained, Metal Polymer, Cemented 21 CFR 888.3560, Knee joint patellofemorotibial polymer/metal/polymer semi-constrained cemented prosthesis. 21 CFR 888.3358 Prosthesis, Hip, Semi-Constrained, Metal/Polymer, Porous, Uncemented 21 CFR 888.3353, Prosthesis, Hip, Semi-Constrained, Metal/Ceramic/Polymer Cemented or Non-Porous, Uncemented
Regulatory Class:	II
Product Code:	JDI/KRO JWH LPH LZO

III. PREDICATE DEVICE

GUARDIAN® Limb Salvage System, K013035
ADVANCE® Modular Tibial Component, K973524
LINEAGE® Acetabular System, K002149
SLT Femoral Heads, K932222

No reference devices were used in this submission.

IV. DEVICE DESCRIPTION

The Onkos Surgical ELEOST™ Limb Salvage System consists of components that are used in the reconstruction of the lower limb. The reconstruction applications are proximal femur, distal femur, total femur, proximal tibia, and hinged knee. The ELEOS Limb Salvage components are femoral head, mid-section, stem, distal hinge femur, tibial hinge assembly, axial pin, tibial poly spacer, tibial sleeve, male-male mid-section, resurfacing hinge femur, and proximal tibia, patella, stem extension, tibial wedges and augments. Instrumentation is provided non-sterile in surgical trays which are to be re-processed per the validated instructions stated below.

Components	RECONSTRUCTION APPLICATIONS				
	Proximal Femur	Distal Femur	Total Femur	Proximal Tibia	Hinged Knee
Femoral head	✓		✓		
Mid-Section	✓	✓	✓	✓	
Segmental Stem	✓	✓		✓	
Distal Femur		✓	✓		
Tibial Hinge Component		✓	✓	✓	✓
Axial Pin		✓	✓	✓	✓
Tibial Poly Spacer		✓	✓	✓	✓
Tibial Baseplate		✓	✓		✓
Male-Male Mid-Section			✓		
Resurfacing Hinge Femur				✓	✓
Proximal Tibia				✓	
Patella		✓*	✓*	✓*	✓*
Wedges and Augments		✓*	✓*	✓*	✓*
Stem Extension		✓*	✓*	✓*	✓*

**These implants are optional for each procedure. The surgeon shall use his/her medical judgement to determine if these implants are necessary based on factor such as patient bone quality, joint stability and pathology.*

V. INDICATIONS FOR USE**ELEOST™ Limb Salvage System Hip Components:**

Indicated for use in total hip arthroplasty for reduction or relief of pain and/or improved hip function in skeletally mature patients with the following conditions:

- 1) Non-inflammatory degenerative joint disease such as osteoarthritis, avascular necrosis, ankylosis, protrusion acetabuli, and painful hip dysplasia;
- 2) Inflammatory degenerative joint disease such as rheumatoid arthritis;
- 3) Correction of functional deformity
- 4) Revision procedures where other treatments or devices have failed; and,
- 5) Treatment of fractures that are unmanageable using other techniques.

Limb salvage system is also indicated for procedures where radical resection and replacement of the proximal, distal and/or total femur is required with the following conditions:

K161520

Onkos Surgical, Inc.
ELEOS™ Limb Salvage Systems

October 27, 2016

- 1) Patients suffering from severe arthropathy of the hip that does not respond to any conservative therapy or better alternative surgical treatment;
- 2) Surgical intervention for severe trauma, revision hip arthroplasties, and/or Oncology indications.
- 3) Metastatic diseases

ELEOS™ Limb Salvage System Knee Components:

Indicated for cemented use in knee arthroplasty for reduction or relief of pain and/or improved knee function in skeletally mature patients with the following conditions:

- 1) Non-inflammatory degenerative joint disease including osteoarthritis, traumatic arthritis, or avascular necrosis;
- 2) Inflammatory degenerative joint disease including rheumatoid arthritis;
- 3) Correction of functional deformity;
- 4) Revision procedures where other treatments or devices have failed; and
- 5) Treatment of fractures that are unmanageable using other techniques.

Limb salvage system is also indicated for procedures where radical resection and replacement of the proximal tibia is required with the following conditions:

- 1) Patients suffering from severe arthropathy of the knee that does not respond to any conservative therapy or better alternative surgical treatment;
- 2) Surgical intervention for severe trauma, revision knee arthroplasties, and/or Oncology indications.
- 3) Metastatic diseases

VI. COMPARISON OF TECHNOLOGICAL CHARACTERISTICS WITH THE PREDICATE DEVICE

The predicate devices and the subject device are the same. The same materials, technology and sterilization methods are used to manufacture the subject device. The predicate device manufacturer is the contract manufacturer of the subject device. There are no technological differences between the two systems.

VII. PERFORMANCE DATA

No performance data is provided. A copy of the full 510(k) for the predicate device (K013035) is provided because there are no technological differences between the subject and predicate devices. Pyrogenicity was assessed using the LAL test which identified an acceptable endotoxin limit. Testing to monitor pyrogens will be performed using periodic testing.

VIII. CONCLUSIONS

Since the subject device is identical with respect to fit, form, function and manufacturing processes as the predicate device, the ELEOS™ Limb Salvage System has the same safety and effectiveness profile as the predicate device.