



January 21, 2022

b-ONE ORTHO, Corp.  
Allison Gecik  
Associate Director, Regulatory Affairs  
3 Wing Drive Suite #259  
Cedar Knolls, New Jersey 07927

Re: K213673

Trade/Device Name: MOBIO Total Knee System  
Regulation Number: 21 CFR 888.3560  
Regulation Name: Knee joint patellofemorotibial polymer/metal/polymer semi-constrained cemented prosthesis  
Regulatory Class: Class II  
Product Code: JWH, OIY  
Dated: November 19, 2021  
Received: November 22, 2021

Dear Allison Gecik:

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](mailto: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

DEPARTMENT OF HEALTH AND HUMAN SERVICES  
Food and Drug Administration

Form Approved: OMB No. 0910-0120  
Expiration Date: 06/30/2020  
See PRA Statement below.

### Indications for Use

510(k) Number (if known)

K213673

Device Name

MOBIO Total Knee System

Indications for Use (Describe)

The MOBIO Total Knee System is intended for total knee arthroplasty due to the following conditions:

- Painful, disabling joint disease of the knee resulting from: noninflammatory degenerative joint disease (including osteoarthritis, traumatic arthritis, or avascular necrosis), rheumatoid arthritis or post-traumatic arthritis
- Post-traumatic loss of knee joint configuration and function
- Moderate varus, valgus, or flexion deformity in which the ligamentous structures can be returned to adequate function and stability.
- Revision of previous unsuccessful knee replacement or other procedure
- Fracture of the distal femur and/or proximal tibia that cannot be stabilized by standard fracture-management techniques

Additional Indications for the PS and PS+:

- Ligamentous instability requiring implant bearing surface geometries with increased constraint
- Absent or nonfunctioning posterior cruciate ligament
- Severe anteroposterior instability of the knee joint

The MOBIO Total Knee System is intended for implantation with bone cement only.

MOBIO Total Knee System components are not intended for use with other knee systems.

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.

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

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**TRADITIONAL  
510(k) SUMMARY  
As required by 21 CFR 807.92**

**Submitter Information:**

Submitter's Name: b-ONE ORTHO, Corp.  
Address: 3 Wing Drive  
Suite 259  
Cedar Knolls, NJ 07927  
Telephone: 866-276-4538  
Contact Person: Allison Gecik  
agecik@b1.co  
Telephone: 973-965-8940

**Date Prepared:** November 10, 2021  
**Proprietary Name:** MOBIO Total Knee System  
**Classification:** Class II  
**Classification Panel:** Orthopedic  
**Common Name:** Total Knee Joint Replacement  
**Product Code(s):** JWH; OIY

Classification Name(s):	Regulation Number
Prosthesis, Knee, Patellofemorotibial, Semi-Constrained, Cemented, Polymer/Metal Polymer. Knee joint patellofemorotibial polymer/metal/polymer semi-constrained cemented prosthesis	888.3560
Prosthesis, knee, patellofemorotibial, semi-constrained, cemented, polymer + additive/metal/polymer + additive. Knee joint patellofemorotibial polymer/metal/polymer semi-constrained cemented prosthesis.	

**Legally Marketed Predicate Devices to Which Substantial Equivalence is Claimed:** b-ONE MOBIO Total Knee System CR Tibial Insert (K202429);  
b-ONE MOBIO Total Knee System CR PLUS Tibial Insert (K210483)

**Legally Marketed Reference Devices Used to Support** b-ONE MOBIO Total Knee System PS/PS+ Tibial Insert Components (K183025; K180446)

## **Substantial Equivalence:**

### **Indications for Use:**

The MOBIO Total Knee System is intended for total knee arthroplasty due to the following conditions:

- Painful, disabling joint disease of the knee resulting from: noninflammatory degenerative joint disease (including osteoarthritis, traumatic arthritis, or avascular necrosis), rheumatoid arthritis or post-traumatic arthritis
- Post-traumatic loss of knee joint configuration and function
- Moderate varus, valgus, or flexion deformity in which the ligamentous structures can be returned to adequate function and stability.
- Revision of previous unsuccessful knee replacement or other procedure
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Additional Indications for the PS and PS+:

- Ligamentous instability requiring implant bearing surface geometries with increased constraint
- Absent or nonfunctioning posterior cruciate ligament
- Severe anteroposterior instability of the knee joint

The MOBIO Total Knee System is intended for implantation with bone cement only.

MOBIO Total Knee System components are not intended for use with other knee systems.

### **Device Description/Technological Characteristics:**

The MOBIO Total Knee System is a modular artificial knee replacement system comprised of symmetric cemented femoral components, symmetric cemented tibial tray, symmetric tibial inserts with locking wires, symmetric patellar resurfacing button, and reusable surgical instruments. The therapeutic effect is replacement of the diseased joint with artificial components to restore joint function. The purpose of this submission is to introduce a line extension to the b-ONE™ MOBIO™ Total Knee System CR/CR PLUS Tibial Insert components and PS/PS PLUS Tibial Insert Components. Compatibility with the b-ONE™ MOBIO™ Total Knee System tibial baseplate and femoral components remains the same. There is no allowed interchangeability with systems manufactured by other companies.

The subject b-ONE™ MOBIO™ Tibial Inserts are made from UHMWPE GUR® 1020-E Crosslinked with .1% tocopherol and conventional UHMWPE GUR® 1020. This material is the same as that used in the reference and predicate devices. The pre-assembled locking wire is the same locking wire used in the existing Tibial Inserts and is manufactured from Cobalt Chrome conforming to ASTM F90.

All system components are supplied sterile and are single use devices.

### **Comparison of Technological Characteristics**

The design features and materials of the subject devices are substantially equivalent to those of the predicate devices. The MOBIO Total Knee System and the predicate devices share the following characteristics:

- Materials of construction

- Manufacturing processes
- Sizes offered
- Product design for shape and macrostructures- only differences to the predicate are dimensions specific to the condylar stabilizing features of the subject devices.
- Sterilization methods

### **Performance Data**

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

#### Non-Clinical Studies

- Locking Mechanism Strength
- Range of Constraint
- Contact Area and Contact Stress
- Tibiofemoral Range of Motion
- Characterization of UHMWPE Insert Material
- Bacterial Endotoxin Testing
- Shelf Life Studies
- Biocompatibility
- Wear Testing

### **Conclusion**

The information provided above supports that the MOBIO Total Knee System is as safe and effective as the predicate devices with the same intended use. The subject and predicate devices only differ in material. However, applicable reference devices have been cited to support the conclusion that these differences do not raise any new questions of safety and effectiveness. The MOBIO Total Knee System is substantially equivalent to the predicate devices.