

August 1, 2023

Blue Belt Technologies, Inc. Corrine Herlinger Senior Principal Regulatory Affairs Specialist 2875 Railroad Street Pittsburgh, Pennsylvania 15222

Re: K231963

Trade/Device Name: REAL INTELLIGENCE<sup>TM</sup> CORI<sup>TM</sup> Regulation Number: 21 CFR 882.4560 Regulation Name: Stereotaxic Instrument Regulatory Class: Class II Product Code: OLO Dated: June 30, 2023 Received: July 3, 2023

Dear Corrine Herlinger:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. Although this letter refers to your product as a device, please be aware that some cleared products may instead be combination products. The 510(k) Premarket Notification Database located at <a href="https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfpmn/pmn.cfm">https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfpmn/pmn.cfm</a> identifies combination product submissions. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration. Please note: CDRH does not evaluate information related to contract liability warranties. We remind you, however, that device labeling must be truthful and not misleading.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the 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 <a href="https://www.fda.gov/combination-products/guidance-regulatory-information/postmarketing-safety-reporting-combination-products">https://www.fda.gov/combination-products/guidance-regulatory-information/postmarketing-safety-reporting-combination-products</a>); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820) for devices or current good manufacturing practices (21 CFR 4, Subpart A) for combination products; and, if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21 CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to <u>https://www.fda.gov/medical-devices/medical-device-safety/medical-device-reporting-mdr-how-report-medical-device-problems</u>.

For comprehensive regulatory information about medical devices and radiation-emitting products, including information about labeling regulations, please see Device Advice (<u>https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance</u>) and CDRH Learn (<u>https://www.fda.gov/training-and-continuing-education/cdrh-learn</u>). Additionally, you may contact the Division of Industry and Consumer Education (DICE) to ask a question about a specific regulatory topic. See the DICE website (<u>https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/contact-us-division-industry-and-consumer-education-dice</u>) for more information or contact DICE by email (<u>DICE@fda.hhs.gov</u>) or phone (1-800-638-2041 or 301-796-7100).

Sincerely,



For

Shumaya Ali, M.P.H.
Assistant Director
DHT6C: Division of Restorative, Repair and Trauma Devices
OHT6: Office of Orthopedic Devices
Office of Product Evaluation and Quality
Center for Devices and Radiological Health

Enclosure

#### Indications for Use

Submission Number (if known)

K231292

Device Name

REAL INTELLIGENCE™ CORI™

Indications for Use (Describe)

CORI is indicated for use in surgical procedures, in which the use of stereotactic surgery may be appropriate, and where reference to rigid anatomical bony structures can be determined. These procedures include:

- unicondylar knee replacement (UKR),
- total knee arthroplasty (TKA),
- · revision knee arthroplasty, and
- total hip arthroplasty (THA).

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

Prescription Use (Part 21 CFR 801 Subpart D)

Over-The-Counter Use (21 CFR 801 Subpart C)

#### CONTINUE ON A SEPARATE PAGE IF NEEDED.

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# 510(k) Summary

510(k) Owner	Blue Belt Technologies, Inc. 2875 Railroad Street Pittsburgh, PA 15222 USA Tel: (412) 683-3844 Fax: (412) 683-6447
Contact Person	Corrine Herlinger Senior Principal Regulatory Affairs Specialist Tel: 412.552.6428 Email: corrine.herlinger@smith-nephew.com
Date of Submission	30-June-2023
Classification Reference	21 CFR 882.4560
Classification Name	Stereotaxic instrument
Product Code	OLO
Supported Codes	HSX, JWH, MBH, NJD
Common/Usual Name	Orthopedic Stereotaxic Instrument
Trade/Proprietary Name	REAL INTELLIGENCE™ CORI™ (CORI)
Predicate Device(s)	REAL INTELLIGENCE™ CORI™ (K221224)
Reason for Submission	This Special 510(k) submission supports an update to CORI to allow the system to be used with porous Unicondylar Knee Replacement (UKR) Implants

### **Intended Use**

Real Intelligence CORI (CORI) is intended to assist the surgeon in providing software-defined spatial boundaries for orientation and reference information to anatomical structures during orthopedic procedures.

### **Indications for Use**

CORI is indicated for use in surgical procedures, in which the use of stereotactic surgery may be appropriate, and where reference to rigid anatomical bony structures can be determined. These procedures include:

- unicondylar knee replacement (UKR),
- total knee arthroplasty (TKA),
- revision knee arthroplasty, and
- total hip arthroplasty (THA).

### **Device Description**

The subject of this Special 510(k) is REAL INTELLIGENCE CORI (CORI), a robotic-assisted orthopedic surgical navigation and burring system. CORI uses established technologies of navigation via a passive infrared tracking camera. Based on intraoperatively-defined bone landmarks and known geometry of the surgical implant, the system aids the surgeon in establishing a bone surface model for the target surgery and planning the surgical implant location. For knee applications, CORI then aids the surgeon in executing the surgical plan by controlling the cutting engagement of the surgical bur.

CORI knee application software controls the cutting engagement of the surgical bur based on its proximity to the planned target surface. The cutting control is achieved with two modes:

- **Exposure control** adjusts the bur's exposure with respect to a guard. If the surgeon encroaches on a portion of bone that is not to be cut, the robotic system retracts the bur inside the guard, disabling cutting.
- **Speed control** regulates the signal going to the tool control unit itself and limits the speed of the drill if the target surface is approached.

Alternatively, the surgeon can disable both controls and operate the robotic drill as a standard navigated surgical drill.

# **Currently Supported Knee Implants**

The following Smith+Nephew knee implants are supported on CORI:

 Table 1: Currently Supported Smith+ Nephew Knee Implants

Implant Model Name	510(k) Number	Classification Product Code
STRIDE Unicondylar Knee	K123380	HSX
ZUK Select Knee System	K160738	HSX
JOURNEY II Unicompartmental Knee System	K191211	HSX
JOURNEY UNI	K102069	HSX
JOURNEY II CR	K121443	JWH
JOURNEY II BCS	K111711	JWH
JOURNEY II XR	К141471, К152726	JWH
LEGION CR/PS	K951987, K962557, K093746	JWH
LEGION Porous CR Femoral Components	К073325, К091543	MBH
LEGION Porous CR Narrow Femoral Components	K210566	MBH
LEGION Porous Tibia	K100897	MBH
Porous Tibia Baseplate	K211221	MBH
GENESIS II CR/PS	К951987, К962557	JWH
ANTHEM	K142807	JWH
SMITH & NEPHEW, INC. REVISION KNEE SYSTEM	K043440	JWH
REVISION KNEE SYSTEM	K041106	JWH
LEGION RK TIBIAL WEDGES (Hemi-Step & Full-Step)	K953274	JWH
LEGION COBALT CHROME REVISION KNEE SYSTEM	K060742	JWH
LEGION Knee System	K180334	JWH, MBH
ENGAGE Partial Knee System	K190439	NJD, HSX
JOURNEY II UK <sup>™</sup> and ENGAGE <sup>™</sup> Cementless Partial Knee System	K222653	NJD, HSX

### **Discussion of Similarities and Differences**

This Special 510(k) submission supports an update to allow for porous UKR implants on Real Intelligence CORI for the following implant combinations:

- ENGAGE<sup>™</sup> Partial Knee System consisting of the ENGAGE Femur and Tibia, intended for cemented or cementless implantation cleared per K190439
- JOURNEY II UK<sup>™</sup> and ENGAGE<sup>™</sup> Cementless Partial Knee System consisting of the Journey II Unicompartmental Cemented Femur and Engage Cemented or Cementless Tibia cleared per K222653

The modifications made to support the change do not impact the system's intended use, indications for use, or fundamental scientific technology.

Blue Belt Technologies believes that CORI is subject to premarket notification requirements under Section 510(k) of the Federal Food, Drug and Cosmetic Act ("FFDCA" or "the Act") and is substantially equivalent to the previously cleared REAL INTELLIGENCE CORI (K221224).

#### Table 1: Predicate Device

Manufacturer	Description	Submission Number	Clearance Date
Blue Belt Technologies, Inc.	REAL INTELLIGENCE CORI	K221224	August 19, 2022

#### Table 2: Summary of Technological Similarities with Predicate

Devices	Subject Device CORI	Primary Predicate CORI – K221224
Intended use	Same as predicate.	REAL INTELLIGENCE CORI (CORI) is intended to assist the surgeon in providing software-defined spatial boundaries for orientation and reference information to anatomical structures during orthopedic procedures.
Indications for Use	Same as predicate.	CORI is indicated for use in surgical procedures, in which the use of stereotactic surgery may be appropriate, and where reference to rigid anatomical bony structures can be determined. These procedures include: • unicondylar knee replacement (UKR), • total knee arthroplasty (TKA), • revision knee arthroplasty, and • total hip arthroplasty (THA).

Devices	Subject Device CORI	Primary Predicate CORI – K221224
Knee Implant Product Codes Supported	HSX, JWH, MBH, NJD	HSX, JWH, МВН
Environment of Use	Same as predicate.	CORI is intended to be used by trained medical professionals in a hospital or clinical setting equivalent to an orthopedic surgery suite.
Technological Characteristics	Same as predicate.	For knee applications, CORI uses established technologies to prepare bone for attachment of UKR and TKA implant components. In the case of a total knee arthroplasty, the bone surface may also be prepared to receive the femoral and tibial cutting guides.
		CORI uses intraoperative data collection (image- free or non-CT data generation) to create a model of the patient's femur and/or tibia, dependent on the procedure being performed, and allows the surgeon to prepare a surgical plan.
		The system uses predefined boundaries generated during the planning process to control the motion of the surgical bur and limit the amount of bone removed in order to shape the condyles or tibial plateau in preparation for placement of the surgical implant. Bur cutting is controlled either by retracting the bur in a guard, or by controlling the speed of the bur as the target surface is approached.
		To support the hip application, CORI uses a Virtual Machine with hypervisor to enable the Windows-based HIP7 software to run on CORI.

# **Non-Clinical Testing (Bench)**

Verification and validation testing demonstrated the safety and efficacy of CORI when the system is used to place porous UKR implants intended for use with or without bone cement. Summative usability testing (including labeling validation) demonstrated that participating surgeons were able use the subject device safely and effectively in a simulated use environment.

Blue Belt Technologies has concluded that all design inputs have been met and that the verification and validation testing performed did not raise any new questions of safety or effectiveness.

## Conclusions

The subject device, CORI, described in this submission has the same intended use, indications for use, and the same technological characteristics as the predicate device, CORI (K221224). The difference between the two systems is an update to the CORI User Manual and an update to the Implant Database to include the ENGAGE<sup>™</sup> Partial Knee System and the JOURNEY II UK<sup>™</sup> and ENGAGE<sup>™</sup> Cementless Partial Knee System.

Comprehensive verification testing demonstrated that the system meets required design inputs. Summative usability testing results demonstrate that representative users can use the subject device safely and effectively in a simulated use environment. The information presented in this 510(k) premarket notification demonstrates that CORI may be used for the placement of porous UKR implants, and that CORI is as safe and effective as the predicate CORI system (K221224). Blue Belt Technologies believes that FDA can find CORI to be substantially equivalent to the predicate device.