



Aibolit Technologies, LLC
Howard Schrayer
Official Correspondent
9616 Moritz Way
Delray Beach, Florida 33446

January 7, 2022

Re: K211443

Trade/Device Name: Aibolit 3D+
Regulation Number: 21 CFR 892.2050
Regulation Name: Medical Image Management And Processing System
Regulatory Class: Class II
Product Code: QIH, LLZ
Dated: November 30, 2021
Received: December 2, 2021

Dear Howard Schrayer:

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

Thalia T. Mills, Ph.D.
Director
Division of Radiological Health
OHT7: Office of In Vitro Diagnostics
and Radiological Health
Office of Product Evaluation and Quality
Center for Devices and Radiological Health

Enclosure

INDICATIONS FOR USE

DEPARTMENT OF HEALTH AND HUMAN SERVICES Food and Drug Administration Indications for Use	Form Approved: OMB No. 0910-0120 Expiration Date: 06/30/2020 See PRA Statement below.
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510(k) Number (if known)

TBD **K21143**

Device Name

AIBOLIT 3D+

Indications for Use (Describe)

Aibolit 3D+ is intended as a medical imaging system that allows the processing, review, analysis, communication and media interchange of multi-dimensional digital images acquired from CT imaging devices. Aibolit 3D+ is intended as software for preoperative surgical planning, training, patient information and as software for the intraoperative display of the multidimensional digital images. Aibolit 3D+ is designed for use by health care professionals and is intended to assist the clinician who is responsible for making all final patient management decisions.

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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**Exhibit E - 510(k) Summary
(Per 21 CFR 807.92)**

Contact: Howard Schroyer
Albolit Technologies, LLC
9616 Moritz Way
Delray Beach, FL 33446

Telephone: 609-273-7350
hs.ss@lucidmedical.net

Date Prepared: January 7, 2022

Device Trade Name: AIBOLIT 3D+

Manufacturer: Albolit Technologies, LLC
9616 Moritz Way
Delray Beach, FL 33446

Common Name: Automated Radiological Image Processing Software
Medical image management and processing system

Classification: Class II

Product Code: QIH - LLZ

Regulation: 21 CFR 892.2050

Predicate Devices:

Primary Predicate
Ceevra, Inc.
Ceevra Reveal 2.0
Image Processing System
[510(k) K173274]

Reference Predicate
Intuitive Surgical, Inc.
IRIS 1.0 System
[510(k) K182643]

Indications for Use:

Aibolit 3D+ is intended as a medical imaging system that allows the processing, review, analysis, communication and media interchange of multi-dimensional digital images acquired from CT imaging devices. Aibolit 3D+ is intended as software for preoperative surgical planning, training, patient information and as software for the intraoperative display of the multidimensional digital images. Aibolit 3D+ is designed for use by health care professionals and is intended to assist the clinician who is responsible for making all final patient management decisions.

Device Description:

Aibolit 3D+ is a web-based stand-alone application that can be presented on a computer connected to the internet. Once the enhanced images are created, they can be used by the physician for case review, patient education, professional training and intraoperative reference.

Aibolit 3D+ is a software only device, which processes CT images from a patient to create 3-dimensional images that may be manipulated to view the anatomy from virtually any perspective. The software also allows for transparent viewing of anatomical structures artifacts inside organs such as ducts, vessels, lesions and entrapped calcifications (stones). Anatomical structures are identified by name and differential coloration to highlight them within the region of interest.

The software may help to facilitate the surgeon's decision-making during planning, review and conduct of surgical procedures and, hence, may potentially help them to decrease or prevent possible errors caused by the misidentification of anatomical structures and their positional relationship.

Substantial Equivalence and Predicate Devices:

The device was shown to be substantially equivalent to a previously cleared video image processing system, the Ceevra Reveal 2.0 Image Processing System [510(k) K173274].

Predicate Comparison Table

Manufacturer	Aibolit Technologies, LLC	Ceevra
Trade Name	AIBOLIT 3D+ Image Processing System	Ceevra Reveal 2.0 Image Processing System
510(k) Number	Subject Device - TBD	K173274
Type of Device/ Product Code /	Radiological Image Processing System / QIH - LLZ	Radiological Image Processing System / LLZ
Regulation / Class	21 CFR 892.2050 – Class II	21 CFR 893.2050 – Class II
Indications for Use	Aibolit 3D+ is intended as a medical imaging system that allows the processing, review, analysis, communication and media interchange of multi-dimensional digital images acquired from CT imaging devices. It is also intended as software for preoperative surgical planning, training, patient information and as software for the intraoperative display of the multidimensional digital images. Aibolit 3D+ is designed for use by health care professionals and is intended to assist the clinician who is responsible for making all final patient management decisions.	Ceevra Reveal 2.0 is intended as a medical imaging system that allows the processing, review, analysis, communication and media interchange of multi-dimensional digital images acquired from CT or MR imaging devices. It is also intended as software for preoperative surgical planning, and as software for the intraoperative display of the aforementioned multidimensional digital images. Ceevra Reveal 2.0 is designed for use by health care professionals and is intended to assist the clinician who is responsible for making all final patient management decisions.
Mechanism of Action	Capture and enhancement of (DICOM) digital video images via software-based conversion to 2-D and 3-D anatomical structure images that can be manipulated for viewing	Capture and enhancement of (DICOM) digital video images via software-based conversion to 2-D and 3-D anatomical structure images that can be manipulated for viewing
Intended Users	Health care professionals	Health care professionals
Intended Use Environment	Healthcare facilities such as hospitals and clinics	Healthcare facilities such as hospitals and clinics

Format of Captured Images	DICOM	DICOM
Intended Use	AIBOLIT 3D+ is intended for use as a medical imaging system that allows the processing, review, analysis, communication and media interchange of multi-dimensional digital images acquired from CT imaging devices. It is also intended as software for preoperative surgical planning, and as software for the intraoperative display of multi-dimensional digital images. AIBOLIT 3D+ is designed for use by health care professionals and is intended to assist the clinician who is responsible for making patient management decisions.	Intended as a medical imaging system that allows the processing, review, analysis, communication and media interchange of multi-dimensional digital images acquired from CT or MR imaging devices. It is also intended as software for preoperative surgical planning, and as software for the intraoperative display of the aforementioned multi-dimensional digital images. Ceevra Reveal 2.0 is designed for use by health care professionals and is intended to assist the clinician who is responsible for making all final patient management decisions.
Security	Data coded and HIPAA compliant	Data coded and HIPAA compliant
Form of Device	AIBOLIT 3D+ is a software only device that permits electronic image uploads, provides image conversion and allows viewing on a mobile device or standard computer monitor.	The Ceevra Reveal 2.0 Video Processor is a software only device that permits electronic image uploads, provides image conversion and allows viewing on a mobile device or standard computer monitor.
Image processing	High-definition digital images up to 4K	High-definition digital images
Functions	Generation of 2D and 3D images from DICOM data Organ segmentation and structure identification Dimensional and volume references Multi-axis image rotation Organ transparency Organ retraction animation	Generation of 2D and 3D images from DICOM data Organ segmentation and structure identification Dimensional and volume references Multi-axis image rotation
Body contact	None	None

<p>User Interface and System Work-Flow</p>	<p>Physician uploads DICOM images and specifies desired anatomical segments of interest</p> <p>Radiologist annotates sample (segments) images</p> <p>AI software facilitates annotation of available images under guidance and control by the Radiologist</p> <p>Imaging technician generates multi-axis rotatable image and retraction model</p> <p>Radiologist reviews images generated by imaging technician and returns output file to requesting physician</p>	<p>Physician uploads DICOM images and specifies desired anatomical segments of interest</p> <p>Imaging technician annotates sample (segments) images</p> <p>Imaging technician generates multi-axis rotatable image and returns output file to requesting physician</p>
<p>External / Internet Connections</p>	<p>Web-based software</p>	<p>Web-based software</p>
<p>CT Image Uploading</p>	<p>By requesting physician</p>	<p>By requesting physician</p>
<p>Other User Inputs</p>	<p>List of organ structures to be annotated and displayed, patient ID and demographics</p>	<p>List of organ structures to be annotated and displayed, patient ID and demographics</p>
<p>Image Segmentation</p>	<p>By Radiologist (MD) – Manual annotation is done for all CT slices with optional use of AI/ML algorithms as determined by Radiologist and with Radiologist’s approval</p>	<p>By Imaging Technician – Manual annotation done for all CT slices – No software used for annotation</p>
<p>Organ identification</p>	<p>By Radiologist</p>	<p>Unknown proprietary method used to identify organ structures</p>
<p>3D Image generation</p>	<p>3D image file generated by 3rd party software (3D Slicer) following Radiologist review and approval of annotation</p>	<p>3D image file generated by 3rd party software</p>

Organ structure identification	Proprietary software assigns color coding to each structure identified by Radiologist and displays color-coded image with labeled key to color/structure identity	Proprietary software assigns color coding to each structure identified by imaging technician and displays color-coded image with key to color/structure identity
Image editing permission	Only the radiologist can alter or edit images following review – User physicians cannot edit images - Physicians have option to show or hide organs on display	Imaging technician can edit images generated by the system software – User physicians cannot edit images - Physicians have option to show or hide organs on display
Device Output Devices	3D image can be displayed on standard monitor	3D image can be displayed on standard monitor, smart phone (with separate software) or Virtual Imaging 3D headset
Supplemental outputs	Organ structure dimensions, volume, organ labels, patient ID, CT date and demographics	Organ structure dimensions, volume, organ labels, patient ID and demographics
Output image manipulation by user	Physician user can show or hide individual organ structures, zoom capability, rotational capability and transparency capability	Physician user can show or hide individual organ structures, zoom capability, rotational capability, transparency capability (current version)

Performance Testing:

The following documentation was submitted in the 510(k).

- Hardware Requirements
- Level of Concern Statement
- Software Description
- Architecture Design
- User Manual and Instructions for Use
- Software Design Specification
- Risk Analysis
- Traceability Analysis
- Software Validation Report
- Usability Evaluation
- Software Development Lifecycle
- Unresolved Anomalies
- Cybersecurity

Conclusion

AIBOLIT 3D+ is substantially equivalent to the previously cleared Ceevra Reveal 2.0 Image Processing System with respect to intended use, principle of operation, general technological characteristics and performance.