



Siemens Medical Solutions USA, Inc.
% Cordell Fields, Esq.
Sr. Regulatory Affairs Specialist
40 Liberty Boulevard 65-1A
MALVERN PA 19355

March 5, 2021

Re: K210309

Trade/Device Name: Cios Select (VA21) Flat Panel
Regulation Number: 21 CFR 892.1650
Regulation Name: Image-intensified fluoroscopic x-ray system
Regulatory Class: Class II
Product Code: OWB, OXO, JAA
Dated: January 27, 2021
Received: February 3, 2021

Dear Cordell Fields:

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

DEPARTMENT OF HEALTH AND HUMAN SERVICES Food and Drug Administration <p style="text-align: center;">Indications for Use</p>	Form Approved: OMB No. 0910-0120 Expiration Date: 06/30/2023 See <i>PRA Statement on last page.</i>
510(k) Number (if known) K210309	
Device Name <p style="text-align: center;">Cios Select (VA21) Flat Panel</p>	
Indications for Use (Describe) <p>The Cios Select is a mobile X-ray system intended for use in Operating room, Traumatology, Endoscopy, Intensive Care Station, Pediatrics, Ambulatory patient care and in Veterinary Medicine.</p> <p>The Cios Select can operate in three different modes, Digital Radiography, Fluoroscopy, and Pulsed Fluoroscopy which are necessary in performing wide variety of clinical procedures, such as intraoperative bile duct display, fluoroscopic display of an intra-medullary nail implants in various positions, low dose fluoroscopy in pediatrics, fluoroscopic techniques utilized in pain therapy and positioning of catheters and probes.</p>	
Type of Use (Select one or both, as applicable) <p style="text-align: center;"> <input checked="" type="checkbox"/> Prescription Use (Part 21 CFR 801 Subpart D) <input type="checkbox"/> Over-The-Counter Use (21 CFR 801 Subpart C) </p>	
<p>CONTINUE ON A SEPARATE PAGE IF NEEDED.</p>	
<p style="text-align: center;">This section applies only to requirements of the Paperwork Reduction Act of 1995.</p> <p style="text-align: center;">*DO NOT SEND YOUR COMPLETED FORM TO THE PRA STAFF EMAIL ADDRESS BELOW.*</p> <p>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:</p> <p style="text-align: center;"> Department of Health and Human Services Food and Drug Administration Office of Chief Information Officer Paperwork Reduction Act (PRA) Staff PRAStaff@fda.hhs.gov </p> <p style="text-align: center;"><i>“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.”</i></p>	
<p>FORM FDA 3881 (6/20)</p>	<p style="text-align: center;">Page 1 of 1</p> <p style="text-align: right;"><small>PSC Publishing Services(301)443-6740 EF</small></p>

510(k) Summary: Cios Select

Company: Siemens Medical Solutions USA, Inc.
40 Liberty Boulevard, 65-1A
Malvern, PA 19355

Date Prepared: January 27, 2021

This 510(k) summary of safety and effectiveness information is being submitted in accordance with the requirements of SMDA 1990 and 21 CFR § 807.92.

1. General Information:

Importer / Distributor:

Siemens Medical Systems USA, Inc.
40 Liberty Boulevard, 65-1A
Malvern, PA 19355

Establishment Registration Number: 2240869

Manufacturing Site:

Siemens Shanghai Medical Equipment Ltd.
278 Zhou Zhu Road, Shanghai
201318, China

Establishment Registration Number: 3003202425

2. Contact Person:

Cordell Fields
Technical Specialist, Regulatory Submissions
Siemens Medical Solutions USA, Inc.
40 Liberty Boulevard, 65-1A
Malvern, PA 19355
Phone: (610) 306-3167
Email: cordell.fields@siemens – healthineers.com

3. Device Name and Classification:

Trade Name:	Cios Select
Classification Name:	Image-Intensified Fluoroscopic x-ray System
Classification Panel:	Radiology
Regulation Number:	21 CFR §892.1650
Device Class:	Class II
Product Codes:	OWB, OXO, JAA

4. Legally Marketed Primary Predicate Device

Trade Name:	Cios Select
510(k) Clearance	K181767
Clearance Date	August 17, 2018
Classification Name:	Image-intensified fluoroscopic x-ray System
Classification Panel:	Radiology

Regulation Number: 21 CFR §892.1650
Device Class: Class II
Product Code: OWB
Subsequent Product Codes: JAA, OXO
Total Product Life Cycle: There are no Recalls nor MDR incidents for this cleared device.

Legally Marketed Secondary Predicate Device

Trade Name: **Cios Spin**
510(k) Clearance: K181550
Clearance Date: October 30, 2018
Classification Name: Image-intensified fluoroscopic x-ray System
Classification Panel: Radiology
Regulation Number: 21 CFR §892.1650
Device Class: Class II
Product Codes: OWB
Subsequent Product Codes: JAA, OXO
Total Product Life Cycle: All product Recall incidents are considered during the Design Input phase of development to ensure the latest models will not be affected by any of the applicable issues.

Legally Marketed Predicate Device

Trade Name: **Cios Flow**
510(k) Clearance: K203504
Clearance Date: December 22, 2020
Classification Name: Image-intensified fluoroscopic x-ray System
Classification Panel: Radiology
Regulation Number: 21 CFR §892.1650
Device Class: Class II
Product Code: OWB
Subsequent Product Codes: JAA, OXO
Total Product Life Cycle: There are no Recalls nor MDR incidents for this cleared device.

1. Device Description:

The Cios Select (VA21) Mobile X-ray System is designed for the surgical environment. The Cios Select (VA21) is a modification of the Cios Select originally cleared under Premarket Notification K181767 on August 17, 2018.

The Cios Select consists of two major units:

The Siemens Healthineers Cios Select mobile fluoroscopy C-arm system is an X-ray imaging system consisting of two mobile units: a mobile acquisition unit and a monitor cart as the image display station.

The mobile acquisition unit is comprised of the X-ray control, the C-arm which supports the single-tank high-frequency generator/X-ray tube assembly, the flat panel detector, and user controls.

The monitor cart connects to the acquisition unit by a cable. It integrates the TFT flat panel displays, Digital Imaging Processing System, user controls and image storage devices (DVD, USB).

The following modifications were made to the predicate device the Cios Select Mobile X-ray System cleared under Premarket Notification K181767 on August 17, 2018. Siemens Medical Solutions USA, Inc. submits this Special 510(k) to request clearance for the Subject Device the Cios Select (VA21) in comparison to the Predicate Device.

- 1.) Upgraded software to version VA21.
- 2.) New optional Cios OpenApps
- 3.) New optional Target pointer.
- 4.) New optional tube side green laser aimer
- 5.) New optional flat detector side green laser aimer
- 6.) Enhanced Cybersecurity

2. Indications for Use:

The Cios Select is a mobile X-ray system intended for use in Operating room, Traumatology, Endoscopy, Intensive Care Station, Pediatrics, Ambulatory patient care and in Veterinary Medicine.

The Cios Select can operate in three different modes, Digital Radiography, Fluoroscopy, and Pulsed Fluoroscopy which are necessary in performing wide variety of clinical procedures, such as intraoperative bile duct display, fluoroscopic display of an intra-medullary nail implants in various positions, low dose fluoroscopy in pediatrics, fluoroscopic techniques utilized in pain therapy and positioning of catheters and probes.

3. Substantial Equivalence:

The Cios Select (VA21) system is substantial equivalent to the legally marketed predicates listed in **Table 1** below:

Table 1: Predicate Device Comparable Properties for Subject Device Modifications:

Predicate Device Name and Manufacturer	510(k) Number	Clearance Date	Comparable Properties
Primary Predicate Cios Select (VA20) w/Flat Panel Detector Siemens Shanghai Medical Equipment Ltd.	K181767	8/17/2018	<ul style="list-style-type: none"> • Indications for use • X-ray technology • Image processing • Mechanical design • Cybersecurity • Software
Predicate Device	K181550	10/30/2018	<ul style="list-style-type: none"> • Tube Side Laser

Predicate Device Name and Manufacturer	510(k) Number	Clearance Date	Comparable Properties
Cios Spin (VA30) SIEMENS AG Sector Healthcare			<ul style="list-style-type: none"> FD Side Laser
Predicate Device	K203504	12/22/20	<ul style="list-style-type: none"> Target Pointer Cios OpenApps
Cios Flow (VA30) SIEMENS AG Sector Healthcare			

4. Summary of Technological Characteristics of the Subject Device as Compared with the Predicate Device:

The Cios Select (VA21) System is designed as a set of components (floor stand, C-arm, X-ray tube and housing, flat panel detector, digital imaging system, collimator, generator etc.) that is combined to provide a specialized angiography system. Components used with Cios Select (VA21) System are either commercially available with current Siemens systems or include modifications to existing components. Technological differences between the Subject Device and the Predicate Device is provided in the **Table 2** below for all modifications.

Table 2: Summary of Comparison of Technological Characteristics

Subject Device Cios Select (VA21) System Modifications		Primary Predicate Device Cios Select VA20 (K181767)	Comparison Results
1.	Updated System Software to VA21 with modifications/features	Secondary Predicate Device Cios Flow (VA30) (K203504)	Same: The system software was upgraded to support new software features (1-3): Upgraded software VA21 with Cios Open Apps and Target Pointer. There are no technological differences in these features and the functionality of these features have not changed except for better visualization of moving k-wires. These features are the exact same features cleared in the Secondary Predicate Device Cios Flow (VA30) K1203504 . Software Testing was conducted per Software Guidance.
1.A	New Optional Cios OpenApps	Optional Cios OpenApps	
1.B	New optional Target pointer.	New optional Target pointer.	
1.C	Enhanced Cybersecurity	Primary Predicate Device Cios Select (VA20) (K181767) Enhanced Cybersecurity	Same: Cybersecurity features are the same features cleared in Cios Select VA20 (K181767)
2.	New Optional tube side green laser aimer	Secondary Predicate Device Cios Spin (VA30) (K181550) Optional tube side green laser aimer	Same: The green laser aimers on the side of the tube and flat detector is the same exact laser aimer cleared in the Secondary Predicate
3.	New Optional flat detector side green laser aimer	Optional flat detector side green laser aimer	

			Device Cios Spin (VA30) K181550.
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5. Nonclinical Performance Testing:

Non-clinical tests were conducted for the Cios Select (VA21) during product development. The Siemens Cios Select has been tested to meet the requirements for conformity to multiple industry standards. Performance testing confirmed, that the Siemens Cios Select complies with the following 21 CFR Federal Performance Standards.

Code of Federal Regulations Title 21 Subchapter J- Radiological Health, applicable sections include:

- 1020.30 Diagnostic X-Ray Systems and their major components
- 1020.31 - Radiographic equipment
- 1020.32 Fluoroscopic Equipment
- 1040.10 Laser products

The Cios Select (VA 21) was certified by Siemens Healthcare GmbH Corporate Testing Laboratory to comply with the following standards for Electrical safety, performance and Electromagnetic Compatibility:

- AAMI ANSI ES60601-1:2005/(R)2012
- IEC 60601-1-2:2014
- IEC 60601-1-3:2013
- IEC 60601-1-6:2010/A1:2013
- IEC 60825-1:2007
- TR 60878:2015
- IEC 62304:2015
- IEC 80001-1:2010
- IEC 60601-2-28:2017
- IEC 60601-2-43:2017
- IEC 60601-2-54:2009/A1:2015
- ISO 10993-1:2009
- ISO 14971:2007

Table 3: FDA Guidance Documents

FDA Guidance Document and Effective Date	
1.	Guidance for Industry and FDA Staff – User Fees and Refunds for Premarket Notification Submissions 510(k) Document issued on October 2, 2017
2.	Guidance for Industry and Food and Drug Administration Staff: Refuse to Accept Policy for 510(k)s Document issued on September 13, 2019
3.	Guidance for Industry and FDA Staff: The Special 510(k) Program Document issued on September 13, 2019

FDA Guidance Document and Effective Date	
4.	Guidance for Industry and FDA Staff: Deciding when to submit a 510(k) for a change to an existing device. Document issued on October 25, 2017
5.	Guidance for Industry and Food and Drug Administration Staff: The 510(k) Program: Evaluating Substantial Equivalence in Premarket Notifications [510(k)] Document Issued on July 28, 2014
6.	Guidance for Industry and FDA Staff: Guidance for the Submission Of 510(k)'s for Solid State X-ray Imaging Devices Document issued on September 1, 2016
7.	Guidance for Industry and FDA Staff: Guidance for the Content of Premarket Submission for Software in Medical Devices Document issued on May 11, 2005
8.	Guidance for Industry and FDA Staff: Guidance for Off-The-Shelf Software Use in Medical Devices Document issued on September 27, 2019
9.	Guidance for Industry and FDA Staff: Applying Human Factors and Usability Engineering to Medical Devices. Document issued February 3, 2016
10.	Guidance for Industry and FDA Staff: Pediatric Information for X-ray Imaging Device Premarket Notifications. Document issued on November 28, 2017
11.	Guidance for Industry and FDA Staff: Content of Premarket Submissions for Management of Cybersecurity in Medical devices. Document issued on October 2, 2014
12.	Guidance for Industry and FDA Staff: Appropriate Use of Voluntary Consensus Standards in Premarket Submission for Medical Devices Document issued on September 14, 2018
13.	Guidance for Industry and FDA Staff: Medical Device Accessories - Describing Accessories and Classification Pathways Document issued on December 20, 2017
14.	Guidance for Industry and FDA Staff: Recommended Content and Format of Non-Clinical Bench Performance Testing Information in Premarket Submissions Document issued on December 20, 2019
15.	Guidance for Industry and FDA Staff: Radio Frequency Wireless Technology in Medical Device Document issued on August 14, 2007.
16.	Guidance for Industry and FDA Staff: Information to Support a Claim of Electromagnetic Compatibility (EMC) of Electrically-Powered Medical Devices Document issued on July 11, 2016

The modifications described in this Premarket Notification are supported with verification and validation testing.

Verification and Validation:

Software Documentation for a **Moderate** Level of Concern software per FDA's Guidance Document "Guidance for the Content of Premarket Submissions for Software Contained in Medical Devices" issued on May 11, 2005 and "Off-The-Shelf Software Use in Medical Devices" is also included as part of this submission. The performance data demonstrates continued conformance with special controls for medical devices containing software. Non-clinical tests were conducted on the Subject Device Cios Select software version VA21 during product development.

The Risk analysis was completed and risk control implemented to mitigate identified hazards. The testing results support that all the software specifications have met the acceptance criteria. Testing for verification and validation for the device was found acceptable to support the claims of substantial equivalence.

Bench testing in the form of Unit, Subsystem and System Integration testing was performed to evaluate the performance and functionality of the new features, hardware, and software updates. All testable requirements in the Engineering Requirements Specifications keys, Subsystem Requirements Specifications keys, and the Risk Management Hazard keys have been successfully verified and traced in accordance with the Siemens product development (lifecycle) process. The software verification and regression testing has been performed successfully to meet their previously determined acceptance criteria as stated in the test plans.

Electrical safety and EMC testing were conducted on the Cios Select, consisting of the acquisition unit (C-arm system) and the image processing and display station. The system complies with the IEC 60601-1, IEC 60601- 2-43 and IEC 60601-2-54 standards for safety and the IEC 60601-1-2 standard for EMC.

The Cios Select software (VA21) was tested and found to be safe and effective for intended users, uses and use environments through the design control verification and validation process. The Human Factor Usability Validation showed that Human factors are addressed in the system test according to the operator's manual and in clinical use tests with customer report and feedback form. Customer employees are adequately trained in the use of this equipment.

Siemens conforms to the cybersecurity requirements by implementing a process of preventing unauthorized access, modifications, misuse or denial of use, or the unauthorized use of information that is stored, accessed or transferred from a medical device to an external recipient. Provided in this submission is a cybersecurity statement that considers IEC 80001-1:2010. The responsibility for compliance with IEC 80001-1-2010 is the hospital. Provided in the Software Section is the required cybersecurity information.

Additional engineering bench testing was performed including: the non-clinical testing identified in the guidance for submission of 510(k)s for Solid State X-Ray Imaging Devices (SSXI); demonstration of system performance; and an imaging performance evaluation. X-ray Imaging Devices- Laboratory Image Quality and Dose Assessment, Tests and Standards" was performed with acceptable results.

Summary:

Performance tests were conducted to test the functionality of the Cios Select (VA21). These tests have been performed to assess the functionality of the Subject Device. Results of all conducted testing and clinical assessments were found acceptable and do not raise any new issues of safety or effectiveness.

6. General Safety and Effectiveness Concerns:

Instructions for use are included within the device labeling, and the information provided will enable the user to operate the device in a safe and effective manner.

Risk management is ensured via a hazard analysis, which is used to identify potential hazards. These potential hazards are controlled via software development, verification and validation testing. To minimize electrical and mechanical hazards, Siemens adheres to recognized and established industry practice, and all equipment is subject to final performance testing. Furthermore, the operators are health care professionals familiar with and responsible for the evaluating and post processing of X-ray images.

7. Conclusion as to Substantial Equivalence:

The Cios Select has the same indications for use, operating environment, and mechanical design as the Primary and Secondary predicates. Siemens concludes via the documentation provided in the 510(k) submission that the Cios Select does not introduce any new potential safety risks and is substantially equivalent, and performs as well as the predicate devices.