

May 21, 2021

iRhythm Technologies, Inc.Rey JacintoSr. Manager, Regulatory Affairs699 8th StreetSan Francisco, California 94103

Re: K202527

Trade/Device Name: Zio ECG Utilization Software (ZEUS) System Regulation Number: 21 CFR 870.1425 Regulation Name: Programmable Diagnostic Computer Regulatory Class: Class II Product Code: DQK, DSI, DXH Dated: August 31, 2020 Received: September 1, 2020

Dear Rey Jacinto:

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 <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,

Jennifer Shih Kozen Assistant Director Division of Cardiac Electrophysiology, Diagnostics and Monitoring Devices Office of Cardiovascular Devices Office of Product Evaluation and Quality Center for Devices and Radiological Health

Enclosure

Indications for Use

510(k) Number *(if known)* K202527

Device Name Zio® ECG Utilization Service (ZEUS) System

Indications for Use (Describe)

The device is intended to capture, analyze and report symptomatic and asymptomatic cardiac events and continuous electrocardiogram information for long-term monitoring. After wear, ECG data from compatible monitoring devices is processed and analyzed by the ZEUS System. A final report is generated on the beat-to-beat information from the entire ECG recording. For the Zio AT service, the ZEUS System supports the capture and analysis of automatically-detected arrhythmia event, as well as the analysis of uploaded patient-triggered events.

The ZEUS System is indicated for use on patients 18 years or older who may be asymptomatic or who may suffer from transient symptoms such as palpitations, shortness of breath, dizziness, light-headedness, pre-syncope, syncope, fatigue, or anxiety and patients who are asymptomatic. The reports are provided for review by the intended user to render a diagnosis based on clinical judgment and experience. It is not intended for use on critical care patients.

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) Notification K202527

I. General Information

Applicant:

iRhythm Technologies, Inc.

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San Francisco, CA 94103 USA

Phone: 415-632-5700

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Contact Person:

Rey Jacinto Sr. Manager, Regulatory Affairs Phone: 415-214-7440 Email: <u>rey.jacinto@irhythmtech.com</u> **Date Prepared:** August 31, 2020

II. Device Information

Trade Name:

ZEUS System

Generic/Common Name:

Programmable diagnostic computer

Classification Names:

Programmable diagnostic computer [21CFR§870.1425]

Arrhythmia detector and alarm (including ST-segment measurement and

alarm)[21CFR§870.1025]

Telephone electrographic transmitter and receiver [21CFR§870.2029]

Regulatory Class:

Class II (Special controls)

Product Codes:

DQK, Computer, Diagnostic, Programmable

DSI, Detector and Alarm, Arrhythmia

DXH, Transmitters and Receivers, Electrocardiographic, Telephone

III. Predicate Devices

The following predicate device has been selected:

- iRhythm Technologies, Inc. Zio[®] ECG Utilization Software (ZEUS) System [K190593]

IV. Indications for Use

The Indications for Use statement for the ZEUS System is as follows:

The device is intended to capture, analyze and report symptomatic and asymptomatic cardiac events and continuous electrocardiogram information for long-term monitoring. After wear, ECG data from compatible monitoring devices is processed and analyzed by the ZEUS System. A final report is generated on the beat-to-beat information from the entire ECG recording. For the Zio AT service, the ZEUS System supports the capture and analysis of automatically-detected arrhythmia events, as well as the analysis of uploaded patient-triggered events.

The ZEUS System is indicated for use on patients 18 years or older who may be asymptomatic or who may suffer from transient symptoms such as palpitations, shortness of breath, dizziness, light-headedness, pre-syncope, syncope, fatigue, or anxiety and patients who are asymptomatic. The reports are provided for review by the intended user to render a diagnosis based on clinical judgment and experience. It is not intended for use on critical care patients.

V. Device Description

The Zio® Service consists of single-patient-use monitoring devices and the Zio ECG Utilization Software (ZEUS) System, the subject device of this submission, for analysis and reporting of cardiac information derived from ECG data. The ZEUS System is a software system consisting of a collection of software modules designed to store and analyze data from compatible cardiac monitoring devices to curate a report of preliminary findings intended for use by clinicians as an aid in arrhythmia diagnosis and management. The ECG Analysis Software of the ZEUS System is used by Certified Cardiographic Technicians (CCTs) prior to publishing the reported cardiac information in the Final Report and is not utilized directly by the prescribing clinician or patient. The reported cardiac information includes beats, ectopic runs, ECG segments, rhythms, and heart rate measurements. Recorded ECG is processed by an automated ECG analysis platform; results are quality reviewed by CCTs, findings and associated ECG are captured in a report provided to clinicians via a secure website. For the Zio® AT Patch/Gateway, the ZEUS System provides capabilities to automatically detect clinically actionable arrhythmia during the monitoring period, as well as receive baseline, scheduled, symptomatic and asymptomatic transmissions.

VI. Comparison of Technological Characteristics with Predicate Devices (Substantial Equivalence)

The proposed indications for use statement for the ZEUS System are identical to the indications for use in the predicate device. The differences in the technological characteristics, namely the introduction of ECG segment detection, between the devices are substantially equivalent as the fundamental scientific technology and intended use is unchanged. Thus, the ZEUS System is considered substantially equivalent to the predicate device.

A comparison table outlining the differences and similarities between the subject device and the predicate device is provided in Table 1.

Feature	Subject Device:	Predicate Device:		
	ZEUS System	ZEUS System		
	(K202527)	(K190593)		
General Characteristics				
Classification	Same	Class II (Special controls)		
		21CFR870.1425		
		21CFR870.1025		
		21CFR870.2029		
Product Code	Same	DQK		
		DSI		
		DXH		
Patient Environment	Same	Ambulatory		
Patient Population	Same	Non-pediatric, non-critical		
		care patients		
Technological Characteristics				
Data Input	Same	Digital long-term		
		continuous ECG		
Data Download	Same	Yes		
Data Storage	Same	Yes		
Rhythm Detection	ECGDL	ECGML + ECGDL		
Algorithm + ECG Analysis				
ECG Segments	QT	N/A		
Interoperability	Zio [®] XT Patch	Zio XT Patch		
	Zio [®] AT Patch	Zio AT Patch		
	Zio [®] Monitor			

Table 1. Substantial Equivalence Summary Table

VII. Performance Data

Safety and performance of the ZEUS System has been evaluated and verified in accordance with design specifications and to support a determination of substantial equivalence to the predicate device.

Nonclinical testing summarized in Table 2 demonstrates that the ZEUS System is in conformance with FDA-recognized consensus standards and FDA guidance documents.

FDA #	Body	Number/Version	Title
5-40	AAMI ANSI ISO	14971:2012(R)2010	Medical Devices – Application of risk
		(Corrected 4	management to medical devices
		October 2017)	
13-79	IEC	62304 Edition 1.1	Medical device software - Software
		2015-06	life cycle processes
		CONSOLIDATED	
		VERSION	
3-127	AAMI ANSI IEC	60601-2-47:2012	Medical Electrical Equipment Part
			2-47: Particular Requirements For
			The Basic Safety And Essential
			Performance Of Ambulatory
			Electrocardiographic Systems
3-118	AAMI ANSI	EC57:2012	Testing and reporting performance
			results of cardiac rhythm and ST
			segment measurement algorithms
N/A	U.S. FDA	October 2, 2014	Guidance for Industry and FDA Staff –
			Content of Premarket Submissions
			for Management of Cybersecurity in
			Medical Devices
N/A	U.S. FDA	May 5 <i>,</i> 2005	Guidance for the Content of
			Premarket Submissions for Software
			Contained in Medical Devices

 Table 2. FDA-Recognized Consensus Standards & Guidance Document Summary

The nonclinical verification and performance test results established that the device meets its design requirements and intended use, that the modifications to the cleared device are as safe and effective as the predicate. During the development, potential hazards were evaluated and controlled by the risk management activities, including risk analysis, risk mitigation, verification and benefit-risk analysis. The verification and system-level regression testing demonstrate that the device meets all its specifications.

VIII. Clinical Testing in Support of Substantial Equivalence Determination

No clinical testing was performed in support of this premarket notification.

IX. Conclusion

The results confirm by examination and provision of objective evidence that the design output met the design input requirements. The results of the nonclinical testing performed demonstrate that the ZEUS System meets the requirements of established conformance standards and performance specifications necessary for its intended use and does not raise new questions of safety or effectiveness as compared to the predicate device. The ZEUS System is substantially equivalent to the predicate device.