



May 4, 2023

Linear Health Sciences, LLC
% Jessica Czamanski
Managing Director
MedTactics, LLC
19924 NE 19th PL
Miami, Florida 33179

Re: K230266

Trade/Device Name: Orchid Safety Release Valve™

Regulation Number: 21 CFR 880.5220

Regulation Name: Intravenous Catheter Force-Activated Separation Device

Regulatory Class: Class II

Product Code: QOI

Dated: January 20, 2023

Received: January 31, 2023

Dear Jessica Czamanski:

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,

A handwritten signature in black ink that reads "David Wolloscheck". The signature is written in a cursive style. In the background, there is a large, light blue watermark of the letters "FDA".

David Wolloscheck, Ph.D.
Acting Assistant Director
DHT3C: Division of Drug Delivery and
General Hospital Devices,
and Human Factors
OHT3: Office of GastroRenal, ObGyn,
General Hospital and Urology Devices
Office of Product Evaluation and Quality
Center for Devices and Radiological Health

Enclosure

Indications for Use

510(k) Number (if known)
K230266

Device Name
Orchid Safety Release Valve™

Indications for Use (Describe)

The Linear Health Science™ Orchid Safety Release Valve™ is a tension-activated accessory for single patient use and placed between the existing IV administration set and IV extension set connection. The Orchid SRV™ is intended for use with electronic IV pumps in peripheral IV catheter applications where tension may act on the IV tubing. The Orchid SRV™ is designed to allow flow to an IV catheter. When excessive tension acts on the line, the Orchid SRV™ separates and closes the flow path in both directions. The Orchid SRV™ can be used during intermittent infusion and continuous infusion.

The Orchid SRV™ is intended to aid in reduction of peripheral IV mechanical complications requiring IV replacement.

The Orchid SRV™ is for use with patients two (2) weeks of age and older.

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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K230266 - 510(k) SUMMARY

This 510(k) Summary is provided per the requirements of section 21 CFR 807.92.

I. Submitter

Submitter's Name: Linear Health Sciences, LLC
Contact Person: Mr. Daniel Clark
CEO
Address: 5333 Wisteria Drive
Oklahoma City, OK 73142
Telephone: (310) 721-6222
Email: dan.clark@linearsciences.com
Date Preparation: May 4, 2023

II. Application Correspondent

Contact's Name: MedTactics, LLC
Contact Person: Jessica Czamanski
Managing Director
Address: 19924 NE 19TH PL
Miami, FL 33179
Telephone: (754) 422-9101
Email: jczamanski@medtactics.us

III. Device

Trade Name: Orchid Safety Release ValveTM
Common Name: Quick Disconnect Accessory
Classification Name: Intravenous Catheter Force-Activated Separation Device
Product Classification: Class II
Regulation Number: 21 CFR § 880.5220
Product Code: QOI

IV. Predicate Device

Manufacturer: Linear Health Sciences, LLC
Device Name: Orchid Safety Release ValveTM

510(k) Number: K212064
Product Classification: Class II
Regulation Number: 21 CFR § 880.5220

V. Device Description

The Orchid Safety Release Valve™ or Orchid SRV™ connects via standard luer-locking connection, allowing flow during IV therapy. The Orchid SRV™ is designed to allow the device to separate into two halves when longitudinal tension exceeds the SRV tension window, automatically closing the flow path to both IV extension set and IV administration set. Following separation, a component of the Orchid SRV™ is left attached to each side of the infusion system to protect the intraluminal pathway. Upon separation, replacement of the SRV™ is necessary. Follow institutional policy to replace the SRV™, or at least every seven (7) days.

VI. Intended Use

The Linear Health Sciences™ Orchid Safety Release Valve™ is a tension-activated accessory, provided sterile and for single use, in line with an IV administration set and peripheral IV extension set on a patient. The Orchid SRV™ provides a quick separation feature that allows the device to quickly separate into two halves upon tension, closing the flow path to prevent leakage. The device is intended to reduce the risk of IV catheter failure, requiring IV catheter replacement.

VII. Indications for Use

The Linear Health Sciences™ Orchid Safety Release Valve™ is a tension-activated accessory for single patient use and placed between the existing IV administration set and IV extension set connection. The Orchid SRV™ is intended for use with electronic IV pumps in peripheral IV catheter applications where tension may act on the IV tubing. The Orchid SRV™ is designed to allow flow to an IV catheter. When excessive tension acts on the line, the Orchid SRV™ separates and closes the flow path in both directions. The Orchid SRV™ can be used during intermittent infusion and continuous infusion.

The Orchid SRV™ is intended to aid in reduction of peripheral IV mechanical complications requiring IV replacement.

The Orchid SRV™ is for use with patients two (2) weeks of age and older.

VIII. Comparison of Technological Characteristics with the Predicate Devices

There are no differences in technological characteristics of the device. Only differences are present in the labeling of the Orchid Safety Release Valve, specifically the instructions for use and quick reference guide.

The following table (**Table 5-1**) provides an overview of general technological characteristics in comparison to the predicate device.

Table 5-1: General Technological Characteristics Comparison

Product Features	<u>Proposed</u> Linear Health Sciences' Orchid Safety Release Valve™	<u>Predicate</u> Linear Health Sciences' Orchid Safety Release Valve™ (K212064)	<u>Substantial Equivalence Determination</u>
Classification	Class II	-same-	
Product Code	QOI	-same-	
Regulation Number	§880.5220	-same-	
Device Classification Name	Intravenous Catheter Force-Activated Separation Device	-same-	
Intended Use	The Linear Health Sciences™ Orchid Safety Release Valve™ is a tension-activated accessory, provided sterile and for single use, in line with an IV administration set and peripheral IV extension set on a patient. The Orchid SRV™ provides a quick separation feature that allows the device to quickly separate into two halves upon tension, closing the flow path to prevent leakage. The device is intended to reduce the risk of IV catheter failure, requiring IV catheter replacement.	-same-	
Indications for Use	The Linear Health Sciences™ Orchid Safety Release Valve™ is a tension-activated accessory for single patient use and placed between the existing IV administration set and IV extension set connection. The Orchid SRV™ is intended for use with electronic IV pumps in peripheral IV catheter applications where tension may act on the IV tubing. The Orchid SRV™ is designed to allow flow to an IV catheter. When excessive tension acts on the line, the Orchid SRV™ separates and	The Linear Health Sciences™ Orchid Safety Release Valve™ is a tension-activated accessory for single patient use and placed between the existing IV administration set and IV extension set connection. The Orchid SRV™ is intended for use with electronic IV pumps in peripheral IV catheter applications where tension may act on the IV tubing. The Orchid SRV™ is designed to allow flow to an IV catheter. When excessive tension acts on the line, the Orchid SRV™ separates and	Indications for use are being updated to include pediatric patients 2 weeks old to 17 years old. The device performance of the subject and predicate device are identical and compatible with IV therapy in adult and pediatric patients. The results of pediatric extrapolation supported the similarities in risk profiles across all patient populations. Therefore, the change in indication does not raise new or different

Table 5-1: General Technological Characteristics Comparison

Product Features	<u>Proposed</u> Linear Health Sciences' Orchid Safety Release Valve™	<u>Predicate</u> Linear Health Sciences' Orchid Safety Release Valve™ (K212064)	<u>Substantial Equivalence Determination</u>
	<p>closes the flow path in both directions. The Orchid SRV™ can be used during intermittent infusion and continuous infusion.</p> <p>The Orchid SRV™ is intended to aid in reduction of peripheral IV mechanical complications requiring IV replacement.</p> <p>The Orchid SRV™ is for use with patients two (2) weeks of age and older.</p>	<p>closes the flow path in both directions. The Orchid SRV™ can be used during intermittent infusion and continuous infusion.</p> <p>The Orchid SRV™ is intended to aid in reduction of peripheral IV mechanical complications requiring IV replacement.</p> <p>The Orchid SRV™ is for use with patients eighteen (18) years of age and older.</p>	<p>questions of safety or effectiveness.</p>
Materials	Polycarbonate and silicone	-same-	
Environment of Use	Hospital	-same-	
Provided Sterile	Yes	-same-	
Principle of Operation	<p>The Orchid Safety Release Valve™ has luer lock connections that will lock the device in place during use. The female luer connects to an administration set while the male luer connects to a vascular access device hub or extension set. Once connected the device allows for continuous flow. The Orchid SRV will separate into the male and female subassemblies, upon a tension event, automatically closing the flow path, while maintaining sterility and preventing fluid leakage from the device.</p>	-same-	
User Profile	Physician or clinical personnel with clearance to administer IV sets and related products	-same-	
Separation Force	1-4.2 lbf	-same-	

Table 5-1: General Technological Characteristics Comparison

Product Features	<u>Proposed</u> Linear Health Sciences' Orchid Safety Release Valve™	<u>Predicate</u> Linear Health Sciences' Orchid Safety Release Valve™ (K212064)	<u>Substantial Equivalence Determination</u>
Vascular Access Catheter Type	Peripheral IV Catheters	-same-	
For Use with Electronic Pumps	Yes	-same-	
Single Use	Yes	-same-	
Continuous and Intermittent Infusion	Yes	-same-	
Sterilization	Ethylene Oxide	-same-	
SAL	10 ⁻⁶	-same-	
Shelf-Life	2 years	-same-	

IX. Performance Data

The following performance data was considered in support of the substantial equivalence determination.

Performance Testing - Bench

There is no change in performance data since there are no changes to the device or its manufacturing processes.

Biocompatibility

There is no change in biocompatibility since there are no changes to the device or its manufacturing processes.

Electrical Safety and Electromagnetic Compatibility (EMC)

There are no electrical or metal components in the proposed Orchid Safety Release Valve™; therefore, the proposed device does not require EMC and Electrical Safety evaluation.

Software Verification and Validation Testing

The Orchid Safety Release Valve™ does not contain software; therefore, the proposed device does not require software verification and validation testing.

Performance Testing - Animal

This submission does not include any animal performance testing. It was determined that no such testing was required to demonstrate substantial equivalence.

Clinical Data

Pediatric extrapolation results supported safety and effectiveness of the use of the Orchid SRV in pediatric patients.

X. Conclusion

The Orchid Safety Release ValveTM has the same intended use, environment, operating principle and fundamental technology, manufacturing, and materials as the predicate device. The conducted pediatric extrapolation assessment demonstrates that the differences in the indications for use do not raise any new or different questions of safety or efficacy. The information provided in this submission demonstrates that the device is substantially equivalent to its predicate.