August 7, 2019

Shockwave Medical, Inc.
Ms. Cindy Morrow
Manager, Regulatory Affairs
5403 Betsy Ross Drive
Santa Clara, California 95054

Re: K191840

Trade/Device Name: Shockwave Medical Peripheral Intravascular Lithotripsy (IVL) System
Regulation Number: 21 CFR 870.1250
Regulation Name: Percutaneous Catheter
Regulatory Class: Class II
Product Code: PPN
Dated: July 8, 2019
Received: July 9, 2019

Dear Ms. Morrow:

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/comination-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.


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,

Eleni Whatley

For

Gregory O'Connell
Assistant Director
DHT2C: Division of Coronary and Peripheral Intervention Devices
OHT2: Office of Cardiovascular Devices
Office of Product Evaluation and Quality
Center for Devices and Radiological Health

Enclosure
Indications for Use

Device Name
Shockwave Medical Peripheral Intravascular Lithotripsy (IVL) System

Indications for Use (Describe)
The Shockwave Medical Peripheral Intravascular Lithotripsy (IVL) System is intended for lithotripsy-enhanced balloon dilatation of lesions, including calcified lesions, in the peripheral vasculature, including the iliac, femoral, ilio-femoral, popliteal, infra-popliteal, and renal arteries. Not for use in the coronary or cerebral vasculature.

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

- [x] 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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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:

- Department of Health and Human Services
- Food and Drug Administration
- Office of Chief Information Officer
- Paperwork Reduction Act (PRA) Staff
- PRASStaff@fda.hhs.gov

“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.”
This 510(k) Summary is submitted in accordance with the requirements of the Safe Medical Devices Act (SMDA) of 1990 and 21 CFR 807.92.

510(k) Owner
Shockwave Medical, Inc.
5403 Betsy Ross Drive
Santa Clara, CA 95054
Phone: (510) 279-4262
Fax: (510) 279-5934

Contact Person
Cindy Morrow

Date Prepared
August 7, 2019

Device Name and Classification

<table>
<thead>
<tr>
<th>Trade Name:</th>
<th>Shockwave S4 Intravascular Lithotripsy (IVL) Catheter,</th>
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<tr>
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<td>Shockwave Medical Peripheral Intravascular Lithotripsy System</td>
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<td>Common Name:</td>
<td>Catheter, lithotripsy, peripheral, transluminal</td>
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<td>21 CFR 870.1250</td>
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<td>Classification Name:</td>
<td>Percutaneous catheter</td>
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<td>Product Code:</td>
<td>PPN</td>
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Predicate Device
The predicate device is the Shockwave Medical Peripheral Intravascular Lithotripsy System, K180454, cleared by FDA on June 27, 2018.

Indications for Use / Intended Use
The Shockwave Medical Peripheral Intravascular Lithotripsy (IVL) System is intended for lithotripsy-enhanced balloon dilatation of lesions, including calcified lesions, in the peripheral vasculature, including the iliac, femoral, ilio-femoral, popliteal, infra-popliteal, and renal arteries. Not for use in the coronary or cerebral vasculature.

Device Description
The Shockwave Medical Peripheral Intravascular Lithotripsy (IVL) System has three components: a proprietary IVL Catheter, an IVL Generator, and an IVL Connector Cable. The IVL Catheter has integrated lithotripsy emitters and is designed to enhance percutaneous transluminal angioplasty by enabling delivery of the calcium disrupting capability of lithotripsy prior to full balloon dilatation at low pressures. The application of lithotripsy mechanical pulse waves alters the structure of an occlusive vascular deposit (stenosis) prior to low-pressure balloon dilatation of the stenosis and facilitates the passage of blood.
The IVL Catheter is delivered through the peripheral arterial system of the lower extremities to the site of an otherwise difficult to treat lesion. The balloon is partially inflated and the lithotripsy emitters are energized thereby generating pulsatile mechanical energy within the balloon at the target treatment site and allowing subsequent dilation of a peripheral artery stenosis using low balloon pressure. The IVL Generator delivers energy through the IVL Connector Cable to the pulse emitters located inside the balloon in the IVL Catheter. The IVL Catheter is a single-use device supplied sterile to the customer. The IVL Generator and IVL Connector Cable are non-sterile reusable devices.

**Technological Comparison**

This Special 510(k) Premarket Notification is being submitted for minor modifications to the Peripheral IVL Catheter sizes (2.5mm, 3.0mm, 3.5mm, and 4.0mm in diameter and 40mm length) for use with the IVL Generator and Connector Cable. The minor modifications include small dimensional changes to the balloon and emitter components, and new tooling for improved manufacturability.

The IVL System has the same intended use, principles of operation and has substantially equivalent technological characteristics, including the same fundamental scientific technology, design, energy source, shelf life, and sterilization method as the already 510(k) cleared IVL System. The IVL Catheters are the same design as the predicate; the IVL Catheters with integrated lithotripsy emitters enable the localized delivery of pulsatile mechanical energy to disrupt calcified lesions.

**Summary of Performance Data**

Objective evidence demonstrating that the IVL System design output meets the product design input requirements as well as that device performance characteristics conform to user needs and intended uses as defined in the product specification was provided. Testing was conducted in accordance with Shockwave Medical’s Risk Analysis procedures, all applicable FDA guidance documents and relevant international standards. Testing included:

- IVL Catheter design verification and validation testing:
  - Sheath compatibility
  - Nominal balloon diameter
  - Balloon compliance (at 10 ATM)
  - Nominal balloon working length (at 6 ATM)
  - Inflation (to 6 ATM) / Deflation times (from 10 ATM)
  - Crossing profile
  - Distal tip profile and Tip durability
  - Marker band alignment
  - Distal & Proximal bond strength
  - Catheter torsional strength
  - Emitters and marker band bond strength integrity
  - Pressure leakage during lithotripsy treatment
  - Minimum RBP
Results demonstrated that the performance of the IVL System meets its design specifications and demonstrates substantial equivalence for its intended use; therefore, additional clinical data were not required.

**Basis for Substantial Equivalence**

The IVL System with the minor modifications to the Peripheral IVL Catheter shares the same intended use, principles of operation, overall technical and functional capabilities, and similar design and materials as the identified predicate device. Any differences between the IVL Systems were evaluated through design verification and validation testing which demonstrated device performance. The IVL System with the minor modifications to the Peripheral IVL Catheter is therefore substantially equivalent to the predicate device.