August 14, 2017

Penumbra, Inc.
Mary Rose
Director, Regulatory Affairs
One Penumbra Place
Alameda, California 94502

Re: K171332
Trade/Device Name: Artemis Neuro Evacuation Device
Regulation Number: 21 CFR 882.1480
Regulation Name: Neurological Endoscope
Regulatory Class: Class II
Product Code: GWG
Dated: July 14, 2017
Received: July 17, 2017

Dear Ms. Rose:

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. 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); good manufacturing practice requirements as set forth in
the quality systems (QS) regulation (21 CFR Part 820); and if applicable, the electronic product
radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

If you desire specific advice for your device on our labeling regulation (21 CFR Part 801), please
contact the Division of Industry and Consumer Education at its toll-free number (800) 638-2041
or (301) 796-7100 or at its Internet address
http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm. 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
http://www.fda.gov/MedicalDevices/Safety/ReportaProblem/default.htm for the CDRH's Office
of Surveillance and Biometrics/Division of Postmarket Surveillance.

You may obtain other general information on your responsibilities under the Act from the
Division of Industry and Consumer Education at its toll-free number (800) 638-2041 or (301)
796-7100 or at its Internet address

Sincerely,

Carlos L. Peña -S

Carlos L. Peña, PhD, MS
Director
Division of Neurological
and Physical Medicine Devices
Office of Device Evaluation
Center for Devices and Radiological Health

Enclosure
Indications for Use

Device Name
Artemis™ Neuro Evacuation Device

Indications for Use (Describe)
The Artemis™ Neuro Evacuation Device is used for the controlled aspiration of tissue and/or fluid during surgery of the Ventricular System or Cerebrum in conjunction with a Penumbra Aspiration Pump.

Penumbra Aspiration Pump:
The Penumbra Aspiration Pump is indicated as a vacuum source for the Penumbra Aspiration Systems.

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.

*DO NOT SEND YOUR COMPLETED FORM TO THE PRA STAFF EMAIL ADDRESS BELOW.*

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."
1 510(k) SUMMARY

(as required by 21 CFR 807.92)

1.1 Sponsor/Applicant Name and Address

Penumbra Inc.
One Penumbra Place
Alameda, CA 94502, USA

1.2 Sponsor Contact Information

Mary Rose
Director, Regulatory Affairs
Phone: 510-748-3346
FAX: 510-217-6414
e-mail: mary.rose@penumbrainc.com

1.3 Date of Preparation of 510(k) Summary

July 14, 2017

1.4 Device Trade or Proprietary Name

Artemis™ Neuro Evacuation Device

1.5 Device Common/Usual or Classification Name

Endoscope, neurological (Product Code: GWG)

1.6 Primary Device Classification

Regulatory Class: II
Classification Panel: Neurology
Classification Name: Endoscope, neurological
Regulation Number: 21 CFR 882.1480
Product Code: GWG
## 1.7 Predicate Devices

<table>
<thead>
<tr>
<th>510(k) Number</th>
<th>Clearance Date</th>
<th>Name of Predicate Device</th>
<th>Name of Manufacturer</th>
</tr>
</thead>
<tbody>
<tr>
<td>K132931</td>
<td>01/17/14</td>
<td>Apollo System</td>
<td>Penumbra, Inc.</td>
</tr>
<tr>
<td>K152699</td>
<td>03/17/16</td>
<td>Apollo System</td>
<td>Penumbra, Inc.</td>
</tr>
</tbody>
</table>

### Reference Devices

<table>
<thead>
<tr>
<th>510(k) Number</th>
<th>Clearance Date</th>
<th>Name of Predicate Device</th>
<th>Name of Manufacturer</th>
</tr>
</thead>
<tbody>
<tr>
<td>K160533</td>
<td>05/24/16</td>
<td>Penumbra Embolectomy Aspiration System (INDIGO™ Aspiration System)</td>
<td>Penumbra, Inc.</td>
</tr>
<tr>
<td>K160449</td>
<td>05/25/16</td>
<td>Penumbra System® and Penumbra Pump MAX</td>
<td>Penumbra, Inc.</td>
</tr>
</tbody>
</table>

## 1.8 Predicate Comparison

<table>
<thead>
<tr>
<th>510(k) No.</th>
<th>Apollo™ System</th>
<th>Subject Device: Artemis™ Neuro Evacuation Device</th>
</tr>
</thead>
<tbody>
<tr>
<td>K132931, K152699</td>
<td>K171332</td>
<td>Class II, GWG</td>
</tr>
</tbody>
</table>

### Intended Use

- **Apollo™ System**: Used for the controlled aspiration of tissue and/or fluid during surgery of the Ventricular System or Cerebrum.
- **Subject Device**: SAME

### Aspiration Control

- **Apollo™ System**: Fingertip controlled via vacuum regulator hole on Wand handle.
- **Subject Device**: SAME

### Mechanism of Action

- **Apollo™ System**: Vacuum Aspiration aided by Vibrational Energy
- **Subject Device**: Vacuum Aspiration aided by Rotational Energy

### Power Source for Mechanism of Action

- **Apollo™ System**: Apollo Generator
- **Subject Device**: Primary Cell Lithium Ion Battery, 1.5 Volt DC Motor contained within the handle

### Aspiration Source

- **Apollo™ System**: Apollo Pump
- **Subject Device**: Penumbra Aspiration Pump

### Wand Hypotube

- **Apollo™ System**: 304 Stainless Steel
- **Subject Device**: SAME

### Wand Materials

- **Apollo™ System**: Polymer, Metal
- **Subject Device**: SAME

### Wand Dimensions

- **Apollo™ System**: L: 27.0 cm – 28.0 cm
  ID: 0.060” – 0.088”
  OD: 0.072” – 0.101”
- **Subject Device**: L: 26.1 cm – 27.0 cm
  ID: 0.048” – 0.100”
  OD: 0.058” – 0.109”

### Aspiration Tubing Material

- **Apollo™ System**: Polymer
- **Subject Device**: SAME
Apollo™ System | Subject Device: Artemis™ Neuro Evacuation Device
--- | ---
Aspiration Tubing Dimensions | OD: 0.188” ID: 0.088” | OD: 0.263” ID: 0.163”
Single Use or Reusable | Single Use | SAME
Sterilization | EO | Gamma
Shelf-Life | 12-Months – Wand | SAME
Aspiration Pump | Apollo System Pump | Penumbra Aspiration Pump
IEC 60601-1 Compliance | Yes | SAME
IEC 60601-1-2 Compliance | Yes | SAME
Voltage | 100-115 VAC/230 VAC | SAME
Frequency | 50 Hz/60 Hz | SAME
Maximum Vacuum | 29 inHg | SAME
Maximum Flow Rate | 0.8 SCFM | SAME

1.9 Device Description

The Artemis™ Neuro Evacuation Device, is a surgical instrument designed to aid a physician in the removal of tissue and/or fluid during image-guided neurosurgery. The Artemis Wand has two functions. These functions are control and transfer of aspiration and generation of rotational energy. Aspiration is generated by a Penumbra Aspiration Pump, which the Artemis Wand connects to through its flexible tubing. The Artemis Wand has a rigid hypotube containing a wire to facilitate removing tissue and/or fluid with the assistance of rotational energy and aspiration.

The Artemis Wand is designed to be image-guided, allowing visualization of the procedure. The method of removal is vacuum aspiration, which draws the tissue and/or fluid into the lumen of the Wand hypotube. The integrated wire is fully contained within the lumen of the Wand hypotube, and has rotational capability facilitating movement of any tissue and/or fluid that may otherwise clog the hypotube lumen.

Intended users for this device are physicians who have received appropriate training in image-guided neurosurgery.
1.10 **Indications for Use**

The Artemis™ Neuro Evacuation Device is used for the controlled aspiration of tissue and/or fluid during surgery of the Ventricular System or Cerebrum in conjunction with a Penumbra Aspiration Pump.

**Penumbra Aspiration Pump:**

The Penumbra Aspiration Pump is indicated as a vacuum source for the Penumbra Aspiration Systems.

1.11 **Summary of Non-clinical Data**

Included in this section is a description of the testing, which substantiates the safe and effective performance of the subject Artemis™ Neuro Evacuation Device as well as its substantial equivalence to the predicate device:

- Biocompatibility
- Design Verification (Bench-Top Testing)
- EMC
- Sterilization
- Shelf-life

1.11.1 **Biocompatibility**

Biocompatibility testing was performed on the subject Artemis™ Neuro Evacuation Device. Tests were selected in accordance with EN ISO 10993-1 guidelines (Biological Evaluation of Medical Devices) and FDA Guidance. All studies were conducted pursuant to 21 CFR, Part 58, Good Laboratory Practices.

The following tests were performed based contact type and duration:

<table>
<thead>
<tr>
<th>Test</th>
<th>Method</th>
<th>Conclusions</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>In vitro</em> Cytotoxicity</td>
<td>MEM Elution Test</td>
<td>Non-toxic</td>
</tr>
<tr>
<td>Sensitization (Magnusson-Kligman Method)</td>
<td>ISO Guinea Pig Maximization Sensitization Test</td>
<td>Non-sensitizing</td>
</tr>
<tr>
<td>Test</td>
<td>Method</td>
<td>Conclusions</td>
</tr>
<tr>
<td>-------------------------------------------</td>
<td>------------------------------------------------------------------------</td>
<td>--------------</td>
</tr>
<tr>
<td>Irritation (Intracutaneous Reactivity (ISO))</td>
<td>ISO Tests for Irritation and Skin Sensitization</td>
<td>Non-irritant</td>
</tr>
<tr>
<td><strong>Systemic Toxicity</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Systemic Injection (ISO)</td>
<td>ISO Acute Systemic Injection Test</td>
<td>Non-toxic</td>
</tr>
<tr>
<td>Material Mediated Pyrogen</td>
<td>ISO Materials-Mediated Rabbit Pyrogen Test</td>
<td>Non-pyrogenic</td>
</tr>
<tr>
<td><strong>Hemocompatibility</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Thrombosis (Dog Thrombogenicity)</td>
<td>Thrombogenicity Study in Dogs - ISO</td>
<td>Non-Thrombogenic</td>
</tr>
<tr>
<td>Coagulation (PT)</td>
<td>Prothrombin Time (PT) Test</td>
<td>Non-hemolytic</td>
</tr>
<tr>
<td>Coagulation (PTT)</td>
<td>Partial Thromboplastin Time (PTT) Test</td>
<td>Non-hemolytic</td>
</tr>
<tr>
<td>Hematology (Hemolysis) – Direct Contact</td>
<td>ASTM Hemolysis (Direct Contact Method)</td>
<td>Non-hemolytic</td>
</tr>
<tr>
<td>Hematology (Hemolysis) – Indirect Contact</td>
<td>ASTM Hemolysis (Extract Method)</td>
<td>Non-hemolytic</td>
</tr>
<tr>
<td><strong>Genotoxicity</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ames Mutagenicity</td>
<td>Ames Test</td>
<td>Non-mutagenic</td>
</tr>
<tr>
<td><em>In vivo</em> Mouse micronucleus</td>
<td>ISO In Vivo Mouse Micronucleus Assay</td>
<td>Non-mutagenic</td>
</tr>
</tbody>
</table>

The non-clinical testing found the Penumbra Artemis™ Neuro Evacuation Device to be biocompatible according to EN ISO 10993 requirements. Therefore, Penumbra Artemis™ Neuro Evacuation Device is substantial equivalent to the predicate device.

**1.11.2 Bench-Top Testing**

Design Verification testing was conducted to evaluate the physical and mechanical properties of the Artemis™ Neuro Evacuation Device and demonstrate substantial equivalence to predicate. The following tests were successfully performed:

<table>
<thead>
<tr>
<th>Test</th>
<th>Test Method Summary</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Packaging Inspection</td>
<td>Confirm the packaging outputs meet all product specifications.</td>
<td>100%</td>
</tr>
<tr>
<td>Dimensional / Visual</td>
<td>Confirm the dimensions / visual outputs meet all product specifications.</td>
<td>100%</td>
</tr>
<tr>
<td>Inspection</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dynamic Vacuum Testing</td>
<td>Wand can sustain maximum vacuum and transfer vacuum to distal tip of wand.</td>
<td>100%</td>
</tr>
<tr>
<td>Simulated Use</td>
<td>Evaluate the effectiveness of the device to generate rotational energy to remove simulated clot.</td>
<td>100%</td>
</tr>
</tbody>
</table>
### Tensile/Torque

<table>
<thead>
<tr>
<th>Description</th>
<th>Specification</th>
<th>Pass</th>
</tr>
</thead>
<tbody>
<tr>
<td>All components are tested</td>
<td>to ensure connections/joints meet all product</td>
<td>100%</td>
</tr>
<tr>
<td></td>
<td>specifications.</td>
<td>Pass</td>
</tr>
</tbody>
</table>

### Electric Safety Testing

Compliant to requirements of IEC 60601-1 (3rd Ed.), IEC 60601-1-2, 60601-1-6, and IEC 62366.

Pass

Design Verification (Bench-Top Testing) found the Penumbra Artemis™ Neuro Evacuation Device to meet all design inputs. Therefore, Penumbra Artemis™ Neuro Evacuation Device is substantially equivalent to the predicate device.

1.11.3 Sterilization

The Artemis™ Neuro Evacuation Device was validated for gamma irradiation sterilization to a 6-log sterility assurance utilizing the VDmax25 sterilization validation in accordance with ISO 11137 and ISO 11737.

1.11.4 Shelf Life

The Artemis™ Neuro Evacuation Device has a 12-month Shelf-life.

1.12 Summary of Substantial Equivalence

The subject Artemis™ Neuro Evacuation Device is equivalent to the predicate Apollo System™. The subject device has an identical intended use as the predicate device. The subject device and predicate device differ slightly in regards to technological and material variations. However, these differences do not raise different questions of safety and effectiveness. The device testing described in Section 1.11 demonstrate the subject device is equivalent to the predicate device in regards to operating principle fundamental technology, materials, and device performance.