February 1, 2018

Cook Incorporated
Carly Powell
Regulatory Affairs Specialist
750 Daniels Way, P.O. Box 489
Bloomington, IN 47402

Re: K171662
Regulation Number: 21 CFR § 876.5130
Regulation Name: Urological Catheter and Accessories
Regulatory Class: II
Product Code: KOD
Dated: December 19, 2017
Received: December 20, 2017

Dear Carly Powell:

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.

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](http://www.fda.gov/MedicalDevices/Safety/ReportaProblem/default.htm) for the CDRH's Office of Surveillance and Biometrics/Division of Postmarket Surveillance.

For comprehensive regulatory information about medical devices and radiation-emitting products, including information about labeling regulations, please see Device Advice ([https://www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/](https://www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/)) and CDRH Learn ([http://www.fda.gov/Training/CDRHLearn](http://www.fda.gov/Training/CDRHLearn)). 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 ([http://www.fda.gov/DICE](http://www.fda.gov/DICE)) for more information or contact DICE by email (DICE@fda.hhs.gov) or phone (1-800-638-2041 or 301-796-7100).

Sincerely,

Benjamin R. Fisher -S

Benjamin R. Fisher, Ph.D.
Director
Division of Reproductive, Gastro-Renal, and Urological Devices
Office of Device Evaluation
Center for Devices and Radiological Health

Enclosure
Device Name

Indications for Use (Describe)
Ureteral catheters are indicated for access and catheterization of the urinary tract, including the following applications:

• Delivery of contrast media
• Drainage of fluids from the urinary tract
• Delivery of irrigation fluids to the urinary tract
• Navigation of a tortuous ureter
• Access, advancement, or exchange of wire guides (open-ended catheters only)

The Pediatric Ureteral Catheter is indicated for access and catheterization of the urinary tract in pediatric patients, including the following applications:

• Delivery of contrast media
• Drainage of fluids from the urinary tract
• Delivery of irrigation fluids to the urinary tract
• Navigation of a tortuous ureter

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.*

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Department of Health and Human Services
Food and Drug Administration
Office of Chief Information Officer
Paperwork Reduction Act (PRA) Staff
PRASTaff@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.”
2.0 510(k) Summary

Ureteral Catheters
As required by 21 CFR 807.92
Date Prepared: January 31, 2018

Submitted By:
Submission: Traditional 510(k) Premarket Notification
Applicant: Cook Incorporated
Contact: Carly Powell
Applicant Address: Cook Incorporated
750 Daniels Way
Bloomington, IN 47404
Contact Phone: (812) 339-2235 x104913
Contact Fax: (812) 332-0281

Device Information:
Trade Name: Open-End Ureteral Catheter,
Open-End Ureteral Catheter Sof-Flex,
EchoTip Open-End Ureteral Catheter,
Open-End Flexi-Tip Ureteral Catheter,
Flexi-Tip Ureteral Catheter (Closed End),
Whistle Tip Ureteral Catheter,
Round Tip Ureteral Catheter,
Spiral Tip Ureteral Access Catheter,
Pediatric Ureteral Catheter
Common Name: Catheter, Urological
Classification Name: Urological catheter and accessories
Regulation, Class: 21 CFR §876.5130, Class II
Product Code, Panel: KOD, Gastroenterology/Urology

Predicate Devices:
- Primary predicate device: Bard TigerTail Ureteral Catheter (K033719)
- Secondary predicate device: Porges Ureteral Catheters (K021856)

Device Description:
The Ureteral Catheter family is comprised of nine different types of Ureteral Catheters. The types of catheters include: Open-End Ureteral Catheter, Open-End Ureteral Catheter Sof-Flex, EchoTip Open-End Ureteral Catheter, Open-End Flexi-Tip Ureteral Catheter, Flexi-Tip Ureteral Catheter (Closed End), Whistle Tip Ureteral Catheter, Round Tip Ureteral Catheter, Spiral Tip Ureteral Access Catheter, and Pediatric Ureteral Catheter.
The Ureteral Catheters range in length from 10 to 125 cm. The catheters may be closed or open-ended. The Ureteral Catheters are sterile, single-use devices.

**Indications for Use:**

Ureteral catheters are indicated for access and catheterization of the urinary tract, including the following applications:

- Delivery of contrast media
- Drainage of fluids from the urinary tract
- Delivery of irrigation fluids to the urinary tract
- Navigation of a tortuous ureter
- Access, advancement, or exchange of wire guides (open-ended catheters only)

The Pediatric Ureteral Catheter is indicated for access and catheterization of the urinary tract in pediatric patients, including the following applications:

- Delivery of contrast media
- Drainage of fluids from the urinary tract
- Delivery of irrigation fluids to the urinary tract
- Navigation of a tortuous ureter

**Comparison to Predicate Devices:**

The subject devices have similar indications for use, methods of operation, and fundamental technological characteristics as the predicate devices due to the similarities with their intended uses. Differences between the subject device and the predicate devices include dimensional variations and slight variations in materials. Characteristics of the subject device that differ from the predicate devices are supported by testing. These differences do not raise any new questions of safety and/or effectiveness. The substantial equivalence comparison of the subject device to the predicates is provided in the following table.

**Substantial Equivalence Comparison**

<table>
<thead>
<tr>
<th>PRIMARY PREDICATE DEVICE</th>
<th>SECONDARY PREDICATE DEVICE</th>
<th>SUBJECT DEVICE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bard - TigerTail Ureteral Catheter</td>
<td>Porges Ureteral Catheters</td>
<td>Ureteral Catheters</td>
</tr>
<tr>
<td>510(k) Number</td>
<td>K033719</td>
<td>Subject of Submission</td>
</tr>
<tr>
<td>Manufacturer</td>
<td>C.R. Bard Inc.</td>
<td>Cook Incorporated</td>
</tr>
<tr>
<td>Regulation</td>
<td>21 CFR §876.5130</td>
<td>Identical</td>
</tr>
<tr>
<td>Product Code</td>
<td>KOD</td>
<td>Identical to primary predicate</td>
</tr>
<tr>
<td>Classification Name</td>
<td>Catheter, Urological</td>
<td>Identical to primary predicate</td>
</tr>
<tr>
<td>Classification</td>
<td>II</td>
<td>Identical</td>
</tr>
<tr>
<td>Indications for Use</td>
<td>Ureteral catheterization: -Drainage catheters -Catheters for retrograde ureteropyelography</td>
<td>Ureteral catheters are indicated for access and catheterization of the urinary tract, including the following applications:</td>
</tr>
</tbody>
</table>
**Reference Devices:**

In addition to the predicate devices, the following reference devices are used to help make a determination of substantial equivalence:

- The Rutner Universal Wedge Catheter (Cook Inc., K760858)
- The C-Flex Ureteral Stent Set (Cook Inc., K162104)

**Performance Data:**

The subject devices underwent the applicable testing listed below to ensure reliable design and performance under the testing parameters. Performance and biocompatibility
testing were conducted in accordance with the following applicable FDA guidance documents to confirm the reliable performance of critical device characteristics.

Performance – Testing shows that the subject device conforms to the performance testing requirements based on intended use. All predetermined acceptance criteria were met.

<table>
<thead>
<tr>
<th>Test Performed</th>
<th>Guidance Document</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tensile Strength – Testing shows that there should be no fracture of catheter tips or shafts during proper clinical use.</td>
<td></td>
</tr>
<tr>
<td>Leakage and Lumen Blockage – Testing evaluated lumen blockage and leakage in a pressurized flow test. Lumen patency, dimensional length, inner diameter, and outer diameter were determined.</td>
<td>N/A</td>
</tr>
<tr>
<td>Radiopacity – Testing shows that the mean radiopacity met the acceptance criteria and evaluated radiopacity by subjecting the ureteral catheters to a comparative fluoroscopic evaluation.</td>
<td></td>
</tr>
<tr>
<td>Kink Radius – Testing determined the kink radius of the ureteral catheter tubing.</td>
<td></td>
</tr>
<tr>
<td>Catheter-Hub Bond – Testing determined the tensile strength of the hub-to-shaft bond.</td>
<td></td>
</tr>
</tbody>
</table>

Biocompatibility – Testing shows that the subject device conforms to the biocompatibility requirements based on its intended use. All predetermined acceptance criteria were met.

<table>
<thead>
<tr>
<th>Testing Performed</th>
<th>Guidance Document</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sensitization – Guinea Pig Maximization</td>
<td></td>
</tr>
<tr>
<td>Irritation/Intracutaneous Reactivity – Intracutaneous Study</td>
<td></td>
</tr>
</tbody>
</table>

All predetermined acceptance criteria were met.

**Conclusion:**

The data included in this submission indicate that the subject devices do not raise new questions of safety or effectiveness compared to the predicate device. This supports a determination of substantial equivalence.