510(k) Summary:

This summary is provided as part of this Premarket Notification in compliance with 21CFR, Section 807.92.

Submitters name: B-K Medical
Address: Mileparken 34, DK2730 Herlev, Denmark
Phone: +45 44528100
Fax: +45 44528199
Contact person: Gert Nielsen, Regulatory Manager
Date prepared: August 23, 2013

Trade name: Ultrasound Scanner Pro Focus 2202
Common name: Diagnostic Ultrasound System
Classification names:
- Ultrasonic Pulsed Echo Imaging System (901YO, CFR 892.1560)
- Ultrasonic Pulsed Doppler Imaging System (901YN, CFR 892.1560)
- Diagnostic Ultrasonic Transducer (901TX, CFR 892.1570)

Identification of predicate, legally marketed device:
The predicate device(s) is(are):
- **ZONARE Medical Systems Inc.**:
  - ZONARE Z5 Ultrasound System (K120703)
    - Transducer Model Number: C10-3
- **Philips Healthcare, Inc.**:
  - CX50 Diagnostic Ultrasound System (K111513)
    - Transducer Type: C8-5
- **Toshiba America Medical Systems, Inc.**:
  - Diagnostic Ultrasound System (K121422)
    - Apio 500 TUS-A500 v2.1
    - Apio 400 TUS-A400 v2.1
    - Apio 300 TUS-A300 v2.1
  - Transducer Model Number: PVT-712BT

Device description:
2202 supports the following scanning modes and combinations thereof:
- B-mode, M-mode, CWD-mode, PWD mode and CFM mode, Tissue harmonic imaging, Contrast harmonic imaging.
- An optional ECG signal can be superimposed the ultrasound information in all modes and mode combinations.
- An optional 3-D unit can reconstruct a series of 2-D images into a single 3-D volume and display this on the screen.
The system can perform simple geometric measurements, and perform calculations in the areas of Vascular, Urology, Cardiology and OB/GYN applications.
The system can guide biopsy- and puncture needles.
An optional Vector Flow Imaging (VFI) module: Color Flow Mapping (CFM) imaging mode with the ability to visualize both the axial and the transverse velocity.

Transducers
Transducers are linear and convex phased arrays and mechanical sector.
The patient contact materials comply with ISO10993-1
All transducers used together with 2202 are Track 3 transducers.

**Acoustic output**
The system will assure that the acoustic output always will stay below the pre-amendments upper limits i.e. 
$I_{spt} \leq 720 \text{ mW/cm}^2$ and $MI \leq 1.9$ (Track 3, non ophthalmic).
The Thermal Index values are maximum 6.0, i.e. $TI \leq 6.0$

**Clinical measurement accuracy.**
Clinical measurements and calculations are described and accuracies are provided with the User Guide.

**Thermal, mechanical and electrical safety.**
The scanner 2202 has been tested by a recognized, certified body according to IEC 60601-1.

**Acoustic Output Reporting**
The Acoustic Output Reporting is made according to the standards required by "Information for Manufacturers Seeking Clearance of Diagnostic Ultrasound Systems and Transducers. FDA. CDRH. September 30, 2008."
The acoustic output is measured and calculated according to: "Acoustic Output Measurement Standard for Diagnostic Ultrasound Equipment" (AIUM 1998).

**Intended use.**
See comparison below
Technological characteristics compared to the predicate device(s).
The predicate device(s) have the same major technological characteristics as the subject device, see comparison below.

Comparison with the predicate devices mentioned above from:

<table>
<thead>
<tr>
<th>Supplier</th>
<th>BK Medical Systems</th>
<th>ZONARE Medical Systems</th>
<th>Philips</th>
<th>Toshiba</th>
<th>Comparison</th>
</tr>
</thead>
<tbody>
<tr>
<td>SLIK No.</td>
<td>K041352 &amp; K160919</td>
<td>K120703</td>
<td>K121754</td>
<td>K121422</td>
<td></td>
</tr>
<tr>
<td>Intended use</td>
<td>Diagnostic ultrasound, imaging and fluid flow analysis in the following applications</td>
<td>Diagnostic ultrasound, imaging and fluid flow analysis in the following applications</td>
<td>Diagnostic ultrasound, imaging and fluid flow analysis in the following applications</td>
<td>Vascular imaging of structures and devices present in the human body using ultrasound and in vivo correlate information for diagnosis at the following clinical applications</td>
<td>Comparable</td>
</tr>
<tr>
<td>Indications for use</td>
<td>Abdominal, Pelvic, Pediatric, Traperitoneal, Transabdominal, Transvaginal, Transrectal, Musculoskeletal, (conventional and superficial)</td>
<td>Abdominal, Pelvic, Pediatric, Traperitoneal, Intra-Aortic, Intra-Cardiac, Carotid, Transesophageal, Transesophageal (non-cardiac and superficial), Cardiac, Adult-Pediatric (Transthoracic &amp; Transesophageal), Musculoskeletal, (conventional &amp; superficial), (CVI, Doppler, Carotid, Adult-Pediatric (Trans), Femoral, Intra-Aortic, Bone, Ultrasound, Vascular, Conventional imaging and tissue characterization, Vascular)</td>
<td>Abdominal, Pelvic, Pediatric, Traperitoneal, Intra-Aortic, Intra-Cardiac, Carotid, Transesophageal, Transesophageal (non-cardiac and superficial), Cardiac, Adult-Pediatric (Transthoracic &amp; Transesophageal), Musculoskeletal, (conventional &amp; superficial), (CVI, Doppler, Carotid, Adult-Pediatric (Trans), Femoral, Intra-Aortic, Bone, Ultrasound, Vascular, Conventional imaging and tissue characterization, Vascular)</td>
<td>Abdominal, Pelvic, Pediatric, Traperitoneal, Intra-Aortic, Intra-Cardiac, Carotid, Transesophageal, Transesophageal (non-cardiac and superficial), Cardiac, Adult-Pediatric (Transthoracic &amp; Transesophageal), Musculoskeletal, (conventional &amp; superficial), (CVI, Doppler, Carotid, Adult-Pediatric (Trans), Femoral, Intra-Aortic, Bone, Ultrasound, Vascular, Conventional imaging and tissue characterization, Vascular)</td>
<td>Comparable</td>
</tr>
<tr>
<td>Transducer Specifics</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SLIK No.</td>
<td>K041352 &amp; K160919</td>
<td>K120703</td>
<td>K121754</td>
<td>K121422</td>
<td></td>
</tr>
<tr>
<td>Indication for use</td>
<td>Neonatal, Cephalic</td>
<td>Neonatal, Cephalic</td>
<td>Neonatal, Cephalic</td>
<td>Neonatal, Cephalic</td>
<td>Comparable</td>
</tr>
<tr>
<td>Frequency range</td>
<td>10-3.8 MHz</td>
<td>10-3 MHz</td>
<td>5-5 MHz</td>
<td>10-3 MHz</td>
<td>Comparable</td>
</tr>
</tbody>
</table>

Summary of Clinical Tests:
This submission introduces no new indications for use, modes, features or technologies relative to the predicate devices that require clinical testing. The clinical safety and effectiveness of ultrasound system with these characteristics are well accepted for both predicate and subject devices.
Conclusion: The device ProFocus 2202 and the probe N13C5 type 8862 in this application has similar intended uses, and in particular the subject for the submission, the addition of the new application Neonatal Cephalic, is the same.

B-K Medical ApS therefore considers that 2202 is substantially equivalent to the predicate devices.
January 6, 2014

B-K Medical ApS
% Mr. Gert Nielsen
Regulatory Affairs Manager
Mileparken 34
Herlev DK-2730
DENMARK

Re: K132685
   Trade/Device Name: Ultrasound Scanner Pro Focus 2202
   Regulation Number: 21 CFR 892.1560
   Regulation Name: Ultrasonic pulsed echo imaging system
   Regulatory Class: II
   Product Code: IYO, IYN, ITX
   Dated: December 5, 2013
   Received: December 19, 2013

Dear Mr. Nielsen:

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.

This determination of substantial equivalence applies to the following transducers intended for use with the Ultrasound Scanner Pro Focus 2202, as described in your premarket notification:

   Transducer Model Number

   N13C5 Type 8862

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 Small Manufacturers, International and Consumer Assistance 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" (21CFR 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 Small Manufacturers, International and Consumer Assistance 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.

Sincerely yours,

[Signature]

for

Janine M. Morris
Director, Division of Radiological Health
Office of In Vitro Diagnostics and Radiological Health
Center for Devices and Radiological Health

Enclosure
Indications for Use

510(k) Number (if known):
K132685

Device Name:
Ultrasound Scanner Pro Focus 2202
Ultrasound Transducer N13C5, Type 8862

Indications For Use:
Ultrasound scanner and transducers for B, Tissue and Contrast Harmonic Imaging, M, PWD, CWD, Color Doppler, Vector Flow Imaging and combined mode imaging.
Signal analysis and display.
Guidance of biopsy needles, geometrical measurements and calculation of parameters.
Non monitoring ECG for superimposing the ultrasound information.
An optional 3-D unit can reconstruct a series of 2-D images into a single 3-D volume and display this on the screen.
An optional Vector Flow Imaging (VFI) module: Color Flow Mapping (CFM) imaging mode with the ability to visualize both the axial and the transverse velocity.

Clinical applications:
Ultrasound Scanner Pro Focus 2202:
Fetal, Obstetrics, Abdominal, Intraoperative, Neurosurgery, Small organ, Pediatric, Neonatal Cephalic, Adult Cephalic, Cardiac, Transrectal, Transvaginal, Transurethral, Peripheral Vascular, Musculoskeletal.

Ultrasound Transducer N13C5, Type 8862: Neonatal Cephalic.

Details on specific Indication for Use forms

Prescription Use __X__ AND/OR Over-The-Counter Use ______
(Part 21 CFR 801 Subpart D) (21 CFR 807 Subpart C)

(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED)

Concurrence of CDRH; Office of In Vitro Diagnostics and Radiological Health (OIR)

Page 1 of 1
**Diagnostic Ultrasound Indications for Use Form**

**System:** 2202

Fill out one form for each ultrasound system and each transducer.

**Intended Use:** Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

<table>
<thead>
<tr>
<th>Clinical Application</th>
<th>Mode of Operation</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>A</td>
</tr>
<tr>
<td>Ophthalmic</td>
<td></td>
</tr>
<tr>
<td>Fetal 2) (K0435524)</td>
<td>P</td>
</tr>
<tr>
<td>Abdominal (K0435524)</td>
<td>P</td>
</tr>
<tr>
<td>Intraoperative (specify) (K0435524)</td>
<td>P</td>
</tr>
<tr>
<td>Intraoperative Neurological (K0435524)</td>
<td>P</td>
</tr>
<tr>
<td>Pediatric (K0435524)</td>
<td>P</td>
</tr>
<tr>
<td>Small Organ (specify) (K0435524)</td>
<td>P</td>
</tr>
<tr>
<td>Neonatal Cephalic</td>
<td>N</td>
</tr>
<tr>
<td>Adult Cephalic (K070077)</td>
<td>P</td>
</tr>
<tr>
<td>Cardiac (K0435524)</td>
<td>P</td>
</tr>
<tr>
<td>Transesophageal</td>
<td></td>
</tr>
<tr>
<td>Transrectal (K0435524)</td>
<td>P</td>
</tr>
<tr>
<td>Transvaginal (K0435524)</td>
<td>P</td>
</tr>
<tr>
<td>Transurethral (K0435524)</td>
<td>P</td>
</tr>
<tr>
<td>Intravascular</td>
<td></td>
</tr>
<tr>
<td>Peripheral Vascular (K0435524). (K009919 *)</td>
<td>P</td>
</tr>
<tr>
<td>Laparoscopic</td>
<td></td>
</tr>
<tr>
<td>Musculo-skeletal Conventional (K0435524)</td>
<td>P</td>
</tr>
<tr>
<td>Musculo-skeletal Superficial (K0435524)</td>
<td>P</td>
</tr>
<tr>
<td>Other (specify)</td>
<td></td>
</tr>
</tbody>
</table>

**N** = new indication; **P** = previously cleared by FDA; **E** = added under Appendix E

**Additional Comments:** 1) **B+M, B+D, B+C, B+D+C**

**B** mode includes **Tissue Harmonic Imaging**

**D** is **PWD, C is Color Doppler**

2) **Fetal is often called Obstetrics**

3) **Vector Flow Imaging**

(PLEASE DO NOT WRITE BELOW THIS LINE - CONTINUE ON ANOTHER PAGE IF NEEDED)

Concurrence of CDRH; Office of In Vitro Diagnostics and Radiological Health (OIR)

Prescription Use (Per 21 CFR 801.109)
## Diagnostic Ultrasound Indications for Use Form

### System:
2202

### Transducer:
8862

**Intended Use:** Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

<table>
<thead>
<tr>
<th>Clinical Application</th>
<th>Mode of Operation</th>
</tr>
</thead>
<tbody>
<tr>
<td>General (Track I Only)</td>
<td>Specific (Tracks I &amp; II)</td>
</tr>
<tr>
<td>Ophthalmic</td>
<td>Ophthalmic</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>Abdominal</td>
<td></td>
</tr>
<tr>
<td>Intra-operative (Specify 2)</td>
<td>(K043524)</td>
</tr>
<tr>
<td>Intra-operative (Neuro)</td>
<td>(K043524)</td>
</tr>
<tr>
<td>Laparoscopic</td>
<td></td>
</tr>
<tr>
<td>Pediatric</td>
<td>(K043524)</td>
</tr>
<tr>
<td>Small Organ (Specify)</td>
<td></td>
</tr>
<tr>
<td>Neonatal Cephalic</td>
<td>N</td>
</tr>
<tr>
<td>Adult Cephalic</td>
<td></td>
</tr>
<tr>
<td>Trans-rectal</td>
<td></td>
</tr>
<tr>
<td>Trans-vaginal</td>
<td></td>
</tr>
<tr>
<td>Trans-urethral</td>
<td></td>
</tr>
<tr>
<td>Trans-esoph. (non-Card.)</td>
<td></td>
</tr>
<tr>
<td>Musculo-skel. (Conventional)</td>
<td></td>
</tr>
<tr>
<td>Musculo-skel. (Superficial)</td>
<td></td>
</tr>
<tr>
<td>Intra-luminal</td>
<td></td>
</tr>
<tr>
<td>Other (Specify)</td>
<td></td>
</tr>
<tr>
<td>Cardiac</td>
<td></td>
</tr>
<tr>
<td>Cardiac Adult</td>
<td></td>
</tr>
<tr>
<td>Cardiac Pediatric</td>
<td></td>
</tr>
<tr>
<td>Trans-esoph. (Cardiac)</td>
<td></td>
</tr>
<tr>
<td>Other (Specify)</td>
<td></td>
</tr>
<tr>
<td>Peripheral Vessel</td>
<td>Peripheral vessel</td>
</tr>
<tr>
<td>Other (Specify)</td>
<td></td>
</tr>
</tbody>
</table>

**Notes:**
- N = new indication; P= previously cleared by FDA; E= added under Appendix E
- *Examples may include: A-mode, Amplitude Doppler, 3-D Imaging, Harmonic Imaging, Tissue Motion Doppler, Color Velocity Imaging

**Additional Comments:**
1) Mode combinations: B+M, B+D, B+C, B+D+C. (D is PWD, C is Color Flow mapping Doppler including Amplitude(power)Doppler)
2) Intraoperative: Gall bladder

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(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED)

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Prescription Use (Per 21 CFR 801.109)