

510(k) Summary  
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1/30/2014

**MAR 04 2014**

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**Official Contact:** Renee Thornborough – Director QA/RA

**Proprietary or Trade Name:** Model HEM-7320

**Common/Usual Name:** Noninvasive blood pressure measurement system.

**Classification Name/Code:** DXN – Noninvasive blood pressure measurement system.  
21CFR 870.1130  
Class II

**Device:** Model HEM-7320

**Predicate Devices:** Omron – HEM -780N3 (HEM-7080-Z)- K061822  
Omron – BP742 (HEM-7200-Z) – K121932

**Device Description:**

The device is an automatic non-invasive blood pressure system. The device is battery powered and can also be powered from an IEC 60601-1 compliant AC adaptor. The device inflates a cuff with an integral pump, then deflates the cuff via an electronically controllable valve. During deflation the cuff pressure is monitored and pulse waveform data is extracted. The extracted pulse waveform data is then analyzed by software which determines pulse rate, as well as systolic and diastolic pressure.

The device is intended to be used with an Omron specified cuff to encompass arms ranging from 9 to 17 inches in circumference.

The device also detects the appearance of irregular heartbeats during measurement.

The device has provisions for selecting two users, measurements from these users are stored in memory. The memory stores up to 100 of the latest measurements. It can also display an average of the last three values.

Note that a number of documents in this submission include reference to the Cuff Wrapping Guide (or Cuff Wrapping Check). This feature has been removed from the device and is not part of this submission.

**Intended User**  
Home user

**Patient Population**

This device is intended for use on adults.

**Indications for Use:**

The device is a digital monitor intended for use in measuring blood pressure and pulse rate in adult patient population.

The device detects the appearance of irregular heartbeats during measurement and gives a warning signal with readings.

**Environment of Use:**

Home

**Contraindications:**

There are no known contraindications.

**Predicate Device Comparison:**

The HEM-7320 was compared to the predicates HEM -780N3 (HEM-7080-Z) - K061822 and BP742 (HEM-7200-Z) – K121932 in the device comparison table below.

Device Comparison

	Predicate Device	Predicate Device	New Device
Model Name:	HEM -780N3 (HEM-7080-Z)	BP742(HEM-7200-Z)	HEM-7320
510(k) Number	K061822	K121932	TBD
Indications	The device is a digital monitor intended for use in measuring blood pressure and pulse rate in adult patient population with arm circumference ranging from 9 inches to 17 inches (22 cm - 42 cm). The device detects the appearance of irregular heartbeats during measurement and gives a warning signal with readings. This blood pressure monitor compares average blood pressure results to pre-established AHA (American Heart Association) hypertension guideline of 135/85 mmHg. The Omron 780N3 model is not intended to be a diagnostic device. Contact your physician if hypertensive values are indicated.	The device is a digital monitor intended for use in measuring blood pressure and pulse rate in adult patient population. The device detects the appearance of irregular heartbeats during measurement and gives a warning signal with readings.	The device is a digital monitor intended for use in measuring blood pressure and pulse rate in adult patient population. The device detects the appearance of irregular heartbeats during measurement and gives a warning signal with readings. <b>Identical to K121932</b>
Environment of Use	Home	Home	Home <b>Identical to both predicates</b>
Patient Population	Adult	Adult	Adult <b>Identical to both predicates</b>
<b>Specifications / Features</b>			
Measurement method / Principal of operation	HEM-7080-Z(HEM 780N3) Cuff oscillometric method	BP742(HEM-7200-Z) Cuff oscillometric method	HEM-7320 Cuff oscillometric method. <b>Identical to both predicates</b>
Measurement range	Pressure: 0 to 294 mmHg Pulse Rate: 40 to 180 beats/min.	Pressure: 0 to 299 mmHg Pulse Rate: 40 to 180 beats/min.	Pressure: 0 to 299 mmHg Pulse Rate: 40 to 180 beats/min. <b>Identical to K121932</b>
Pressure sensor	Electrostatic capacitive sensor	Semiconductor pressure sensor	Semiconductor pressure sensor. <b>Identical to K121932</b>

	HEM-7080-Z(HEM 780N3)	BP742(HEM-7200-Z)	HEM-7320
Applicable cuff (Arm Circumference)	22-42cm	22-42cm	22-42cm, identical to both predicates
Accuracy of pressure indicator	Within $\pm 3$ mmHg or 2 % of reading	Within $\pm 3$ mmHg or 2 % of reading	Within $\pm 3$ mmHg or 2 % of reading <b>Identical to both predicates</b>
Accuracy of pulse rate	Within $\pm 5$ % of reading	Within $\pm 5$ % of reading	Within $\pm 5$ % of reading <b>Identical to both predicates</b>
Inflation method	Automatic by electric pump	Automatic by electric pump	Automatic by electric pump. <b>Identical to both predicates</b>
Deflation method	Active electronic control valve	Automatic pressure release valve	Automatic pressure release valve <b>Identical to K121932</b>
Display	LCD digital display	LCD digital display	LCD digital display <b>Identical to both predicates</b>
Power Source	4" AA"batteries or AC adapter	4" AA"batteries or AC adapter	4" AA"batteries or AC adapter <b>Identical to both predicates</b>
Operating conditions	10 to 40 °C (50 to 104 °F) 30 to 85 %RH	10 to 40 °C 15 to 90 %RH	10 to 40 °C 15 to 90 %RH <b>Identical to K121932</b>
Storage conditions	-20 to 60 °C 10 to 95 %RH	-20 to 60 °C 10 to 95 %RH	-20 to 60 °C 10 to 95 %RH <b>Identical to K121932</b>
Dimensions (mm)	131 (W) $\times$ 155 (D) $\times$ 84 (H) mm without cuff and battery	124(W) $\times$ 161(D) $\times$ 90(H)mm	124 (W) $\times$ 161 (D) $\times$ 90 (H) mm, size is not a factor in function of the device
Weight	Approx. 420g (not including cuff and batteries)	Approx. 340g (not including battery)	Approx. 380g (13 3/8 oz) (not including battery), weight is not a factor in function of the device.
Irregular Heart beat Feature	Yes	Yes	Yes, <b>identical to both predicates</b>
Body movement detection	Yes	Yes	Yes, <b>identical to both predicates</b>
Hypertension indicator	No	Yes	Yes, <b>identical to K121932</b>
TruRead™	Yes	No	Yes, <b>identical to K061822</b>

Technology / Features Comparison		HEM-7080-Z(HEM 780N3)	BP742(HEM-7200-Z)	HEM-7320
Power supply	Regulates power voltage regardless of battery voltage.	Regulates power voltage regardless of battery voltage.	Regulates power voltage regardless of battery voltage.	Regulates power voltage regardless of battery voltage. <b>Identical to both predicates</b>
Microprocessor	<ul style="list-style-type: none"> <li>determines blood pressure and pulse rate</li> <li>controls the pump, the valve, and the display</li> <li>detects switch operations</li> <li>stores measurement results</li> <li>manages date and time</li> </ul>	<ul style="list-style-type: none"> <li>determines blood pressure and pulse rate</li> <li>controls the pump, the valve, and the display</li> <li>detects switch operations</li> <li>stores measurement results</li> <li>manages date and time</li> </ul>	<ul style="list-style-type: none"> <li>determines blood pressure and pulse rate</li> <li>controls the pump, the valve, and the display</li> <li>detects switch operations</li> <li>stores measurement results</li> <li>manages date and time</li> </ul>	<ul style="list-style-type: none"> <li>determines blood pressure and pulse rate</li> <li>controls the pump, the valve, and the display</li> <li>detects switch operations</li> <li>stores measurement results</li> <li>manages date and time</li> </ul> <b>Identical to both predicates</b>
Pressure sensor	Electrostatic capacitive sensor	Semiconductor pressure sensor	Semiconductor pressure sensor	Semiconductor pressure sensor, <b>identical to K121932</b>
Rapid exhaust Valve	Active electronic control valve that performs cuff air bleeding and release	Automatic rapid air release valve.	Automatic rapid air release valve.	Active electronic control valve that performs cuff air bleeding and release, <b>identical to K061822</b>
Deflation Valve	Automatic pressure release valve	Automatic pressure release valve	Automatic pressure release valve	Automatic pressure release valve, <b>identical to K061822</b>
Inflation source	DC rolling pump	DC rolling pump	DC rolling pump	DC rolling pump <b>Identical to both predicates</b>
Display	LCD (Liquid Crystal Display) displays; <ul style="list-style-type: none"> <li>current cuff pressure</li> <li>systolic blood pressure</li> <li>diastolic blood pressure</li> <li>pulse rate</li> <li>error messages</li> <li>measurement results in the memory</li> </ul>	LCD (Liquid Crystal Display) displays; <ul style="list-style-type: none"> <li>current cuff pressure</li> <li>systolic blood pressure</li> <li>diastolic blood pressure</li> <li>pulse rate</li> <li>error messages</li> <li>measurement results in the memory</li> </ul>	LCD (Liquid Crystal Display) displays; <ul style="list-style-type: none"> <li>current cuff pressure</li> <li>systolic blood pressure</li> <li>diastolic blood pressure</li> <li>pulse rate</li> <li>error messages</li> <li>measurement results in the memory</li> </ul>	LCD (Liquid Crystal Display) displays; <ul style="list-style-type: none"> <li>current cuff pressure</li> <li>systolic blood pressure</li> <li>diastolic blood pressure</li> <li>pulse rate</li> <li>error messages</li> <li>measurement results in the memory</li> </ul> <b>Identical to both predicates</b>
Controls	<ul style="list-style-type: none"> <li>START/STOP Button</li> <li>Setting Button</li> <li>Memory button</li> <li>User ID Selections Button</li> <li>Morning/Evening Average Buttons</li> </ul>	<ul style="list-style-type: none"> <li>START/STOP Button</li> <li>Date/Time setting Button</li> <li>Up/Down Button</li> <li>User ID Selections Button</li> </ul>	<ul style="list-style-type: none"> <li>START/STOP Button</li> <li>Date/Time setting Button</li> <li>Up/Down Button</li> <li>User ID Selections Button</li> </ul>	<ul style="list-style-type: none"> <li>START/STOP Button</li> <li>Date/Time setting Button</li> <li>Up/Down Button</li> <li>User ID Selections Button</li> </ul> <b>Identical to K121932</b>

	<b>HEM-7080-Z(HEM 780N3)</b>	<b>BP742(HEM-7200-Z)</b>	<b>HEM-7320</b>
Cuff	The cuff is applied to the arm circumference between 9-17 inches(22- 42 cm). The cuff has an elasticity board to hold the arm.	Wide Range Cuff, Standard Adult Arm Cuff Large Cuff	ComFit Cuff(HEM-FL31) 22-42cm, new cuff to reduce dependence on bladder placement relative to the brachial artery
Materials	Patient contact materials of the cuff have been cleared in the referenced 510(k)	Patient contact materials of the cuff have been cleared in the referenced 510(k)	Patient contact materials of the cuff have been tested in accordance with ISO 10993 and FDA guidance

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**Differences Between Other Legally Marketed Predicate Devices:**

The Omron HEM-7320 is viewed as substantially equivalent to the predicate devices because: The HEM-7320 uses the exact same technology and has identical indications for use. The differences that exist between the devices are insignificant in the terms of safety or effectiveness.

**Indications –**

The device is a digital monitor intended for use in measuring blood pressure and pulse rate in adult patient population.

The device detects the appearance of irregular heartbeats during measurement and gives a warning signal with readings.

Discussion – These indications are identical to the predicate Omron BP742 (K121932).

**Prescriptive –** The HEM-7320 and predicates are all OTC.

**Design and Technology –** The HEM-7320 has equivalent design and features when compared to the predicates and has the identical technology to the predicate.

**Performance and Specifications –** The HEM-7320 has equivalent specifications of performance when compared to the predicate.

**Compliance with standards –** The predicate devices declare compliance with SP10, IEC 60601-1 and IEC 60601-1-2. The HEM-7320 complies with AAMI ANSI ES6060-1 (which replaced IEC 60601-1), IEC 60601-1-2 and ANSI/AAMI/ISO 81060-2 (which replaced SP10). The HEM-7320 also complies with IEC 80601-2-30 and IEC 60601-1-11 for home healthcare.

**Materials –**

The patient contacting materials of the cuffs has been tested in accordance with ISO 10993-1 and FDA Guidance. The tests included Cytotoxicity, Sensitization, and Intracutaneous Reactivity.

**Patient Population –**

The HEM-7320 and predicates are indicated for adults

**Environment of Use –** Home, Identical to the predicates

**Non-Clinical Testing Summary:**

We have performed bench tests and found that the HEM-7320 met all requirements specifications and standards requirements and were found to be equivalent in comparison to the predicate. Testing includes the following:

- Verification Testing to insure the device meets its specifications

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- Testing of hazard mitigations
- Testing for compliance to AAMI ANSI ES60601-1
- Testing for compliance to IEC 60601-1-2
- Testing for compliance to IEC 80601-2-30
- Testing for compliance to IEC 60601-1-11
- Comparative Testing to the predicates

**Clinical Testing Summary:**

Testing to insure clinical accuracy of the device in accordance with ANSI/AAMI/ISO 81060-2 as documented in **Section 20**.

Eighty five patients (40 males and 45 females) were recruited for the study.

Standard auscultation method was used as the reference blood pressure (BP) measuring in the left upper arm. BP measurements were repeated alternatively with the device and auscultation in the same arm according to the sequence in AAMI.

**Substantial Equivalence Conclusion**

Omron maintains that the HEM-7320 is substantially equivalent to the predicate device in indications for use, patient population, and environment for use, technology characteristics, specifications / performance and compliance with international standards



DEPARTMENT OF HEALTH & HUMAN SERVICES

Public Health Service

Food and Drug Administration  
10903 New Hampshire Avenue  
Document Control Center - WO66-G609  
Silver Spring, MD 20993-0002

March 4, 2014

Omron Healthcare, Inc.  
c/o Paul Dryden  
24301 Woodsage Dr.  
Bonita Springs, FL 34134 US

Re: K133383  
Trade/Device Name: HEM-7320  
Regulation Number: 21 CFR 870.11307  
Regulation Name: Noninvasive Blood Pressure  
Regulatory Class: Class II  
Product Code: DXN  
Dated: January 31, 2014  
Received: November 6, 2013

Dear Paul Dryden:

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 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" (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 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,

The logo of the U.S. Food and Drug Administration (FDA), featuring the letters 'FDA' in a stylized, bold font with a graphic element resembling a caduceus or a similar medical symbol behind the letters.

for Bram D. Zuckerman, M.D.

Director

Division of Cardiovascular Devices

Office of Device Evaluation

Center for Devices and

Radiological Health

Enclosure

**Indications for Use**

510(k) Number (if known)  
K133383

Device Name  
Omron HEM-7320

Indications for Use (Describe)

The device is a digital monitor intended for use in measuring blood pressure and pulse rate in adult patient population.  
The device detects the appearance of irregular heartbeats during measurement and gives a warning signal with readings.

K133383

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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)

PLEASE DO NOT WRITE BELOW THIS LINE – CONTINUE ON A SEPARATE PAGE IF NEEDED.

FOR FDA USE ONLY

Concurrence of Center for Devices and Radiological Health (CDRH) (Signature)

 Date: 2014-03-04  
14:53:17-05'00'  
for Bram Zuckerman