

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

July 23, 2015

Grandway Technology (shenzhen) Limited Patrick Chow Block 6 And 7, Zhu Keng Industrial Zone, Ping Shan, Long Gang District, Shenzhen, 518118 CN

Re: K143735

Trade/Device Name: Digital Automatic Blood Pressure Monitor MD2300

Regulation Number: 21 CFR 870.1130

Regulation Name: Noninvasive Blood Pressure Measurement System

Regulatory Class: Class II Product Code: DXN Dated: June 24, 2015

Received: June 24, 2015

Dear Patrick Chow,

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 <u>Federal Register</u>.

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

http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm.

Sincerely yours,

for Bram D. Zuckerman, M.D.

Director

Division of Cardiovascular Devices

Office of Device Evaluation

Center for Devices and Radiological Health

Enclosure

DEPARTMENT OF HEALTH AND HUMAN SERVICES Food and Drug Administration

Indications for Use

Form Approved: OMB No. 0910-0120 Expiration Date: January 31, 2017 See PRA Statement below.

alt's arterial blood flatable cuff wrapped ags together with the

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.

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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."

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1. Submitter Identification

510(k) Submitter	GRANDWAY TECHNOLOGY (SHENZHEN) LIMITED				
Address	Block 6 and 7, Zhu Keng Industrial Zone, Ping Shan, Long Gang				
	District, Shenzhen, Guang Dong, People's Republic of China				
Phone Number	(00852)-2851-6789				
Fax Number	(00852)-2851-6278				
Contact Person	Mr. Patrick Chow				
Date of Submission	29-Dec-2014				

2. Device Identification

Trade Name	Digital Automatic Blood Pressure Monitor MD2300
Common Name	Non-invasive Blood Pressure Measurement System
Classification Name	Non-invasive Blood Pressure Measurement System
	(CFR 870.1130, Class II, Product Code DXN)

3. Predicate Device

Predicate Device	Digital Automatic Blood Pressure Monitor BPM18 Series	
Manufacturer	GRANDWAY TECHNOLOGY (SHENZHEN) LIMITED	
510(k) Number	K133619	

4. Device Description

Digital Automatic Blood Pressure Monitor MD2300 is a non-invasive blood pressure measurement system for use by medical professional or at home. It is designed to measure the systolic and diastolic blood pressure, and pulse rate of an individual in each measurement and then display the readings on a digital panel.

The device utilizes the oscillometric methodology, in which an inflatable cuff is wrapped around the upper arm of an individual, for blood pressure measurement. This means the monitor detects your blood's movement through your brachial artery and converts the movement into a digital reading.

The table below illustrate the feature presence in Digital Automatic Blood Pressure Monitor MD2300.

Model	Blood Pressure	Pulse Rate	WHO	Irregular	LCD Type	Backlight	User ×
Model	Measurement	Measurement	Classification	Heartbeat	LCD Type	Illumination	Memory
MD2300	~	•	~	>	Positive Transmissive	>	2 × 60

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5. Indication for Use

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The upper arm blood pressure monitor is used for non-invasive measurement and monitoring for adult's arterial blood pressure. You can use it to measure your systolic and diastolic pressure, and pulse rate through an inflatable cuff wrapped around the upper arm. Quickly and easily, storing the results and displaying the progression of readings together with the average.

6. Comparison of Technological Characteristics between New Device and Predicate Devices

Digital Automatic Blood Pressure Monitor MD2300 is compared to the predicate device, BPM18 Series (K133619) in the device comparison table below.

Comparison between BPM18 Series and Predicate device				
Item	Predicate Device	MD2300	Comment	
Indication for Use	Digital Automatic Blood Pressure Monitor	The upper arm blood pressure monitor is	Equivalent	
	BPM18 Series is for use by medical	used for non-invasive measurement and		
	professional or home user. The BPM18 Series	monitoring for adult's arterial blood		
	is intended to measure the systolic and	pressure. You can use it to measure your		
	diastolic blood pressure, and pulse rate of an	systolic and diastolic pressure, and pulse		
	adult individual by using a non-invasive technique, in which an inflatable cuff is	rate through an inflatable cuff wrapped		
La de la companya de	wrapped around the upper arm.	around the upper arm. Quickly and easily,		
	wrapped around the upper arm.	storing the results and displaying the		
		progression of readings together with the		
		average.		
Measurement	Non-invasive, Oscillometric	Non-invasive, Oscillometric	Identical	
Method			·	
IHB Detection	Yes	Yes	Identical	
Patient Population	Adult	Adult	Identical	
BP Measurement	Cuff Pressure: 0 - 300 mmHg	Cuff Pressure: 0 - 300 mmHg	Identical	
Range	Systolic Pressure: 50 - 250 mmHg	Systolic Pressure: 50 - 250 mmHg		
	Diastolic Pressure: 30 - 200 mmHg	Diastolic Pressure: 30 - 200 mmHg		
Number of User	2 independent users	2 independent users	Identical	
Memory Space	2 users × 120 memory space	2 users × 60 memory space	Equivalent	
Resolution of	Blood Pressure: 1 mmHg or 0.1kPa	Blood Pressure: 1 mmHg	Identical	
Measurement	Pulse Rate: 1 beat/ min	Pulse Rate: 1 beat/ min		
Blood Pressure	± 3 mmHg or 2% of reading	± 3 mmHg	Equivalent	
Measurement				
Accuracy				
Pulse Rate	40 - 180 beats/min	40 - 180 beats/min	Identical	
Measurement				
Range				
Pulse Rate	\pm 5 % of the reading	\pm 5 % of the reading	Identical	
Measurement				
Accuracy				
Display Type	LCD	LCD	Identical	
Power Source	4 × 1.5 V AAA-batteries and/or AC	4 × 1.5 V AA-batteries and/or AC	Equivalent	
	Adaptor	Adaptor		
Pressurization	Automatic Inflation	Automatic Inflation	Identical	
Mode				
Deflation Mode	Automatic Exhaust/ Deflation	Automatic Exhaust/ Deflation	Identical	
Operating	Temperature: +5 to +40 °C	Temperature: +5 to +40 °C	Identical	
Condition	Humidity: 15 to 93 % R.H. max	Humidity: 15 to 93 % R.H. max		
	Atmospheric Pressure: 700 - 1060 hPa	Atmospheric Pressure: 700 - 1060 hPa		

Comparison between BPM18 Series and Predicate device					
Item	Predicate Device	MD2300	Comment		
Storage and	Temperature: -25 to +70 °C	Temperature: -25 to +70 °C	Identical		
Transportation	Humidity: up to 93% R.H. max	Humidity: up to 93% R.H. max			
Condition	Atmospheric Pressure: 700 - 1060 hPa	Atmospheric Pressure: 700 - 1060 hPa			
Material	Resistances, capacitance, transistors, amplifiers, pressure sensor, CPU, PCB, cuff ABS button, ABS cabinet, batteries	Resistances, capacitance, transistors, amplifiers, pressure sensor, CPU, PCB, cuff ABS button, ABS cabinet, batteries	Identical		
	and packaging	and packaging			
Compatibility with	No influence with environment and other	No influence with environment and other	Identical		
Environment and	device	device			
Other Devices					
Applicable	- EN 1060-1:1995+A2:2009	- EN 1060-1:1995+A2:2009	Equivalent		
Standard	- EN 1060-3:1997+A2:2009	- EN 1060-3:1997+A2:2009			
	- IEC 60601-1:2012	- IEC 60601-1:2012			
	- EN 60601-1-2:2007	- IEC 60601-1-2:2007			
	- FCC Part 15 Subpart B	- FCC Part 15 Subpart B	**		
	- ISO 10993-5:2009	- ISO 10993-5:2009			
	- ISO 10993-10:2010	- ISO 10993-10:2010			
	- IEC 62304:2006	- IEC 62304:2006			
	- IEC 81060-2:2009	- IEC 81060-2:2013			

Digital Automatic Blood Pressure Monitor MD2300 is a non-invasive measuring device and utilizes the oscillometric methodology to measure the blood pressure reading. The key components of device are: a pressure sensor, a electric valve and an electronic control module together with an electric pump. The electric pump inflate (and deflate) the inflatable cuff automatically according to our designed architecture. The predicate device adopts exactly same methodology and key components for measuring blood pressure.

7. Clinical and Non-clinical Tests

Clinical Test Summary

Testing to insure clinical accuracy of the device in accordance with ISO 81060-2:2013 as documented in Clinical Test report.

One hundred patients (49 males and 51 females) were invited for the study. Standard auscultation method was used as the reference blood pressure monitor measuring in the left arm. Blood pressure measurements were repeated alternatively with the device and auscultation in the same arm according to the sequence in ISO 81060-2:2013.

Non-Clinical Test Summary

Digital Automatic Blood Pressure Monitor MD2300 has performed several non-clinical tests to show that all requirement specifications and standard requirements are met. The tests includes the follows:

- ♦ EN 1060-1:1995+A2:2009
- ♦ EN 1060-3:1997+A2:2009
- ♦ IEC 60601-1:2012
- ♦ IEC 60601-1-2:2007
- ♦ FCC Part 15 Subpart B
- ♦ ISO 10993-5:2009
- ♦ ISO 10993-10:2010

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♦ IEC 62304:2006

All of the clinical and non-clinical testing performed on Digital Automatic Blood Pressure Monitor MD2300 are same as the predicate device.

Also, bench testing, IEC 80601-2-30, is conducted to show the performance of Digital Automatic Blood Pressure Monitor MD2300 is equivalent to the predicate device.

8. Conclusion

Digital Automatic Blood Pressure Monitor MD2300 has the same intended use and same technological characteristics as the predicate device, Digital Automatic Blood Pressure Monitor BPM18 Series (K133619). Moreover both clinical and non-clinical testing has demonstrated that no differences in the technological characteristics and questioning on safety or effectiveness to be raised. Thus, Digital Automatic Blood Pressure Monitor MD2300 is substantially equivalent to the predicate device.