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Safety Notice

satisfactory use

Thank you for purchasing the DBP-1319b Blood Pressure Monitor. The unit has been constructed using reliable circuitry and durable materials. Used properly, this unit will provide yeas of

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Measure blood pressure(systolic and diastolic) and pulse rate of adults and adolecents age 12 through 21 years of age. All functions can be used safely and values can be read out in one LCD DISPLAY. Measurement position is on adult upper arm only. The PATIENT is an intended OPERATOR.

Blood pressure measurement determined with this device are equivalent to those obtained by a trained observer using the cuff/ stethoscope auscultation method, within the limits prescribed by the Recognized Consensus Standard (IEC 81060-2-30) for electronic sphygmomanometers.

Precautions to Ensure Safe, Reliable Operation

- 1. Do not drop the unit. Protect it from sudden jars or shocks.
- 2. Do not insert foreign objects into any openings.
- 3. Do not attempt to disassemble the unit.
- 4. Do not crush the pressure cuff.
- 5. If the unit has been stored at temperatures below 0 oC, leave it in a warm place for about 15 minutes before using it. Otherwise, the cuff may not inflate properly.
- 6. If the unit has been stored at temperatures above 40 oC, leave it in a cool place for about 15 minutes before using it. Otherwise, the cuff may not inflate properly.
- 7. Do not store the unit in direct sunlight, high humidity or dust.
- 8. To avoid any possibility of accidental strangulation, keep this unit away from children and do not drape tubing around your neck.
- 9.Ensure that children do not use the instrument unsupervised; some parts are small enough to be swallowed.
- 10.Some may get a skin irritation from the cuff taking frequent readings over the course of the day, but this irritation typically goes away on its own after the monitor is removed.

Safety Notice

- Important Instructions Before Use
- 1. Do not confuse self-monitoring with self-diagnosis. Blood pressure measurements should only be interpreted by a health professional who is familiar with your medical history. 2. Contact your physician if test results regularly indicate abnormal readings.
- 3. If you are taking medication, consult with your physician to determine the most appropriate time to measure your blood pressure. NEVER change a prescribed medication without first consulting with your physician.
- 4. Individuals with serious circulation problems may experience discomfort. Consult your physician prior to use.
- 5. For persons with irregular or unstable circulation resulting from diabetes, liver disease, arteriosclerosis or other medical conditions, there may be variations in blood pressure values measured at the wrist versus at the upper arm. Monitoring the trends in your blood pressure taken at either the arm or the wrist is nevertheless useful and important.
- 6. People suffering from vascular constriction, liver disorders or diabetes, people with cardiac pacemakers or a weak pulse, and women who are pregnant should consult their physician before measuring their blood pressure themselves. Different values may be obtained due to their condition.
- 7. People suffering from arrhythmias such as atrial or ventricular premature beats or atrial fibrillation only use this blood pressure monitor in consultation with your doctor. In certain cases oscillometric measurement method can produce incorrect readings
- 8. Too frequent measurements can cause injury to the patient due to blood flow interference. 9. The cuff should not be applied over a wound as this can cause further injury.
- 10.DO NOT attach the cuff to a limb being used for IV infusions or any other intravascular access, therapy or an arterio-venous (A-V) shunt. The cuff inflation can temporarily block blood flow, potentially causing harm to the patient.
- 11. The cuff should not be placed on the arm on the side of a mastectomy. In the case of a double mastectomy use the side of the least dominant arm
- 12. Pressurization of the cuff can temporarily cause loss of function of simultaneously used monitoring equipment on the same limb.
- 13.A compressed or kinked connection hose may cause continuous cuff pressure resulting in blood flow interference and potentially harmful injury to the patient.
- 14. Check that operation of the unit does not result in prolonged impairment of the circulation of the patient.
- 15. Product is designed for its intended use only. Do not misuse in any way.
- 16. Product is not intended for infants or individuals who cannot express their intentions.
- 17. Prolonged over-inflation of the bladder may cause ecchymoma of your arm.
- 18. Do not disassemble the unit or arm cuff. Do not attempt to repair.
- 19. Use only the approved arm cuff for this unit. Use of other arm cuffs may result in incorrect measurement results.
- 20. The system might produce incorrect readings if stored or used outside the manufacturer's specified temperature and humidity ranges. Make sure to store the blood pressure monitor, children, pets and pests are outside of accessible range.
- 21. Do not use the device near strong electrical or electromagnetic fields generated by cell phones or other devices, they may cause incorrect readings and interference or become interference source to the device.
- 22. Do not mix new and old batteries simultaneously.

Safety Notice

Federla Commulcation Commission (FCC) Interference Statement

1. This device complies with part 15 of the FCC Rules. Operation is subject to the condition that this device does not cause harmful interference.

2. This device is verified to comply with part 15 of the FCC Rules for use with cable television service.

3. This device complies with part 15 of the FCC Rules. Operation is subject to the following two conditions:

(1) This device may not cause harmful interference, and

(2) this device must accept any interference received, including interference that may cause undesired operation. Please note that changes or modifications not expressly approved by the party responsible for compliance could void the user's authority to operate the equipment. 4. This equipment has been tested and found to comply with the limits for a Class B digital device, pursuant to part 15 of the FCC Rules. These limits are designed to provide reasonable protection against harmful interference in a residential installation. This equipment generates, uses and can radiate radio frequency energy and, if not installed and used in accordance with the instructions, may cause harmful interference to radio communications. However, there is no guarantee that interference will not occur in a particular installation. If this equipment does cause harmful interference to radio or television reception, which can be determined by turning the equipment off and on, the user is encouraged to try to correct the interference by one or more of the following measures:

-Reorient or relocate the receiving antenna.

-Increase the separation between the equipment and receiver.

-Connect the equipment into an outlet on a circuit different from that to which the receiver is connected.

-Consult the dealer or an experienced radio/TV technician for help.

Safety Notice

- 23. Replace batteries when Low Battery Indicator "ix appears on screen. Replace both batteries at the same time.
- 24. Do not mix battery types. Long-life alkaline batteries are recommended.
- 25. Remove batteries from device when not in operation for more than 3 months. 26. Dispose batteries properly; observe local laws and regulations.
- 27. Only use a recommended AC adaptor double-insulated complying with EN 60601-1 and EN 60601-1-2. An unauthorized adapter may cause fire and electric shock.



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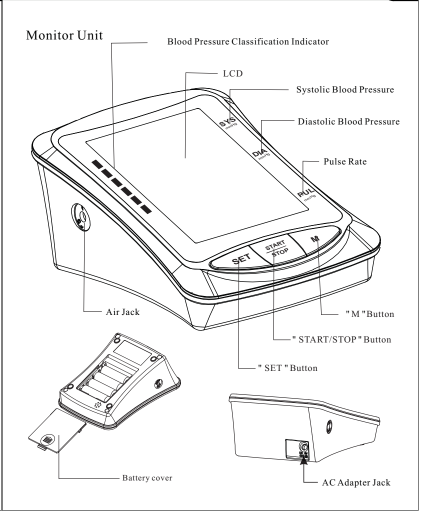
5

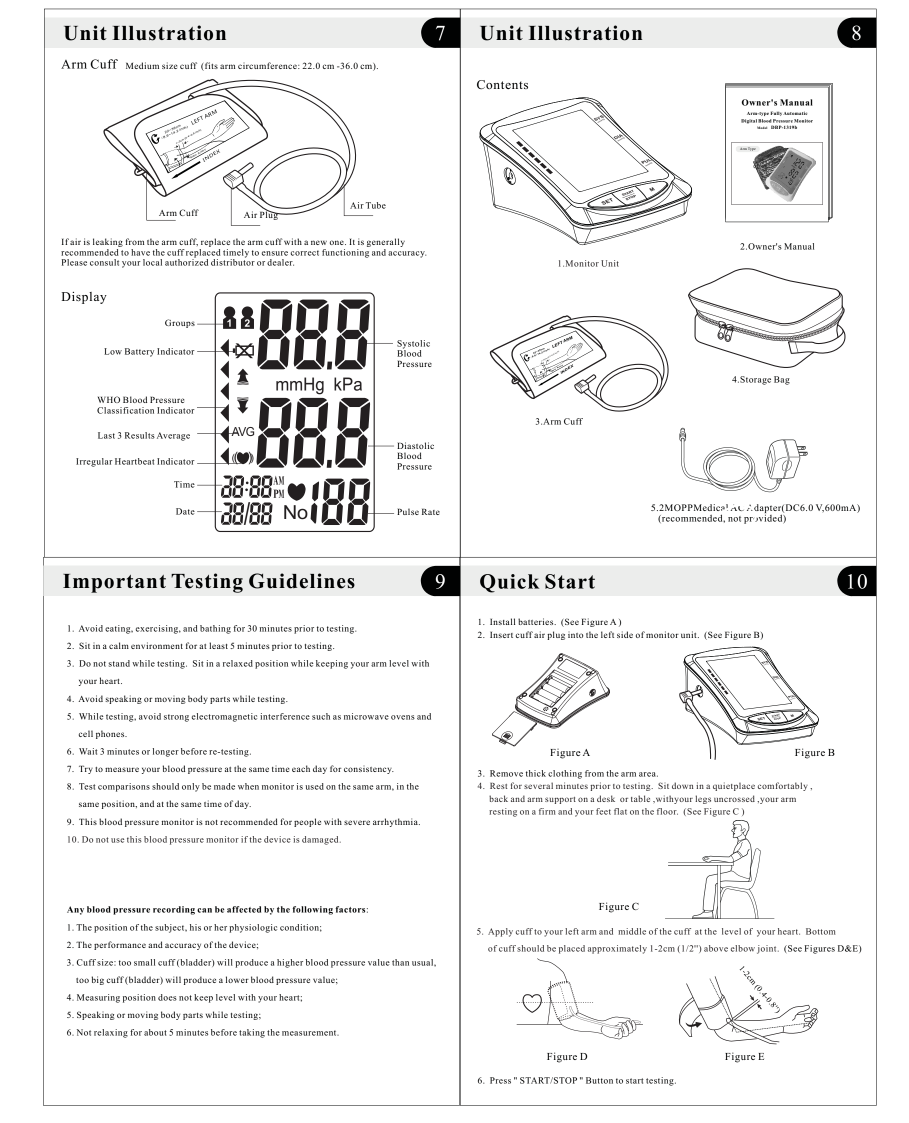
Advising operator that Instruction manual/ Booklet must be consulted. 29. Do not use the device during transport vehicles for influencing measurement accuracy,

- such as patient transport in an ambulance or helicopter.
- 30. Contains small parts that may cause a chocking hazard if swallowed by infants. 31. Please align the polarities of each battery with the +ve and -ve signs
- imprinted on the battery housing when you replace the batteries
- 32. This ME equipment or ME systems should be only used in shielded location. 33. Use of this equipment adjacent to or stacked with other equipment should be avoided because it could result in improper operation. If such use is necessary, this equipment and the other equipment should be observed to verify that they are operating normally.
- 34.Portable RF communications equipment (including peripherals such as antenna cables and external antennas) should be used no closer than 30 cm (12 inches) to any part of the unit, including cables specified by the manufacturer. Otherwise, degradation of the performance of this equipment could result.

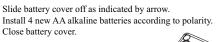
WARNING SIGNS AND SYMBOLS USED				
Ť	Keep Dry			
No.	Keep off Sunlight			
Type BF Equipment				
Instructions For Use MUST be Consulted				
Discard the used product to the recycling collection point according to local regulations				
Bluetooth	The Bluetooth® Smart word mark and logos are registered trademarks owned by Bluetooth SIG, Inc. and any use of such marks by JOYTECH Healthcare Co.,Ltd.			

Unit Illustration





Battery Installation

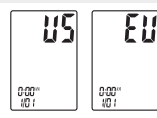


AC Adapter jack is on the back side of the monitor. Medical AC adapter(DC 6.0 V,600mA) can be used with the device (recommended, not provided). The adapter connect pin should be positive inside and negtive outside with a 2.1mm coaxial joint. Do not use any other type of AC adapter as it may harm the unit.

AC Adapter Jack

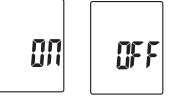
Note: Power supply is specified as part of ME EQUIPMENT.

Unit Operation



4. Voice Setting

Press "SET" button to enter voice setting mode. Set voice format ON or OFF by pressing the "M" button.



5. Volume Setting Press "SET" button to enter volume setting mode. Set the voice volume by adjusting the "M" button . There are six volume levels.



6. Saved Settings

While in any setting mode, press " START/STOP " button to turn the unit off. All information will be saved.

Note: If unit is left on and not in use for 3 minutes, it will automatically save all information and shut off

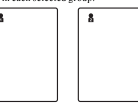
Unit Operation

System Settings

With power off, press "SET" button to activate System Settings. The Memory Group icon flashes

1. Select Memory Group

While in the System Setting mode, you may accumulate test results into 2 different groups. This allows multiple users to save individual test results (up to 60 memories per group.) Press " M " button to choose a group setting. Test results will automatically store in each selected group.



2. Time/Date setting Press "SET" button again to set the Time/Date mode. Set the year first by adjusting the "M" button.

Press "SET" button again to confiom curret month:Continue setting the date, hour and minute in the same way Every time the "SET" button is pressed, it will lock in your selection and continue in succession (month, day, hour, minute)



3. Time Format setting Press "SET" button again to set the time format setting mode.Set the time format by adjusting the "M" button. EU means European Time US means U.S Time.

Unit Operation

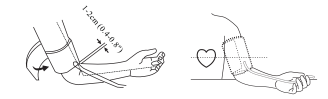
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Applying the Arm Cuff

1. Firmly insert air plug into opening located on left side of monitor unit.



- 2. With sticky nylon section facing outward, insert end of cuff underneath metal ring of cuff.
- 3. Fasten cuff about 1-2cm (0.4-0.8") above the elbow joint. For best results apply cuff to bare arm and keep level with heart while testing.



Note: Do not insert air plug into opening located on right side of monitor unit. This opening is designed for an optional power supply only.

Testing

ready for testing.

1. Power On

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2. Pressurization

Unit Operation

The unit will automatically inflate to the proper pressure value and stop inflating. During this time, please keep quiet.



Note: Pressurization will gradually subside and ultimately stop when cuff is not properly applied to the arm. If this occurs, press "START/STOP" button to turn the unit off.

Unit Operation

3. Testing

After cuff inflation, air will slowly subside as indicated by the corresponding cuff pressure value. A flashing " \clubsuit " will appear simultaneously on screen signaling heart beat detection.

Press and hold "START/STOP" button to turn the unit on. The LCD screen will appear for one second as unit performs a quick diagnosis. A voice tone will indicate when unit is

Note: Unit will not function if residual air from previous testing is present in cuff.

The LCD will flash " 🍸 " until pressure is stabilized.



Note: Keep relaxed during testing. Avoid speaking or moving body parts.

4. Result Display

The screen will display measurements for systolic and diastolic blood pressure with voice broadcast. An indicator representing the current measurement will appear next to the corresponding WHO Classification.



Note: Refer to Page 23~24 for detail WHO Blood Pressure Classification Information.

Unit Operation

Irregular Heartbeat Indicator

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If the monitor detects an irregular heart rhythm two or more times during the measuring process, the Irregular Heartbeat Symbol " (() "appears on screen along with measurement results. Irregular heartbeat rhythm is defined as rhythm that is either 25% slower or faster than the average rhythm detected while measuring systolic blood pressure and diastolic blood pressure. Consult your physician if the Irregular Heartbeat Symbol " () ")" "requently appears with your test results.

5. Deleting/Storing Test Results

User may delete their current test result due to unfavorable testing conditions or for any other reason. To delete the last test result, press the "SET" button after result is displayed. If result is not deleted, it will automatically store by date within the previously configured Memory Group.

Note: Be sure the appropriate Memory Group selection is made prior to testing.

If the number of tests surpasses the allotted 60 memories per group, the most recent tests will appear first, thus eliminating the oldest readings.

Power Off

The "START/STOP" button can be pressed to turn off the unit in any mode. The unit can turn off the power itself about 3 minutes no operation in any mode.

Safety Precaution: If pressure in arm cuff becomes too extreme while testing, press the "START/STOP" button to turn power off. The cuff pressure will rapidly dissipate once the unit is off.

Last 3 Tests Average

With power off, press the "M" button to activate screen display. After the unit performs a self-diagnosis, the screen will display the average test results from the last 3 readings of the last group used. The "AVG" symbol will appear along with the corresponding WHO Blood Pressure Indicator. The Memory Check mode can be accessed by pressing "M" buttons. To check the average results from other groups, select the desired group first prior to activating "SET" button in the off position.(See "Select Memory Group" on Page 10)

8 mmHg AVG 8:30^M • 15

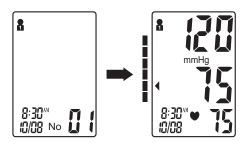
Unit Operation

Memory Check

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With power off, you may check past test results by using the "M" buttons. The most recent test result and oldest test result in memory can be viewed by pressing and holding the "M" button. Upon activating test results. you can press the "M" button to scroll through all test results stored in memory.

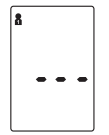


Note: Previous test will only be displayed from the most recently used memory group. To check previous test results in other memory groups, you must first select the desired group and then turn monitor off. (See "Select Memory Group" on Page 10)

Unit Operation

Memory Deletion

Memory for a selected group may be deleted while in Memory Check mode. Press and hold the "SET" button for approximately 3 seconds to delete all memory records from the selected groupwith voice broadcast "Memory Clear" and then transfer into testing mode. Press the "START/STOP" button to turn the unit off.



Note: Memory cannot be recovered once it has been deleted.

Low Battery Indicator

The unit will broadcast "Low Battery" when battery life is depleting and unable to inflate cuff for testing. The "



Unit Operation

Static Pressure Measurement

In the power down state, press and hold the "START/STOP" button, and theninstall the batteries. Until the LCD screen is full, release the "START/STOP" button. When the LCD screen displays the double zero, the bloodpressure meter is in static state. Software version is displayed at the heart rate.



Note: Only Service personnel permitted to access to this mode, the mode unavailable in normal use.

Bluetooth requirements

The monitor requires a device with:

- . Bluetooth 4.0 or later
- . Android 5.0 or later
- . IOS 9.0 or later
- And works with:
- . iphone , iPod, iPad
- . Android Phones and Tablets

Wireless communication

Frequency range : 2.4 Ghz (2400-2483.5 MHz)

Modulation : GFSK

Antenna gain:0.5dBi

Bluetooth connection

Using for the first time

1. Download the free "JOYTECH healthcare" App: On your mobile phone or table

go to www.sejoy.com.

2.Open the App on your phone or tablet. If requested, you should enable Bluetooth on your

device. You can enable Bluetooth under the Settings menu on your smart phone or table.

3. Create a new user login, or login with your existing user name and password.

4. Selection device "Blood pressure monitor".

Pairing your monitor with a Smart Device

The date and time on your monitor will automatically be set when you pair it with your

smart device.

- 1. Make sure the Bluetooth in your smart device is turned on.
- 2. Open the app on your smart device and follow the set-up and pairing instructions.
- 3. Confirm the monitor is connected successfully.

If the monitor is connected successfully to the smart device, "bLE on" will appear on the monitor display.



Blood Pressure Information

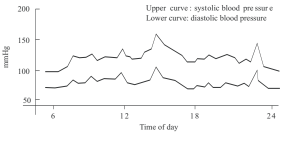
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Blood Pressure

Blood pressure is the force of blood pushing against the walls of arteries. It is typically measured in millimeters of mercury (mmHg.) Systolic blood pressure is the maximum force exerted against blood vessel walls each time the heart beats. Diastolic blood pressure is the force exerted on blood vessels when the heart is resting between beats.

An individual's blood pressure frequently changes throughout the course of a day. Excitement and tension can cause blood pressure to rise, while drinking alcohol and bathing can lower blood pressure. Certain hormones like adrenaline (which your body releases under stress) can cause blood vessels to constrict, leading to a rise in blood pressure.

If these measuring numbers become too high, it means the heart is working harder than it should.



Example: fluctuation within a day (male, 35 years old)

Unit Operation

Troubleshooting

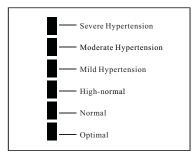
23

Problem	Possible Cause	Solution	
	Cuff is too tight or not properly positioned on the arm	Firmly reposition cuff approximately1-2cm (1/2") above the elbow joint (See Page 12)	
Blood pressure results are not within typical range	Inaccurate test results due to body movement or monitor movement	Sit in a relaxed position with arm placed near heart. Avoid speaking or moving body parts while testing. Make sure the monitor unit is placed in a stationary position throughout the testing period. (See Page 7)	
	Cuff fails to inflate properly	Make sure hose is properly fastened to cuff and monitor unit	
" Err "displayed	Improper operation	Read user manual carefully and re-test properly.	
	Pressurization is over cuff rated pressure 300mmHg	Read user manual carefully and re-test properly.	
Connection failure./ Data is not being	The blood pressure monitor might not be porperly placed within the smart device's tranmission range and is too far from the smart device.	If there are no causes of data transmission interference found near the blood pressure monitor, move the blood pressure monitor within 16ft.(5m) of the smart device and try again	
transmitted	The blood pressure did not pair successfully to the smart device	Try to pair the devices once again	
	The application on the smart device is not ready.	Check the application then try sending the data again.	

Blood Pressure Information

WHO Blood Pressure Classification Indicator

The DBP-1319b is equipped with a classification indicator based on established guidelines from the World Health Organization. The chart below (color coded on monitor unit) indicates test results.





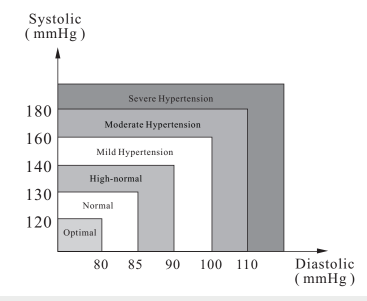
S: Blood Pressure Classification Indicator

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Blood Pressure Information

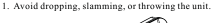
Health Reminder

Hypertension is a dangerous disease that can affect the quality of life. It can lead to a lot of problems including heart failure, kidney failure, and cerebral hemorrhaging. By maintaining a healthy lifestyle and visiting your physician on a regular basis, hypertension and relative diseases are much easier to control when diagnosed in their early stages.



Note: Do not be alarmed if an abnormal reading occurs. A better indication of an individual's blood pressure occurs after 2-3 readings are taken at the same time each day over an extended period of time. Consult your physician if test results remain abnormal.

Maintenance





2. Avoid extreme temperatures. Do not expose unit directly under sunshine.



3. When cleaning the unit, use a soft fabric and lightly wipe with mild detergent.

Use a damp cloth to remove dirt and excess detergent.



Blood Pressure Q&A

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- Q: What is the difference between measuring blood pressure at home or at a professional healthcare clinic?
- A: Blood pressure readings taken at home are now seen to give a more accurate account as they better reflect your daily life. Readings can be elevated when taken in a clinical or medical environment. This is known as White Coat Hypertension and may be caused by feeling anxious or nervous.

Note: Abnormal test results may be caused by:

- Improper cuff placement Make sure cuff is snug-not too tight or too loose. Make sure bottom of the cuff is approximately 1-2cm (1/2") above the elbow joint.
 Improper body position Make sure to keep your body in an upright position.
- Feeling anxious or nervous Take 2-3 deep breaths, wait a few minutes and resume testing.

Q: What causes different readings?

- A: Blood pressure varies throughout the course of a day. Many factors including diet, stress, cuff placement, etc. may affect an individual's blood pressure.
- Q: Should I apply the cuff to the left or right arm? What is the difference?
- A: Either arm can be used when testing, however, when comparing results, the same arm should be used. Testing on your left arm may provide more accurate results as it is located closer to your heart.
- Q: What is the best time of day for testing?
- A: Morning time or any time you feel relaxed and stress free.

Maintenance

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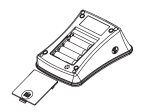
- 4. Cuff Cleaning and Disinfection:
- A) Spread the cuff (skin-contact surface) upwards onto a clean table. Use a damp clean cloth (water-based) to wipe the skin-contact surface with a force.
- B) Soak the cloth clean with drinking water and wring it dry. Repeat A) with the damp cloth (water-based) for 3 times.

C) Apply 70%-80% alcohol to a new cloth (or 75% alcohol cotton-ball), use it to wipe the skin-contact surface with a force. Then soak the cloth with the alcohol again (or change a new 75% alcohol cotton-ball), repeat the disinfection procedure for 3 times.
D) When the disinfection towards the skin-contactsurface is finished, wipe thenon-skin contact surface with a cloth (alcohol-based) or alcohol cotton-ball thoroughly for

- 3 times. E) Leave the cuff naturally dry, then it is ready for reuse.
- Notice: Do not soak in water or splash water on it.
- 5. Do not use petrol, thinners or similar solvents



- 6. Remove batteries when not in operation for an extended period of time.
- 7. Do not disassemble product.





- 8. It is recommended the performance should be checked every 2 years.
- 9. Expected service life: Approximately three years at 10 tests per day.
- 10.No service and maintenance while it is in use and maintenance only be performed by service personnel. Service and maintenance require parts, repair, technical support will be provided.

Specifications

Product Description	Arm-type Fully Aut	omatic Blood Pressure Monitor		
Model	DBP-1319b			
Display	LCD Digital Display	y Size: 102mm×68.9mm (4.02" x 2.71")		
Measurement Method	Oscillometric Meth	od		
	Systolic Pressure	Systolic Pressure 60mmHg~280mmHg		
	Diastolic Pressure	30mmHg~200mmHg		
Maaaaaa	Pressure	0mmHg~300mmHg		
Measurement Range	Pressure	±3mmHg		
	Pulse	30 ~ 180 Beats/Minute		
	Pulse	±5%		
Pressurization	Automatic Pressuri	zation		
Memory	120 Memories in Tw	o Groups with Date and Time		
	Irregular Heartbeat	Detection		
	WHO Classification Indicator			
	Last3 Tests Average			
Function	Low Battery Detect	ion		
runction	Automatic Power-C	ff		
	Backlight			
	Voice			
	Bluetooth			
Power Source	4 AA batteries or M (recommended, not	edical AC Adapter(DC6.0V, 600mA) provided)		
Battery Life	Approximately 2 months at 3 tests per day			
Unit Weight	Approx.480g (16.93 oz.) (excluding battery)			
Unit Dimensions	Approx.166 x 114 x 72mm (6.54" x 4.49" x 2.83")(L x W x H)			
Cuff Circumference	Medium cuff: Fits arm circumference 22-36 cm			
Operating Environment	Temperature	10°C ~ 40°C (50°F~104°F)		
Operating Environment	Humidity	15% ~ 93% RH		
	Pressure	700hPa~1060hPa		

Specifications

Storage Environment	Temperature:	-25°C~70°C (-13°F~158°F)	
Storage Environment	Humidity	≪93% RH	
Classification:	Internal Powered Equipment, Type BF 📩 . Cuff is the Applied Part		
Ingress Protection Rating:	IP20, Indoor Use Only		
Battery Shelf life:	60 months		
Battery Storage Temperature:	-25°C~55°C (-13°F~131°F)		

Specifications are subject to change without notice.

- 1. IEC 80601-2-30, medical electrical equipment part 2-30: particular requirements for the basic safety and essential performance of automated noninvasive sphygmomanometers. (Cardiovascular)
- 2. ANSI/AAMI ISO 81060-2, non-invasive sphygmomanometers part 2: clinical validation of automated measurement type. (Cardiovascular)
- 3. AAMI/ANSIES60601-1:2005/(R)2012 and C1:2009/(R)2012 and, a2:2010/(r)2012 (consolidated text) medical electrical equipment -- part 1: general requirements for basic safety and essential performance
- 4. AAMI/ANSI/IEC 60601-1-2, Medical Electrical Equipment -- Part 1-2: General Requirements For Basic Safety And Essential Performance -- Collateral Standard: Electromagnetic Disturbances -- Requirements And Tests (General II (ES/EMC)).
- 5. IEC 60601-1-11, medical electrical equipment part 1-11: general requirements for basic safety and essential performance collateral standard: requirements for medical electrical equipment and medical electrical systems used in the home healthcare environment.

Correct Disposal of This Product (Waste Electrical & Electronic Equipment)



This marking shown on the product indicates that it should not be disposed with other household waste at the end of its life. To prevent potential harm to the environment or to human health, please separate this product from other types of wastes and recycle it responsibly. When disposing this type of product, contact the retailer where product was purchased or contact your local government office for details regarding how this item can be disposed in an environmentally safe recycling center.

Business users should contact their supplier and check the terms and conditions of the purchasing agreement. This product should not be mixed with other commercial wastes for disposal. This product is free of hazardous materials.

Warranty

The Blood Pressure Monitor is guaranteed for 2-year from the date of purchase. If the Blood Pressure Monitor does not function properly due to defective components or poor workmanship, we will repair or replace it freely. The warranty does not cover damages to your Blood Pressure Monitor due to improper handling. Please contact local retailer for details.

33 Electromagnetic Compatibility Information **34**

The device satisfies the EMC requirements of the international standard IEC 60601-1-2. The requirements are satisfied under the conditions described in the table below. The device is an electrical medical product and is subject to special precautionary measures with regard to EMC which must be published in the instructions for use. Portable and mobile HF communications equipment can affect the device. Use of the unit in conjunction with non-approved accessories can affect the device negatively and alter the electromagnetic compatibility. The device should not be used directly adjacent to or between other electrical equipment. **Table 1**

Guidance and	declaration o	of manufacturer-o	alactromagnetic	amissions
Ouruance and	ucciaration 0	i manufacturer=v	rectionagnetic	cimissions

The device is intended for use in the electromagnetic environment specified below. The customer or the user of the device should assure that it is used in such an environmen

Emissions test	Compliance	Electromagnetic environment -guidance
Radiated emission CISPR 11	Group 1, class B.	The device uses RF energy only for its internal function. Therefore, its emissions are very low and are not likely to cause any interference in nearby electronic equipment.
Conducted emission CISPR 11	Group 1, class B.	The device is suitable for use in all establishments, including domestic establishments and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.
Harmonic emissions IEC 61000-3-2	Class A	
Voltage fluctuations/ flicker emissions IEC 61000-3-3	Complies	

Electromagnetic Compatibility Information 35 Electromagnetic Compatibility Information 36

Table 3

Table 2

Guidance and declaration of manufacturer-electromagnetic immunityThe device is intended for use in the electromagnetic environment specified below. The customer or the user of the device should assure that it is used in such an environment.IMMUNITY testIEC 60601 test levelCompliance levelElectromagnetic environment -guidanceElectrostatic discharge (ESD) IEC 61000-4-2is 8 kV ± 8 kV, ± 15 kV airElectromagnetic environment -guidanceElectrostatic transient/burst IEC 61000-4-4± 8 kV, ± 15 kV airK and ± 2 kV, ± 4 kV, ± 15 kV airFloors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30 %.Surge IEC 61000-4-5± 0 kV, ± 1 kV (differential mode)Mains power quality should be tha of a typical commercial or hospita environment.Voltage dips, short interrupti- ons and voltage variations on p- ower supply in- put lines0 % UT; 0 % UT; 25/30 cycles Single phase: at 10°0 % UT; 1 cycle and and of o'Mains power quality should be tha of a typical commercial or hospita environment.	ble 2			
The customer or the user of the device should assure that it is used in such an environment. IMMUNITY test IEC 60601 test level Compliance level Electromagnetic environment -guidance Electrostatic discharge (ESD) IEC 61000-4-2 $\pm 8 kV$ contact $\pm 2 kV, \pm 4 kV, \\ \pm 15 kV air \pm 8 kVcontact\pm 2 kV, \pm 4 kV, \\ \pm 15 kV air Floors should be wood, concreteor ceramic tile. If floors arecovered with synthetic material,the relative humidity should beat least 30 %. Electrostatictransient/burstIEC 61000-4-5 \pm 2 kV, \\ 100 kHz, forAC power port \pm 0.5 kV, \pm 1 kV(differentialmode) Mains power quality should be thaof a typical commercial or hospitaenvironment. SurgeIEC 61000-4-5 \pm 0.5 kV, \pm 1 kV(differentialmode) \pm 0.5 kV, \pm 1 kV(differentialmode) Mains power quality should be thaof a typical commercial or hospitaenvironment. Voltage dips,short interrupti-ons and voltagevariations on p-ower supply in-put lines 0 % UT; 1 cycleandand25/30 cyclesSingle phase: 0 % UT; 1 cycleandand30 cyclesSingle phase: Mains power quality should be thaof a typical commercial or hospitaenvironment. $	Guidance	and declaration of	of manufacturer-	electromagnetic immunity
IMMUNITY testtest levellevelcertoinagitte christiante rguidanceElectrostatic discharge (ESD) IEC 61000-4-2 $\pm 8 kV$ $\pm 8 kV, \pm 4 kV, \pm 2 kV, \pm 4 kV, \pm 8 kV, \pm 15 kV$ air $\pm 8 kV$ $\pm 15 kV$ airFloors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30 %.Electrostatic transient/burst IEC 61000-4-4 $\pm 2 kV, \pm 1 kV, \pm 15 kV$ air $\pm 2 kV, \pm 1 kV, \pm 15 kV$ airMains power quality should be tha of a typical commercial or hospita environment.Surge IEC 61000-4-5 $\pm 0.5 kV, \pm 1 kV$ (differential mode) $\pm 0.5 kV, \pm 1 kV$ (differential mode)Mains power quality should be tha of a typical commercial or hospita environment.Voltage dips, short interrupti- ons and voltage variations on p- put lines $0 \% UT;$ $25/30 cyclesSingle phase:0 \% UT;25/30 cyclesSingle phase:0 \% UT;25/30 cyclesSingle phase:0 \% UT;25/30 cyclesSingle phase:0 \% UT;25/30 cyclesSingle phase:$	The customer or th			
Electrostatic discharge (ESD) IEC 61000-4-2contact contact $\pm 2 kV, \pm 4 kV,$ $\pm 15 kV aircontactcontact\pm 2 kV, \pm 4 kV,\pm 15 kV aircontactcoreramic tile. If floors arecovered with synthetic material,the relative humidity should beat least 30 %.Electrostatictransient/burstIEC 61000-4-4\pm 2 kV,\pm 2 kV,100 kHz, forAC power port\pm 2 kV,100 kHz, forAC power portMains power quality should be thaof a typical commercial or hospitaenvironment.SurgeIEC 61000-4-5\pm 0.5 kV, \pm 1kV(differentialmode)\pm 0.5 kV, \pm 1kV(differentialmode)Mains power quality should be thaof a typical commercial or hospitaenvironment.Voltage dips,short interrupti-ons and voltagevariations on p-put lines0 % UT;0 % UT; 1 cycle0 % UT; 1 cycleandand70 % UT;25/30 cyclesSingle phase:0 % UT;0 % UT;25/30 cyclesSingle phase:0 % UT;0 % UT;0 % UT;0 % UT;0 % UT;$	IMMUNITY test			
transient/burst IEC 61000-4-4100kHz, for AC power port100kHz, for AC power portof a typical commercial or hospita environment.Surge IEC 61000-4-5±0.5kV, ±1kV (differential mode)±0.5kV, ±1kV (differential mode)±0.5kV, ±1kV (differential mode)Mains power quality should be tha of a typical commercial or hospitaVoltage dips, short interruptions and voltage variations on power supply in- ower supply in- put lines0 % UT; 1 cycle 0 % UT; 1 cycle 25/30 cycles Single phase:0 % UT; 25/30 cycles Single phase:Mains power quality should be tha of a typical commercial or hospita environment.	discharge (ESD)	contact $\pm 2 \text{ kV}, \pm 4 \text{ kV},$ $\pm 8 \text{ kV},$	contact $\pm 2 \text{ kV}, \pm 4 \text{ kV},$ $\pm 8 \text{ kV},$	or ceramic tile. If floors are covered with synthetic material, the relative humidity should be
Surge IEC 61000-4-5 (differential mode) (differential mode)<	transient/burst	100kHz, for	100kHz, for	of a typical commercial or hospita
Voltage dips, short interrupti- ons and voltage variations on p- put lines $\begin{bmatrix} 0,5 \text{ cycle} \\ 0,5 \text{ cycle} \\ At 0^\circ, 45^\circ, 90^\circ, At 0^\circ, 45^\circ, 90^\circ, 135^\circ, 180^\circ, 225^\circ \text{Mains power quality should be tha} \\ ,270^\circ \text{ and } 315^\circ, 270^\circ \text{ and } 315^\circ, 270^\circ, 2$		(differential	(differential	of a typical commercial or hospita
IEC 61000-4-11	short interrupti- ons and voltage variations on p- ower supply in- put lines	0,5 cycle At 0°, 45°, 90°, 135°, 180°, 225 , 270° and 315° 0% UT; 1 cycle and 70% UT; 25/30 cycles Single phase:	0,5 cycle At 0°, 45°, 90°, 135°, 180°, 225 , 270° and 315° °0 % UT; 1 cycle and 70 % UT; 25/30 cycles Single phase:	of a typical commercial or hospita environment.

	v. The custome	er or the user o	omagnetic environment f the device should assure that it
IMMUNITY test	IEC 60601 test level	Compliance level	Electromagnetic environment -guidance
Conducted RF IEC 61000-4-6		3V for 0.15- 80MHz; 6V in ISM and amate -ur radio bands between0.15- 80MHz	Portable and mobile RF communications equipment should be used no closer to any part of the device, including cables, than the recommended separation distance calculated
Radiated RF IEC 61000- 4-3	385MHz, 27V /m	385MHz, 27V /m	from the equation applicable to the frequency of the transmitter. Recommended seperation distance
	450MHz, 28V /m	450MHz, 28V /m	$d = [\frac{3.5}{E_1}]\sqrt{P} \text{ 80 MHz to 800 MHz}$ $d = [\frac{7}{E_1}]\sqrt{P} \text{ 800 MHz to 2.7 Ghz}$
	710MHz,745 MHZ,780MHz 9V/m	710MHz,745 MHZ,780MHz 9V/m	$d = \left[\frac{i}{E_1}\right]\sqrt{P} 800 \text{ MHz to } 2.7 \text{ Ghz}$ where P is the maximum output power rating of the transmitter in watts (W) according to the transm-
	810MHz,870 MHZ,930MHz 28V/m	810MHz,870 MHZ,930MHz 28V/m	itter manufacturer and d is the recommended separation distance in metres (m).
	1720MHz,1845 MHZ,1970MHz 28V/m	1720MHz,1845 MHZ,1970MHz 28V/m	
	2450MHz, 28V /m	2450MHz, 28V /m	Interference may occur in the vicinity of equipment marked with the following symbol:
	5240MHz,5500 MHZ,5785MHz 9V/m	5240MHz,5500 MHZ,5785MHz 9V/m	

Guidance and declaration of manufacturer-electromagnetic immunity

Electromagnetic Compatibility Information 37

30 A/m; 50Hz or 60Hz Power frequency magnetic fields should be at levels charactertic of a

typical location in a typical commercial or hospital environment.

0 % UT; 250/300 cycle 0 % UT; 250/300 cycle

30 A/m; 50Hz or 60Hz

Table 4

Power frequency (50/60 Hz)

magnetic field

IEC 61000-4-8

Recommended separation distances between portable and mobile RF communications equipment and the device The device is intended for use in an electromagnetic environment in which radiated therefore disturbances are controlled. The customer or the user of the device can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the device as recommended below, according to the maximum output power of the communications equipment.

Rated maximum	Separation distance according to frequency of transmitter		
output power of	I	n	
transmitter	80 MHz to 800 MHz	800 MHz to 2.7 GHz	
W	$d = [\frac{3.5}{E_1}]\sqrt{P}$	$d = \left[\frac{7}{E_1}\right]\sqrt{P}$	
0.01	0.12	0.23	
0.1	0.38	0.73	
1	1.2	2.3	
10	3.8	7.3	
100	12	23	

For transmitters rated at a maximum output power not listed above, the recommended separation distance d in metres (m) can be estimated using the equation applicable to the frequency of the transmitter, where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer.

NOTE1 At 80 MHz and 800 MHz, the separation distance for the higher frequency range applies.

NOTE2 These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.