HOMEDICS

Automatic Arm Blood Pressure Monitor



Instruction Manual



BPA-2000-EU

Before use

Before the blood pressure monitor is taken into use, a number of basic steps should be observed. Please read this manual thoroughly before the blood pressure monitor is used for the first time and keep the manual for future use.

The blood pressure monitor is intended for home use only. Any measurement is solely for your information and can under no circumstances replace a visit to the doctor. The blood pressure measurements should always be assessed by a qualified person (a doctor, nurse or similar) who is familiar with the user's general medical condition. By regular use and by keeping a record of the results, this person can be kept well informed about developments in the user's blood pressure. Measurements must never be interpreted by the user with the aim of changing levels in the use of medicine as prescribed by a doctor. The doctor's instructions must be followed AT ALL TIMES.

Use of appropriate cuff size is crucial to correct measurement. Follow the instructions presented in this manual and printed on the cuff to ensure that the correct cuff size is used.

The blood pressure monitor is not suited for users who suffer from arrhythmia (heart rhythm disturbances) and errors may occur in the measurement results if the user has suffered a stroke, suffers from cardiovascular diseases, has a very low blood pressure, or suffers from other symptoms such as circulatory diseases (diabetes, kidney diseases, arterial sclerosis (deposits in the arteries), or poor peripheral blood circulation (e.g. in hands and feet).

Electromagnetic disturbances: The device contains sensitive electronic components. Keep it clear of strong electric or electromagnetic fields in the immediate surroundings (e.g. mobile telephones, microwave ovens) as these may temporarily reduce measurement accuracy.

Use the blood pressure monitor only for its original purpose.

The device is intended for measurement of blood pressure and pulse with adult persons. Do not apply the monitor to infants nor to persons who cannot express their permission. The device must not be operated by children.

The blood pressure monitor functions in accordance with the oscillometric measurement principle. These blood pressure changes are measured in the cuff on the basis of the arm pulse and are converted into a numerical value of the current blood pressure. Simultaneously, the number of pulse beats are registered and calculated as number of pulse beats per minute.

There are two kinds of discernible blood pressures. Both are expressed in mmHg (millimetres on the mercury column): The systolic and the diastolic. The systolic ('the high blood pressure') states the pressure when the heart chambers contract to send blood through your body, and the diastolic ('the low blood pressure') states the pressure when the heart is at ease and filling with blood before the next muscle contraction. The connection between values (mmHg) is expressed as e.g. 130 over 85 which means that the systolic pressure is 130 and the diastolic pressure is 85.

Why is it important to keep an eye on your blood pressure?

Many of today's most common diseases are connected to hypertension, also referred to as 'high blood pressure'. Hypertension is closely connected to cardiovascular diseases, and for people in the danger zone the blood pressure provides an important tool to keep an eye on the development.

What are systolic pressure and diastolic pressure?

When ventricles contract and pump blood out of the heart, blood pressure reaches its maximum value, the highest pressure in the cycle is known as systolic pressure. When the heart relaxes between heartbeats, the lowest blood pressure is diastolic pressure.

What is the standard blood pressure classification?

Below illustrates the blood pressure classification mode by World Health Organization (WHO) and International Society of Hypertension(Isa) in 1999.



Level Blood Pressure (mmHg)	Optimal Optimal	Normal Norm.	High-normal Norm.	Mild G1	Moderate G2	Severe G3
SYS	<120	120~129	130~139	140~159	160~179	≥180
DIA	<80	80~84	85~89	90~99	100~109	≥110

Only a physician/GP can tell you your normal blood pressure range and the point at which you are at risk. Consult your physician to obtain these values. If the measurements taken with this product fall outside the range, consult your physician/GP.

SAFETY INFORMATION

(3)	Caution: consult accompanying documents	*	Type B applied part
C € º120	CE mark: conforms to essential requirements of the Medical Device Directive 93/42/EEC.	X	Disposal: do not dispose of this product as unsorted municipal waste. Collection of such waste separately for special treatment is necessary.
	Manufacturer		Direct current
SN	Specifies serial number		Direct current

- · This device is intended for adult use only.
- This device is intended for non-invasive measuring and monitoring of arterial blood pressure. It is not intended for use on extremities other than the arm.
- Do not confuse self-monitoring with self-diagnosis. This unit allows you to monitor your blood pressure. Do not begin or end medical treatment based solely on the readings from this unit. Always consult your physician first.
- If you are taking medication, consult your physician to determine the most appropriate time to measure your blood pressure. Never change a prescribed medication without consulting your physician.
- This unit is not suitable for continuous monitoring during medical emergencies or operations.
- If the cuff pressure exceeds 40 kPa (300 mmHg), the unit will automatically deflate. Should the cuff not deflate when pressures exceeds 40 kPa (300 mmHg), detach the cuff from the arm and press the 🕐 button to stop inflation.
- To avoid measurement errors, carefully read this manual before using the product.
- The equipment is not AP/APG equipment and not suitable for use in the presence of a flammable anesthetic mixture with air of with oxygen or nitrous oxide.
- The operator shall not touch output of AC adapter and the patient simultaneously.
- To avoid measurement errors, please avoid the condition of strong electromagnetic field radiated interference signal or electrical fast transient/burst signal when using the AC adaptor.
- The user must check that the equipment functions safely and see that it is in proper working condition before being used.
- Please note that Luer lock connectors are not used on the product and please DO NOT change any provided connectors.

LCD display signal



Symbol	Description	Explanation
SYS	Systolic blood pressure	High pressure result
DIA	Diastolic blood pressure	Low pressure result
Pul/min	Pulse	Pulse/minute
	Deflating	CUFF is deflating
Ĩ:38	Time (hour:minute)	Current time
18	Memory	The displayed measurement values is from the memory.
kPa	kPa	Measurement Unit of the blood pressure
mmHg	mmHg	Measurement Unit of the blood pressure
	Low battery	Batteries are low and need to be replaced
	Arrhythmia	Irregular heartbeat
	Grade	The grade of the blood pressure.

MONITOR COMPONENTS



List

1. Blood Pressure Monitor



3.4*AA batteries



2. Cuff



4. User manual

Inserting and Changing Batteries



- 1. Slide off the battery cover.
- 2. Install the batteries by matching the correct polarity, as shown.
- 3. Replace the cover.

Replace the batteries whenever the below happen

- The L 🛈 + 📼 shows
- The display dims
- The display does not light up



Alternatively an AC adaptor powered mode is available: 100-240V~, 50-60HZ, 400mA (You can purchase the AC adaptor PP-ADPR2K3K-EU from the HoMedics website).

CAUTION

8

Remove batteries if the device is not likely to be used for some time.

This product uses the Oscillometric Measuring method to detect blood pressure. Before every measurement, the unit establishes a "zero pressure" equivalent to the air pressure. Then it starts inflating the arm cuff, meanwhile, the unit detects pressure oscillations generated by beat-to-beat pulsatile, which is used to determine the systolic and diastolic pressure, and also pulse rate.

The device also compares the longest and the shortest time intervals of detected pulse waves to mean time interval then calculates standard deviation. The device will display a warning signal with the reading to indicate the detection of irregular heartbeat when the difference of the time intervals is over 25%.

SETTING DATE, TIME AND MEASUREMENT UNIT

It is important to set the clock before using your blood pressure monitor, so that a time stamp can be assigned to each record that is stored in the memory. (year: 2000–2050, time: 24 H)

- 1. When the unit is off, hold pressing "SET" for 3 seconds to enter the mode for year setting.
- 2. Press the "MEM" to change the [YEAR].
- **3.** When you get the correct year, press "SET" and it will turn to the next step automatically.



- 4. Repeat step 2 and 3 to set the [MONTH] and [DAY].
- 5. Repeat step 2 and 3 to set the [HOUR] and [MINUTE].





6. Repeat step 2 and 3 to set the [UNIT].





7. After the unit is set , done will show, then turn off automatically.



HOW TO USE THE BLOOD PRESSURE DEVICE

Fitting the cuff

Correct cuff size is critical to accurate measurement. This monitor comes with a standard size 22-32cm arm cuff. A large size cuff 32-42cm is available from the HoMedics website www.homedics.co.uk

- 1. If the cuff is not assembled at first use, put the end of the cuff which is furthest away from the tube through the D-ring to form a noose.
- 2. Apply the tube to the left side of the device.
- **3.** Remove tight-fitting clothing from your left upper arm and position the cuff at approximately 2-3 cm above the elbow so that the tube is centred at the middle of the arm. The tube must face downward towards the hand.
- **4.** Pull gently at the end of the cuff so that it tightens easily and smoothly around the arm. The cuff is secured by pulling the Velcro fastener towards the cuff.
- **5.** You should be able to fit two fingers between the cuff and your arm. The cuff should not be fitted any tighter than this.
- 6. Relax your body and rest your arm on a table with your palm up so that the cuff is positioned at the same height as the heart. Make sure that the tube does not bend.

If it is not possible to fit the cuff on the left arm, it can be placed on the right arm instead. However, it is important that all measurements are carried out on the same arm. If the measurement is to be taken on the right arm, the artery symbol on the cuff should be positioned above the main artery. Press the arm's arteries with two fingers at approximately 2 cm above the elbow on the inside of the arm to find out where the pulse is strongest. This is the main artery.

















Worth knowing about blood pressure measurement

- Blood pressure measurements should be performed in quiet and comfortable surroundings as measurements can be affected by high or low temperatures. The blood pressure should be measured at normal body temperatures.
- Do not move or speak during measurement. Keep your feet flat on the floor.
- Do not touch the cuff or the blood pressure device during measurement.
- To obtain the best possible measurement, the blood pressure should be measured at approximately the same time every day, and on the same arm.
- If you are to perform several successive measurements, please wait 5 minutes between each measurement.
- The measurements are for guidance only. If you have doubts about your blood pressure, consult your doctor.
- The blood pressure device automatically deflates when the pressure exceeds 300 mmHg.
- The product is not suited for persons suffering from arrhythmia (heart rhythm disturbances).
- Measurement errors may occur if the user suffers from diabetes, poor blood circulation, kidney problems, or has suffered a stroke.
- The blood pressure changes with every heartbeat and is consequently changing constantly 24 hours a day.
- The measurement of blood pressure can be influenced by the user's position, condition, or other factors. To obtain the highest degree of accuracy it is recommended that the measurement is taken 1 hour after workout, bathing, finishing a meal, consuming alcoholic or caffeine beverages, or after smoking.
- Before measurement it is recommended that you sit still and relax for 15 minutes. The user should not be physically tired or exhausted when the measurement is taken.
- The user must not talk or move arm or hand during measurement.

How to measure blood pressure

- 1. Press the 🔱 to turn on the monitor, and it will start and finish the whole measurement automatically.
- 2. When the measurement is completed, the cuff automatically deflates and the display will show the systolic pressure, the diastolic pressure, and the pulse. The measurement is simultaneously stored in the monitor's memory.
- **3.** The device automatically switches off approximately 1 minute after use but can also be switched off manually by pressing the **U** button.

NOTE

If necessary, a measurement can be interrupted by pressing the 🕑 button. The cuff will subsequently deflate immediately.

The cuff will automatically re-inflate if the device determines that the body requires higher pressure to complete the measurement.

Recalling records

1. Please press the "MEM" to show the recent record.



2. Press the "MEM" or "SET" to get the UP record you want. DOWN SE1 The number of the record. 117 R date and time will be shown accordingly. The current record is The month and day The corresponding time will show and alternate shown with date.

The most recent record (1) is shown first. Each new measurement is assigned to the first (1) record. All other records are pushed back one digit (e.g., 2 becomes 3, and so on), and the last record (60) is dropped from the list.

Deleting your records

If you did not get the correct measurement, you can delete all results using the following steps.

- 1. When the device is off, hold "MEM" for 3 seconds. The flash display will show dEL ALL.
- **2**. Press "SET" to confirm deleting and the monitor will turn off automatically.





- 3. If you do not want to delete the record, press 🕐 to cancel.
- **4**. If there is no record the right display will show.



Error search

This section includes a list of error messages and frequently asked questions for problems you may encounter with your blood pressure monitor. If irregularities occur during use, please check the following items.

PROBLEM	SYMPTOM	CHECK THIS	REMEDY
	Display is dim or will	Batteries are exhausted.	Replace with new batteries
No power	not light up.	Batteries are inserted incorrectly	Insert the batteries correctly
Low batteries	LO+ C Show on the display	Batteries are low.	Replace with new batteries
	E 1 shows	The cuff is not secure.	Refasten the cuff and then measure again
	E 2 shows	The cuff is very tight	Refasten the cuff and then measure again
	E 3 shows	The pressure of the cuff is excess.	Refasten the cuff and then measure again
Error Message	E 10 or E 11 shows	The monitor detected motion while measuring	Movement can affect the measurement. Relax for a moment and then measure again
	E 20 or E 21 shows	Measure incorrectly	Relax for a moment and then measure again
	Eexx, shows on the display.	System error.	Retake the measurement. If the problem persists, contact the retailer or our customer service department for further assistance. Refer to the warranty for contact information and return instructions.

If the device still does not function, please contact the store where the device was purchased. Do not attempt to open the device or adjust any of the inner parts yourself.

Specifications

Power supply	Battery powered mode: 6VDC 4*AA batteries AC adaptor powered mode: 100-240V~, 50-60HZ, 400mA (You can purchase the AC adaptor PP-ADPR2K3K-EU from the HoMedics website).
Display mode	Digital LCD V.A.93*61mm
Measurement mode	Oscillographic testing mode
Measurement range	Pressure: 0~40kpa (0~300mmHg) pulse value: (40~199) times/minute
Accuracy	Pressure: 15~25°C within ± 0.4kpa (3mmHg) 10°C~40°C (out of 15°C~25°C) within ± 0.8kpa (6mmHg) pulse value: ±5%
Normal working condition	Temperature: 5°C~40°C Relative humidity ≤80%
Storage & transportation condition	Temperature: -20°C~60°C Relative humidity: 10%~93%
Measurement perimeter of the upper arm	About 22cm~32cm
Weight	Approx. 353g (Excluding the batteries)
External dimensions	Approx.140*120*70mm
Attachment	4*AA batteries, user manual, cuff
Mode of operation	Continuous operation
Degree of protection	Type B applied part
Protection against ingress of water	IPX-0

Cleaning and maintenance

The blood pressure monitor and the cuff can be wiped with a soft cloth moistened with a mild detergent and wiped dry with a soft, dry cloth or paper towel. Never use strong, caustic detergents as these may cause damage to the synthetic parts. Also, never use diluents, alcohol or kerosene to clean the device.

Safety and electrical devices

- When the device is not used, store it in a safe place.
- Only use the device on the upper arm, not on other body parts.
- When the device is not used for extended periods of time, it is recommended that you remove the batteries to avoid leakage which may cause damage to the device.
- If the device has been stored at very low temperatures near the freezing point, please allow it to obtain room temperature before using it again.
- Neither device nor cuff must be dissembled as this may cause serious damage to the device. If repair is necessary, the device must be returned to the store where it was purchased. Do not attempt to open the device or adjust the inner parts yourself.
- Do not submerge the device in water or other liquids as this will cause damage to the device.
- Do not expose the device nor the cuff to extreme temperatures, moisture or direct sunlight. Protect the device against dust.
- The cuff and the tube should not be folded up tightly. The device must not be activated if it is not appropriately fitted to the upper arm.
- Avoid dropping the device and do not subject it to heavy pressure.
- The device must not be exposed to extreme shocks such as being dropped on the floor.
- The device is intended for use in private households only and is only suited for use with adults (over 18 years of age).
- If the highly unlikely incident should occur that the cuff continues to inflate and does not stop, immediately remove the cuff from the upper arm.

Battery Directive



This symbol indicates that batteries must not be disposed of in the domestic waste as they contain substances which can be damaging to the environment and health. Please dispose of batteries in designated collection points.

WEEE explanation



This marking indicates that this product should not be disposed with other household wastes throughout the EU. To prevent possible harm to the environment or human health from uncontrolled waste disposal, recycle it responsibly to promote the sustainable reuse of material resources. To return your used device, please use the return and collection systems or contact the retailer where the

product was purchased. They can take this product for environmental safe recycling.

Table 1 Guidance and manufacture's declaration – electromagnetic emissions for all EQUIPMENT and SYSTEMS

Guidance and manufacture's declaration - electromagnetic emission

The BPA-2000-EU is intended for use in the electromagnetic environment specified below. The customer of the user of the BPA-2000-EU should assure that it is used in such and environment.

Emission test	Compliance	Electromagnetic environment – guidance	
RF emissions CISPR 11	Group 1	The BPA-2000-EU uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.	
RF emission CISPR 11	Class B		
Harmonic emissions IEC 61000-3-2 Not applicable		The BPA-2000-EU is suitable for use in all establishments other than domestic and those directly connected to the public low-voltage power supply network that	
Voltage fluctuations/ flicker emissions IEC 61000-3-3	Not applicable	supplies buildings used for domestic purposes.	

Table 2 Guidance and manufacture's declaration – electromagnetic immunity – for all ME EQUIPMENT and ME SYSTEMS

Guidance and manufacture's declaration - electromagnetic immunity

The BPA-2000-EU is intended for use in the electromagnetic environment specified below. The customer or the user of BPA-2000-EU 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	±6 kV contact ±8 kV air	±6 kV contact ±8 kV air	Floors should be wood, concrete or ceramic tile. If floor are covered with synthetic material, the relative humidity should be at least 30%.	
Electrical fast transient/burst IEC 61000-4-4	±2 kV for power supply lines	±2kV for power supply lines	Mains power quality should be that of a typical commercial or hospital environment.	
Surge IEC 61000-4-5	±1 kV line(s) to line(s)	±1 kV differential mode	Mains power quality should be that of a typical commercial or hospital environment.	
Voltage dips, short interruptions and voltage variations on power supply input lines IEC 61000-4-11	<5% UT (>95% dip in UT) for 0.5 cycle	<5% UT (>95% dip in UT) for 0.5 cycle	Mains power quality should be that of a typical commercial or hospital environment If the user of the BPA-2000-EU requires continued operation during power mains interruptions, it is recommended that the BPA-2000-EU be powered from an uninterruptible power supply or a battery.	
	40% UT (60% dip in UT) for 5 cycles	40% U _T) (60% dip in U _T) for 5 cycles		
	70% UT (30% dip in UT) for 25 cycles	70% U _T (30% dip in U _T) for 25 cycles		
	<5% U _T (>95% dip in U _T) for 5 sec	<5% U _T (>95% dip in U _T) for 5 sec	-	
Power frequency (50Hz) magnetic field IEC 61000-4-8	3A/m	3A/m	Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.	
NOTE: U _T is the a.c. mains voltage prior to application of the test level.				

Table 3 Guidance and manufacture's declaration – electromagnetic immunity – for ME EQUIPMENT and ME SYSTEMS that are not LIFE-SUPPORTING

Guidance and manufacture's declaration - electromagnetic immunity

The BPA-2000-EU is intended for use in the electromagnetic environment specified below. The customer or the user should assure that it is used in such an environment.

Immunity test	IEC 60601 test level	Compliance level	Electromagnetic environment - guidance
			Portable and mobile RF communications equipment should be used no closer to any part of the YS-6100, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter. Recommended separation distance $d = 1.167 \sqrt{P}$ $d = 1.167 \sqrt{P}$ 80 MHz to 800 MHz
Conducted RF	3 Vrms	3 Vrms	d = 2.333 \sqrt{P} 800 MHz to 2.5 GHz
Radiated RF	3 V/m 80 MHz	3.V/m	where <i>P</i> is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and <i>d</i> is the recommended
IEC 61000-4-3	to 2.5 GHz		separation distance in metres (m).
			Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey, a should be less than the compliance level in each frequency range.
			Interference may occur in the vicinity of equipment marked with the following symbol: $((\cdot, \cdot))$
NOTE 1 At 80 MHz and 800 MHz, the higher frequency range applies. NOTE 2 These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.			
a Field strengths fro	om fixed transmitters	, such as base station	s for radio (cellular/cordless) telephones and

a Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy.

To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the ELE007839V1 is used exceeds the applicable RF compliance level above, the ELE007839V1 should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as reorienting or relocating the ELE007839V1.

b Over the frequency range 150 kHz to 80 MHz, field strengths should be less than 3 V/m.

Table 4 Recommended separation distances between portable and mobile RF communications equipment and the EQUIPMENT or SYSTEM – for ME EQUIPMENT or ME SYSTEM that are not LIFE-SUPPORTING

Recommended separation distances between portable and mobile RF communications equipment and the ELE007839V1 Fitness Equipment.

The BPA-2000-EU is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the BPA-2000-EU can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the BPA-2000-EU as recommended below, according to the maximum output power of the communications equipment.

Rated maximum output	Separation distance according to frequency of transmitter m			
power of transmitter W	$150 \text{ kHz to } 80 \text{ MHz}$ $d = 1.167 \sqrt{P}$	80 MHz to 800 MHz d = 1.167 \sqrt{P}	800 MHz to 2,5 GHz d = 2.333 \sqrt{P}	
0,01	0.167	0.167	0.233	
0,1	0.369	0.369	0.738	
1	1.167	1.167	2.333	
10	3.690	3.690	7.338	
100	11.67	11.67	23.330	

For transmitters rated at a maximum output power not listed above, the recommended separation distance *d* in meters (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.

NOTE 1 At 80 MHz and 800 MHz, the separation distance for the higher frequency range applies. NOTE 2 These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

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Register your product today at: http://www.homedicsgroup.com/register

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