FA 48

Leading Health Trend



SYSTOLIC BLOOD PRESSURE

DIASTOLIC BLOOD PRESSURE

MEAN BLOOD PRESSURE

PULSE

PULSE PRESSURE

MOTION STATE



The device bears the CE label in accordance with the provisions of Medical Device Directive 93/42/EEC.

THE PERSONS RESPONSIBLE FOR PLACING DEVICES ON THE EC MARKET UNDER MDD 93/42/EEC



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INTRODUCTION

1. OVER VIEW

The FA 48 is a compact, non-invasive Ambulatory Blood Pressure Monitoring (ABPM) system. The appropriate use is 24-hour blood pressure monitoring and pulse wave analysis (PWA). The FA 48 should be used only under medical supervision.

INDICATION FOR USE

The FA 48 Ambulatory Blood Pressure Monitor is designed to measure systolic and diastolic blood pressure and pulse rate of adults who are eighteen (18) years and older using the oscillometric method on a cuffed arm.

INCORRECT USE



Because of the risk of strangulation from the tube and cuff, the FA 48 may not be used for patients who are legally incompetent.

The FA 48 must not be used for blood pressure monitoring during surgeries. It has not been designed for alarm-giving monitoring in intensive care units.

It is not recommended for people with serious arrhythmia to use this Wireless Blood Pressure Monitor.

PROPER USE OF THE FA 48

- 1. The FA 48 ABPM is intended for use by a trained medical practitioner. Interpretation of blood pressure measurements should always be performed by a physician.
- 2. The reliability of the monitor and the software depend upon conformance with the operation and maintenance instructions in this manual.
- 3. The FA 48 ABPM is designed for use with patients with normal sinus rhythms.
- 4. Measurement accuracy for the FA 48 ABPM may be affected by position of the subject, physical conditions, movement, and use outside of the operating instructions contained in this manual.

USER RESPONSIBILITY

This product is designed to perform in conformity with the description thereof contained in this manual and accompanying labels and inserts, when assembled, operated, maintained and repaired in accordance with the instructions provided. A defective product should not be used. Parts that are broken, plainly worn, missing or incomplete, distorted or contaminated should be replaced immediately. Should any repair or replacement become necessary, we recommend that service be performed at the nearest approved service center. The user of the product shall have the sole responsibility for any malfunction, which results from improper use, faulty maintenance, improper repair, damage or alteration by anyone other than Jawon Medical Co., Ltd. or their authorized service personnel.

The FA 48 System

The FA 48 ABPM system is lightweight, quiet, comfortable, and does not interfere with daily activities. It is easily configured and individually-fitted for each patient by a physician or nurse. The 24-hour measurements for systolic and diastolic blood pressure, the mean arterial pressure, and the pulse rate are automatically obtained without medical supervision, and later downloaded to a computer for analysis and interpretation.

Blood pressure measurements 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 ISO 81060-2.

SAFETY and EFFECTIVENESS WARNINGS

Make sure that you are familiar with all Safety and Effectiveness considerations before using the ABPM (model - FA 48).

ACCESSORIES

The Jawon Medical Co., Ltd. warranty can only be honored if you use Jawon Medical Co., Ltd. approved accessories and replacement parts.

Caution

Use of accessories other than those recommended by Jawon Medical Co., Ltd. may compromise product performance.

2. WORD DEFINITIONS

SYS Systolic Blood Pressure
DIA Diastolic Blood Pressure

Exhaust This means "releasing the cuff air as soon as possible".

3. CLASSIFICATION AND COMPLIANCE

- 1) This device is classified as;
 - Battery powered and the type-BF against electric shock
 - Ordinary equipment without protection against ingress of water
- Equipment not suitable for use in presence of a flammable anesthetic mixture by standard of IEC 60601-1:2005(Safety of Medical Electrical Equipment)
- 2) This device is in compliance with Class A for Noise-Emission, Level B for Noise-immunity, by standard of IEC 60601-1-2:2007(Electromagnetic Compatibility Requirements).
- 3) This device complies with:
 - IEC 60601-1:2005 Medical electrical equipment Part 1: General requirements for basic safety and essential performance
 - IEC 60601-2-30 Medical electrical equipment -- Part 2-30: Particular requirements for basic safety and essential performance of automated non-invasive sphygmomanometers
 - ISO 81060-1: 2007 Non-invasive sphygmomanometers Part 1: Requirements and test methods for non-automated measurement type
 - ISO 81060-2:2009 Non-invasive sphygmomanometers -- Part 2: Clinical validation of automated measurement type
 - ISO 10993-1: Biological evaluation of medical devices

4. SAFETY PRECAUTIONS



This device is designed and manufactured with consideration of safety of the operator and patient and also to the reliability of the unit.

The following precautions must be observed for additional safety;

- 1) The unit must be operated only by, or under supervision of a qualified person.
- 2) The operator and patient must not touch or handle inner side of the equipment at any time.
- 3) Do not modify the unit. If any modification is needed, ask our company or its authorized dealer for service.

- 4) The unit has previously been adjusted in the factory for optimum performance. Do not attempt to adjust switches or any other things except those specified in this manual for operation.
- 5) If you have experienced any trouble with the unit, switch it off immediately, and contact our company or its authorized dealer for assistance.
- 6) Continuous cuff pressure due to connection tubing kinking can cause the blood flow interference and resulting harmful injury to the patient.
- 7) Too frequent measurements can cause injury to the PATIENT due to blood flow interference.
- 8) The application of the CUFF over a wound can cause further injury.
- 9) The application of the CUFF and its pressurization on any limb where intravascular access or therapy, or an arterio-venous (A-V) shunt, is present because of temporary interference to blood flow and could result in injury to the patient;
- 10) CUFF and its pressurization on the arm on the side of a mastectomy should not be applied.
- 11) The pressurization of the CUFF can temporarily cause loss of function of simultaneously used monitoring ME EQUIPMENT on the same limb;
- 12) It is necessary to check that operation of the automated sphygmomanometer does not result in prolonged impairment of PATIENT blood circulation
- 13) If you plan to connect any device of other manufacturers electrically or mechanically to the unit, contact our company or its authorized dealer for instructions before doing so.

When you connect computer or other system to the unit (RS-232C), the attached systems should be those certified by IEC 950 or equivalent standards for data processing equipment.

Configurations shall comply with the system standard IEC 60601-1:2005.

Everybody who connects additional equipment to the signal input part or signal output part configures a medical system standard IEC 60601-1:2005.

If in doubt, consult the A/S department of local distributor.

- 14) Avoid the following environments for storage;
- Where the ambient temperature falls -20°C or exceeds 60°C.
- Where the atmospheric pressure falls below 70kPa (700mbar) or exceeds 106kPa (1060mbar).
- Where the humidity is over 95% non-condensing.
- Where the unit is exposed to spray or splashing water.
- Where the unit is exposed to dust.
- Where the unit is exposed to water vapor.
- Where the unit is exposed to salty atmosphere.
- Where the unit is exposed to explosive gas.
- Where the unit is exposed to excessive shocks or vibrations.
- Where the angle of inclination of mounting surface exceeds 10 degrees.
- Where the AC power line voltage heavily fluctuates.
- Where the AC power line voltage changes heavily when this unit is in operation.
- Where the unit is exposed to direct sunlight.
- 15) This equipment has been tested and found to comply with the limits for medical devices to the IEC 60601-1-2:2007. These limits are designed to provide reasonable protection against harmful interference in a typical medical 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 other devices in the vicinity. However, there is no guarantee that interference will not occur in a particular installation. If this equipment does cause harmful interference to other devices, 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 device.
- Increase the separation between the equipment.
- Connect the equipment into an outlet on a circuit different from that to which the other device(s) are connected.
- Consult the manufacturer or field service technician for help.
- 16) Do not to touch signal input, signal output or other connectors, and the patient simultaneously.
- 17) a statement that MEDICAL ELECTRICAL EQUIPMENT needs special precautions regarding EMC and needs to be installed and put into service according to the EMC information provided in the ACCOMPANYING DOCUMENTS;
- 18) a statement that portable and mobile RF communication equipment can affect MEDICAL ELECTRICAL EQUIPMENT.
- 19) Please consult a physician or a trained health professional for interpretation of measurement results.
- 20) No phthalates are used for this product and its container.
- 21) The cuff is not made with natural rubber latex

5. SAFETY SYMBOLS AND INFORMATION

The International Electrotechnical Commission (IEC) has established a set of symbols for medical electrical equipment which classifies a connection or warning of any potential hazard.

The classifications and symbols are shown below. Save these instructions for your safety.

†	DEFIBRILLATION-PROOF TYPE BF APPLIED PART
	General Warning Sign. Failure to turn off the power before you begin this procedure can lead to product damage and result in serious injury or death.
<u> </u>	Caution. Failure to follow instructions could result in damage to product or minor injury
0	General mandatory action sign. Important to follow instructions. Not a caution.
	Waste Electrical and Electronic Equipment (WEEE) The device could be sent back to the manufacturer for recycling or proper disposal after their useful lives. Alternatively the device shall be disposed in accordance with national laws after their useful lives.

<u> </u>	identifies the battery
\odot	Turning on the recorder
Ċ	Turning off the recorder
~~ <u></u>	Date of manufacture
	Manufacturer
$((\bullet))$	Non-ionizing radiation
(€ ⁰197	CE mark
SN	Serial No
	Refer to instruction manual/ booklet.
EC REP	Authorized representative in the European community.
	Keep dry
₩oHS2	RoHS2

6. Guidance for Electromagnetic compatibility (EMC)

Details about the electromagnetic compatibility (EMC) of the FA 48 are given below. Before using the FA 48, be sure to read and understand the following information.

1) Guidance and manufacturer's declaration – electromagnetic emissions

The FA 48 is intended for use in the electromagnetic environment specified below. The customer or the user of the FA 48 should assure that it is used in such an environment.

Emissions test	Compliance	Electromagnetic environment – guidance
RF emissions CISPR 11	Group 1	The FA 48 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 emissions CISPR 11	Class B	The EA 49 is suitable for use in all establishments, including
Harmonic emissions IEC 61000-3-2	Class A	The FA 48 is suitable for use in all establishments, including domestic establishments and those directly connected to the public low-voltage power supply network that supplies
Voltage fluctuations/ flicker emissions IEC 61000-3-3	Compliance	buildings used for domestic purposes.

2) Guidance and manufacturer's declaration – electromagnetic immunity

The FA 48 is intended for use in the electromagnetic environment specified below. The customer or the user of the FA 48 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	±6kV: Contact ±8kV: Air	±6kV: Contact ±8kV: Air	Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30 %.
Electrical fast transition/burst IEC 61000-4-4	±2kV: Power supply lines ±1kV: Input/output lines	±2kV: Power supply lines ±1kV: Input/output lines	Mains power quality should be that of a typical commercial or hospital environment.
Surge IEC 61000-4-5	±1 kV differential mode ±2kV common mode	±1 kV differential mode ±2kVcommon mode	Mains power quality should be that of a typical commercial or hospital environment.
Voltage drops, dips, and fluctuations of input power supply line IEC 61000-4-11	<5 % <i>U</i> T (>95 % dip in <i>U</i> T) for 0,5 cycle 40 % <i>U</i> T (60 % dip in <i>U</i> T) for 5 cycles 70 % <i>U</i> T (30 % dip in <i>U</i> T)	<5 % <i>U</i> T (>95 % dip in <i>U</i> T) for 0,5 cycle 40 % <i>U</i> T (60 % dip in <i>U</i> T) for 5 cycles 70 % <i>U</i> T (30 % dip in <i>U</i> T)	Mains power quality should be that of a typical commercial or hospital environment. If the user of the FA 48 requires continued operation during power mains interruptions, it is recommended that the FA 48 be powered from an uninterruptible power supply or a battery.

	for 25 cycles <5 % <i>U</i> T (>95 % dip in <i>U</i> T) for 5 sec	for 25 cycles <5 % <i>U</i> T (>95 % dip in <i>U</i> T) for 5 sec	
Magnetic field of commercial frequency (50/60Hz) IEC 61000-4-8	3 A/m	3 A/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.

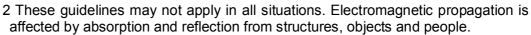
3) Guidance and manufacturer's declaration – electromagnetic immunity 2
The FA 48 is intended for use in the electromagnetic environment specified below. The customer or the user of the FA 48 should assure that it is used in such an environment.

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 FA 48, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter.
			Recommended separation distance $d = 1.2\sqrt{P}$
Conducted RF IEC 61000-4-6	3 Vrms 150 kHz to 80 MHz	3 Vrms	$d = 1.2\sqrt{P}$ 80 MHz to 900 MHz $d = 2.3\sqrt{P}$ 900 MHz to 2,5 GHz
Radiated RF IEC 61000-4-3	3 V/m 80 MHz to 2,5 GHz	3 V/m	where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and d is the recommended separation distance in meters (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:



Note

1. At 80 MHz and 900 MHz, the higher frequency range applies.





- 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 FA 48 is used exceeds the applicable RF compliance level above, the FA 48 should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as reorienting or relocating the FA 48.
- Over the frequency range 150 kHz to 80 MHz, field strengths should be less than 3 V/m.

4) Recommended separation distances between portable and mobile RF communications equipment and the FA 48

The FA 48 is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the FA 48 can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the FA 48 as recommended below, according to the maximum output power of the communications equipment.

Rated maximum	Separation distance according to frequency of transmitter m			
output power	150 kHz to 80 MHz	80 MHz to 900 MHz	900 MHz to 2,5 GHz	
of transmitter W	$d = 1.2\sqrt{P}$	$d = 1.2\sqrt{P}$	$d = 1.2\sqrt{P}$	
0.01	0.12	0.12	0.23	
0.1	0.38	0.38	0.73	
1	1.2	1.2	2.3	
10	3.8	3.8	7.3	
100	12	12	23	

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 900 MHz, the separation distance for the higher frequency range applies.



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

TERMS OF EACH PART AND FUNCTION

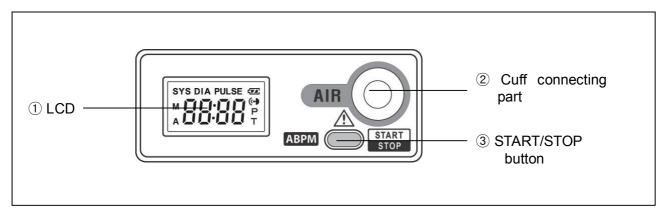
1. DISPLAY

① LCD (Display)

It displays time, battery status and measurement process.

Also, systolic blood pressure, diastolic blood pressure and pulse result will be indicated after measurement.

- Connecting part of cuff
 It connects main unit and connecting hose which connected on cuff.
- START / STOP button If measurement is set with MANUAL mode, measurement can be started with START / STOP button.





SYS: Systolic blood pressure

DIA: Diastolic blood pressure

PULSE: Pulse

 : Battery remains

Data transmitting

M: This icon will be indicated when wake-up time is set.

A: This icon will be indicated when wake-up time automatic measurement is set.

P: This icon will be indicated when bed-time is set.

T: This icon will be indicated when bed-time automatic measurement is set.

12:00: Time

Pressure value will be indicated when measuring and systolic/diastolic will be indicated after measuring.

2. FRONT AND REAR PART

MENU button

This button is for setting the device such as time, measurement interval and etc.

② UP button

Number is going up when setting the device.

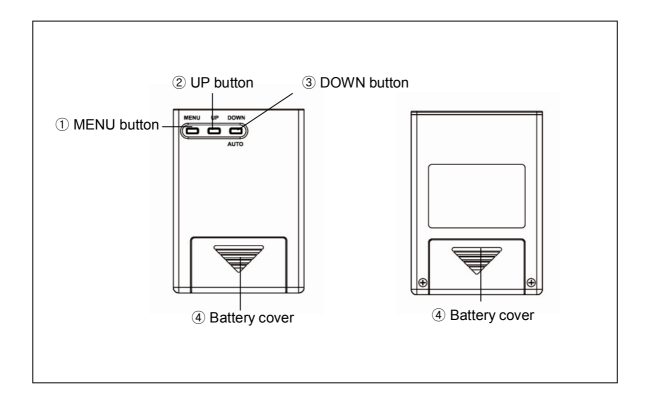
3 DOWN button

Number is going down when setting the device.

Battery cover

It is a cover for protecting battery inlet.

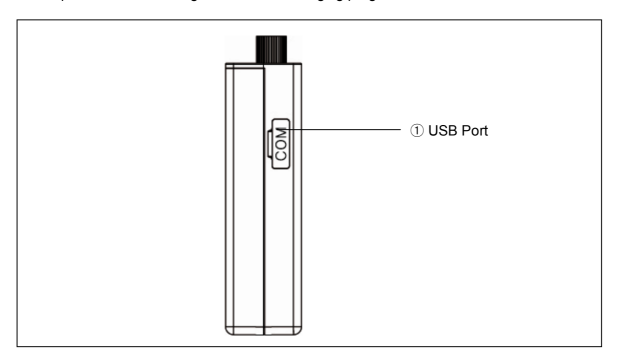
When replace the battery, replace 4 batteries each 2 batteries on front and rear part.



3. SIDE PART

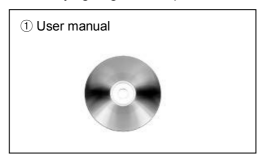
O USB Port

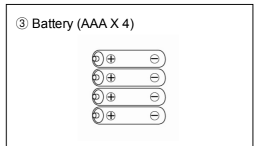
This port is for transmitting data to data managing program on PC.



4. Accessories

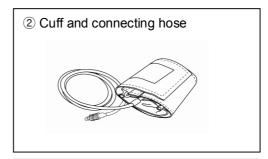
- User manual (CD)
- 3 Battery (AAA X 4)
- S Carrying bag and strap

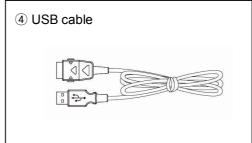






- ② Cuff and connecting hose(Standard)
- 4 USB cable
- Data managing program

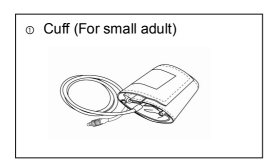




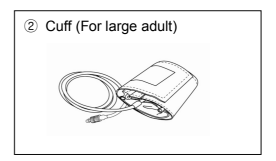


5. Options

O Cuff (For small adult)



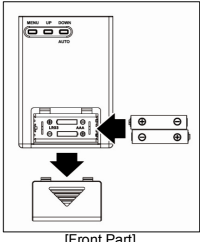
② Cuff (For large adult)

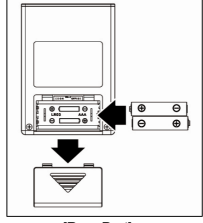


INSTALLATION

1. How to put battery

Put batteries on front and rear part on the device for turning on the device. Open the battery cover and put each 2 batteries (AAA type battery) on front and rear part.

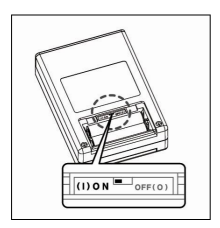




[Front Part]

[Rear Part]

After putting batteries on rear part, turn on the ON/OFF switch as shown in the picture.



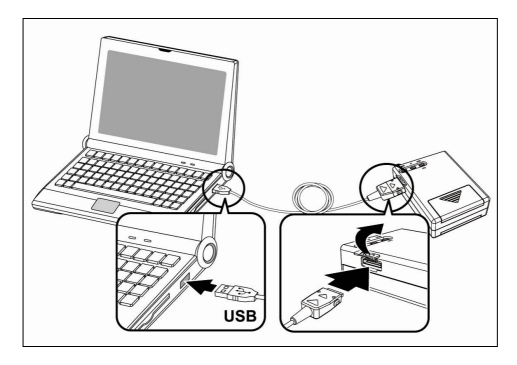
Note



Replace all batteries (4 batteries) when replacing battery. When renting the device, please be sure to put new batteries.

2. How to connect communication port

Connect the USB port which located on side part of main unit and computer USB port by USB cable.



3. Setting

Time will be blinked when turning off the power and you can set up in this state.

If you want to change the setting, press MENU button for 5 seconds then system will enter to setting mode.

Date / Time setting

Setting sequence is hour/minute \rightarrow month/day \rightarrow year.

Hour/minute setting



When you turn ON the power or press the MENU button for few seconds while using, time will be blinked. Set the hour by UP and DOWN button.

When you press the MENU button once again, system moves to minute setting. Press UP and DOWN button for setting minute.

Month/day setting



After setting minute, press MENU button once again for entering month setting mode. Press UP and DOWN button for setting month. Press MENU button once again for entering date setting mode. Press UP and DOWN button for setting day.

Year setting.



After setting day, press MENU button once again for entering year setting mode. Press UP and DOWN button for setting year. After setting day, press MENU button for few seconds for entering 'Alarm setting' mode.

Note



If you want to discontinue the date and time setting, press STOP switch. Then, device will be turning to measurable state. Hour, minute, month, day, year needs to be set together. If you discontinue the process in the middle of setting, time display will be returned to the state before you started setting.

② Alarm setting

Ring the alarm when measured systolic blood pressure is higher or lower than setting value.

Setting maximal blood pressure alarm



After setting the year, press and hold the MENU button, and the device will show 'SYS' and 'H' on the LCD screen. You may set the maximal blood pressure alarm. Set the desired value using the UP and DOWN buttons. When the measured blood pressure is higher than the preset value, the alarm will go off for 90 seconds and the SYS icon will blink.

Setting minimal blood pressure alarm



After setting the maximal value, press the MENU button once again, and the device will show 'SYS' and 'L' on the LCD screen. You may also set the minimal blood pressure alarm. Set the desired value using the UP and DOWN buttons. When the measured blood pressure is lower than the preset value, the alarm will go off for 90 seconds and the SYS icon will blink.

Alarm ON/OFF setting



After setting the minimal blood pressure alarm, press the MENU button once again, and the device will show a bell icon on the top right corner, and the screen will shift to the alarm ON/OFF setting. Adjust alarm on or off using the UP and DOWN buttons. When the alarm is set off, the device does not ring any alarms regardless of the measured value.

③ Pressure value setting

It is required to set the initial pressure for blood pressure measurement. Once the initial pressure is set, repeated pressurization is prevented for hypertension patients.

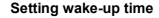


Setting initial pressure

After setting the alarm function, press the MENU button and the LCD screen will display 'PULSE' and enters the initial pressure setting menu. Set the desired value using the UP and DOWN buttons.

Wake-up time setting

Set the measuring intervals at wake-up and after wake-up times.





After setting the desired initial pressure, press and hold the MENU button, and the LCD screen will display 'M' and enter the wake-up time setting menu. Set desired wake-up time using UP and DOWN buttons.

Once the wake-up time is set, press the MENU button again and the minute will blink, and enter the minute setting menu. Set the desired wake-up time (minute) using the UP and DOWN buttons.



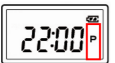


After setting desired wake-up time, press the MENU button again, and the LCD screen will display 'M' and 'A', and enters measuring intervals menu. Set desired interval using UP and DOWN button. The device automatically measures blood pressure at every preset interval, and the device operates manually when the value is set as '0'.

Sed-time setting

Setting the measuring intervals at bed-time and after bed-time

Setting bed-time



After setting the desired measuring intervals, press and hold the MENU button, and the LCD screen will display 'P', and enter the bed-time setting menu. Set the desired bed-time using the UP and DOWN button.

Once the bed-time is set, press the MENU button again and the minute will blink, and enter the minute setting menu. Set the desired bed-time (minute) using the UP and DOWN buttons.

Setting measuring intervals after bed-time



After setting the desired bed-time, press the MENU button again, and the LCD screen will display 'P' and 'T', and enter measuring intervals menu. Set the desired interval using the UP and DOWN button. The device will automatically measure the blood pressure at every preset interval, and the device operates manually when the value is set at '0'.

Note

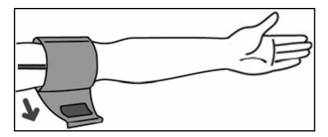


It is recommended to set the bed-time measuring interval wider than that of wake-up time.

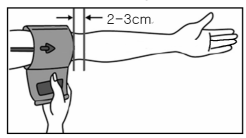
MEASUREMENT

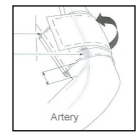
1. Wrapping the cuff

① Wrap the cuff around your upper arm with palm facing up.
Following the directions which are printed directly on the cuff as it should be applied brachial artery. (inside forearm)

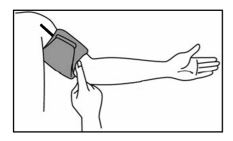


2 Position the lower edge of the cuff is 2~3cm above the elbow joint as shown in the picture.





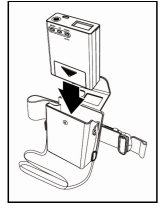
3 Strain the cuff with 1~2 finger space between the cuff and the arm.



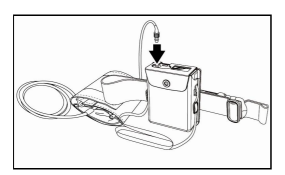
Note This device is only for adult.

2. Measurement preparation

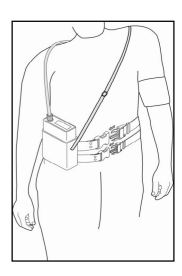
- Insert the cuff-end into the ring so it can be put around the upper arm.
- ② Align the arrow-head (▼) at the upper arm artery (brachial artery), and make the lower-end of the cuff positioned 2-3cm above the elbow joint.
- Put the cuff around the arm, two-finger loose, and make sure the cuff is not folded or misplaced.
- Hang the air-hose around the neck, and put the belt on the flank so that the device is placed on the
 other side of the cuff.
- Place the device in the bag, and make sure the LCD screen is visible from above.



Insert the hose that is connected to the cuff into the cuff-insertion hole, and turn it clock-wise to lock.

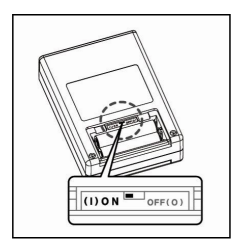


O Adjust the shoulder and waist straps comfortable for the users.



3. Measurement

Turn the power on.



- When the device is turned on, the LCD screen will show the current time, and will be ready for measurements.
- If measuring intervals are preset, the device automatically measures blood pressure according to the preset value. If not, press the START button on the device to begin measuring.

Note



If the cuff pressure is too high and it causes pain, press the STOP button and take the cuff off immediately.

Caution



If the zipper is not completely zipped, the internal robber bag might come out during pressurization, and it may cause damage or rupture the bag.

Note

- If the measuring pressure goes too high when the air-hose is blocked;
 - The device will display 'E1' on the screen.

 If the same message is repeatedly displayed, please request for a repair

service.

- If the air-hose leaks, the connection to the cuff is defected, or the pressure does not go up;
 - The device will display 'E2' on the screen.

If the same message is repeatedly displayed, please request for a repair service.

- If the patient moves or talks during the measurement;
 - The device will display 'E3' on the screen.

Please take a rest, and redo the measurement with accurate measuring techniques.

If the same message is repeatedly displayed even with correct measuring conditions, request for a repair service.

• When an error is detected, the device will make a buzzing sound, and turn back to the measurement-ready status.

4. Measurement result

Display systolic / diastolic blood pressure and pulse on LCD. Reply 3 times.

If you need more detail information about measurement result (Graph, list of 24 hour measured data), Use the data management program FA 48 manager. (Program version: FAMN.E.1.0.00) Reference the manager's user manual about detail contents of data manage program FA 48 manager. (Rev 1.1)



STORAGE AND MAINTENACE

- Do not operate or store the device with below environments.
 - Store where the ambient temperature falls -20°C or exceeds 60°C.
 - Operate where the ambient temperature falls under 10°C or exceeds 40°C.
 - Where the atmospheric pressure falls below 70kPa (700 mbar) or exceeds 106kPa (1060 mbar).
 - Operate where the humidity is over 30~75% non-condensing.
 - Store where the humidity is over 95%.
 - Where the unit is exposed to vapor or water.
 - Where the unit is exposed to dust.
 - Where the unit is exposed to high concentration of oil vapor.
 - Where the unit is exposed to salty atmosphere.
 - Where the unit is exposed to explosive gas or dust.
 - Where the unit is exposed to excessive shocks or vibrations.
- Clean the recorder, cuff and accessories for the next use.
- The recorder is a precision instrument. Please check all functions (every year) periodically. Contact your nearest A/S center for this inspection.
- If this unit is not being used for a long period of time, please remove the batteries.
- Do not drop the unit.
- Do not disassemble the main body or the cuff.
- Do not fold or bend the cuff forcibly.
- Do not wash or dip the cuff in the water.

SAFETY PRECAUTION

- The recorder should be operated by a physician who knows it well.
- Confirm that the recorder works correctly and measurement values are proper.
- Clear the old data before starting a new measurement.
- Cover the USB port using the rubber cap, to avoid dust.
- Do not mix new and used batteries in the recorder.
- Use the recorder only during the time of diagnosis or medical treatment.
- Confirm that the cuff and air hose are connected properly.
- Confirm parts of contact with patient directly. Keep clean for these parts.
- Instruct the patient on how to stop operation if there is an abnormal measurement, and how to remove the cuff if there is excessive arm pain.
- The recorder may not make a measurement when a patient has continuous arrhythmia or the recorder senses noise due to the patients movement.
- Please check measurement values by other methods, if you suspect an value.
- Do not use this recorder on a person who is using a heat lung machine and a defibrillator.
- Avoid strong magnetic field and static electricity.
- Please transfer the data as soon as possible.

TROUBLE SHOOTING

1. Measurement error and solution

ERROR	CAUSE	SOLUTION	
Err 0	 When inflation lever is not sufficient. When air hose is not connected well. When air leaks from the cuff. When battery power is low. 	 Please set higher inflation level. Check if there is any 'CAUSE'. After settle it, monitor again. 	
Err 1	 When a cuff is not connected to a monitor. When a cuff is positioned incorrectly 	Please check if the cuff is connected well to the unit and monitor again.	
Err 2	When the pressurization is over than 300mmHg.	Please turn the power off then on and monitor again.	
Err 3	 When monitoring is carried out the wrong way. When a readed data is not output as a result If noise or movement detected. 	 Please read 'wrapping the cuff' and 'correct measuring posture' in 'Measurement' part. Monitor again following the way instructed. 	
Err 4	When moving or speaking is sensed during monitoring.	Please do not move or speak during monitoring.	
Err 5	When battery is not fixed well. When the polarities are positioned incorrectly.	 Check if the polarities of the batteries are positioned correctly. After disconnecting batteries in unit and press [I/O] button, then replace them. 	

2. Trouble shooting

SYMPTOM	CAUSE	COUNTERMEASURE
¢	The battery power is low. (When batteries are exhausted completely, the mark does not appear on the LCD any more.)	Replace all four "AA" type batteries at the same time. Before change transmit the data to your computer first.
The blood pressure value seems too low.	When your arm positioned higher than your heart during monitoring.	Make sure that your arm position at the same
The blood pressure value seems too high	When your arm positioned lower than your heart during monitoring.	level as your heart, then monitor it again.
Re- pressurization	 In case of insufficient inflation. The movement of the arm or body during inflation is sensed. (In case of excessive hypertension or a feeble pulse, it is possible to occur repressurization over 2 times.) 	 Relax for a few minutes then monitor it again with correct posture. Do not move during monitoring.
The unit does not start monitor again after error occurs.	The cuff may not be positioned incorrectly.	Monitor it again after wearing the cuff correctly and positioning your arm as your heart level.

3. After service

If there is any problem with the unit, please follow the steps below;

- ** Contact our company's Overseas Service Department immediately.
 After gathering the model name, Serial Number, date of purchase and description of the problem, contact our company with information shown below.
- * Try to solve the problem over the phone with the personnel of local service department. If the problem cannot be solved over the phone, just return to service department directly.
- We our company or local distributor will make available on-request circuit diagrams, component part list, descriptions, calibration or other information which will assist your appropriately qualified technical personnel to repair those parts of unit which are designated by our company as repairable.
- * Calibration interval for this device is 2 years.

To ensure the device's proper and sage operation, please contact our company or distributor periodically for calibration.

How to contact our company Write us at:

JAWON MEDICAL CO., LTD

29, Gongdan 4-ro, Jillyang-eup, Gyeongsan-si, Gyeongsangbuk-do, Korea

TEL: 82-53-856-0993 FAX L 82-53-856-0995

(You can also contact the following representative or your local distributor)

4. Packing and transport

Our company wraps this device up with the most suitable method to protect it from any impact or damage during shipping and transporting. This device can be damaged during delivery if it is packed with other ways except the one our company uses. Please handle this device carefully without any impact in packing and delivering it.

If this device needs to be transported wrap this device up again and transport it as follows.

- 1) Turn off the power of this device.
- ② If peripherals are connected, turn off the power of them and then disconnect each device.
- 3 Disassemble this device in reverse order to assembly.
- 4 Wrap up this device with original packing materials.
- (5) Transport it carefully in order not to give a shock to this device.

SPECIFICATION

Model	FA 48
Measuring method	Oscillometric
Display mode	TN LCD display
Result Contents	Systolic / Diastolic / Mean Blood Pressure, Pulse, Pulse Pressure
Measuring range	Pressure: 30~300mmHg, Pulse: 30~240bpm
Accuracy	Pressure: ±3mmHg, Pulse: ±3%
Resolving Power	1mmHg
Pressurizing method	DC Motor
Cuff type	Belt type
Pressurizing time	Approx. 10 seconds
Measuring time	Approx. 1 minute
Power	DC 6V (AAA type 1.5V X 4)
Memory	576 memory
Ambience for operation	Temperature: 10~40 ℃, Homidity: 30~75%
Ambience for storage	Temperature: -20~60 $^{\circ}\mathrm{C}$, Humidity: Less than 95%
Data transmission	USB
Dimension	Main unit: 72(W) × 28(D) × 101(H) mm Cuff: standard - 140(W) X 611(H) mm (±2mm) Small adult - 135(W) X 535(H) mm (±2mm) Large adult - 160(W) X 795(H) mm (±2mm) Cuff Range: standard - 200~310 mm Small adult - 150~220 mm Large adult - < 450 mm Air bag: standard - 120(W) X 240(H) mm Small adult - 110(W) X 210(H) mm Large adult - 140(W) X 350(H) mm
Weight	Approx. 155g (Main + Battery)

WARRANTY and REPAIR

WARRANTY

Item	Ambulatory Blood Pressure Monitor		Warranty period	
Model	FA 48		1year (main unit only)	
Serial NO.				Tyear (main unit only)
Date of purchase		Month	Day	Year
Customer	Name:		•	TEL:
Customer	Address:			
Dealer	Name:	TEL:		
Dealei	Address:			
Date		Defection		Confirmation

All repairs on products under warranty must be performed or approved by Jawon Medical Co., Ltd. Unauthorized repairs will void the warranty. In addition, whether or not covered under warranty, any product repair shall exclusively be performed by Jawon Medical Co., Ltd. certified service personnel.

Repairs

If your product requires warranty, extended warranty, or non-warranty repair service, please call first the nearest Jawon Medical Technical Support Center. A representative will assist you troubleshooting the problem and will make every effort to solve it over the phone, avoiding potential unnecessary return. In case the return cannot be avoided, the Customer Service Representative will guide you through the Return Authorization process.

Limited Warranty Statement

Jawon Medical Co., Ltd. warrants that the FA 48 you have purchased meets the labeled specifications of the Product and will be free from defects in materials and workmanship that occur within 1 year after the date of purchase. Accessories used with the Product are warranted for 90 days after the date of purchase.

The date of purchase is: 1) the date specified in our records, if you purchased the product directly from us, 2) the date specified in the warranty registration card that we ask you to send to us, or 3) if you don't return the warranty registration card, 120 days after the date on which the Product was sold to the dealer from whom you bought the product, as documented in our records.

This warranty does not cover damage caused by: 1) handling during shipping, 2) use or maintenance contrary to labeled instructions, 3) alteration or repair by anyone not authorized by Jawon Medical, and 4) accidents.

If a Product or accessory covered by this warranty is determined to be defective because of defective materials, components, or workmanship, and the warranty claim is made within the warranty period described above, Jawon Medical Co., Ltd. will, at its discretion, repair or replace the defective product or accessory free of charge.

You must obtain a return authorization from Jawon Medical Co., Ltd. to return your product before you send it to Jawon's designated service center for repair.

This warranty is in lieu of all other warranties, express or implied, including but not limited to the implied warranties of merchantability and fitness for a particular purpose. Jawon's obligation under this warranty is limited to repair or replacement of products containing a defect. Jawon Medical Co., Ltd. is not responsible for any indirect or consequential damages resulting from a product defect covered by the warranty.

ACTIVITY RECORD (24 Hour Blood Pressure Measurement) Please read the Instructions and Caution on the back of this sheet.

Name :		Address:	
Home TEL. :			
Office TEL. :			
Patient No :		Chart No :	
Doctor :		Referred By:	
Hospital :		Unit :	
Recorder I.D.#	Date Out :		Date Due In :
·	·	· ·	

MEASUREMENT PERIOD	From:	To:

INTERVAL

No.1	to	(every	min.)	No.4	to	(every	min.)
No.2	to	(every	min.)	No.5	to	(every	min.)
No.3	to	(every	min.)	No.6	to	(every	min.)

ACTIVITY RECORD

TIME	ACTION	SYMPTOMS(CONDITIONS)
:	Starting measurement	
:		
:		
:		
:		
:		
:		
:		
:		
:		
:		
:		
:		
:		
:		

EXAMPLE

TIME	ACTION	SYMPTOMS(CONDITIONS)
11:00	Measurement beginning.	
11:40	Lunch is taken.	
14:25	I talked standing with the friend.	I have gotten excited.

Ambulatory Blood Pressure Monitor FA 48

<Information>

Even for one day, blood pressure is varying depending on the measurer's position and physical state. Therefore, the data about measurer's position, meal, medicine dosage, etc during the measurement is very important for a doctor to analyze blood pressure data for 24 hours.

Motion detecting sensor is embedded in FA 48 and it detects the patient's motion without recording it manually. However, the information of meal, medicine dosage, urination, excretion and how well you sleep and wake/asleep time is as important as the measurer's activity status. So record these data attentively.

Also read the safety caution and operation manual carefully as stated below.

<How to operate ABPM FA 48>

Basically, FA 48 automatically measures the patients according to the set measurement period.

However, it is hard to have a measurement when the patients are changing the clothes or taking a shower. In this case, convert it to manual mode. Please refer to the content mentioned below.



M: It is indicated when the awake time is set.

A: It is indicated when the measurement period is set for activity period. (Automatic measurement after awake time.)

Press 'START/STOP' button for 2 sec. and LCD will be turned off. It removes the Automatic mode and 'A' is not indicated on LCD anymore.

Press 'START/STOP' button again and the Automatic measurement starts.

P: It is indicated when the asleep time is set.

T: It is indicated when the measurement period is set for sleeping period. (Automatic measurement after asleep time.)

Press 'START/STOP' button for 2 sec. and LCD will be turned off. It removes the Automatic mode and 'T' is not indicated on LCD anymore.

Press 'START/STOP' button again and the Automatic measurement starts.

: It indicates the remaining balance of battery.

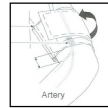


START/STOP Button

When you press this button during the manual mode set up, the measurement starts. During the measurement, press it again to stop it. Set the measurement mode. (AUTO or MANUAL)

<How to wear cuff>

- O Put the ring at the end of the cuff and make it like the shape of arm circumference.
- As shown in the picture at cuff, the mark "ARTERY" has to be on an artery where the inner part
 of pulse is beating.
- As shown in the picture, pull the end of the cuff and wind it around your arm so that cuff stays 2~3cm above your elbow.
- Oheck that there's space for one or two fingers between the cuff and arm.



<Measurement Precautions >

It is an ambulatory blood pressure monitor, but it also measures blood pressure during the rest. Do not move your arm too much during the measurement.

Try to rest during the measurement. Remove the Automatic mode if you have to concentrate on the exercise or driving.

*Press 'STOP' button to cease the measurement and unfasten the cuff if you feel severe pain or the device doesn't work properly.

Periodic Check List

Management No	
---------------	--

Item		Inspection Subject		Requirements		Judgment	Remarks	
Visual Check	(
Mainframe	1	Enclosure		No scratch, crack,		Pass/Fail		
				defo	rmation and ru	st		
	2	Labels and pa	anels	No peel	ing and dust		Pass/Fail	
	3	Keys		No dam	age		Pass/Fail	
	4	Cuffs		No scratch and damage		Pass/Fail		
Accessories	1	Battery		No scratch and damage		Pass/Fail		
	2 User manual		Kept in proper place		Pass/Fail			
Mechanical C	hec	k						
Mainframe	1	Keys		Smooth	operation		Pass/Fail	
	2	Recorder		Smooth	operation with	no	Pass/Fail	
				abno	ormal sound			
	3	Cuffs		Smooth	operation		Pass/Fail	
Accessories 1 Battery		Smooth operation and		Pass/Fail				
				removal				
Electrical Ch	eck							
Performance	1	Battery		Screen	display upon		Pass/Fail	
				pow	er-on			
	2	Display		No abno	rmality and		Pass/Fail	
				flickering				
	3	Connect PC		Connecting possible		Pass/Fail		
	4	Measurement	t	Proper measurement		Pass/Fail		
General Judgment					Pass/Fail			
Model FA 48						Serial No.		
Installation pla	Installation place			Date		of purchase		
Check date Check		Check	ced by		Аррі	roved by		

Copy this sheet for use

If repair is required, write down so in the Remarks column.

Daily Check List

Management No.

Visual Check 1 Enclosure No scratch, crack, deformation and rust Pass/Fail 2 Labels and panels No peeling and dust Pass/Fail 3 Keys No damage Pass/Fail 4 Cuffs No scratch and damage Pass/Fail Accessories 1 Battery No scratch and damage Pass/Fail Mechanical Check Kept in proper place Pass/Fail Mainframe 1 Keys Smooth operation Pass/Fail Accessories 1 Battery Smooth operation and removal Pass/Fail	Remarks
Mainframe 1 Enclosure deformation and rust Pass/Fail	
3 Keys No damage Pass/Fail 4 Cuffs No scratch and damage Pass/Fail Accessories 1 Battery No scratch and damage Pass/Fail Wechanical Check Smooth operation Pass/Fail Mainframe 2 Recorder Smooth operation with no abnormal sound Pass/Fail Accessories 1 Battery Smooth operation and removal Pass/Fail	
4 Cuffs No scratch and damage Pass/Fail Accessories 1 Battery No scratch and damage Pass/Fail 2 User manual Kept in proper place Pass/Fail Mechanical Check Smooth operation Pass/Fail Mainframe 2 Recorder Smooth operation with no abnormal sound Pass/Fail Accessories 1 Battery Smooth operation and removal Pass/Fail	
Accessories 1 Battery No scratch and damage Pass/Fail	
Accessories 2 User manual Kept in proper place Pass/Fail Mechanical Check 1 Keys Smooth operation Pass/Fail Mainframe 2 Recorder Smooth operation with no abnormal sound Accessories 1 Battery Smooth operation and removal Pass/Fail	
2 User manual Kept in proper place Pass/Fail	
Mainframe 1 Keys Smooth operation Pass/Fail Smooth operation with no abnormal sound Pass/Fail Accessories 1 Battery Smooth operation and removal Pass/Fail	
Mainframe 2 Recorder Smooth operation with no abnormal sound Pass/Fail Accessories 1 Battery Smooth operation and removal Pass/Fail	
2 Recorder abnormal sound Pass/Fail Accessories 1 Battery Smooth operation and removal Pass/Fail	
Accessories 1 Battery removal Pass/Fail	
Electrical Check	
1 Battery Screen display upon power-on Pass/Fail	
Performance 2 Display No abnormality and flickering Pass/Fail	
3 Connect PC Connecting possible Pass/Fail	
4 Measurement Proper measurement Pass/Fail	
Other 1 Clock Present date/time Pass/Fail	
General Judgment Pass/Fail	
Model FA 48 Serial No.	
Installation place Date of purchase	
Check date Checked by Approved by	

Copy this sheet for use

If repair is required, write down so in the Remarks column.

Service center
TEL: 82 - 2 - 587 - 4056
82 - 53 - 856 - 0993

If the problems continue, call the service center. When you ask for service, the manufacturer's label, serial number, date of original purchase and explanation of malfunction will be required.

For purpose of improvement, specifications and subject to change without notice.



* Our company name will be changed to JANEX MEDICAL.

HEAD OFFICE: 29, Gongdan 4-ro, Jillyang-eup, Gyeongsan-si,

Gyeongsangbuk-do, Korea

TEL: 82-53-856-0993, FAX: 82-53-856-0995

EUROPEAN REPRESENTATIVE: VITAKO Sp. z o.o.

UL. MALEJ SYRENKI 2 71-790 SZCZECIN, POLAND TEL: 91 8522900 / 901 FAX: 91 8522910