Professional 24-hour ambulatory blood pressure monitor with Atrial Fibrillation (AFIB) detection

Instruction Manual
The Microlife WatchBP O3 AFIB is designed to provide reliable and unbiased ambulatory blood pressure measurement results, and strictly follows European Society of Hypertension (ESH) and American Heart Association (AHA) recommendations for ambulatory blood pressure measurement. The WatchBP O3 AFIB device has been clinically validated according to the ESH protocol. In addition, this ambulatory blood pressure monitor has an implemented algorithm that allows the detection of atrial fibrillation (AFIB) during 24-hour blood pressure measurement.
### Table of Contents

**Product description**
- Name of parts and display ............................................ 6-7

**Before using WatchBP O3 AFIB for the first time**
- Activating the device ....................................................... 8
- Confirm the cuff size ........................................................ 9

**Selecting operation mode** .................................................... 10

**In «AMBULATORY» mode** .............................................. 10-11
- Measurement programming ............................................. 12-14
- Taking blood pressure measurements .......................... 15-19

**In «CASUAL» mode** ....................................................... 20
- Taking blood pressure measurements .......................... 21

**Recording intake medication**
- Medication record .................................................... 22-23

**Special Function: AFIB detection**
- Screening for atrial fibrillation during blood pressure measurement ........................................ 24

### Atrial fibrillation detector
- Important facts about atrial fibrillation .......................... 25

**Viewing, transferring and deleting measurement data**
- Viewing measurement data .................................................. 26-27
- Transferring measurement data ....................................... 28
- Deleting measurement data ............................................ 29

**Appendix**
- Battery indicator and how to replace batteries ............... 30
- Safety, care, accuracy test and disposal ....................... 31-33
- Error messages ........................................................ 34-36
- Technical specifications .................................................. 37
Product description

Name of parts

Cuff socket
USB port
Mode switch
ON/OFF button
M button (Memory)
Display
Medication button
Battery compartment

Battery compartment

Battery compartment

Battery compartment

Battery compartment

Battery compartment

Battery compartment
Display

- Mode indication
- Battery display
- Stored value
- Time left to next measurement
- Morning data
- Evening data
- Date/Time
- Systolic value
- Diastolic value
- Pulse indicator
- Pulse rate
- Number of stored data
Before using the WatchBP O3 AFIB for the first time

Activating the device
Pull out the protective strip from the battery compartment.

Setting the date and time
Upon removing the protective strip or installing new batteries, the year number flashes in the display.

1) Set the year – Use the M button to select the year. Press the ON/OFF button to confirm your selection.

2) Set the month – Use the M button to set the month. Press the ON/OFF button to confirm.

3) Set the day – Press the M button to set the day. Press the ON/OFF button to confirm.

4) Set the time – Once you have set the hour and minutes and pressed the ON/OFF button the date and time are set, and the current time is displayed.
5) To change the date and time, briefly take out and put back one battery from the battery compartment. The Year number will flash. Complete the process as described above.

Confirm the cuff size

A variety of different cuff sizes are available.

⚠️ Only use Microlife cuff!

✔️ Washable cuffs are available in both nylon and cotton.

S

M

L

XL

S, XL size cuffs can be ordered optionally.

✔️ M, L size cuffs are standard delivered.
Selecting the operation mode

Prior to each measurement, use the mode switch on the side of the device to select the preferred measurement mode. The WatchBP O3 AFIB offers two measurement modes: «AMBULATORY» and «CASUAL».

«AMBULATORY» mode
Select «AMBULATORY» mode for fully programmable 24-hour ambulatory blood pressure measurement.
Programmable measurement intervals

The device automatically takes measurements at fixed intervals of 15, 20, 30, or 60 minutes, as can be programmed.

Two measurement periods

The awake (day) measurement period can be programmed to start at any hour between 00 to 23, and the asleep (night) measurement period can be set to any hour between 00 to 23. Measurement intervals can be set to 15, 20, 30, or 60 minutes for both awake and asleep time periods.

The default measurement interval is 30 minutes for the awake hours and 60 minutes for the asleep hours.
Measurement programming in «AMBULATORY» mode

Installing the software program
1) Put the CD in the CD-ROM drive of your computer. Alternatively click on «setup.exe» in the CD’s directory.
2) Simply follow the instructions provided in the installation window on the computer screen.
3) When the installation is finished, restart the computer before working with the program for the first time.

Connecting the device to a computer
1) Connect the device to the PC. A successful connection is displayed by «PC» on the device.
2) Start the software program.
3) Enter name, identity number, and date of birth to create a new record (if required).

Software can also be downloaded from the website www.watchbp.com.

User manual can be found in the software by clicking on «Help» in the main window.
Programming the measurement procedure

1) With “Ambulatory settings” the 24-hour measurement procedure can be programmed. Choose at what time the Day Period and Night Period starts. Using the drop-down menu on the lower-left of the screen, choose from 00 to 23 as the start of day and night periods. For instance the awake time is 06 AM and asleep time is 10PM.

2) Assign the measurement intervals for the awake and asleep hours by using the drop-down menu to choose from 15, 20, 30, and 60 minutes as the measurement interval.
3) Setting ambulatory options.

<table>
<thead>
<tr>
<th>Setting ambulatory options.</th>
</tr>
</thead>
<tbody>
<tr>
<td>☑️ Hiding BP data</td>
</tr>
<tr>
<td>☑️ Silent mode</td>
</tr>
</tbody>
</table>

On default the blood pressure data are hidden and the device is in silent mode. In order to show the measured blood pressure values on the LCD screen and to provide a beep before each measurement, click on the squares to the left.

4) For setting the highest inflation pressure click on the circle left to “Enable” and click on the arrow to determine the height of the inflation pressure. “Disable” means that the device will automatically search for the right inflation pressure (default).

5) Once you have completed the settings, press “Program” to program the schedule into the device.
Taking blood pressure measurements in «AMBULATORY» mode

Be sure the device is set to «AMBULATORY» mode.

1) **Preparing measurement arm** – Remove all clothing covering or constricting the measurement arm.

- Avoid rolling up long sleeves, as this may lead to constriction of blood flow to the measurement arm.

2) **Positioning the cuff and device** – Fit the cuff closely, but not too tight. Make sure that the cuff is 2~3 cm (1 inch) above the elbow with the tube on the inside of the arm. The cuff tube should point upward and be mounted over the patient’s shoulder.

- Adjust the tube holder on the belt of the pouch so it is properly positioned on the shoulder.
Taking blood pressure measurements in «AMBULATORY» mode (cont.)

3) **Proper positioning of the cuff using the anchor strap**

**Option 1:** Use the Shoulder sling ① and the Anchor strap ② to position the cuff as demonstrated in the diagram below.

![Diagram showing proper positioning of the cuff using the Anchor strap.]

Instructions: Place the Shoulder sling over the shoulder. Insert the device into the pouch. Position the device at front. Apply the cuff to the arm. Buckle up the Anchor strap with the Shoulder sling. Snap the connector of the Anchor strap into the D-ring of cuff. Adjust length for fit and comfort.

**Option 2:** Use the Shoulder strap ③ and the Anchor Strap ② to position the cuff as demonstrated.

![Diagram showing proper positioning of the cuff using the Anchor strap.]

Instructions: Place the Shoulder strap over both shoulders as demonstrated in the diagram. Connect the device to a belt. Position the device at front. Apply the cuff to the arm. Buckle up the Anchor strap with the Shoulder strap. Snap the connector of the Anchor strap into the D-ring of the cuff. Adjust the length for fit and comfort.
4) **Next measurement indicator** – The device will display the next measurement time, indicated by a countdown in minutes on the display.

5) One minute before the next scheduled measurement, the device will partially inflate and immediately deflate the cuff to remind the patient of the upcoming measurement.

*The screen will display X countdown minutes to indicate the time to the next measurement.*

*Measurement reminders will be disabled for asleep hours.*
Taking blood pressure measurements in «AMBULATORY» mode (cont.)

6) **Measurement reminder** – Five seconds before the next measurement, the device can emit a short series of beeps to notify the patient of the coming measurement (optionally). The beep will be off on default.

7) **During measurements** – the patient should be reminded to remain still, refrain from talking, and to breathe normally during the measurement. If the patient is occupied when a measurement begins, the patient should, when safe, relax the measurement arm.

8) **Repeat measurements in case of an error** – The device will automatically repeat the measurement after a two minute countdown should an error occur during measurement.

- The beeper of «AMBULATORY» mode can be enabled via software setting. (see P.14)

- The device will not beep prior to asleep measurement in «AMBULATORY » mode.

- If the repeat measurement encounters an error again, the device will take an extra measurement after a four minutes countdown. If the extra measurement is not successfully either, the device will record an error message.

- If the device stops taking measurements, the patient should return the device to the physician to determine the cause of error.

- The patient can stop any single measurement at anytime by pressing the ON/OFF button.
9) **Storing measurement data** – The WatchBP O3 AFIB device can store up to 250 single measurements in «AMBULATORY» mode.

When the memory is full, each new measurement will automatically overwrite the earliest measurement.

The display of blood pressure values on the LCD screen can be disabled in «Ambulatory» mode by means of software setting.
Taking blood pressure measurements in «CASUAL» mode

In «CASUAL» mode, the device functions like a regular blood pressure monitor – single measurements are automatically stored and can be reviewed at a later stage. The casual mode can be used to perform a single blood pressure measurement to test if the device is functioning properly. Do not forget to put the device back in «Ambulator» mode before sending away the patient for 24-hour blood pressure measurement.

Taking blood pressure measurements

Be sure the device is set to «CASUAL» mode.

1) **Applying the cuff** – The cuff should fit closely around the arm, but not too tight, and be placed 2~3cm above the elbow with the tube on the inside of the arm. The patient should support the arm at the same height as the heart during the measurement.

Additional instructions on taking proper blood pressure measurement can be found on [www.watchbp.com](http://www.watchbp.com).
2) **Start the measurement** – Press the ON/OFF button to start the measurement.

3) **During the measurement** – The cuff will inflate automatically. A single measurement will be performed by each push on the ON/OFF button.

4) **Finishing the measurement** – Once the measurement is complete, the measurement value is automatically stored for later reference. If an error displays after the measurements, the process should be repeated.

5) **250 measurements safely stored** – The WatchBP O3 AFIB device can store up to 250 single measurements in «CASUAL» mode.

ⓘ *When the memory is full, each new measurement will automatically overwrite the earliest measurement.*
Recording the taking of medication

Medication record
The patient can record the time of medication intake by pressing the medication button.

1) Press and hold the medication button for 2 seconds and the Pill icon will appear.

Press the pill button and hold two seconds until “Pill” appears.

2) Release the Medication button and the Pill icon will flash in alternation with the recorded time.

3) Medication record is saved after the beep sound.

☞ Medication record can be used in both modes.

☞ The beeper can be disabled via software setting.
50 records safely stored
The WatchBP O3 AFIB device can store up to 50 medication records.

 Quando la memoria è piena, l'apparecchio visualizzerà “Full”.

Delete stored medication records
To clear the memory of all medication records, press and hold the medication button for 7 seconds. The “CL” symbol will flash. Press the M button to delete the memory or the ON/OFF button to cancel the deletion.

Pressing the M button to delete data will erase all medication recordings in the selected mode.
Special Function: atrial fibrillation (AFIB) screening

Screening for atrial fibrillation during blood pressure measurement
The device is designed to screen atrial fibrillation during blood pressure measurements in the «AMBULATORY» mode. This device is able to detect atrial fibrillation with high accuracy: a sensitivity of 97% and a specificity of 89%*. If atrial fibrillation is detected this will be shown in the report.

AFIB Detector
The device can screen for atrial fibrillation during blood pressure measurement.
Occasionally, the device might falsely detect atrial fibrillation which can have two causes:
1) The arm has moved during blood pressure measurement. For this reason it is of essential importance to keep the arm still during the measurement.
2) Some arrhythmia (irregular heart beat) other than atrial fibrillation might be present.


This device detects atrial fibrillation, a major cause of stroke. Not all risk factors for stroke, including atrial flutter, may be detected by this device.
This device may not detect atrial fibrillation in people with pacemakers or defibrillators.
Important facts about atrial fibrillation

Atrial fibrillation is a common heart rhythm problem and a common cause of major strokes. It affects 8% of those 65 years and older and about 20% of all strokes are caused by atrial fibrillation.

Atrial fibrillation is a rhythm problem that can last from a few minutes, to days or weeks and even years. Atrial fibrillation can lead to the formation of blood clots in the heart. These clots can break off and flow to the brain causing stroke.

One sign of atrial fibrillation is palpitations. However, many people have no symptoms and therefore may remain undetected whereas diagnosing atrial fibrillation early followed by adequate treatment can largely reduce the chance of getting a stroke.
Viewing, transferring and deleting measurement data

Viewing measurement data
1) Use the mode switch to first select the type of measurements to be viewed.

2) Then press the M button

In «AMBULATORY» mode
1) When the M button is pressed, it briefly displays the total number of measurements stored, e.g. N=40, followed immediately by the average of all measurement data stored in the memory.

«A» is displayed when the number shown is the average of all data.

«- -» will display when the number of measurements is less than 12.
2) Pressing the M button briefly displays the total number of awake measurements stored, e.g. N=20, followed immediately by the average of all measurements during awake hours.

3) Pressing M button once again briefly displays the total number of nighttime measurements stored, e.g. N=20, followed immediately by the average of all measurements during night hours.

In “CASUAL” mode

1) When the M button is pressed, it briefly displays the total number of measurement data stored, e.g. N=63, followed immediately by the average of all measurement data stored in memory.

2) All individual readings can be viewed by repeatedly pressing the M button.

All individual readings can be viewed by repeatedly pressing the M button.

The display of BP reading and memory of Ambulatory Mode can be disabled via software setting.
Viewing, transferring and deleting measurement data on the PC. (cont.)

Transferring measurement data

1) Connect the device to the PC. A successful connection is displayed by «PC» on the device.
2) Start the software program.
3) To transfer the data in both modes and the medication compliance record, click «Download BP data to PC».

Software commands

<table>
<thead>
<tr>
<th>Store data</th>
<th>Click «Save Excel report» or «Save PDF report», the file name is formed automatically from the patient’s identity number and the suffix «WatchBPO3 AFIB_(date). xls». or WatchBPO3 AFIB_(date). pdf&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>View the data</td>
<td>Click «Open file»</td>
</tr>
<tr>
<td>Delete the memories</td>
<td>Click «Clear Memory»</td>
</tr>
<tr>
<td>Close the program</td>
<td>Click «Exit»</td>
</tr>
</tbody>
</table>

Please unplug and re-plug the USB cable to the PC if the software instructs to do so.

The program will prompt to synchronize the date/time of the device to that of the PC.

Before using «Clear Memory» or «Exit» make sure the data are stored.

Press «Help» to open user manual of software

System Requirements: 550MHz CPU, 256MB Memory, 1024x768 pixel resolution, 256 color, CD-ROM drive, 1 free USB port, 40MB free hard disk space, Microsoft Windows XP / Vista / Win7 / Win8.
Deleting measurements
Data from «AMBULATORY» and «CASUAL» mode measurements can be deleted independent of each other.

1) Use the mode switch to first select the type of measurements you wish to delete.

2) Press the M button and hold it until the CL symbol flashes.

3) Release the M button and press it once more while the delete symbol flashes.

☞ Each deletion will erase the entire records of each mode.
Appendix

New batteries should be used for each patient monitoring session. “AAA” alkaline batteries are the main power source of WatchBP O3 AFIB.

How to Replace batteries
1) Open the battery compartment at the back of the device.
2) Replace the batteries – ensure correct polarity as shown by the symbols in the compartment.

Battery indicator
When the device is connected to a computer, the pump runs for 1 second instantaneously and “Pc --“ is displayed together with voltage of the batteries on the PC screen. If the voltage is low, the buzzer of the device beeps. The battery icon and voltage number are also displayed on the screen of the device. It is a reminder for replacing batteries. The buzzer keeps beeping until the batteries are replaced.

Caution: do not use rechargeable batteries.
Use 4 new, long-life 1.5V, size AAA alkaline batteries.
Do not use batteries out of expiration date.
Remove batteries if the device will not be used for a prolonged period.
Safety, care, accuracy test and disposal

Safety and protection
This device may only be used for the purposes described in these instructions. The device comprises of sensitive components and must be treated with caution. The manufacturer cannot be held liable for damage caused by incorrect application.

Read the instruction manual carefully before using this device, especially the safety instructions, and keep the instruction manual for future use.

- Ensure that children do not use the device unsupervised; some parts are small enough to be swallowed.
- Only activate the pump when the cuff is installed.
- Do not use the device if you think it is damaged or if anything appears unusual.
- Read the further safety instructions in the individual sections of the instruction manual.
- Do not connect the device to a computer until prompted to do so by the computer software.

Observe the storage and operating conditions described in the “Technical specifications” section of this manual.

- Protect the device from water and moisture
- Protect the device from direct sunlight
- Protect the device from extreme heat and cold
- Avoid proximity to electromagnetic fields, such as those produced by mobile phones
- Never open the device
- Protect the device from impact and drops
Device care
Use a soft cloth with one of the following recommended cleaning solutions to wipe the exterior of the device:

- Mild soap and water
- Hydrogen peroxide solution (3% diluted with water)
- Sodium hypochlorite solution
  (1:10 dilution of household chloride bleach in water)
- Isopropyl alcohol (70% solution)

Then wipe the exterior of the device with a soft, dry cloth.

Cleaning the cuff
Take out the bladder. Fold and place the cuff cover inside a washing bag. Wash the cuff cover with warm water (43°C; 110°F) and a mild detergent in the washing machine.

Pasteurization: wash the cuff cover in 75°C (167°F) hot water for 30 minutes.

Do not iron the cuff!
Accuracy test
We recommend the WatchBP O3 AFIB device to be tested for accuracy every 2 years or after mechanical impact (e.g. being dropped). Please contact Microlife to arrange an accuracy test.

Disposal
Batteries and electronic instruments must be disposed of in accordance with the locally applicable regulations, and not as domestic waste.
Error messages

If an error occurs during measurement, the measurement is interrupted and an error message «Er» is displayed.

<table>
<thead>
<tr>
<th>Error</th>
<th>Description</th>
<th>Potential cause and remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td>«Er 1»</td>
<td>Signal too weak</td>
<td>The pulse signals on the cuff are too weak. Reposition the cuff and repeat the measurement.</td>
</tr>
<tr>
<td>«Er 2»</td>
<td>Error signal</td>
<td>During the measurement, error signals were detected by the cuff, caused for instance by movement or muscle tension. Repeat the measurement, keeping your arm still.</td>
</tr>
</tbody>
</table>

- The device will take another measurement automatically when an error occurs.
- Please consult your doctor, if this or any other problem occurs repeatedly.
- If you think the results are unusual, please read through the information in this instruction manual carefully.
| «Er 3» | No pressure in the cuff | An adequate pressure cannot be generated in the cuff. A leak may have occurred. Replace the batteries if necessary. Repeat the measurement. |
| «Er 5» | Abnormal result | The measuring signals are inaccurate and no result can therefore be displayed. Read through the checklist for performing reliable measurements and then repeat the measurement. |
| «HI» | Pulse or cuff pressure too high | The pressure in the cuff is too high (over 300 mmHg) OR the pulse is too high (over 200 beats per minute). Relax for 5 minutes and repeat the measurement. |
| «LO» | Pulse too low | The pulse is too low (less than 40 beats per minute). Repeat the measurement. |
Technical Specifications

Operating temperature: • 10 - 40 °C / 50 - 104 °F

Storage temperature: • -20 - +55 °C / -4 - +131 °F
• 15 - 90 % relative maximum humidity

Weight: • 260g (including batteries)

Dimensions: • 115 x 80 x 35 mm

Measuring procedure: • oscillometric, corresponding to Korotkoff

Measurement range: • 30 - 280 mmHg – blood pressure
• 40 - 200 beats per minute – pulse

Cuff pressure display: • Range: 0 - 299 mmHg
• Resolution: 1 mmHg
• Static accuracy: pressure within ±3 mmHg
• Pulse accuracy: ±5 % of the readout value

Power source: • 4X1.5 V Batteries; size AAA

Reference to Standards:
• Device corresponds to the requirements of the standard for non-invasive blood pressure monitor.
  EN 1060-1
  EN 1060-3
  EN 1060-4
  IEC 60601-1
  IEC 60601-1-2

Electromagnetic compatibility:
• Device fulfils the stipulations of the standard IEC 60601-1-2.

The stipulations of the EU Directive 93/42/EEC for Medical Devices Class all have been fulfilled.

Type BF applied part

Microlife reserves the right to alter technical specifications without prior written notice.
This device is covered by a two-year guarantee from the date of purchase. This guarantee is valid only on presentation of the guarantee card completed by the owner confirming date of purchase or purchase receipt. Batteries and wearing parts are not covered by this guarantee.

Name: ____________________________
Address: __________________________
Date: ____________________________
Telephone: ________________________
Email: ____________________________

Product: WatchBP O3 AFIB
Product Number: BP3MZ1-1A
Serial Number:
Date: 
<table>
<thead>
<tr>
<th>Region</th>
<th>Company Name</th>
<th>Address</th>
<th>Phone</th>
<th>Fax</th>
<th>Email</th>
<th>Website</th>
</tr>
</thead>
<tbody>
<tr>
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