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The information contained in this manual is subject to change without notice. All changes will be in compliance with regulations governing manufacture of medical equipment.

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 Welch Allyn or their authorized service personnel.

Accessories
The Welch Allyn warranty can only be honored if you use Welch Allyn approved accessories and replacement parts.

Caution
Use of accessories other than those recommended by Welch Allyn may compromise product performance.
Warranty, Service, and Spare Parts

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

Assistance and Parts
If the product fails to function properly or if assistance, service, or spare parts are required, contact the nearest Welch Allyn Technical Support Center.

<table>
<thead>
<tr>
<th>Location</th>
<th>Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>USA</td>
<td>1-800-535-6663</td>
</tr>
<tr>
<td>Canada</td>
<td>1-800-561-8797</td>
</tr>
<tr>
<td>Latin America</td>
<td>(+1) 305-669-9003</td>
</tr>
<tr>
<td>South Africa</td>
<td>(+27) 11-777-7555</td>
</tr>
<tr>
<td>European Call</td>
<td>(+353) 46-90-67790</td>
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<td>Australia</td>
<td>(+61) 2-9638-3000</td>
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<td>United Kingdom</td>
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<td>Singapore</td>
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<td>(+33) 1-55-69-58-49</td>
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<tr>
<td>Japan</td>
<td>(+81) 42-703-6084</td>
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<td>Germany</td>
<td>(+49) 695-098-5132</td>
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<tr>
<td>China</td>
<td>(+86) 21-6327-9631</td>
</tr>
<tr>
<td>Netherlands</td>
<td>(+31) 202-061-360</td>
</tr>
<tr>
<td>Sweden</td>
<td>(+46) 85-853-65-51</td>
</tr>
</tbody>
</table>

Before contacting Welch Allyn it is helpful to attempt to duplicate the problem and to check all accessories to ensure that they are not the cause of the problem.

When calling, please be prepared to provide:
- Product name and model number and complete description of the problem
- The serial number of your product (if applicable)
- The complete name, address and phone number of your facility
- For out-of-warranty repairs or spare parts orders, a purchase order (or credit card) number
- For parts order, the required spare or replacement part number(s)

Repairs
If your product requires warranty, extended warranty, or non-warranty repair service, please call first the nearest Welch Allyn 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 representative will record all necessary information and will provide a Return Material Authorization (RMA) number, as well as the appropriate return address. A Return Material Authorization (RMA) number must be obtained prior to any return.

Note: Welch Allyn does not accept returned products without an RMA.

Packing Instructions
If you have to return goods for service, follow these recommended packing instructions:
- Remove all hoses, cables, sensors, power cords, and ancillary products (as appropriate) before packing, unless you suspect they are associated with the problem.
- Wherever possible use the original shipping carton and packing materials.
- Include a packing list and the Welch Allyn Return Material Authorization (RMA) number.

It is recommended that all returned goods be insured. Claims for loss or damage to the product must be initiated by the sender.
Limited Warranty Statement
Welch Allyn, Inc. warrants that the Welch Allyn CardioPerfect Workstation computer based product you have purchased meets the labelled 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 labelled instructions, 3) alteration or repair by anyone not authorized by Welch Allyn, and 4) accidents.

You assume all responsibility for use of the Product with any hardware or software that does not meet the system requirements described in the Product documentation.

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, Welch Allyn will, at its discretion, repair or replace the defective Product or accessory free of charge.

You must obtain a return authorization from Welch Allyn to return your Product before you send it to Welch Allyn’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. WELCH ALLYN'S OBLIGATION UNDER THIS WARRANTY IS LIMITED TO REPAIR OR REPLACEMENT OF PRODUCTS CONTAINING A DEFECT. WELCH ALLYN IS NOT RESPONSIBLE FOR ANY INDIRECT OR CONSEQUENTIAL DAMAGES RESULTING FROM A PRODUCT DEFECT COVERED BY THE WARRANTY.
# ABPM 6100 Hardware Manual

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1. Overview

The purpose of this manual is to provide an overview of information that you will need to safely and effectively use the ABPM 6100 monitor. Information about the CardioPerfect Workstation Software is available in the CardioPerfect Workstation software manual. Make sure that you familiarize yourself with all of the safety precautions listed in this chapter before attempting to use either device or the software.

Intended Use

PROPER USE OF THE ABPM 6100 MONITORS:
1. The ABPM monitors are intended for use by a trained medical practitioner. Interpretation of blood pressure measurements should always be performed by a physician.
2. The reliability of all the monitors and the software depend upon conformance with the operation and maintenance instructions in this manual.
3. The ABPM 6100 monitors are designed for use with patients with normal sinus rhythms.
4. Measurement accuracy for the ABPM 6100 monitors may be affected by position of the subject, physical conditions, movement, and use outside of the operating instructions contained in this manual.

Conventions

- Caution: consult accompanying documents
- Identifies information within the manual to avoid injury
- Serial Number
- Defibrillation-proof type BF applied part
- Meets essential requirements of European Medical Device Directive 93/42/EEC
- Manufacturer
- No protection against the ingress of liquids
- Identifies the battery
- Do not dispose of this product as unsorted municipal waste. Prepare this product for reuse or separate collection as specified by Directive 2002/96/EC of the European Parliament and the Council of the European Union on Waste Electronic and Electrical Equipment (WEEE). If this product is contaminated, this directive does not apply.
- For more specific disposal information, see www.welchallyn.com/weee, or contact Welch Allyn Customer Service.
EMC Framework of Australia

Recycle
Safety and Effectiveness Warnings and Cautions

Safety and Effectiveness Warnings

Make sure that you are familiar with all Safety and Effectiveness considerations before using the ABPM 6100 monitor.

IMPORTANT WARNINGS

The following warnings apply to the ABPM 6100 Ambulatory Blood Pressure Monitors:

- **Warning**
  - Patient Injury risk. Do not place the cuff on any area where circulation is compromised. Instruct the patient to remove the cuff immediately and notify the doctor if they are experiencing pain, swelling, redness or numbness in the limb where the cuff is placed as this may indicate compromised circulation.
  - The ABPM 6100 has not been designed for use with high frequency (HF) surgical equipment and does not protect against hazards to the patient.
  - Ensure the location of the ABPM 6100 provides maximum separation away from all sources of high-frequency energy.
  - DO NOT use the ABPM 6100 monitor in the presence of flammable anaesthetics due to risk of explosion.
  - DO NOT immerse the ABPM 6100 monitor in any fluid, place any fluid on the monitor, or attempt to clean either monitor with liquid detergents or cleaning agents. If any of these instances occurs, return the unit to an authorized Welch Allyn Service Center. The ABPM 6100 may be cleaned with a damp cloth only.
  - DO NOT remove the ABPM 6100 covers. Neither unit contains any serviceable parts.
  - DO NOT use the monitor if it has failed any diagnostic self-test.
  - DO NOT use the unit if it displays a pressure greater than zero when no cuff is attached. This could lead to inaccurate measurements.
  - DO NOT attach the cuff to a limb being used for intravenous infusions. This may cause the infusion to be blocked and cause the patient harm.
  - DO NOT substitute any component for those supplied by Welch Allyn.
  - DO NOT attempt to repair the unit yourself. Repairs should be performed only by authorized Welch Allyn Service Centers.
  - DO NOT attach the cuff to a patient while the PC Serial Connector is attached to the unit.
Caution

The ABPM 6100 is intended to be used as an ambulatory blood pressure recorder that is sent home with the patient to record their blood pressure over a 24 hour period. It is used to gather trending information in non-critical settings. It is not intended to provide clinicians with real time monitoring of patients in an ER or critical care environment.

The ABPM 6100 monitors are not intended for use on pregnant women or neonates.

The ABPM 6100 units may not yield accurate blood pressure measurements for patients experiencing moderate to severe arrhythmias.

Check to ensure that the operation of the unit does not result in prolonged impairment of the patient's circulation. Instruct the patient to manually remove the cuff if it fails to deflate within three minutes.

Avoid compression or restriction of pressure tubes.

The ABPM 6100, the cuff and tubing are defibrillator protected. The ABPM 6100 has no specific precautions during defibrillation, and defibrillation has no effect on the ABPM 6100 monitor.
2. ABPM 6100 Monitor

2.1 Introduction

The ABPM 6100 Unit is worn by the patient in a belt or shoulder strap and is connected to a cuff, which is wrapped around the non-dominant arm. ABPM 6100 inflates the cuff at pre-programmed intervals throughout the day and measures blood pressure using the oscillometric method, which senses the cessation of pressure waves in the artery when occluded by pressure in the cuff. Heart rate can also be measured using the frequency of pressure waves.

Blood pressure measurements made by the ABPM 6100 are equivalent to those made by a trained observer using the cuff/stethoscope auscultation method within the limits prescribed by the American National Standard, Electronic or Automated Sphygmomanometers.

To get the most out of the ABPM 6100, you should read this section of the manual thoroughly. You will also need to read and understand the CardioPerfect Workstation software manual to properly interface the ABPM 6100 monitor with the software.

Checklist

Check to make sure that the ABPM 6100 package contains the following:
- ABPM 6100 Monitor, PC Interface Cable, Belt and Shoulder Strap,
- Warranty Card, Patient Diary and Cuff Anchors pads
- Adult Size Cuff
- 4 AA Batteries
- CardioPerfect Workstation Software Manual CD
- Monitor Pouch

⚠️ Caution

Substitution of a component different from that supplied might result in measurement error!

Remember to fill out your warranty registration card and send it to Welch Allyn as soon as possible. Report any damaged or missing components to your authorized Welch Allyn representative.
2.2 Operation

This section provides a brief overview of the ABPM 6100 unit, how to load the batteries, and the unit controls.

The ABPM 6100 features a simple design. One Start/Stop button functions as the primary control. The LCD (liquid crystal display) displays easy-to-read information. The CardioPerfect Workstation Software allows you to program the unit before the study and retrieve data after the study. A single air hose connector allows you to connect the cuff to the unit.

The Start/Stop button allows you to perform the following functions:
- Turn the unit On when it is Off.
- Put the unit into Study Mode and take a reading when the unit is in Normal Mode. The time will flash on the LCD when the unit is in Normal Mode.
- Turn the unit Off from Normal or Study mode when the button is pressed and held until the unit beeps five times (approximately five seconds)
- Initiate a reading when the unit is in Study Mode.
- Abort a reading and deflate the cuff if pressed when the unit is taking a reading.

On the back of the ABPM 6100 monitor, a label lists the model and serial number of the unit. The first four digits of the serial number correspond to the year that the unit was manufactured. The battery compartment is on the back of the unit. The connector on the bottom of the unit allows you to connect the unit to a PC using the PC Interface Cable.

Batteries

The ABPM 6100 monitor requires 2 AA batteries. Batteries are installed into the ABPM 6100 unit in the battery bay located at the back of the unit. If rechargeable batteries are used, please refer to the manufacturer’s guidelines for safe use and adequate maintenance. When batteries are properly loaded and first installed, the unit will display the following:
- Incrementing dashes for 2 seconds.
- Two sets of numbers, the first set of three being the Software Version
- Battery voltage displayed for 2 seconds (prior to the voltage, the letter “b” will present)
- Three quick audible beeps.
- The number of BP readings (if any) in memory along with a flashing printer symbol for 3 seconds (the number of readings may not be displayed if the batteries were removed before the unit was turned off)
- One long audible beep.
- Displays flashing time for 20 seconds (after 20 seconds the units turns itself off and goes into Sleep Mode to conserve battery life)
At this point the unit will be ready to upload a BP Study. When the unit is turned on subsequently, the unit will display the following:
- Three quick audible beeps.
- The number of BP readings (if any) in memory along with a flashing printer symbol for 3 seconds.
- One long audible beep.
- Displays flashing time for 20 seconds (after 20 seconds the units turns itself off and goes into Sleep Mode to conserve battery life)

The LCD Display

The LCD displays the following information depending upon the state that the unit is in:
- The time is displayed whenever the unit is in Normal Mode and ready for an action.
- A sun symbol is displayed whenever the buzzer is on (usually during the day).
- A Crescent Moon symbol is displayed when the buzzer is off (usually during programmed sleeping periods).
- A clock is displayed whenever the unit is in Study Mode.
- A battery symbol is displayed whenever the battery voltage is low and the batteries need to be replaced.
- A printer symbol indicates that the unit contains readings in the memory.
3. Interface

This section describes how to connect the ABPM 6100 monitor to your PC and how to set up patients for monitoring sessions.

To set up the ABPM 6100 for communication to your PC for the first time:

**Windows XP**

1. Right-click on My Computer.
2. Select Properties.
3. Click the Hardware tab.
4. Click the Device Manager.
5. Double-click Ports (COM & LPT)
6. Plug the ABPM 6100 PC Interface Cable into a USB port on your PC. Allow Windows to install the driver.

**Windows 7/ Windows 8**

1. Right-click on Computer.
2. Select Properties.
3. Click the Device Manager.
4. Double-click Ports (COM & LPT)
5. Plug the ABPM 6100 PC Interface Cable into a USB port on your PC. Allow Windows to install the driver.
To configure Welch Allyn CardioPerfect Workstation for use with the recorder:

2. In the File menu, click Settings and click Ambulatory blood pressure.
3. Click the Advanced tab.
4. Select device.
5. Set the COM-port number to the number of the serial port to which the ABP recorder is connected.
6. Click OK to save the settings.

### 3.1 Preliminary Directive

Advise the patient to:
- Wear a loose fitting blouse or shirt.
- Avoid wearing long sleeved sweaters or dresses during monitoring period.
- Avoid swimming, showering, or bathing during monitoring.
- Avoid operating heavy equipment or power tools, as vibrations may functionally disrupt the monitor.

### 3.2 Prior to Hook-up

Before hooking the ABPM 6100 unit up to the patient, make sure you have performed the following tasks:
- Be sure the patient name and ID are on the diary and logged separately to avoid confusion with other patient data.
- Make sure that the ABPM 6100 monitor contains new or charged batteries.
- For new studies, make sure that all old data has been downloaded to the CardioPerfect Workstation Software and the ABPM 6100 monitor’s memory has been cleared out.
- Make sure that the proper Study parameters have been written from the CardioPerfect Workstation Software to the unit.
- Seat the patient comfortably. If the patient has long sleeves, have the patient remove his or her shirt. Ask the patient which is the non-dominant arm. You will place the cuff on this arm.

### 3.3 Sizing

Since correct cuff sizing is vital to the accuracy of monitoring data, make sure that you read this section carefully and understand all of the information contained herein before sizing the cuff to the patient.

#### USING THE CUFF RANGE MARKS

Each cuff has two range marks. To size using the range marks, wrap the cuff around the patient’s non-dominant arm. If the edge of the cuff falls within the range marks, the cuff is the correct size for the patient. If the cuff edge does not fall within the range marks, try a different size cuff.

#### USING THE SIZING TABLE

For proper cuff selection, find the circumference of the patient’s upper arm and refer to this cuff sizing chart.
<table>
<thead>
<tr>
<th>Cuff Size</th>
<th>Arm Circumference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pediatric Cuff</td>
<td>NA</td>
</tr>
<tr>
<td>Small Adult Cuff</td>
<td>Part 101340</td>
</tr>
<tr>
<td></td>
<td>18-27 cm</td>
</tr>
<tr>
<td>Adult Cuff</td>
<td>Part 101341</td>
</tr>
<tr>
<td></td>
<td>25-35 cm</td>
</tr>
<tr>
<td>Adult Plus Cuff</td>
<td>Part 101342</td>
</tr>
<tr>
<td></td>
<td>33-40 cm</td>
</tr>
<tr>
<td>Large Adult Cuff</td>
<td>Part 101343</td>
</tr>
<tr>
<td></td>
<td>39-46 cm</td>
</tr>
</tbody>
</table>

### 3.4 Traditional Cuff Placement

Proper cuff placement is very important to achieve accurate blood pressure measurements. Follow these instructions to ensure the cuff is accurately placed on the patient.

**Traditional cuff**

1. Wrap the cuff snugly around the patient’s non-dominant arm as shown in the picture, making sure that the air hose leading from the cuff is not crimped or damaged. The cuff may be worn over a thin shirt without compromising the readings.
2. Attach an adhesive cuff anchor to the clear tab on the cuff, and fasten the cuff anchor to the patient. Note: Make sure that the parameters have been written to the ABPM 6100 monitor before proceeding. The monitor should be in sleep mode.
3. Attach the tubing from the ABPM 6100 monitor to the cuff tubing and insert the ABPM 6100 monitor into the monitor pouch. Attach the pouch to the shoulder strap or the belt, depending upon which is more comfortable to the patient. Be sure to allow enough slack for the patient to move comfortably.
3.5 Sleeve Cuff Placement

Using an incorrect cuff size could result in erroneous and misleading blood pressure measurements.

Sleeve cuff

Step 1: To determine the correct cuff size for your patient, follow these simple steps:

1. To find the right sized cuff, wrap the cuff around the patient’s upper arm without sliding the arm through the sleeve.
2. Use the color-coded RANGE indicator on the inside of the cuff and the bold INDEX marker to check that the arm circumference falls within the cuff range.
3. If the arm is within range, this cuff size is correct for your patient. If the measurement is outside the RANGE indicator, select a new cuff size as indicated by color.

Step 2: Applying the Cuff

1. To apply the Welch Allyn ABPM 6100 cuff, simply slide the sleeve up the patient’s arm, ensuring the color size indicator is at the top of the cuff. The cuff should be midway between the elbow and shoulder.
2. Be sure the ARTERY indicator is over the patient’s brachial artery, between the biceps and triceps muscles (see illustration showing left arm placement).
3. Wrap the cuff snugly around the patient’s upper arm.
4. Take the initial BP reading and ensure hookup is working.
5. Refer to figures 1, 2 and 3 on inside flap for an illustrated overview.

Step 3: Prepare the Patient

1. Preparing the patient is the most important step in obtaining an accurate, reliable blood pressure measurement.
2. Review the following instructions with your patient:
   - Avoid excess movement during readings
   - Relax the instrumented hand, slightly away from the body
   - Avoid hand movement
Avoid flexing muscles during the reading
Do not remove the cuff between readings

3.6 Office Readings

Once the ABPM 6100 monitor has been properly attached to the patient and the PC interface cable has been removed from the unit, it is important and necessary to perform office readings in order to make sure that the unit is functioning properly and does not cause discomfort to the patient. The ABPM 6100 monitor will not begin regular runs until at least one manually initiated reading has been taken. Follow these instructions to perform an Office Reading:

1. If the unit is in Sleep Mode (no display on the LCD), press the Start/Stop button to “wake” the unit.
2. Press the Start/Stop button again to initiate a manual reading. This will cause the cuff to inflate and a reading to be taken.
3. Record the readings that are taken in the office as office readings into the patient diary, so that information can be taken into account in the analysis of the study. This is also a good opportunity to instruct the patient about the use of the patient diary.

If the Start button is disabled when the parameters are written to the unit (thus disallowing initiation of manual readings), the ABPM 6100 monitor still allows you to manually initiate measurement for up to 30 minutes after parameters have been written to the unit and the unit has been “woken” from Sleep Mode. This allows you to initiate manual office readings even if you do not wish the patient to be able to initiate readings.

3.7 Regular Runs

Once at least one manual office reading has been taken, regular readings will initiate based upon the interval configured for the first time period.

BEFORE THE PATIENT LEAVES

Make sure of the following before the patient leaves the office:
• The belt or shoulder strap and pouch are positioned comfortably.
• The Start/Stop button is accessible to the patient. (Even if the Start button is disabled in the Study Parameters, the Start/Stop button will still allow the patient to cancel a reading.)
• The ABPM 6100 monitor is concealed according to the patient’s wishes.
• Remember the patient’s comfort and ability to perform normal tasks can significantly impact the relevance of monitoring data.
• If the display is on, briefly review with the patient how to read the data.
• Explain to the patient the kind of information required in the Patient Diary.
• If the Start button is enabled, explain to the patient that he or she has the option to initiate readings using the Start/Stop button.

OTHER INSTRUCTIONS FOR PATIENTS

The patient should also be advised of the following:
• Pressing the Start/Stop button during a reading will cancel that reading and deflate the cuff regardless of whether the Start button is set to On or Off in the Study Parameters.
• Advise the patient to undress carefully at bedtime, using caution not to disconnect the hose from the monitor.
• Placing a pillow over the monitor during sleep will reduce any electrostatic hum from the ABPM 6100 unit.
• All data is stored internally after deactivation.
PATIENT DO’S AND DON’TS:

Finally, familiarize the patient with the following list of do’s and don’ts:

Do:
- Wear a loose fitting blouse or shirt.
- Record time, symptom/mood and activity/position in the patient diary.
- Activate a reading (if the Start/Stop button is activated) at the first sign of symptoms.
- Remain motionless during readings.
- Keep vehicle driving and travel to a minimum.
- Bring diary upon return.

Don’t:
- Remove the cuff.
- Get the monitor wet.
- Use power tools or heavy equipment during a reading.
- Remove the batteries from the unit.
- Wear long-sleeved sweaters or dresses during monitoring period.
- Swim, shower, or bathe during monitoring.
- Operate heavy equipment or power tools, as vibrations may functionally disrupt the monitor.

3.8 Data

After the ABPM 6100 monitor is disconnected from the patient, you will need to reconnect the monitor to the PC in order to read data from the unit. To reattach the monitor to the PC, simply locate the monitor (smaller) end of the PC Interface Cable and insert it into the communications port located at the bottom of the unit (the PC end of he PC Interface Cable should still be attached to the PC).

Reading Data from the Unit

Whenever data is contained in the unit, it can be read from the unit into the ABPM 6100 software. The CardioPerfect Workstation software will then allow you to view the data and configure reports. See the CardioPerfect Workstation software manual for more information.

To read data from the unit:
1. Make sure that the ABPM 6100 monitor is properly connected to the PC.
2. Open the CardioPerfect Workstation software.
3. Select the Patient Study that you wish to read data into by clicking the representative date under the appropriate patient name. If no Patient Study has been created yet, create a Patient Study by following the instructions in the Creating a New Patient Study section of the CardioPerfect Workstation software manual.
4. Follow the instructions in the CardioPerfect Workstation Software Manual chapter 2.3
5. Within a few seconds, the data will be retrieved from the unit. If the Patient ID in the Unit does not match the Patient ID in the CardioPerfect Workstation software Patient Study, a message will appear stating, “Patient ID in unit and study do not agree. Use ID in unit?” Select Yes to use the Patient ID currently stored in the unit; select No to use the Patient ID in the CardioPerfect Workstation software.

The data from the unit will now be stored in the CardioPerfect Workstation software. If you receive an error message when you try to read data from the unit, make sure the PC Interface Cable is properly attached to both the monitor and the PC and repeat the Read Data from Unit command.

See the CardioPerfect Workstation software manual for instructions on using the software and the data handling and reporting options it offers.
4. Maintenance and Cleaning

Preventative maintenance should be routinely performed to ensure the safe and efficient operation of the ABPM 6100 monitor. In addition, the monitor should be cleaned after each use.

Maintenance
The following inspections of the ABPM 6100 unit should be performed on a regular basis:
- The PC connection cable should be inspected for any cracks, exposed wires or other damage.
- The monitor itself should be visibly inspected for any signs of damage.
- Pneumatic tubing should be inspected for any cracks, fraying, or kinks.
- Do not remove any covers or break the warranty seal while inspecting the unit.

If any signs of damage are detected, do not use the ABPM 6100 monitor. It should be sent to an authorized Welch Allyn service center. See the Service and Warranty section of this manual for a list of authorized Welch Allyn service centers.

Cleaning

**CLEANING THE ABPM 6100 UNIT**

| Importance: The ABPM 6100 cannot be sterilized. DO NOT immerse the monitor in any fluid or use liquid detergents, cleaning agents, or solvents to clean it. If the unit is immersed in any liquid, do not use the unit again. The unit should be sent to an authorized Welch Allyn service center. |

The ABPM 6100 unit should be cleaned after every use. Use a soft, damp cloth to remove any dirt and dust from the unit.

**CLEANING AND DISINFECTING THE CUFF**

1. **Medical Disinfectant Spray**
   - The cuff may be sprayed with a mild disinfectant solution (e.g. Cidezyme®, ENZOL®, or 10% bleach solution. Follow the direction on the solution), rinsed with distilled water, and line dry. Ensure that no liquid enters the bladder tubing.

2. **Medical Disinfect Wipe**
   - Use a mild disinfectant wipe (e.g. Sani-Cloth®) and thoroughly wet cuff surface and line dry.

3. **Machine Wash**
   - Remove bladder to machine wash the cuff shell. Machine wash warm (50 - 130° F, 10 - 54° C) with a mild detergent and line dry.
5. Calibration Check

Welch Allyn recommends that the calibration of the ABPM 6100 monitor be verified annually by the user using the following procedure:

1. With the ABPM 6100 unit already powered up, remove one of the AA batteries from the battery compartment. Then immediately replace the battery back in its proper orientation, which will result in the unit to start the power up cycle.
2. While the LCD is displaying dashes, press and hold down the Start/Stop button. The unit will display the software version, the battery voltage, followed by a click as the valves are closed. When the process is finished, a pressure value will be displayed on the LCD and the unit is ready to have the calibration checked.
3. Disconnect the ABPM 6100 monitor cuff assembly from the unit.
4. Attach the appropriate end of ABPM 6100 Y-connector (Welch Allyn Part #6100-25) to the monitor. Attach a cuff to the appropriate end of the Y-connector, and wrap around a suitably sized can or bottle. This acts as the reservoir for the unit. Connect the third leg of the Y-connector to a high quality, known pressure standard\(^1\). Refer to the calibration figure below for a sketch of the test set up.
5. Pressurize gauge to 250 mmHg and compare against pressure standard (see Note below). If the unit does not meet the required calibration, unit needs to be returned to Welch Allyn for calibration or repair.
6. Bleed pressure down no faster than 10 mmHg per second, stopping to check the pressure at 250, 200, 150, 100, and 50 mmHg.
7. When finished, remove one of the AA batteries from the battery compartment. Then immediately replace the battery back in its proper orientation, which will cause the unit to start the power up cycle.

Note: Your ability to measure the accuracy of the ABPM 6100 depends upon the sensitivity of the pressure standard you use for the calibration procedure.

- If using a manometer (mercury column or aneroid gauge) rated at ± 3.0 mm Hg, you will be able to determine the accuracy of the unit being tested to within ± 6.0 mm Hg.
- If using a device (e.g., digital pressure standard) rated at ± 1.0 mm Hg, you will be able to determine the accuracy of the unit being tested to within ± 4.0 mm Hg.
- \(^1\)Welch Allyn recommends using the most sensitive pressure standard possible when performing calibration checks. A Setra Pressure Meter 2270-01, which is calibrated for ± 0.1 mm Hg, works well for this application.
6. Specifications

This section provides the unit specifications for the ABPM 6100 ambulatory blood pressure monitor.

**Power Requirements:**
- Two "AA" alkaline batteries or high capacity rechargeable batteries (NiMh).

**Dimensions:**
- 124 x 70 x 33 mm (approximate).

**Weight:**
- 270g (including batteries, approximate).

**Environmental Operating Condition:**
- Atmospheric Pressure Range of 700 HPA to 1060 HPA
- Protection against electric shock: Internally powered, Type BF

**Operating Ambients:**
- Temperature: 10°C (50°F) to 50°C (122°F).
- Humidity: 20% to 95% RH non-condensing.

**Storage Ambients:**
- Temperature: -20°C (-68°F) to 70°C (158°F).
- Humidity: 15% to 95% RH non-condensing.
- Altitude: -170 to 1,700 meters

**Gauging Ranges:**
- Systolic Blood Pressure: 60 to 250 mmHg
- Diastolic Blood Pressure: 25 to 200 mmHg
- Maximum Inflation: 270 mmHg
- Heart Rate: 40 to 200 bpm

**Memory:**
- Up to 250 readings using Alkaline batteries
- Up to 110 reading using rechargeable NiMh

**Method of Measurement:**
- Oscillometric with step deflation.

**Accuracy:**
- ± 3 mmHg

**International Standards:**
- EN 60601-1:2001 ; Medical electrical equipment – Part 1-1: General requirements for safety. Collateral standard: Safety requirements for medical electrical systems
- Specific requirements for the protection against hazards of ignition of flammable anesthetic mixtures and to control the risk of fire (Fire prevention)
- EN 60601-2-30:2000 ; Medical electrical equipment – Part 2-30: Particular requirements for the safety, including essential performance, of automatic cycling non-invasive blood pressuring monitoring equipment
- EN 1060-3:1997 ; Non-invasive sphygmomanometers – Part 3: Supplementary requirements for electro-mechanical blood pressure measuring systems
- AAMI SP 10 ES1 Category C: 1992 (battery powered) Electronic or automated sphygmomanometers

**Calibration:**
- Minimally, once per year

**Safety System:**
- Maximum Inflation pressure limited to 300 mmHg.
- Auto safety release valve for power failure.
- Maximum BP measurement time limited to less than 180 seconds.

**Data Connector:**
• Stereo mini-plug headphone type connector.

**Operator Control:**
• 1-button control and LCD.

**Automatic Measurement Intervals:**
• Programmable up to four separate time periods at 5 to 120 minute intervals.

## 7. Classification

**Protection against electric shock:**
• Internally powered, Type BF

**Protection against the ingress of liquids:**
• IPX0, per IEC 60529

## 8. Discarding the Equipment

Discard the old battery appropriately.
• In the USA, call 1800-SAV-LEAD for instructions on recycling it.
• International users, contact your local authorities concerning recycling.
• Discard the ABPM 6100 and accessories according to local laws.

Do not dispose of this product as unsorted municipal waste. Prepare this product for reuse or separate collection as specified by Directive 2002/96/EC of the European Parliament and the Council of the European Union on Waste Electronic and Electrical Equipment (WEEE). If this product is contaminated, this directive does not apply. For more specific disposal information, see www.welchallyn.com/weee, or contact Welch Allyn Customer Service.
9. Troubleshooting and Error Codes

9.1 Error Codes

The ABPM 6100 monitor displays error codes whenever an error situation is encountered. Error codes will display on the unit’s LCD. Error codes that apply to a specific reading will also display in the CardioPerfect Workstation Software when the data is read from the unit. The following table explains error codes generated by the ABPM 6100 monitor along with possible solutions for each error code.

<table>
<thead>
<tr>
<th>Error Code</th>
<th>Description</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Weak or no oscillometric signal</td>
<td>Check cuff position and cuff tightness</td>
</tr>
<tr>
<td>2</td>
<td>Artifact/Erratic Oscillometric Signal</td>
<td>Instruct patient to remain still during reading. Try reading again.</td>
</tr>
<tr>
<td>3</td>
<td>Exceeded retry count (4 inflate attempts)</td>
<td>Instruct patient to remain still during reading. Try reading again.</td>
</tr>
<tr>
<td>4</td>
<td>Exceeded measurement time limit</td>
<td>Check air hose connections and make certain cuff is tight enough.</td>
</tr>
<tr>
<td>85</td>
<td>Reading aborted (blocked valves or pneumatics)</td>
<td>Check air hose connections and make certain air tubing is not crimped.</td>
</tr>
<tr>
<td>86</td>
<td>Reading aborted (manual abort)</td>
<td>Push Start/Stop button to restart reading</td>
</tr>
<tr>
<td>87</td>
<td>Reading aborted (inflate time-out or air leak)</td>
<td>Check air hose and cuff</td>
</tr>
<tr>
<td>88</td>
<td>Reading aborted (safety time-out)</td>
<td>Push Start/Stop button to restart reading. If problem persists, return for servicing.*</td>
</tr>
<tr>
<td>89</td>
<td>Reading aborted (cuff overpressure)</td>
<td>Check air hose for blockage or kinking</td>
</tr>
<tr>
<td>90</td>
<td>Service Required (power supply out-of-range or other hardware problem)</td>
<td>Replace batteries. If problem persists, return for servicing.*</td>
</tr>
<tr>
<td>91</td>
<td>Service Required (safety override fitted or autozero out-of-range)</td>
<td>Push Start/Stop button to retry reading. If problem persists, return for servicing.*</td>
</tr>
<tr>
<td>97</td>
<td>Service Required Transducer out-of-range</td>
<td>Return for servicing*</td>
</tr>
<tr>
<td>98</td>
<td>Service Required (A/D out-of-range)</td>
<td>Return for servicing*</td>
</tr>
<tr>
<td>99</td>
<td>Service Required (EEPROM calibration data CRC failure)</td>
<td>Unit needs to be recalibrated. Return for servicing.*</td>
</tr>
</tbody>
</table>

The codes mentioned above, are the codes as shown on the device display. Please refer to the CPWS ABP Software manual for the codes as used by the software.
9.2 Troubleshooting

The following table contains a list of troublesome scenarios accompanied by suggestions for problem solving.

<table>
<thead>
<tr>
<th>Trouble</th>
<th>Response to Trouble</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cycle starts but cuff will not fully inflate</td>
<td>Ensure a connection with the ABPM 6100 monitor is secure and check bladder for leak; replace if needed. Replace batteries and try again.</td>
</tr>
<tr>
<td>Blood Pressure readings are not displayed during Regular Runs.</td>
<td>Verify that the display setting is ON in the Basic Parameters menu.</td>
</tr>
<tr>
<td>Patient activation button does not initiate readings while in the Regular Run mode.</td>
<td>Verify that Start Button is ON in the Basic Parameters menu.</td>
</tr>
<tr>
<td>Time Regular Runs do not initiate</td>
<td>Make sure that the time period is not set for MAN (manual) operation.</td>
</tr>
<tr>
<td>Blood Pressure readings are failing with error codes being displayed</td>
<td>Refer to the Error Codes section CardioPerfect Workstation Software manual.</td>
</tr>
</tbody>
</table>

* Always return ABPM 6100 monitor to an authorized service center for repairs. Unauthorized service will void all warranties.
10. Guidance and Manufacturer’s Declarations

**CAUTION**

The ABPM6100 needs special precautions regarding EMC and needs to be installed and put into service according to the following EMC information provided.

Portable and mobile RF communications equipment can affect the ABPM6100.

---

### Electromagnetic Emissions

The ABPM6100 is intended for use in the electromagnetic environment specified below. The customer or the user of the ABPM6100 should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Emissions test</th>
<th>Compliance</th>
<th>Electromagnetic environment – guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF emissions CISPR 11</td>
<td>Group 1</td>
<td>The ABPM6100 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.</td>
</tr>
<tr>
<td>RF emissions CISPR 11</td>
<td>Class B</td>
<td></td>
</tr>
<tr>
<td>Harmonic emissions IEC 61000-3-2</td>
<td>n.a.</td>
<td>The ABPM6100 is suitable for use in all establishments, including domestic establishments and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.</td>
</tr>
<tr>
<td>Voltage fluctuations/ flicker emissions IEC 61000-3-3</td>
<td>n.a.</td>
<td></td>
</tr>
</tbody>
</table>
The ABPM6100 is intended for use in the electromagnetic environment specified below. The customer or the user of the ABPM6100 should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Immunity test</th>
<th>IEC 60601 test level</th>
<th>Compliance level</th>
<th>Electromagnetic environment guidance</th>
</tr>
</thead>
</table>
| 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 floors 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  
±1 kV for input/output lines | n.a. | Mains power quality should be that of a typical commercial or hospital environment.

| Surge IEC 61000-4-5 | ±1 kV differential mode  
±2 kV common mode | n.a. | 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 % \(U_T\)  
(>95 % dip in \(U_T\)) for 0,5 cycle  
40 % \(U_T\)  
(60 % dip in \(U_T\)) for 5 cycles  
70 % \(U_T\)  
(30 % dip in \(U_T\)) for 25 cycles  
<5 % \(U_T\)  
(>95 % dip in \(U_T\)) for 5 sec | n.a. | Mains power quality should be that of a typical commercial or hospital environment. If the user of the ABPM6100 requires continued operation during power mains interruptions, it is recommended that the ABPM6100 be powered from an uninterruptible power supply or a battery.

| Power frequency (50/60 Hz) magnetic field 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.
### Electromagnetic Immunity

The ABPM6100 is intended for use in the electromagnetic environment specified below. The customer or the user of the ABPM6100 should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Immunity test</th>
<th>IEC 60601 test level</th>
<th>Compliance level</th>
<th>Electromagnetic environment — guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conducted RF</td>
<td>3 Vrms 150 kHz to 80 MHz</td>
<td>n.a.</td>
<td>Portable and mobile RF communications equipment should be used no closer to any part of the ABPM6100, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter.</td>
</tr>
</tbody>
</table>
| Radiated RF   | 3 V/m 80 MHz to 2.5 GHz | 3 V/m           | **Recommended separation distance**<br>
|               |                      |                 | \[d = 1.2 \cdot \sqrt{P}\] |
|               |                      |                 | \[d = 1.2 \cdot \sqrt{P}\] 80 to 800 MHz |
|               |                      |                 | \[d = 2.3 \cdot \sqrt{P}\] 800 MHz to 2.5 GHz |

where \(P\) is the maximum output power rating of the transmitter in watts (W) and \(d\) is the recommended separation distance in metres (m).

Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey, should be less than the compliance level in each frequency range.¹

Interference may occur in the vicinity of equipment marked with the following symbol: \(\Box\)

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.

¹ 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 ABPM6100 is used exceeds the applicable RF compliance level above, the ABPM6100 should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as reorienting or relocating the ABPM6100.

² Over the frequency range 150 kHz to 80 MHz, field strengths should be less than 3 V/m.
The ABPM6100 is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the ABPM6100 can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the ABPM6100 as recommended below, according to the maximum output power of the communications equipment.

<table>
<thead>
<tr>
<th>Rated maximum output power of transmitter W</th>
<th>Separation distance according to frequency of transmitter m</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>150 KHz to 80 MHz</td>
</tr>
<tr>
<td>0.01</td>
<td>0.12</td>
</tr>
<tr>
<td>0.1</td>
<td>0.37</td>
</tr>
<tr>
<td>1</td>
<td>1.2</td>
</tr>
<tr>
<td>10</td>
<td>3.7</td>
</tr>
<tr>
<td>100</td>
<td>12</td>
</tr>
</tbody>
</table>

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

**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.