SunTech® Oscar 2™
24-HR ABPM

24-Hour Ambulatory BP Monitoring System
User Manual
Changes
This manual is identified as Part number: 80-0062-00. An updated version may be available for download from the SunTech Medical website. Should you notice errors or omissions in this manual, please notify us at:
SunTech Medical, Inc. 507 Airport Boulevard, Suite 117
Morrisville, NC 27560 USA
Tel: 800.421.8626 Fax: 919.654.2301
Email: CustomerService@SunTechMed.com Web: www.SunTechMed.com

Additional contact information is listed on page 50.

This manual is for the Oscar 2, Model 250 Ambulatory Blood Pressure Monitor System.

Caution: Federal law restricts this device to sale by or on the order of a licensed practitioner

Copyright Information
All content in this manual is the proprietary information of SunTech Medical and is provided solely for purposes of operation, maintenance or service of the Oscar 2 ABPM System. This manual and the Oscar 2 ABPM System described in it are protected under copyright law under which they may not be copied, in whole or in part, without written consent of SunTech Medical. SunTech and Oscar are registered trademarks of SunTech Medical, Inc. SphygmoCor is a registered trademark of AtCor Medical Pty. Ltd., Sydney Australia. All other trademark names are the trademarks of their respective holders. The information in this manual is furnished for guidance only, is subject to change without notice, and should not be construed as a commitment by SunTech Medical. SunTech Medical assumes no liability for errors or inaccuracies that may appear in this manual. © 2017 SunTech Medical. All rights reserved.
1. Introduction to Ambulatory Blood Pressure Monitoring

Ambulatory blood pressure monitoring (ABPM) is an accepted clinical tool for collecting multiple blood pressure measurements to better assist clinicians with the diagnosis and management of hypertension by providing data related to: blood pressure variability, estimation of true blood pressure, overnight changes in blood pressure, Blood Pressure Load, Sleep Dip, and morning surge in blood pressure. In-clinic and home blood pressure measurements cannot provide the same depth of information that a 24-hour study provides. Several studies have shown that ambulatory blood pressure monitoring, when compared to clinic or home blood pressure measurement, is superior in predicting target organ damage, morbid events, or cardiovascular risk.\(^1\)\(^\text{2}\)\(^3\)

The data obtained from ambulatory blood pressure monitors is accurate and useful for managing a wide variety of hypertensive situations including:

- White-coat hypertension
- Resistant hypertension
- Masked hypertension
- Childhood hypertension
- Efficacy of anti-hypertensive drug therapy on a 24-hour basis
- Nocturnal hypertension
- Episodic hypertension and/or anxiety disorders
- Hypotensive symptoms
- Changes in diet and daily routine designed to reduce hypertension

2. Introduction to Central Blood Pressure Monitoring

Central blood pressure measurement derives the central aortic pressure waveform from cuff pulsations recorded non-invasively at the brachial artery. Analysis of the waveform provides key parameters including central systolic pressure, central pulse pressure and indices of arterial stiffness such as augmentation pressure and augmentation index.

Increased central systolic pressure and augmentation index have been shown to be markers of cardiovascular risk.\(^4\)

Additionally, research has shown its significance as a biomarker for guiding assessment of drug safety and efficacy and, ultimately, patient treatment.\(^5\)
# 3. Symbols Used in Labeling

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
<th>Standard/Source</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>General Warning Sign</td>
<td>ISO 7010-W001</td>
</tr>
<tr>
<td></td>
<td>Batch Code</td>
<td>ISO 7000-2492</td>
</tr>
<tr>
<td></td>
<td>Caution</td>
<td>ISO 7000-0434A</td>
</tr>
<tr>
<td></td>
<td>Refer to Instruction Manual</td>
<td>ISO 7010-M002</td>
</tr>
<tr>
<td></td>
<td>Authorized representative in the European Community</td>
<td>SunTech Design</td>
</tr>
<tr>
<td></td>
<td>USB-A or USB-B</td>
<td>Industry</td>
</tr>
<tr>
<td></td>
<td>Consult Instructions for Use</td>
<td>ISO 7000-1641</td>
</tr>
<tr>
<td></td>
<td>This product meets the requirements of the applicable Directives</td>
<td>EU Directive</td>
</tr>
<tr>
<td></td>
<td>Disposal in compliance with WEEE Directive</td>
<td>WEEE Directive</td>
</tr>
<tr>
<td></td>
<td>Manufacturer</td>
<td>ISO 7000-3082</td>
</tr>
<tr>
<td></td>
<td>Date of Manufacture</td>
<td>ISO 7000-2497</td>
</tr>
<tr>
<td></td>
<td>Serial Number</td>
<td>ISO 7000-2498</td>
</tr>
<tr>
<td></td>
<td>Caution: Federal (U.S.A.) law restricts this device to sale by or on the</td>
<td>FDA</td>
</tr>
<tr>
<td></td>
<td>order of a physician</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cuff index line must fall within range markings</td>
<td>SunTech Design</td>
</tr>
<tr>
<td></td>
<td>Arrow should be placed over artery</td>
<td>SunTech Design</td>
</tr>
<tr>
<td></td>
<td>Symbol indicating limb circumference</td>
<td>SunTech Design</td>
</tr>
<tr>
<td></td>
<td>Index line</td>
<td>SunTech Design</td>
</tr>
<tr>
<td></td>
<td>Not made with natural rubber latex</td>
<td>SunTech Design</td>
</tr>
</tbody>
</table>
4. The Oscar 2 ABPM System

**Indications for Use**

The Oscar 2 System is a non-invasive oscillometric ambulatory blood pressure monitor that is intended to be used with AccuWin Pro™, a PC-based computer program for the recording and displaying of up to 250 measurements of systolic and diastolic blood pressure and heart rate. It is intended for use as an aid or adjunct to diagnosis and treatment when it is necessary to measure an adult and pediatric (> 3yrs.) patient’s systolic and diastolic blood pressures over an extended period of time. The system is only for measurement, recording, and display. It makes no diagnoses. Optionally, the Oscar 2 will provide a derived ascending aortic blood pressure waveform and a range of central arterial indices. These measurements are provided non-invasively through the use of a brachial cuff.

It is to be used on those patients where information related to ascending aortic blood pressure is desired but the risks of cardiac catheterization procedure or other invasive monitoring may outweigh the benefits (excludes pediatric subjects).

Bluetooth wireless connectivity is available as an option.

**Device Operation**

The Oscar 2 monitor is worn by the patient on a waist belt and is connected to a cuff around the non-dominant upper arm. The cuff is inflated automatically at intervals which can be programmed during setup. Blood pressure is measured by the oscillometric method which senses pressure waves in the artery when occluded by pressure in the cuff. Heart rate is determined by the frequency of the pressure waves detected.

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 American National Standard, Electronic or Automated Sphygmomanometers®. The Korotkoff sounds heard over the artery below the compression cuff vary in character as the pressure in the cuff is reduced from above systolic toward zero, or atmospheric pressure. They are divided into phases. Phase 1 (K1) or systolic begins with the sudden appearance of a faint, clear tapping or thumping sound that gradually increases in intensity. Phase 5 (KS) or diastolic begins when silence develops, and was used to determine overall efficacy of the Oscar 2.
The Oscar 2 meets or exceeds all requirements for validation by the International Protocol of the European Society of Hypertension (ESH)\(^7\) and the British Hypertension Society (BHS)\(^8\). To obtain results of these studies please send a written request to:

SunTech Medical®
507 Airport Boulevard, Suite 117
Morrisville, NC 27560-8200 USA
or visit our website to review the abstracts: [www.suntechmed.com/bp-devices-and-cuffs/ambulatory-blood-pressure-monitoring/oscar-2#Research_Study_Library](http://www.suntechmed.com/bp-devices-and-cuffs/ambulatory-blood-pressure-monitoring/oscar-2#Research_Study_Library).

### Products and Accessories

The Oscar 2 System should contain the following items. If you are missing any item, please contact SunTech Medical immediately (see Limited Warranty for contact information).

For additional information on how these accessories fit together consult the Oscar 2 Quick Start Guide (82-0403-00)

<table>
<thead>
<tr>
<th>Oscar 2 ABPM System (Included Accessories)</th>
<th>Part Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oscar 2 ABP Monitor*</td>
<td>See System Table below</td>
</tr>
<tr>
<td>AccuWin Pro™ 4 Software</td>
<td>27-0144-A1</td>
</tr>
<tr>
<td>Micro USB Cable</td>
<td>91-0130-00</td>
</tr>
<tr>
<td>Oscar 2 Pouch</td>
<td>98-0032-07</td>
</tr>
<tr>
<td>Oscar 2 Belt</td>
<td>98-0037-00</td>
</tr>
<tr>
<td>Oscar 2 User Guide</td>
<td>80-0062-00</td>
</tr>
<tr>
<td>Oscar 2 Quick Start Guide</td>
<td>82-0403-00</td>
</tr>
<tr>
<td>ABPM Cuff, Size 2 (25-35 cm)</td>
<td>98-0239-02</td>
</tr>
<tr>
<td>ABPM Cuff, Size 3 (32-44 cm)</td>
<td>98-0239-03</td>
</tr>
</tbody>
</table>

* See table "Oscar 2 ABPM System (Monitor Options)" to determine the part number and description for your Oscar 2 ABP Monitor.

<table>
<thead>
<tr>
<th>Oscar 2 ABPM Optional Accessories</th>
<th>Part Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>ABPM Cuff, Size 1 (18-26 cm)</td>
<td>98-0239-01</td>
</tr>
<tr>
<td>ABPM Cuff, Size 4 (42-55 cm)</td>
<td>98-0239-04</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Oscar 2 ABPM System (*Monitor Options)</th>
<th>Part Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oscar 2 ABP Monitor - Standard</td>
<td>99-0133-00</td>
</tr>
<tr>
<td>Oscar 2 ABP Monitor - Bluetooth</td>
<td>99-0133-01</td>
</tr>
<tr>
<td>Oscar 2 ABP Monitor w/ SphygmoCor Inside</td>
<td>99-0133-02</td>
</tr>
<tr>
<td>Oscar 2 ABP Monitor - Bluetooth w/ SphygmoCor Inside</td>
<td>99-0133-03</td>
</tr>
<tr>
<td>Oscar 2 ABP Monitor – Standard-Without AWP4 Software</td>
<td>99-0133-10</td>
</tr>
<tr>
<td>Oscar 2 ABP Monitor - Bluetooth - Without AWP4 Software</td>
<td>99-0133-11</td>
</tr>
<tr>
<td>Oscar 2 ABP Monitor - w/ SphygmoCor Inside - Without AWP4 Software</td>
<td>99-0133-12</td>
</tr>
<tr>
<td>Oscar 2 ABP Monitor - w/ SphygmoCor Inside - BT - Without AWP4 Software</td>
<td>99-0133-13</td>
</tr>
</tbody>
</table>

### About AccuWin Pro™ 4

AccuWin Pro™ 4 is a simple software application designed for exclusive use with the Oscar 2 monitor and which allows for maximum flexibility in the configuration, analysis, interpretation, and reporting of ABPM studies.

Personal computer is needed as an accessory but it is not provided.
**PC System Requirements**
- Windows 7 or later
- SVGA or compatible display adapter and monitor. Minimum 1280 x 1024 recommended resolution
- One available USB port
- Minimum 4GB of RAM
- Minimum of 30GB of HDD space for patient database

**A Note About HIPAA**
The regulations set forth by the Health Insurance Portability and Accountability Act of 1996 (HIPAA) cover a number of topics, two of which present possible compliancy issues for health care providers using a software application, such as AccuWin Pro™ 4: privacy and security.

Password-controlled user access and encrypted patient information capabilities of AccuWin Pro™ 4 assist health care providers in maintaining a HIPAA compliant environment. Since HIPAA compliancy is ultimately the responsibility of the provider, please be aware that report printouts, report PDF files, and exported data contain unencrypted patient information and should be handled appropriately.

**Biocompatibility and Applied Parts**
The Orbit ABPM cuff is the only Applied Part (AP) of the Oscar 2 system. All AP have been evaluated for biocompatibility in conjunction with the applicable standards.

**Specifications**

<table>
<thead>
<tr>
<th>Specifications</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Method of Measurement</td>
<td>Oscillometric with step deflation</td>
</tr>
<tr>
<td>Blood Pressure Range</td>
<td>Systolic: 40-260 mmHg</td>
</tr>
<tr>
<td></td>
<td>Diastolic: 25-200 mmHg</td>
</tr>
<tr>
<td>Heart Rate Range</td>
<td>40-200 bpm</td>
</tr>
<tr>
<td>Maximum Inflate Pressure</td>
<td>280 mmHg</td>
</tr>
<tr>
<td>Accuracy</td>
<td>Heart Rate accurate within +/-2% or +/-3 bpm, whichever is greater.</td>
</tr>
<tr>
<td></td>
<td>Blood Pressure results meet or exceed ANSI/AAMI/ISO 81060-2:2013 standards for non-invasive accuracy: ±5 mmHg mean error &amp; 8 mmHg standard deviation.</td>
</tr>
<tr>
<td>Validations</td>
<td>Clinically validated to ESH International Protocol, BHS (A/A), and ANSI/AAMI/ISO 81060-2:2013</td>
</tr>
<tr>
<td>Operating Conditions</td>
<td>10°C (50°F) to 50°C (122°F)</td>
</tr>
<tr>
<td></td>
<td>20-95% RH non-condensing</td>
</tr>
<tr>
<td>Shipping/Storage</td>
<td>-20°C (°F) to 70°C (°F)</td>
</tr>
<tr>
<td>Conditions</td>
<td>15-95% RH non-condensing</td>
</tr>
<tr>
<td>Ingress Protection</td>
<td>Ordinary Equipment, No ingress protection, IPX0.</td>
</tr>
<tr>
<td>Classification</td>
<td>Continuous operation</td>
</tr>
<tr>
<td></td>
<td>Power: Two (2) AA batteries,</td>
</tr>
<tr>
<td>Data Memory</td>
<td>Flash memory stores up to 250 readings</td>
</tr>
</tbody>
</table>
Calibration Recommendation: Minimally, once every two years

Safety Systems
Maximum inflation pressure limited to 300 mmHg; Auto safety release valve for power failure; Maximum measurement time limited to less than 140 seconds

Sampling Periods
24 independently programmable time periods (Time interval options: none, 5, 10, 15, 20, 30, 45, 60, 90, and 120 minutes)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Standard</th>
<th>EU Norm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood Pressure</td>
<td>IEC 80601-2-30 :2009 + A1:2013</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td>IEC 62366:2015 Ed. 1.0</td>
<td>EN 62366:2008</td>
</tr>
<tr>
<td>Home Use</td>
<td>IEC 60601-1-11:2010 Ed. 1.0</td>
<td>EN IEC 60601-1-11:2010</td>
</tr>
<tr>
<td></td>
<td>ISO 10993-10:2010</td>
<td>N/A</td>
</tr>
<tr>
<td>Size</td>
<td>Approximately 120 x 70 x 35 mm</td>
<td></td>
</tr>
<tr>
<td>Weight</td>
<td>Approximately 275 g, including batteries</td>
<td></td>
</tr>
<tr>
<td>Storage Conditions</td>
<td>-20° C to +70° C, 15%-95% RH non-condensing</td>
<td></td>
</tr>
<tr>
<td>Data</td>
<td>USB 2.0 (Micro USB) with optional Bluetooth™ 4.0</td>
<td></td>
</tr>
</tbody>
</table>

**Safety and Effectiveness Considerations**

The following safety and effectiveness issues are to be considered prior to the usage of the Oscar 2 monitor.

- This device is defibrillator protected.

  **NOTE:** No precautions specific to the Oscar 2 are required during defibrillation, and defibrillation discharge has no effect on the Oscar 2.

- The monitor is intended for use following consultation and instruction by a physician.
- The reliability of the device is dependent upon conformance with the operation and service instructions, as detailed in this manual.
- This device has been designed for use on patients with normal sinus rhythms.
- The interpretation of blood pressure measurements should only be made by a physician. The accuracy of any blood pressure recording may be affected by the position of the subject, his or her physical condition, and use outside the operating instructions detailed in this manual.
- Safety and effectiveness of central blood pressure measurements on children under the age of 18 years of age, pregnant women and neonates have not been tested.

**Disposal**

This symbol indicates that the monitor contains materials which may be hazardous to human health. This product complies with the WEEE Directive. Please return the Oscar 2 monitor to SunTech Medical for proper disposal. Please dispose of other materials according to local regulations.

**Potential Adverse Reactions**

Allergic exanthema (symptomatic eruption) in the area of the cuff may result, including the formation of urticaria (allergic reaction including raised edematous patches of skin or mucous membranes and intense itching) caused by the fabric material of the cuff.

Petechia (a minute reddish or purplish spot containing blood that appears in the skin) formation or Rumple-Leede phenomenon (multiple petechia) on the forearm following the application of the cuff, which may lead to Idiopathic thrombocytopenia (spontaneous persistent decrease in the number of platelets associated with hemorrhagic conditions) or phlebitis (inflammation of a vein) may be observed.

**Cautions for Use**

This monitor is designed to perform in conformity with the description thereof contained in this operation manual when operated, maintained and repaired in accordance with the instructions provided. The monitor should not be modified in any way. Ensure pressure compatibility to all patients. If any abnormality occurs in the monitor, suspend the operation immediately and disconnect it from the patient. If the monitor has been used or stored outside its acceptable range (see Specifications page), it may not meet performance specifications. If the cuff fails to deflate, the patient should be instructed on its proper and safe removal.

**Warnings**

The general warning sign indicates a potentially hazardous situation which could result in serious injury.

**WARNING: Do not use in the presence of flammable anesthetics; this could cause an explosion. This device is not suitable for use in an oxygen enriched environment.**

**WARNING: Do not immerse the monitor in any fluid, place fluids on top, or attempt to clean the monitor with any liquid detergents, cleaning agents, or solvents. This may cause an electrical hazard. Do not use the monitor if accidental wetting occurs; please return to SunTech Medical® (see Limited Warranty). Refer to Maintaining and Cleaning the Oscar 2 ABP System, for care instructions.**

**WARNING: Too frequent measurements can cause injury to the patient due to blood flow interference.**

**WARNING: The cuff should not be applied over a wound as this can cause further injury.**

**WARNING: The cuff should not be placed on the arm on the side of a mastectomy. In the case of a double mastectomy use the side of the least dominant arm.**

**WARNING: Pressurization of the cuff can temporarily cause loss of function of simultaneously used monitoring equipment on the same limb.**
WARNING: Do not use if device is dropped and/or is damaged. Have a qualified service representative check the monitor before using again.

WARNING: Do not attach the cuff to a limb being used for IV infusions or any other intravascular access, therapy or an arterio-venous (A-V) shunt. The cuff inflation can temporarily block blood flow, potentially causing harm to the patient.

WARNING: Use only with the cuffs supplied by SunTech Medical. Different cuffs have not been validated with Oscar 2 and measurements with non-validated components may not be accurate.

WARNING: Use of an ACCESSORY, transducer or cable with ME EQUIPMENT and ME SYSTEMS other than those specified may result in increased EMISSIONS or decreased IMMUNITY of ME EQUIPMENT or ME SYSTEM.

WARNING: The Oscar 2 may be interfered with by other equipment even if the other equipment complies with CISPR Emission Requirements.

WARNING: Performance can be affected by extremes of temperature, humidity and altitude.

WARNING: Do not use the monitor during magnetic resonance imaging (MRI) or in an MRI environment.

Cautions
The caution symbol indicates a potentially hazardous situation which may result in minor or moderate injury. It may also be used to alert against unsafe practices.

CAUTION: When downloading data from the monitor’s communications USB port, the device should not be in use with a patient.

CAUTION: Do not remove monitor covers, except to replace batteries. The monitor does not contain any user serviceable components. Return monitor if service is required.

CAUTION: Do not use on neonates, pediatric patients less than 3 years old, or patients known to be readily susceptible to bruising.

CAUTION: Do not use the monitor if it has failed its diagnostic self test, or if it displays a greater than zero pressure with no cuff attached. The values displayed by such a monitor may be inaccurate.

CAUTION: Substitution of a component different from that supplied may result in measurement error. Repairs should be undertaken only by personnel trained or authorized by SunTech Medical.

CAUTION: The Oscar 2 does not contain any user serviceable internal parts and should only be repaired by an authorized SunTech Medical service representative. Do not service the product while in use.

CAUTION: If cuff fails to deflate within two and a half minutes, instruct patient on manual removal of cuff.

CAUTION: Check that operation of the monitor does not result in prolonged impairment of the circulation of the patient.

CAUTION: Remove batteries when device is not in use for long periods of time to prevent possible battery leakage and product damage.
**CAUTION:** A compressed or kinked connection hose may cause continuous cuff pressure resulting in blood flow interference and potentially harmful injury to the patient.

**CAUTION:** Using an incorrect cuff size could result in erroneous and misleading blood pressure measurement results.

**CAUTION:** Do not machine wash the cuff bladder.

**CAUTION:** On hypotensive patients, the device should be used with caution.

---

**Contraindications**

The Oscar 2 ABPM system should be used in conjunction with all other available medical histories and diagnostic test information about the patient. The following are reasons to withhold use of the Oscar 2 ABPM system from a patient:

**CONTRAINDICATION:** Do not use on patients with erratic, accelerated or mechanically controlled irregular heart rhythms, including patients with arrhythmias.

**CONTRAINDICATION:** Do not use on patients with carotid or aortic valve stenosis.

**CONTRAINDICATION:** The system is not applicable in generalized constriction or localized spasm of muscular conduit arteries such as seen immediately after hypothermic cardiopulmonary bypass surgery or accompanying Raynaud’s phenomena or intense cold.
Oscar 2 At a Glance

- **Sun**: Indicates AWAKE portion of study.
- **Time**: Shown when the monitor is not taking a reading.
- **Battery**: Indicates low battery; REPLACE BATTERIES.
- **Clock**: ABP study in progress.
- **Moon**: Indicates ASLEEP portion of the study.
- **BP Reading**: During a reading, displays the pressure of the cuff in mmHg. Immediately after a reading, shows the BP results in mmHg followed by heart rate in beats per minute.
- **Printer**: Indicates number of BP readings in memory.

**Start/Stop Button**

- **To Power On**: Press the Start/Stop button.
- **To Power Off**: When not taking a reading, press and HOLD the Start/Stop button until you hear 5 beeps (approx. 5 seconds), then release.
- **To Abort a Measurement**: Press the Start/Stop button any time during a reading.
- **To Start a Programmed ABP Study**: When time is flashing, press the Start/Stop button to take 1st reading.
- **To Start a Manual BP Reading**: When clock is displayed, press the Start/Stop button.

**Day/Night Button**

- Allows patient to switch to ASLEEP or AWAKE schedule (if allowed).

**Systolic/Diastolic**

- **Dose Response Event Marker**: Allows patient to mark event or dose.
### Buttons and Functions

<table>
<thead>
<tr>
<th>Buttons</th>
<th>Functions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Start/Stop</td>
<td>TO POWER ON: Press the Start/Stop button.</td>
</tr>
<tr>
<td></td>
<td>TO POWER OFF: When the monitor is not taking a measurement, press and hold the Start/Stop button until you hear five quick beeps, then release.</td>
</tr>
<tr>
<td></td>
<td>TO STOP A MEASUREMENT IN PROGRESS: Press the Start/Stop button.</td>
</tr>
<tr>
<td></td>
<td>TO START A PROGRAMMED STUDY: When the time is flashing, press the Start/Stop button.</td>
</tr>
<tr>
<td></td>
<td>TO START A SINGLE BP READING: When the time is displayed, press the Start/Stop button.</td>
</tr>
<tr>
<td>Day/Night</td>
<td>Toggles between day (AWAKE) mode and night (ASLEEP) mode.</td>
</tr>
<tr>
<td>Event</td>
<td>Marks an event or starts a dose response sequence.</td>
</tr>
</tbody>
</table>

### Display Symbols and Description

<table>
<thead>
<tr>
<th>Display Symbols</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time</td>
<td>Indicates current time. When flashing, the monitor will turn off in 20 seconds unless an ABPM study is in progress.</td>
</tr>
<tr>
<td>Pressure</td>
<td>Indicates the pressure of the cuff in mmHg during a measurement.</td>
</tr>
<tr>
<td>Reading Result</td>
<td>Immediately after a measurement is complete, the display shows the results, if enabled. BP in mmHg is shown first, followed by HR in beats per minute.</td>
</tr>
<tr>
<td>CBP Check</td>
<td>Indicates the CBP measurement function is operating properly. This will only appear during the first 30 minutes of the study.</td>
</tr>
<tr>
<td>Clock</td>
<td>Denotes that a programmed ABP study is in progress.</td>
</tr>
<tr>
<td>Sun</td>
<td>Denotes the monitor is collecting readings according to the AWAKE program of the study.</td>
</tr>
<tr>
<td>Moon</td>
<td>Denotes the monitor is collecting readings according to the ASLEEP program of the study.</td>
</tr>
<tr>
<td>Battery</td>
<td>Indicates low battery voltage; BATTERIES NEED TO BE REPLACED.</td>
</tr>
<tr>
<td>Printer</td>
<td>Indicates the number of readings in memory.</td>
</tr>
</tbody>
</table>

---

### 5. Setting Up The Oscar 2 System

Setting up the Oscar 2 System involves powering the Oscar 2, installing AccuWin Pro™ 4 on a personal computer (PC), and connecting the Oscar 2 to the PC.

The Oscar 2 ABPM system is packaged with everything you need to start. See Product and Accessories for complete contents.

#### Powering the Oscar 2 for Use

Install two (2) AA batteries in the bay located at the back of the monitor. The bay shows the orientation in which the batteries should be placed. When batteries are properly loaded, the monitor’s display will show the following:

1. Incrementing dashes
2. Software and safety version of the monitor
3. Battery voltage followed by three quick audible beeps
4. The number of BP readings in memory followed by one long audible beep
5. Time flashing

The monitor is now ready to be used.
NOTE: Ensure batteries are inserted with the correct polarity. Improper installation will prevent the monitor from functioning. Batteries are required for Oscar use at all times including programming, scheduling and retrieving.

NOTE: Install batteries before connecting Oscar to PC via USB.

NOTE: Device will not re-charge batteries through via USB connection.

⚠️ CAUTION: Remove batteries when device is not in use for long periods of time to prevent possible battery leakage and product damage.

**Installing AccuWin Pro™ 4**

**PC System Requirements**
- Windows 10, 8 or 7 (32 or 64 bit)
- SVGA or compatible display adapter and monitor. Minimum 1280 x 1024 recommended resolution
- One available USB A port
- Minimum 4GB of RAM
- Minimum of 30GB of disk space for patient database

NOTE: Standard install of AccuWin Pro™ 4 does not support a server license. If you would like to support multiple users in a network system, this will require a custom install. Please visit [www.suntechmed.com](http://www.suntechmed.com) or contact SunTech Medical customer support for assistance.

NOTE: Install AccuWinPro™ 4 before connecting the USB cable to the computer.

Visit [www.suntechmed.com/software](http://www.suntechmed.com/software) to register the Oscar 2 device. After registration, download the installation file for AccuWin Pro™ 4 on to your personnel computer (PC). Installation includes the required software for use with Oscar 2 Model 250. After download is complete, run the install file on your PC. If you do not have internet access, please contact SunTech Medical customer support for assistance. After installing AccuWin Pro™ 4, connect the Oscar 2 to the computer.

1. Connect the Oscar 2 USB cable to the micro-USB connector at the bottom of the ABP monitor (Figure 1; part A).
2. Connect the USB end of the cable to the USB port on your PC (Figure 1; part B).

*Figure 1: Connecting the Oscar 2 to a PC*

NOTE: The cable can be left connected when the PC is off.
**Communicating with the Oscar 2**

To successfully complete an ambulatory blood pressure (ABP) study: First connect the Oscar 2 ABP monitor to the PC then program the study in the downloaded SunTech application from your PC. When the patient returns you will again use the SunTech application on your computer to retrieve the collected data.

**Wireless Communications**

The Oscar 2 may be equipped with an optional Bluetooth™ module, that will allow communications with wireless home network devices, mobile phones, and other appropriate devices. For details on how to connect to these devices, please contact your authorized SunTech Medical service representative.

**True24 Mobile Application**

The True24™ ABPM Patient Diary mobile app is designed to work with the SunTech Medical Oscar 2™ ambulatory blood pressure monitor during a physician prescribed ABPM study.

Following each daytime awake blood pressure measurement, the True24™ mobile app prompts the patient to enter information about their activity, posture or any symptoms experienced while the BP measurement was taken. This diary information can assist the clinician with understanding the ambulatory blood pressure study data and making hypertension treatment decisions.

The True24™ Mobile Application works only with the Apple iOS platform, and must be downloaded by the physician onto the patient’s phone from the iTunes App store using the search word ‘True24’

After the mobile application has been downloaded, the physician can access training information that can be found on the application to learn how to:

- Connect the phone to Oscar 2
- Use the True24 Home Screen
- Use the True24 Diary Entries
- Prepare the patient for Sleep
- Tips for a Successful Study
- Using the Oscar 2
Pairing Process

Once the True24™ Application is downloaded from the Apple App Store, launch the application, and follow the instructions below to establish proper pairing.

To enter Bluetooth Pairing mode on the Oscar 2:
1. Simultaneously press the two buttons indicated in Figure 2 until the 5 long beep pattern is heard, and the LCD display shows “bt Pr”.
2. Upon successful pairing with a Bluetooth host, the Oscar 2 will emit one long beep, and the LCD Display will briefly show “bt con”. The Oscar 2 will exit the pairing mode and connect to the paired Bluetooth host.
3. The Oscar 2 will remain in pairing mode for approximately 3 minutes.
   • If no host is paired at the end of 3 minutes, the Oscar 2 will stop the process and exit pairing mode.
   • The pairing process may be restarted by performing step 1, above.

*(The doctor has to train the user on the proper use of the app, including this next step.)*

While using the mobile True 24 app, if the app does not prompt the user for a diary entry after a BP reading is completed, it is suggested that the user check pairing using the instructions on the True24 Mobile App.

It is recommended that True24 mobile app is not used during dosage sequence due to interference with the dosage schedule.

Logging in to AccuWin Pro™ 4

If your AccuWin Pro™ 4 administrator enables login security, you must enter a valid user name and password when you open AccuWin Pro™ 4 or change the current user. See Admin Tools for more information. If login security is not enabled, no login is necessary.

*NOTE: The AccuWin Pro™ 4 administrator can program AccuWin Pro™ 4 to automatically log a user off after a specified time of inactivity elapses. If you are logged off, you must log back in again.*

To log in enter a valid user name and password then click OK.

Changing Your Password

If you know your current password, you can change it. If you are logging in for the first time, you must change the default password assigned to you. You must be logged in under your name before you can change your password.

1. To change your password:
2. From the Configure menu, select Admin Tools.
3. Click Change Password.
4. Enter your current password.
5. Enter your new password. It can be alphanumeric and it must be at least 6 characters.
6. Re-enter your new password to confirm it.
7. Click OK.

*NOTE: If you forget your password, you must ask the AccuWin Pro™ 4 administrator to reset it.*
Changing the Current User

To change the current user:
1. From the Configure menu, select Admin Tools.
2. Click Change User.
3. Enter a valid user name.
4. Enter a valid password.
5. Click Login.

6. AccuWin Pro™ 4 Layout

AccuWin Pro™ 4 gives you the flexibility to program the ABP monitor the way you want and the simplicity to collect and retrieve important test data. The toolbar provides easy, one-click access to frequently used functions. The menu bar allows you to access all functions of the software.

Toolbar Buttons

- **Program**: Opens the Program Monitor window
- **Retrieve**: Initiates data retrieval from the monitor
- **BP Data**: Opens the Open ABP Study – Select Patient window
- **Print Preview**: Previews a report for the open ABP study
- **Print**: Prints a report for the open ABP study
- **Print Page**: Prints the report page of the displayed data
- **PDF**: Saves the report for the open ABP study in PDF format
- **Settings**: Opens the Configuration options window
- **Help**: Opens the online help window
- **Exit**: Closes AccuWin Pro™ 4
- **Languages**: Enables the user to run the program in 8 different languages
7. Conducting an Ambulatory Blood Pressure Study

Programming the Oscar 2 for an ABP Study

Preparation for an ABP study involves filling out an on-screen form to set the parameters for your study to be programmed into the monitor. You can also use a template to fill out the form. Templates help to ensure consistent programming and adherence to specific protocols. AccuWin Pro™ 4 provides default templates, or you can create your own. See also Default Templates.

To program the monitor:

1. From the Monitor menu, select Program study. Alternatively, click the Program button on the toolbar.

2. Enter the desired parameter settings in the form (Figure 3). (See Parameter settings.) If the Oscar 2 has SphygmoCor Technology, the SphygmoCor can be turned on or off depending if CBP data is desired.

3. Click one of the following:
   - **Program** to transfer the information to the Oscar 2. An indicator bar shows the progress and disappears when programming is successfully completed.
   - **Cancel** to close the dialog box.
   - **Help** to start the online Help.

To create patient details before conducting a study:

1. In the Program Monitor window, click the button for Select/Create Patient. Next, click Create New Patient.

2. A new window Patient Information (Figure 4) will appear. Enter patient information.

3. Click OK to save the patient information to be used on this programmed study. Click Cancel to close the dialog box.

   *Note: A Patient ID number and a letter in the last name is required to create a new patient.*

To add measurement periods:

1. Click the Add special period button.

2. Select a start time, an end time, and intervals from the pull-down menus. You can add up to twenty-four (24) additional time periods. A pie chart shows the time periods created for the study.

3. Clicking the X will delete the time period from the list.

To program the monitor using a template:

1. From the open Program Monitor window, click the Open Template button at the bottom of the Program Monitor window.

---

*Figure 3: Programming the Oscar 2 (Standard Tab)*

*Figure 4: Programming the Oscar 2 (Patient Information Window)*
2. From the Available Templates dialog box (Figure 5), select the template name, and then click Open. The form automatically populates with the template's settings.

3. Click one of the following:
   - Program to transfer the information to the Oscar 2. An indicator bar shows the progress and disappears when programming is successfully completed.
   - Cancel to close the dialog box.
   - Help to start the online Help.

**To create a template:**
1. In the open Program Monitor window, enter the desired parameter settings in the form. See Parameter settings.
2. Click the Save Template button at the bottom of the Program Monitor window.
3. In the Assign Template Name dialog box, type a name for the template and click Save.

**Figure 5: Available Templates Window**

**To open a template:**
1. From the open Program Monitor window, click the Open Template button at the bottom of the Program Monitor window.
2. From the Available Templates dialog box, select the template name, and then click Open.
3. The template information will populate the corresponding information in the Program Monitor window.

**To delete a template:**
1. From the open Program Monitor window, click the Open Template button at the bottom of the Program Monitor window.
2. From the Available Templates dialog box, select the template name, and then click Delete.
3. At the prompt, click Yes to confirm deletion.

**To email a template:**
An email server must be specified before a Programming Template can be emailed. See Email Settings on page 45 for setup instructions.

1. From the File menu, select Email>Programming Template.
2. The Programming Templates window opens. Select the template(s) you want to email and click OK.
3. The Email dialog box will appear. Type in the recipient email address(es) with multiple email addresses separated by a semi-colon, a subject for your message, and a message.
4. Click Send to email the files. Click Attach to change the files you want to email. Click Configure to configure the email settings.
Parameter Settings
The parameters can be adjusted as follows:

Standard Tab

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patient ID</td>
<td>Patient ID for reporting and referencing data.</td>
</tr>
<tr>
<td>Patient name</td>
<td>Enter patient name (first, middle, last).</td>
</tr>
<tr>
<td>Select/Create Patient</td>
<td>Click to open the Patient Information window. Use this window to use existing patient information or to create a new patient entry.</td>
</tr>
<tr>
<td>Measurement Schedule:</td>
<td>Specifies when and how often the monitor takes readings. For <strong>Awake time</strong> and <strong>Sleep time</strong>, select from the <strong>Hour</strong> pull-down menu to establish the start time for these periods. From the <strong>Brachial BP Interval</strong> and <strong>Central BP Interval</strong> pull-down menus, select the desired interval between readings (5, 10, 15, 20, 30, 45, 60, 90 or 120 minutes). Please note that the available option for <strong>Central BP Intervals</strong> will only be multiples of the selected <strong>Brachial BP Interval</strong>.</td>
</tr>
<tr>
<td>Start study in 5 minutes:</td>
<td>Check denotes that the study will start automatically after programming; unchecked denotes that the study will be started with the first press of the Start/Stop button when the monitor is powered ON.</td>
</tr>
<tr>
<td>Quality Control</td>
<td>Shows the Quality Control tab that provides more control on the number of minimum readings in a time period.</td>
</tr>
</tbody>
</table>

Advanced Tab

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Max Pressure</td>
<td>Establishes the maximum inflation pressure for the monitor (options between 160 and 280 mmHg). Suggested setting is 30 mmHg above the highest expected systolic BP. <strong>NOTE:</strong> The ABP monitor will not inflate to Max Pressure with each reading; instead it inflates to 30 mmHg above the previous systolic reading.</td>
</tr>
<tr>
<td>Intervals</td>
<td>Set interval type. Select <strong>Fixed</strong> to set the intervals to exact times. Select <strong>Standard</strong> for +/- 5 minutes around the selected intervals.</td>
</tr>
<tr>
<td>Display results</td>
<td>When on, allows the patient to view the results immediately after a measurement. <strong>NOTE:</strong> Display Results is always on for the first 30 minutes of study. Only brachial BP results will be displayed.</td>
</tr>
<tr>
<td>Manual readings</td>
<td>When on, allows the patient to take measurements outside of the scheduled program using the Start/Stop key. If manual readings are disabled/off, Study in 5 minutes <strong>MUST</strong> be enabled/on. If Start study in 5 minutes is enabled/on, user can elect to enable/on or disable/off manual readings <strong>NOTE:</strong> Manual Readings is always ON for the first 30 minutes of the study.</td>
</tr>
</tbody>
</table>
### Day/night button
When on, enables the Day/Night button on the monitor, allowing the patient to start the Awake and Sleep periods according to their daily schedule. A period can be started up to four hours before the programmed period begins.

### Event marking
When enabled, allows the patient to signal up to 30 events during the study. **NOTE: If Event Marking is on, Dose Response Sequences is unavailable.**

### Does Response
Allows Dose Response Sequences to be created to provide greater measurement granularity after a drug dose is administered.

### Audible alerts
Play an alert sounds at the beginning and upon completion of each reading, during the awake period only.

### Retry attempts
The monitor will reattempts a measurement that initially fails.

### Note 1, Note 2
Enter up to 20 alphanumeric characters.

### Time zone difference
Adjust the monitor’s clock to the time zone that the patient is in relative to your time zone. (The PC Time and Monitor Time fields are populated automatically.)

### Dose Response Sequences
Specify the duration, brachial BP interval, and CBP interval for up to four dose response sequences. These sequences are useful for tracking BP after a drug dosage is administered. A sequence begins when the patient presses the Event button. To clear fields, click **Reset**.

Once dosage sequence is initiated, the dosage button should only be pressed to start the next sequence. Manual readings are not included in the dose response analysis. **NOTE: If Event Marking is on, Dose Response Sequences is unavailable.**

### Quality Control Tab

<table>
<thead>
<tr>
<th>QC Review Period:</th>
<th>Select start and end times for a quality control review period for the study program.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Minimum Reading Requirements:</td>
<td>Enter the requirements for a minimum percentage of scheduled readings captured during the defined QC period. The minimum number of readings captured per hour during the defined QC period (between zero and 12). The minimum number of hours to contain readings for the entire study. <strong>NOTE: To access the Quality Control tab you must first turn on the “Quality Control” dial at the bottom of the Program Monitor window.</strong></td>
</tr>
</tbody>
</table>

### Fitting a Patient with the Oscar 2 and SunTech ABPM Cuff
After you have successfully programmed the Oscar 2 using AccuWin Pro™ 4, you may begin fitting the patient with the monitor and a blood pressure cuff. Cuffs may be used on either arm.

1. Choose the proper cuff size
To determine the correct cuff size for your patient, wrap the cuff around the patient’s upper arm without sliding the arm through the sleeve. 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. If the arm is within range, this cuff size is correct for your patient. If the marker is outside the RANGE indicator, select a new cuff size as indicated by color.

⚠️ **CAUTION:** Using an incorrect cuff size could result in erroneous and misleading blood pressure measurements.

2. Apply the SunTech ABPM cuff

To apply the SunTech ABPM 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. Be sure the ARTERY indicator is over the patient’s brachial artery, between the bicep and tricep muscles. Wrap the cuff snugly around the patient’s upper arm.

![Figure 6: Instructions for Applying the SunTech ABPM Cuff](image)

3. Connect the hoses

Connect the hoses from the cuff and monitor by twisting the fittings together until you hear a snap. Drape the hose over the patient’s shoulder, behind the neck and across to the opposite side of the body.

4. Attach to patient

Insert the Oscar 2 into its pouch with the display showing through the window. Attach the pouch to the patient using the belt (Figure 7).

![Figure 7: Fitting the ABPM Cuff to the Patient](image)

**Preparing and Educating The Patient**

When conducting blood pressure measurements, with an oscillometric NIBP device, it is important to follow suitable procedures to ensure valid, accurate results. Preparing your patient for the ABP study is the most important step to achieving a successful test. Review the following instructions with your patient.

- When the pressure in the cuff increases, the patient should avoid excess movement during measurements. Let the cuffed arm hang loosely, slightly away from the body with the middle of the cuff at heart level. Avoid flexing the muscles or moving the hand and fingers of the cuffed arm.
- The patient can stop a measurement in progress by pressing the Start/Stop button.
• If the Manual Readings setting is on, the patient can start a measurement at any time by pressing the Start/Stop button.
• The cuff should not be removed between BP measurements.
• Before sleeping, the patient should make sure that the hose is not kinked and will not become kinked.
• The batteries can be replaced during a study without the data being lost or interrupting the monitor’s program. Alternatively, the monitor can be turned off without losing its data.
• Instruct the patient on how and when to fill out the patient diary.
• If Dose Response Sequences are set up, instruct the patient on how and when to start a sequence. Bluetooth pairing should be connected prior to start of first sequence.
• If the Day/Night button is on, instruct the patient on how to set day and night modes.
• If Event Marking is on, instruct the patient on how and when to mark events.
• Ensure the patient knows how to care for the monitor. Keep the monitor dry and do not drop it.
• If the monitor or cuff causes extreme pain, or pain not normally associated with blood pressure measurement, the patient should remove the cuff and turn off the monitor.
• The patient should not talk during BP measurements. The patient should be seated, standing or lying down. If seated, the patient should have legs uncrossed, feet flat on the floor with back and arms supported.

**Starting the Study**

Before the patient leaves with the monitor and cuff, verify that the monitor operates correctly. To verify proper monitor operation, ensure that the monitor is on and start a BP reading by pressing the Start/Stop button. The cuff will inflate and complete the brachial BP measurement. If CBP measurement has been programmed, before the cuff is fully deflated, the monitor will hold pressure in the cuff for ten seconds to capture the waveform data required to derive CBP values. The cuff will then be fully deflated. The clock icon should appear on the display of the Oscar 2 indicating that the study is in progress. If problems occur, review the setup and fitting of the system.

Replace the batteries (2 AA’s) for every study with new, alkaline batteries. Failure to do so may result in incomplete 24-hour studies.

**To record an event:**

Press the Event button.

The monitor sounds a long audible beep for confirmation, and "rcd : 01" appears on the display (Subsequent recordings use 02, 03, and so on). The monitor can record up to 30 events. If the patient tries to record more than 30 events, the monitor beeps four times, and "No rcd" appears on the display.

*Note: If Event Marking is enabled, then the Dosage Response feature will be disabled.*

**To start a dose response sequence:**

Press the Event button.

The monitor sounds an audible beep for confirmation and begins the first reading in the sequence. The Event button is disabled until all readings in the sequence are complete.

*Note: If Dosage Response is enabled, then the Event Marking feature will be disabled.*

**To manually set day (Awake) or night (Asleep) mode:**

Press the Day/Night button.

The monitor beeps and displays either the sun or moon icon depending on the mode that it was switched to. If switching to Night mode, the monitor will display a moon icon.
If switching to Day mode, the monitor will display a sun icon.

**Finishing the Study**

If you wish to finish the study before the patient returns, instruct the patient to turn off the monitor by holding down the Start/Stop button for five (5) seconds. The Oscar 2 will beep five (5) times and the display will turn off.

When the patient returns, take the cuff, monitor, and belt off and download the captured data to AccuWin Pro™ 4 for review.

### 8. Notes on Blood Pressure Data

Any blood pressure reading can be affected by the measurement site, the position of the patient, motion, or the patient’s physiologic condition. Environmental or operational factors which can affect the performance of the device and/or its blood pressure reading are common arrhythmias such as atrial or ventricular premature beats or atrial fibrillation, arterial sclerosis, poor perfusion, diabetes, age, pregnancy, preeclampsia, renal diseases, patient motion, trembling, and shivering.

**Retrieving Data from the ABP Monitor**

To retrieve the data:

1. Connect the Oscar 2 to the computer.
2. In AccuWin Pro™ 4, click the **Retrieve** toolbar button, or select **Retrieve Data** from the **Monitor** menu. The dialog box on the screen shows the progress as the data is transferred. After completion, a dialog box appears to confirm patient details.
3. In the dialog box, do one of the following:
   1. Click **No**. A list of current patient files appears. Select the patient file to save the retrieved data.
   2. Click **Yes**. The **Patient information** dialog box appears. Enter patient information in the data fields.
4. Click **OK** to save the data.
5. If the alert for adjusting Awake and Asleep times appears, click **Yes** or **No** according to your preference.
6. If the study contains quality control settings, review the results in the **Quality Control Results** dialog box that appears, and then click **Close**.

**NOTE:** A green checkmark in the results indicates that the study meets that requirement. A red x indicates that the study failed to meet that requirement.

The BP file will now open automatically and the patient’s BP data is now saved to the AccuWin Pro™ 4 patient database.

⚠️ **CAUTION:** If you do not retrieve data from the monitor, this data will be lost when you program it for the next study.

### 9. Reviewing and Editing an ABP Study

**Opening A Patient File**

AccuWin Pro™ 4 will automatically open a patient file directly after the data is retrieved.

1. From the **File** menu, select **Open**. Alternatively, from the toolbar, click the **BP Data** button.
2. The **Open ABP Study - Select Patient** dialog box appears, allowing you to select the patient file to open. Select the patient whose BP data you wish to retrieve by clicking on the patient’s name.

3. Click **OK**.

4. If the patient has more than one ABP study, a second dialog box, **Choose Date**, appears. Select a study date and click **OK**. The patient’s data appears in the display area. You can now review the BP study or print a report.

Each study in a patient file contains data that is displayed in the views accessed by the tabs at the bottom of the application. The study information bar located toward the top of the view identifies the patient name, patient ID, patient age (at the time of the study) and the study date of the displayed file.

Tabs and their contents

- **ABP Data**: ABP measurement data from the monitor and relevant graphs.
- **Patient Info**: Patient name, ID, contact information, and physical description.
- **Physician Info**: Patient history, clinical information, and interpretation.
- **Statistics**: Statistical analysis of the ABP study.
- **Dose Response Statistics**: Statistical analysis for dose response sequences.
- **Time-Slice Statistics**: Statistical analysis of specific user-defined time windows.
- **Hourly Averages**: Analysis of ABP study data including study comparison.
- **Summary**: Interpretative summary settings and results for current study.

**Viewing an Ambulatory Blood Pressure Study**

In the upper portion of the ABP Data tab, a table displays the results for each of the BP measurements taken or attempted during the study. The table also includes events recorded by the patient and dose response sequences. To display only these events in the table, select Events Only from the Show pull-down menu. Events and Readings displays all records.

Below the table are three tabs offering different graphical representations of the BP data: Overview Graph, Central BP Data, or AASI Graph.

---

**Figure 8: ABP Data Graph, Threshold tab**

To adjust the size of the data table:

1. Position the mouse pointer over the bottom boundary of the table.
2. When the pointer turns to a double arrow, drag it up or down.

**Overview Graph**

The scale of the vertical axis represents blood pressure (mmHg) and heart rate (bpm). The horizontal axis displays the time in clock hours. Clicking on any point, measurement or event, in the graph highlights the corresponding row in the table.
There are three tabs to the right of the graph: Threshold, Time-Slice, and Legend. The Threshold tab allows the user to change the Threshold settings on the graph. The Time-Slice tab allows users to define specific time windows in which to perform a statistical analysis. The Legend tab provides details on the configuration of colors for the Overview Graph.

Shading on the graph indicates various periods of the study. The default colors (see Display Settings) and definitions are:

- **Blue shading**: Indicates the asleep period of the study.
- **Yellow shading**: Indicates the white coat period, which is the first hour of the study (appears only if enabled).
- **Light green shading**: Indicates a dose response period (appears only if programmed).
- **Light teal shading**: Indicates the periods used to calculate Morning Surge (appears only if enabled).

**Central BP Data**: The Central BP Data tab provides a detailed view of individual CBP measurements taken during the study. If your Oscar 2 is configured with the SphygmoCor Inside technology, you will be able to collect, display, and analyze these parameters. The tab will display the number, time and date of the selected reading along with the following:

**Average Central Pressure Waveform**: A graphical representation of the derived average central pressure waveform. The shape of the aortic pressure pulse is a result of the ventricular ejection and physical properties of the arterial system. The waveform shape changes with changes in arterial stiffness.

**Central Systolic Pressure (cSYS)**: The maximum pressure during aortic ejection. High cSYS indicates high cardiovascular load. High arterial stiffness increases the reflected pressure wave in the arterial system, and augments or increases cSYS pressure. With aging, the arteries become stiffer. Consequently, wave reflection increases, leading to increased cSYS and resulting in increased risk of cardiovascular disease or organ damage.

**Central Diastolic Pressure (cDIA)**: The minimum pressure during aortic ejection.

**Central Pulse Pressure (cPP)**: Represents the height of the aortic pressure waveform. cPP can also be described as the difference between the maximum and minimum of the central pressure waveform, or cSYS minus cDIA. Aortic cPP greater than 50 mmHg has been shown to predict cardiovascular disease.

**Central Augmentation Pressure (cAP)**: cAP is a measure of the pressure wave reflected back from lower body. It is calculated as the difference between the two pressure peaks during ejection (systole). The first peak is related to cardiac ejection and the second peak is related to wave reflection due to arterial stiffness. The cAP value is affected by both the magnitude and speed of the reflected wave, which is an indicator of arterial stiffness.

**Central Augmentation Index (cAIx, cAIx@75)**: The ratio of cAP to cPP, expressed as a percentage. Studies have shown that patients with diabetes tend to have high cAIx, indicating stiffer arteries and higher risk of organ damage. cAIx is also corrected for a heart rate of 75 beats per minute.

**Measured Brachial Waveform**: A graphical representation of the brachial waveform captured during the displayed measurement. The brachial waveform data is used to derive the average central pressure waveform.

The **Central BP Data** tab also offers the user the option to print or save the details for any single CBP measurement from the study. The printed page or saved file will be a single page document containing the same information as is presented on the Central BP Data tab for the selected measurement along with the patient name, patient ID, study date, brachial systolic and diastolic values, and heart rate.

The following diagram shows the central aortic pressure waveform.
**AASI Graph**

The AASI Graph tab is optional, and can be enabled by selecting **Ambulatory Arterial Stiffness Index** on the Report>Configuration>Data (see Customizing and Configuring AccuWin Pro™ 4). The graph on this tab plots the diastolic on systolic BP values for each measurement captured in the study. Additionally, this tab lists the calculated AASI value, diastolic as a function of systolic, systolic as a function of diastolic, the correlation coefficient of the linear regression equation, and the coordinate of the average systolic and diastolic. The calculated AASI value will also be added to the Overall results on the Statistics tab. The user can specify which BP values to use in the graph and calculations by selecting **Brachial BP Values** or **Central BP Values** near the bottom of the legend.

**Reviewing an Ambulatory Blood Pressure Study**

In the ABP Data view, you can review an ABP study for accuracy and context. When the data is retrieved by AccuWin Pro™ 4, all readings in an ABP study that have event codes will be "tagged" in the first column of the table. Tagged records are omitted from the analysis of the ABP study displayed in the Statistics and Hourly Averages views, but these readings can be printed in the report.

The monitor tags data with an asterisk based on criteria used to determine the validity of the data. The exclamation point (!) is a permanent tag and is used when an accurate reading cannot be determined by the ABP monitor. This tag cannot be changed and associated data will not be used in data analysis. The asterisk (*) is a tag that you can edit. Tags numbered r01 to r30 indicate events entered by the patient, and the comments can be changed. Data can be tagged or un-tagged based on a number of factors, including patient history, patient diary information, or other factors.

There are two methods to remove or insert an asterisk (*) tag on a measurement.
To remove or insert an asterisk (*) tag using the table:
1. Select the reading you want to tag (omit) or un-tag (include). It will be highlighted.
2. Click in the first, or left-most, cell under the column labeled “Tag”.

To remove or insert an asterisk (*) tag from the graph:
1. Use the cursor on the graph to select the reading you want to tag (omit) or un-tag (include). Consequently, this action highlights the reading in the table.
2. Right-click the mouse and select Toggle Tag from the menu.

NOTE: All changes made to the ABP study are saved immediately.

Entering Comments
Use the Comments column, the right-most column in the table, to keep track of patient activity during a BP reading. While activity is only one of the many factors that can affect blood pressure, it can be helpful in understanding a BP reading within the context of the study.

To assist the patient with tracking their activities, a patient diary template is available for printing from the Download Library on the SunTech Medical website.

To enter a patient comment from the table:
1. Highlight the reading to which you want to add a comment, and select the corresponding cell in the Comments column.
2. Left-click the mouse over the comment box to open the pull-down menu of predefined entries.
3. Select a comment from the list or type a new comment.
4. Press Enter.

To enter a comment from the graph:
1. Select the reading in the ABP data graph to which you want to add a comment. This causes the corresponding row in the table to be highlighted.
2. Right-click the mouse and select Set comment from the menu.
3. Add a comment by typing or selecting from predefined list.
4. Click OK.

You can add or delete entries appearing in the Comments pull-down list by going to the Diary Comments section of the Configuration window. (For instructions on configuring Diary Comments, see Customizing and Configuring AccuWin Pro™ 4.)

To change the comment for an event marked by the patient:
Click the event's checkmark above the graph to cycle through the following selections (symbol: comment):

✓: “Event marked”
Rx: “Dosage”
Sx: “Symptom”

The selected symbol replaces the checkmark, and the comment appears in the table.

To change time period start times:
1. Display an ABP study and choose the Threshold tab.
2. Click Edit Awake/Asleep Times.
3. In the Times tab (Figure 12), reset the patient’s asleep and awake times to the nearest half-hour by moving the appropriate sliders right or left. The new times are displayed on the clock face, with the black portion representing time asleep and the white portion, time awake. The clock is displayed in military time (i.e. 12:00 is noon, 18:00 is 6 p.m., and 00:00 or 24:00 is midnight).
4. Click Apply to keep the new settings.

**Setting BP Threshold Limits**

Users may select the thresholds used to calculate the blood pressure load and above threshold values for brachial and central BP parameters. There are two settings for thresholds: global and custom. Global thresholds apply to all patient files; custom thresholds apply to all studies for a particular patient.

**To define thresholds:**
1. Open a study in the ABP Data tab. Within the Overview Graph tab, navigate to the Threshold tab located to the right of the graph view.
2. Click Edit Thresholds.
3. From either the Brachial Thresholds or Central Thresholds tab, click Use custom or Use global thresholds.
4. Move the sliders to change the settings according to your preferences.
5. Click OK to apply the changes.

**Brachial Thresholds**

Brachial thresholds can be set for Awake and Asleep systolic and diastolic BP. The default setting is a global threshold set to 140/90 mmHg for Awake periods and 120/80 mmHg for Asleep periods. If desired, you can set thresholds to match a published standard using the corresponding buttons: JNC7/AHA or ESH.

- JNC79 recommends 135/85 mmHg for Awake periods and 120/75 mmHg for Asleep periods.
- The American Heart Association (AHA)10 recommends a 24-hour average BP of 130/80 mmHg.
- The ESH11 recommends 135/85 mmHg for Awake periods and 120/70 mmHg for Asleep periods.

**Central Thresholds**

Central thresholds can be set for Awake and Asleep central systolic BP (cSYS), central pulse pressure (cPP), augmented pressure (cAP), and augmentation index (cAIx). The default setting for the Awake thresholds is a global threshold set to calculated values dependent on gender and age of the patient. If no gender or age is provided, the default values will be set to 130 mmHg for central systolic BP and 50 mmHg for central pulse pressure. No settings will be established for augmented pressure or augmentation index. There is also the option to set the central pulse pressure (cPP) according to the published STRONG standard, 50 mmHg.

The default setting for the Asleep thresholds is no selected values. The user does have the option of automatically setting Asleep thresholds to match the selected Awake values. If no Asleep thresholds are defined, all graphical representations and related statistics will be left blank.
**Setting Pediatric Threshold Limits**

The American Heart Association has a published statement supporting ABP monitoring in children, as hypertension is being diagnosed with increasing frequency in pediatric patients.\(^8\) The guidelines for hypertension differ from those for adults; hypertension in pediatric patients is generally defined as a blood pressure values above the 95th percentile for that gender and height. Specific ABPM thresholds based on published recommendations are programmed into AccuWin Pro\textsuperscript{™} 4.12,13,14

**To apply a pediatric threshold:**
1. Open a study in the ABP Data tab. Within the Overview Graph tab, navigate to the Threshold tab located to the right of the graph view.
2. Click Pediatric Thresholds.
3. Enter the patient’s gender and height. The height may be entered in centimeters (cm), inches, or feet/ inches. (Figure 15)
5. Click Apply.

*NOTE: The pediatric threshold is applied as a custom brachial BP threshold, which applies to a single patient's file.*

**Defining Time-Slice Periods**

Users may define up to twelve specific time windows for which to perform a statistical analysis. The statistical analysis corresponding to time-slice periods will appear in the Time-Slice Statistics tab to the right of the Statistics tab.

**To create a time-slice:**
1. Open a study in the ABP Data tab. Within the Overview Graph tab, navigate to the Time-Slice tab located to the right of the graph view. (Figure 16).
2. Click the New button. On the graph left-click and hold the mouse to select the starting time of the time-slice period then drag the mouse to the end time. Releasing the mouse button determines the ending time of the time slice. The selected time-slice will be highlighted on the Overview graph.
3. Once the time-slice is selected, a Time-Slice Settings box (Figure 17) will appear and allow the user to name the time-slice and edit the start and end times, if necessary.
4. Click Save to create the new time-slice.
To edit a time-slice:
1. Open an ABP study and click the Time-Slice tab (Figure 16).
2. Choose the time-slice from the drop-down menu. The time-slice period will be highlighted on the Overview graph.
3. Click Edit. The Time-Slice Settings box (Figure 17) will appear.
4. Edit the name or the start and end times.
5. Click Save to save the settings.

To delete a time-slice:
1. Display the ABP study. Choose the Time-Slice tab.
2. Choose a time-slice from the drop-down menu. The time-slice period will be highlighted on the ABP data graph.
3. Choose Delete. Click Yes to delete the time-slice (Fig. 18).

Editing Patient Info and Physician Info

Patient and physician information may be entered when data is retrieved from the monitor or after the patient file is saved.

To edit Patient Info (Fig. 19):
1. Click the Patient Info tab at the bottom of the screen.
2. Click on the Edit button at the bottom of the screen.
3. Update any patient information and click Save. Alternatively, you may click Cancel to discard all changes.

To edit Physician Info (Fig. 20):
1. Click the Physician Info tab at the bottom of the screen.
2. Click on the Edit button.
3. Update any information and click Save. Alternatively, you may click Cancel to discard all changes.
Users are able to enter and edit Patient History, Reason for Test, Current Medications, and Physician Interpretation. As an alternative to typing in the free text field for Current Medications, users may add Medication, Dosage, and Frequency to dictionaries that are stored within AccuWin Pro™.

To Add Medications (Fig. 21):
1. Type a medication into the drop-down titled Medication. (See Fig. 22 for abbreviations)
2. Once the medication is typed, press Enter or Tab on the keyboard or use your mouse to move the cursor to the Dosage column.
3. Enter Dosage and Frequency in the same manner. Predefined entries in the Frequency column are listed below.
4. Once the Medication, Dosage, and Frequency are entered into their respective columns, click the Add button to add the information to the Current Medications field above and to the dictionary.

To locate and choose a previously stored entry:
1. Either start to type an entry or choose the down arrow with your mouse. As more characters of the desired entry are typed, a list of matching entries will be displayed.
2. To choose the stored entry, use the down arrow on the keyboard, highlight the desired entry, and press Enter on the keyboard. Alternatively, use the mouse to highlight the entry, and then double-click the entry.
3. When each of the columns are populated with the correct information, click Add.

Viewing Statistics
To view the statistical analysis for the displayed ABP study, click on the Statistics tab. This window initially shows the following tables:

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>ac</td>
<td>Before meals</td>
</tr>
<tr>
<td>bid</td>
<td>Twice daily</td>
</tr>
<tr>
<td>pc</td>
<td>After meals</td>
</tr>
<tr>
<td>prn</td>
<td>As needed</td>
</tr>
<tr>
<td>qam</td>
<td>Every day before noon</td>
</tr>
<tr>
<td>qd</td>
<td>Every day</td>
</tr>
<tr>
<td>qh</td>
<td>Every hour</td>
</tr>
<tr>
<td>qhs</td>
<td>Every night at bedtime</td>
</tr>
<tr>
<td>qid</td>
<td>Four times a day</td>
</tr>
<tr>
<td>qod</td>
<td>Every other day</td>
</tr>
<tr>
<td>qpm</td>
<td>Every day after noon</td>
</tr>
<tr>
<td>qwk</td>
<td>Every week</td>
</tr>
<tr>
<td>tid</td>
<td>Three times a day</td>
</tr>
</tbody>
</table>
- **Overall**: Data for the complete ABPM study
- **Awake Period**: Data collected while the patient is awake
- **Asleep Period**: Data collected while the patient is asleep
- **White Coat Period**: Data collected during the first hour (if enabled).

The Statistics tab can also show additional time periods or BP load charts.

**To show additional time periods or BP load charts:**
Under **Show**, select one of the following:
- **Additional Time Periods** to show tables with data collected during other time periods, as programmed.
- **BP Load Charts** to show pie charts for the Overall, Awake, and Asleep periods, illustrating values above and below established thresholds for brachial and central systolic BP, and brachial diastolic BP.

**NOTE: All data in these tables is included in exports, except white coat analysis values and BP load charts.**

In the **Statistics** tab, each table includes the mean, standard deviation, coefficient of variation (CV), minimum value, and maximum value for the following parameters: brachial and central systolic, brachial and central diastolic, brachial and central MAP, brachial and central pulse pressure, central augmentation index, central augmentation index at 75 bpm, and central augmentation pressure.

Additionally, the following calculations are also provided on the Statistics tab:

- **BP Load**: Blood pressure load characterizes the distribution of measurements in an ABPM study relative to brachial systolic and diastolic thresholds indicative of hypertension. The percentage of readings above this threshold is the BP load which has been shown to predict target organ involvement.\(^{14}\)
- **Asleep Dip**: The nocturnal fall of blood pressure expressed as a percentage of the mean awake values. The expected nocturnal dip for brachial BP is 10-20%. The absence of the expected dip may indicate end-organ damage or risk for cardiovascular events.\(^ {7,16}\)

The Statistics tab may include any of the following additional calculations if they are enabled under **Report>Configuration>Data>**:

- **Ambulatory Arterial Stiffness Index (AASI)**: AASI is calculated as one minus the regression slope of the diastolic on systolic blood pressure. (The regression slope is calculated by plotting the diastolic against the systolic values.) This index has been used to measure arterial stiffness and has been shown to help predict target organ damage, cardiovascular mortality, and stroke.\(^ {17}\) The user can specify which BP values to use when calculating this value by selecting **Brachial BP Values or Central BP Values** from the AASI Graph tab.
- **Morning Surge**: Morning Surge is calculated as the difference between the mean systolic BP during the two hours after waking, minus the systolic BP during the Asleep hour containing the lowest BP value measured. This calculation can help to identify cases in which the morning BP increase is unusually high and presents a potential cardiovascular risk or a risk of stroke.\(^ {14}\)
- **Coefficient of Variation**: The Coefficient of Variation (CV) can be used as a measure of BP variability, which can be a determinant of end-organ damage.\(^ {18}\) CV is calculated as the standard deviation divided by the mean blood pressure of a time period.
- **Smoothness Index**: Smoothness Index (SI) is the ratio between the average of the blood pressure changes computed for each hour of the recording and its standard deviation. SI is generally used to evaluate the effectiveness of antihypertensive treatment as it shows the averaged trend of blood pressure reduction between two studies.\(^ {19}\)
- **White Coat Analysis**\(^ {18}\): White coat syndrome refers to abnormally elevated blood pressure when the patient is in a medical setting. If selected, this option does the following:
• Adds a white coat period to the statistical analysis.
• Displays the first hour of the study, or white coat period, with a light blue background in the graphs.
• Evaluates for and reports on the presence of white coat syndrome on the interpretive summary, when present.

NOTE: Either brachial or central BP values can be used to calculate Morning Surge, Coefficient of Variation, and Smoothness Index. To specify which blood pressure values to use, see Data Settings.

Viewing Dose Response Statistics
To view the statistical analysis for dose response sequences, click on the Dose Response Statistics tab. This tab is present only when dose response sequences have been recorded. You can specify which sequence to display by selecting its number from the Show Sequence menu. The window shows seven scatter plots for ten parameters measured within the dosage response sequence. Results are also provided for BP load and maximum change from the baseline (the first reading of the sequence) for the systolic (SYS), diastolic (DIA), and central systolic (cSYS) parameters.

Viewing Time-Slice Statistics
To view the statistical analysis for user defined time-slice periods, click on the Time-Slice Statistics tab (Figure 23).

Viewing Hourly Averages
To display trends in a patient’s ABP study, click the Hourly Averages tab at the bottom of the window (Figure 24). The table shows the average BP readings for each hour the patient was tested. In addition to brachial systolic (SYS) and diastolic (DIA), and the heart rate (HR), the following statistical averages are provided:

• Mean Arterial Pressure (MAP) - This is the average pressure in an artery over the period of one heartbeat. In the brachial artery, it is calculated by adding the diastolic to one-third of the difference between the systolic and diastolic readings.
- **Pulse Pressure (PP)** - This is calculated by subtracting the diastolic from the systolic reading. It is another hemodynamic parameter that may serve as an indicator for cardiovascular risk. Pulse pressure is provided using both brachial (PP) and central (cPP) values.

- **Pressure Rate Product (PRP)** - This is the product of the average systolic reading multiplied by the average heart rate. PRP strongly correlates to a patient’s activity level and may be a key indicator of cardiovascular risk. As typically reported in research and clinical applications, PRP data is divided by 1000.

Use the check boxes at the top of each column to select the statistics you would like to see displayed in the graph at the bottom of the page. You can resize the table and zoom in on the graph with your mouse pointer. (For instructions on resizing and zooming, see Viewing an Ambulatory Blood Pressure Study.)

![Figure 24: Hourly Averages](image)

**Viewing the Interpretive Report Summary**

This window (Fig. 25) shows an interpretation of the ABPM data based on published guidelines. Because traditional BP guidelines might not apply to the 24-hour ABPM readings, AccuWin Pro™ 4 includes the recommendations specifically for evaluation of ABPM levels. The summary provides normal or hypertensive results for the 24-hour average, awake, and asleep systolic and diastolic pressure readings, asleep “dipping” status, and an optional white coat analysis.

![Figure 25: Summary Tab](image)
To generate an interpretive summary for the displayed ABP study, click the Summary tab on the bottom of the screen and select one of the following options for interpretation:

- Joint National Committee Seventh Report (JNC)\(^7\) and American Heart Association (AHA)\(^8\) guidelines.
- European Society for Hypertension (ESH)\(^6\).
- Pediatric – AHA\(^1\)
- Pediatric – Soergel Tables\(^1\)

Based on the calculated brachial dip percentage, AccuWin Pro™ 4 determines whether the patient is one of the following:

- Dipper (normal): A patient shows a decrease of 10% or more in brachial systolic or diastolic blood pressure values during sleep compared to readings taken when awake.
- Non-dipper (abnormal): A patient shows no decrease or less than a 10% decrease in brachial systolic or diastolic blood pressure values during sleep compared to readings taken when awake.

**Comparing Two Studies**

To further enhance the utility of ABPM, you may want to measure the progress of a patient’s blood pressure by conducting multiple studies. AccuWin Pro™ 4 allows for the comparison of the hourly averages and statistics for two studies conducted on the same patient.

To compare two studies:

1. Click on the **Hourly Averages** tab at the bottom of the window.
2. Click on the **Get Comparison Study** button. (If the button is grayed out, the patient file you are viewing contains only one study.)
3. The **Choose Date** window will appear. Select the study you want to use as a reference for comparison by highlighting it and clicking **OK**.

The statistics generated during the previous study are now incorporated into the Hourly Averages window. The table expands to include the hourly average values for the reference study, and the differences between the displayed and the reference studies.

On the graph at the bottom of the window, both studies’ data are displayed. The dotted lines represent data from the reference study, and solid lines represent data from the displayed study. Select Difference and the graph will display one line representing the difference between the two studies. The same navigation tools for the Hourly Average and ABP Data windows also apply to the comparison window.

---

**10. Creating Reports**

**Configuring and Customizing the Report**

To document a study and its findings, you can create a customized report (Figure 26).

AccuWin Pro™ 4 provides the following preconfigured report formats:

- One page report: Summary page only.
- Standard report: Summary, Patient Information (all information from the Physician Info tab), Statistics, ABP Data, and Monitor Configuration pages.
- Full report: All report pages.
NOTE: All report pages include the SunTech Medical logo, patient demographics (all information from the Patient Info tab), and test date. The footer of each page identifies the type of monitor used and the page number.

To configure or customize your report (Fig. 26):
1. From the Report menu, select Configure, or click on the Settings button on the toolbar then navigate to the Report tab in the Configuration window.
2. On the Formats tab, do one of the following:
   - Choose one of the formats listed in the left pane. When you select a format, the pages included in that format are selected in the Report Pages box.
   - Create a new custom report format by clicking New Report. Enter the report format name. Select the pages from the Report Pages list on the right.
3. To include an additional page in your report, click the respective page in the Report Pages box.
4. Under Report Title, enter up to five lines of text that will appear in the header of every report page.
5. Select either a single-page or multi-page view for the Print Preview window on the Details tab (Figure 27).

You can choose from the following pages to customize your report (listed in the order the pages will print, see Figure 26):

**Summary**: The Interpretive Summary appears above the data graph and summary statistics. Summary brachial and peak central BP statistics for the overall, awake, asleep, and white coat (if chosen) periods include: mean systolic and diastolic, mean heart rate, BP load, and asleep dip.

**Patient Information**: Includes patient history, reason for test, patient medications, and the physician interpretation.

**Statistics**: Displays detailed statistics for brachial and central BP parameters according to the time periods included in the study. Overall, awake and asleep time period results will be displayed first, followed by the white coat period (if enabled), and finally any additional time periods programmed by the user.

**Time-Slice Statistics**: Displays statistics for brachial and central BP parameters for each user defined time-slice period.

**Reviewed Data**: Shows all the BP readings captured during the study in table format as it is shown on the ABP Data tab within AccuWin pro™ 4, excluding the event codes (EC column). The vertical black line between the # and Time columns indicates the asleep time period. A “>” symbol adjacent to the BP values indicates that it has exceeded the threshold.

**Omitted Data**: Shows the BP readings that were excluded from the statistics calculations. This page appends to the Reviewed Data page, if printed.

**Events Marked**: Shows events marked by the patient during the course of the study. This page appends to the Reviewed Data page, if printed.

**Dosage Response**: Displays the statistics and scatter plots for each of the dosage response sequences captured during the study, as they are displayed in the Dose Response Statistics tab.

**AASI**: Displays the data and graph for the AASI parameter, as they are displayed in the ABP Data>AASI tab.

**BP Load**: Shows the BP load pie charts.

**Hourly Averages**: Displays hourly average data for the study in graph and table formats, as displayed on the Hourly Averages tab.

**Study Comparison**: Displays study comparison data in graph and table formats, as displayed on the Hourly Averages tab.

**Histograms**: Contains frequency histograms for awake and asleep time periods for brachial and central systolic and diastolic BP, heart rate, MAP, PRP, and PP.
Monitor Configuration: Shows the programmed time period, dosage response, and monitor settings for the study. The page also displays the quality control results for the study (if applicable) and definitions for event codes.

To configure report details (Figure 27):
1. Click the Details tab. From here you can control the following parameters for printing reports:
   - **Overview Graph Type**: Select line, bar, or the type of graph currently displayed according to the type you want to print on the Summary page of the report.
   - **Overview Graph Scale**: Select time scale for the graph that prints on the Summary page of the report.
   - **Summary Page User Fields**: Check desired fields for medications, patient history, test reason, and physician interpretation.
   - **Horizontal Sight Lines**: Select preferred frequency of sight lines in the Reviewed ABP Data table when printing the Reviewed Data Page, or Omitted Data Page in the report.
   - **Report Viewing Options**: Select either a Single-page or multi-page view of the report which will appear in the Print Preview window.
2. Click OK to apply the changes.

Previewing the Report
1. From the Report menu, select Preview. Alternatively, click the Print Preview toolbar button. The Print Preview window is displayed (Figure 28).
2. To scroll through the report pages, click on the numbered icons on the left side of the window, or select the page you want to view using the drop-down menu.
3. To change the format and details, click Configure.
4. To print the report, click Print Report. To print the currently viewed page, click Print Page
**Printing the Report**

AccuWin Pro™ 4 provides you with flexibility in printing the report. You can either:

- Print the configured report by clicking the **Print Report** button on the toolbar or by going to the Report menu and selecting **Print**. The Print dialog box is displayed, where you can choose the preferred printer.
- Print the report page corresponding to the current tab that is being displayed by clicking **Print Page**.

**Saving the Report as a PDF**

To save the report as an Adobe Portable Document Format (PDF), from the Report menu, select **Create PDF**. Alternatively, click the PDF toolbar button. The system creates the PDF and saves the file to the location you specify.

*NOTE: You must have a printer installed. If you do not have a printer installed, see the online Help to install a driver to print to a file.*

---

**11. Overview of AccuWin™ Pro 4 Report Symbols**

The following symbols are used in the Reviewed and Omitted BP data report pages.

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td>+</td>
<td>Signifies the reading was initiated manually by the patient pressing the Start/Stop key on the monitor.</td>
</tr>
<tr>
<td>-</td>
<td>Indicates that the pressure to which the cuff inflated in dynamic inflation mode may have been too low to obtain an accurate systolic reading. Readings marked with “-” are automatically omitted from the study. Compare the tagged reading with a “-” to the BP readings above and below it. If the omitted reading seems to be acceptable, un-tag it from the ABP Data tab; if it does not seem acceptable, leave it “tagged”.</td>
</tr>
<tr>
<td>r</td>
<td>Indicates that this reading was a retry reading. Retries are readings that are automatically obtained 4 minutes after a measurement attempt fails or a questionable reading is obtained by the monitor.</td>
</tr>
<tr>
<td>!</td>
<td>The exclamation point is a permanent tag and is used when an accurate reading cannot be determined by the Oscar 2.</td>
</tr>
<tr>
<td>&gt;</td>
<td>This symbol will appear to the left of the measurement if the reading results were above the set threshold.</td>
</tr>
<tr>
<td>r01 to r30</td>
<td>These numbered tags identify a series of patient-marked events which occurred during the study.</td>
</tr>
</tbody>
</table>

---

**12. Managing Patient Studies**

**Opening a Patient Study**

This feature allows you to retrieve and view a patient study.

1. From the toolbar, click the **BP Data** button or from the **File** menu, select **Open**. The **Open ABP Study – Select Patient** dialog box appears listing all available patient files.
2. Select the patient whose BP data you wish to retrieve by clicking on the patient’s name. A search bar is available to locate the patient by any information relevant to the patient’s file. Click **OK**.
3. If the patient has more than one ABP study, a second dialog box, **Choose Date**, will be displayed. Select a date and click **OK**.
4. The patient's data appears in the Display Area. You can now review the BP data or print a report.

**Exporting a Patient Study**
You can export BP data files to ASCII, GDT or XML format. XML can be used to create an HL7 compatible file.

1. From the File menu, select Export.
2. Choose either to export the Displayed File or select files From Database.
3. If exporting files from the database, select the files you want to export and click OK.
4. The Export dialog box appears. Click Export to export the file(s). Click Edit to change the list of files you want to export. Click Configure to configure the details of the exported file(s). (For instructions on configuring export, see Export in Customizing and Configuring AccuWin Pro™ 4).

The exported files will be exported to the selected file location as either a *.ASC, *.GDT or *.XML file depending on the format chosen.

**Emailing a Patient Study**

To email AccuWin Pro™ 4 data files as an attachment:
1. From the File menu, select Email.
2. Choose either to email the Displayed File or select files From Database. If displayed File is chosen, the data currently displayed on the screen is emailed. If From Database is chosen, the patient database window opens and prompts you to Select file(s) to email. Check the files you want to email and click OK.
3. The Email dialog box will appear. Type in the recipient email address(es), a subject for your message, and a message.
4. Click Send to email the file(s). Click Attach to change the files you want to email. Click Configure to configure the email settings. (For instructions on configuring email, see Email in Customizing and Configuring AccuWin Pro™ 4).

**Deleting a Patient Record**
This feature allows you to delete one or more patient records at a time. Each patient record contains ALL ABPM studies for that patient; therefore ALL ABPM studies for that patient will be deleted when selected.

To Delete Selected Patient Study(s):
1. From the file menu click Patient Management.
2. In the Patient Management window, select the patient(s) that you'd like to delete.
3. Click Delete.
4. A dialog box Select Date may appear. You may select one study date by clicking on it. Alternatively, you may use the checkboxes to select more than 1 patient file.
5. Click OK to delete the selected study(s). A confirmation window Delete Study will appear. Select Yes to confirm deletion of selected studies. Click No to cancel and return to the Select Date window.

**Importing a Patient Record**
You may import patient studies that are saved on your computer, so that the files are stored in the main AccuWin Pro™ 4 patient database.

To import patient studies:
1. Click File then select Import > Import Patient Study.
2. Click Browse Files. A Windows Explorer Window will appear to allow you to select the patient study files that you would like to import.
3. Select the patient study files that you would like to import. You may select multiple files using standard windows shortcuts, if you prefer.
4. Click Open. The patient study files will populate in the Import Patient Study window.
5. Click Import. A progress bar will show the status of the import process.
**Grouping Patient Records**

Grouping patient records can be useful for organizing patient data based on certain demographics, templates, or studies.

**To create a new group:**
1. Click File then select **Group Management**.
2. Click **Add**. A dialog box appears with a text entry box for the new group name.
3. Type the new group name in the text entry box.
4. Click **OK**.
5. A dialog box appears **Group added successfully**. Click **OK**.

**To move patients between groups:**
1. Click File then select **Patient Management**.
2. Using the checkboxes to the left of the dialog box, select the patients that you would like to manage.
3. Click **Move Patient**.
4. Select the Group Name using the drop-down menu from the dialog box.
5. Click **OK**. A confirmation window will appear once the patient(s) have been moved successfully.

---

**13. Customizing and Configuring AccuWin Pro™ 4**

It may be helpful to have multiple configuration profiles for your users, for example, one for each physician in the practice.

**To create or edit configuration profiles in AccuWin Pro™ 4:**
1. From the **Configure** menu, select **Preferences**.
2. Create a new profile or edit an existing profile. Click on the name of the profile you want to edit and click **Edit**. To create additional profiles, click **New**, name the profile, click **Enter**, and then select **Edit**.
3. Click **OK** to choose a configuration.

If you are editing a configuration profile, a window opens to display a number of configuration options which are described in detail on the following pages.

*NOTE: On each tab, clicking a Default button returns the settings of that tab to the default factory settings*
**Hardware Settings**

The Hardware Settings window (Figure 29) allows you to configure your computer to communicate with your ABP monitor via a specified port. You may also test the settings that you have selected.

1. On the tool bar, click on Monitor, then choose Configuration Port.
2. Under ABP device, select your monitor type.
3. Under Serial port, select the port into which you have plugged the device.
4. Click Test port. The unit will beep once, and you will receive a message confirming successful communication.

If you receive an error message, “Cannot communicate with ABP device,” one of the following applies:
- You have not selected the appropriate port or device.
- The device is not connected properly.

**Data Settings**

The Data Settings tab (Figure 30) allows you to configure the following settings:

**Data Source**: Select how heart rate, Morning Surge, and Smoothness Index are displayed or calculated (i.e. using brachial or central BP values). A lowercase “c” will designate which values are set to use central BP values (e.g. HR becomes cHR).

**Additional calculations**: Select which additional parameters will be displayed when viewing a patient file. Options include:
- AASI
- Smoothness Index
- Morning Surge
- White Coat Analysis
- Coefficient of Variation

Selecting any of these options will make them visible on corresponding tabs and graphs.
Display Settings
This feature allows you to customize the color scheme of the graphs and charts (Figure 31). Customizable palettes allow you to fine-tune the color scheme for optimum reproduction by your printer. Shading on the graph indicates the period of the study. The default colors (which can be changed using the Configuration>Display tab) are the following:

- Blue shading: Indicates the asleep period of the study.
- Yellow shading: Indicates the white coat period, which is the first hour of the study (appears only if enabled).
- Light green shading: Indicates a dose response period (appears only if programmed).
- Light teal shading: Indicates the periods used to calculate Morning Surge (appears only if enabled).

To customize the color palette:
1. Click on the tab for the attribute you want to change.
2. To change a color, click on it. A color palette window will appear, showing 48 basic colors. You may select one of these, or you may create your own custom colors by clicking on the Define Custom Colors button.
3. Configure the color then click Add to Custom Colors.
4. Repeat this process for each color you want to change.
5. Click OK to save changes.

Diary Comment Settings
You can use this feature to keep track of entries from patient diaries describing the patient’s activities during BP measurements. The entries in this window (Figure 32) are a default list which can be customized by adding, editing, or deleting entries. The entries in this list are incorporated into a list that appears in the pull-down menu under the Diary Comments column located on the ABP Data tab.

To add a new diary comment:
1. In the Diary Comments tab of the Configuration window, Click New.
2. Type the text for the new diary comment.
3. Click anywhere to exit the text entry window and store the diary comment.
4. Click OK to save your changes and exit the Configuration window.

To delete a diary comment:
1. In the Diary Comments tab of the Configuration window, select the diary comment that you want to delete.
2. Click Delete.
3. Click OK.

To edit a diary comment:
1. In the Diary Comments tab of the Configuration window, select the diary comment that you want to edit.
2. Edit the text, as desired.
3. Click anywhere to exit the text entry window and store the diary comment.
4. Click OK to save your changes and exit the Configuration window.
**Email Settings**

The Email feature creates a new email message, attaches selected patient data files or programming templates, and sends the email to a specified destination.

**To set up email for AccuWin Pro™ 4 (Figure 33):**

1. Choose the type of connection you are using by selecting LAN or Dial up.
2. Fill in the Email server settings. This information is available from your Internet service provider (ISP) or office network administrator. Your ISP may require you to check the Authorize box in order for you to send email.
3. The Email settings you fill in will appear in the header of your email: the name of the recipient, your return email address, the subject of the email and a message of your choosing. All bolded categories on the tab are required.

Encryption Options allow you to encrypt your emails with a key of your choosing. Patient files are always encrypted. If the key is included in the file, any copy of AccuWin Pro™ 4 can open the file. To increase security, uncheck the box for Include key in file and type in a unique encryption key. The recipient of the emailed file must have the key to open the attached file.

**Report Settings**

This feature allows you to format your reports and to choose what information is included in them.

To format a report:

1. From the Configure menu, select Preferences. Select a profile and click Edit.
2. Click the Reports tab and then the Formats tab.
3. Select which items will be included in your One Page report, Full Report, and Standard Report. Select or de-select the checkboxes on the right to make these changes (Figure 26).
4. To create a new report, click New Report, enter a name, and select the pages you want to appear in the report.
5. To delete any of these reports, select the report and click Delete Report.
6. Enter a Report Title by typing the title of your choice in the box provided. The title appears as a header on each page of the report. Each report format allows for a different report title.
7. To change the graph type, graph scale, and horizontal sight line settings, Click Details (Figure 27).
**Export Settings**
This feature allows you to export reports either ASCII or GDT output (Figure 34), which is useful when the data will be merged with a database. GDT is a format commonly used to share computer medical records.

To change export settings, navigate to report>configure, then click on the Export tab within the Configuration window.

**To configure export to ASCII output,** make selections in these areas:
1. **Delimiters:** Choose which character will separate field items in the database.
2. **Export:** Choose exactly what information will be exported.
3. **To configure export to GDT output,** do the following:
4. Select the **Export GDT** tab.
5. Click the **GDT Summary** checkbox.

![Configuration Window, Export](image)

**14. Overview of Event Codes**
Event Codes (EC) are used during the review of ABPM data. The codes are displayed in the table on the ABP Data tab and in the Reviewed and Omitted BP data report pages under the columns labeled EC. Event Codes describe the conditions under which the BP measurement was taken. They are also accompanied by solutions, if applicable, that can be taken to avoid their occurrence in the future.

**NOTE:** Codes do not automatically indicate that a reading is invalid; they serve only as guides to help you review the data.

**Event Code Definitions**

<table>
<thead>
<tr>
<th>Code</th>
<th>Description in AccuWin Pro™ 4</th>
<th>Solution or Response</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Weak or no oscillometric signal</td>
<td>Check position and tightness of the cuff.</td>
</tr>
<tr>
<td>2</td>
<td>Artifact/erratic oscillometric signal</td>
<td>Patient should remain still during the BP reading.</td>
</tr>
<tr>
<td>3</td>
<td>Exceeded retry count: 4 attempts</td>
<td>Patient should remain still during the BP reading.</td>
</tr>
<tr>
<td>4</td>
<td>Exceeded measurement time limit: 140 seconds</td>
<td>Check the air hose connections and make certain the cuff is tight.</td>
</tr>
<tr>
<td>5</td>
<td>Results outside of published range: BP: 25-260 mmHg HR: 40-200 bpm</td>
<td>Retry the reading by pushing the Start/Stop button. If the problem persists, return the unit for service.</td>
</tr>
<tr>
<td>6</td>
<td>Release interval violation</td>
<td>Retry the reading by pushing the Start/Stop button. If problem persists, return the unit for service.</td>
</tr>
<tr>
<td>85</td>
<td>Reading aborted – blocked valves or pneumatics</td>
<td>Check the air hose connections and make certain the air tubing is not crimped.</td>
</tr>
<tr>
<td>Code</td>
<td>Description in AccuWin Pro™ 4</td>
<td>Solution or Response</td>
</tr>
<tr>
<td>------</td>
<td>--------------------------------</td>
<td>----------------------</td>
</tr>
<tr>
<td>86</td>
<td>Reading aborted – user abort</td>
<td>Restart the reading by pushing the Start/Stop button.</td>
</tr>
<tr>
<td>87</td>
<td>Reading aborted – inflate time-out or air leak</td>
<td>Check the air hose and cuff.</td>
</tr>
<tr>
<td>88</td>
<td>Reading aborted – safety time-out</td>
<td>Retry the reading by pushing the Start/Stop button. If the problem persists, return the unit for service.</td>
</tr>
<tr>
<td>89</td>
<td>Reading aborted – cuff over-pressure</td>
<td>Check for blocks or kinks in the air hose.</td>
</tr>
<tr>
<td>90</td>
<td>Service required – power supply out of range or other hardware problem</td>
<td>Replace the batteries. If the problem persists, return the unit for service.</td>
</tr>
<tr>
<td>91</td>
<td>Service required – safety override fitted or auto-zero out of range</td>
<td>Retry the reading by pushing the Start/Stop button. If the problem persists, return the unit for service.</td>
</tr>
<tr>
<td>94</td>
<td>Low battery warning</td>
<td>Replace the batteries. If the problem persists, return the unit for service.</td>
</tr>
<tr>
<td>97</td>
<td>Service required – transducer out of range</td>
<td>Return the unit for service.</td>
</tr>
<tr>
<td>98</td>
<td>Service required – A/D out of range</td>
<td>Return the unit for service.</td>
</tr>
<tr>
<td>99</td>
<td>Service required – EEPROM calibration data CRC failure</td>
<td>The unit needs to be recalibrated. Return for service.</td>
</tr>
<tr>
<td>109</td>
<td>Dose response sequence end</td>
<td>Review full dose response sequence on Dose Response Statistics tab</td>
</tr>
<tr>
<td>110</td>
<td>Event marked by patient</td>
<td>In ABP table, select desired comment.</td>
</tr>
<tr>
<td>111</td>
<td>Day to night mode switch</td>
<td>If desired, adjust ABP Data graph to match the time marked by the patient.</td>
</tr>
<tr>
<td>112</td>
<td>Night to day mode switch</td>
<td>If desired, adjust ABP Data graph to match the time marked by the patient.</td>
</tr>
</tbody>
</table>

15. Administrative Tools

AccuWin Pro™ 4 includes administrative tools that implement additional security-related features. With AccuWin Pro™ 4 Admin Tools, you can create new user accounts, change user access levels, reset user passwords, enable the login process, and enable automatic logoff. The available user roles are as follows:

1. Administrative: Full access to entire functionality of AccuWin Pro™ 4, including editing Admin Tool settings.
2. User: Full access to entire functionality of AccuWin Pro™ 4, except for editing Admin Tool settings.
To display the Admin Tools window:
1. From the Configure menu, select Admin Tools.
2. If the Enter Name and Password dialog window appears, enter your administrative name and password, and click OK. (The window appears if you have not logged in as an administrator or if login security is not enabled.)
3. The first time you access Admin Tools, you must use the default user name and password provided when the AccuWin Pro™ 4 was downloaded. For assistance, please visit www.suntechmed.com or contact SunTech Medial customer support. See section 18, Technical Assistance, for contact information.

Specifying Login Options
AccuWin Pro™ 4 includes the flexibility of enabling or disabling login security, which requires users to log in with a password. Enabling this option supports the HIPAA access protection requirements. If your facility uses an Electronic Medical Record (EMR) system or another system that already provides login security, you may not want to enable login security in AccuWin Pro™ 4.

When login security is enabled, you can also set the software to automatically logoff users after a specified period of inactivity in AccuWin Pro™ 4.

To enable login security:
1. In the Admin Tools window, click the Use Passwords checkbox. A check indicates that the option is enabled.
2. Click Done.

To enable automatic logoff:
1. In the Admin Tools window, click the Auto Logoff Enabled checkbox.
2. In the Auto Logoff Minutes box that appears, enter the number of minutes that must elapse before an inactive user is automatically logged off. Alternatively, you can click the Up or Down arrow to select a number. You can set the number of minutes between 1 and 240; the default is 20 minutes.
3. Click Done.

Adding User Accounts
An AccuWin Pro™ 4 administrator can create a user at either an administrative or one of two user levels. Each new user is assigned a default password of “user”. When new users first log in, they must change the default password before they can perform any tasks in AccuWin Pro™ 4. This applies to both administrative and user accounts.

To add a new user:
1. In the Admin Tools window, click New User.
2. In the User Name field that appears, enter the new user login name.
3. In the Select Role field, use the drop-down menu to select the level/role for the user.
4. Click OK to create the new user.
5. Click OK to save changes.

Changing User Level
You can change user account levels, but you cannot change user names.

To change user level:
1. In the Admin Tools window, click the user account you want to change.
2. Click Change Role. This option toggles the user level that appears in the Role column.
3. Click Done.
Deleting User Accounts
You can delete both administrative and user-level accounts; however, you cannot delete an administrator account if it is the only one. You also cannot delete yourself. Deleting a user account prevents the user from logging into AccuWin Pro™ 4, if the login security feature is enabled.

To delete a user account:
1. In the Admin Tools window, click the user account you want to delete.
2. Click Delete User.
3. A confirmation message appears. Click Yes to continue deleting the user account.
4. Click OK.

Resetting User Passwords
When a user forgets his or her password, that user must contact the AccuWin Pro™ 4 administrator, who will reset the user’s password to the default password of “user.” When he or she first logs in again, the user must change the default password before being able to perform any tasks in AccuWin Pro™ 4.

To reset a user password:
1. In the Admin Tools window, click the user account you want to change.
2. Click Reset Password.
3. Click Yes to confirm resetting the password.
4. In the confirmation message that appears, click OK.
5. Click OK.

16. Maintaining and Cleaning the Oscar 2

The Oscar 2 system is designed to perform in conformity with the description contained in this user manual and accompanying labels and inserts, when assembled, operated, maintained, and repaired in accordance with the instructions provided. After use, it is important to perform preventative maintenance to ensure the safe and efficient operation of the monitor. It is your responsibility to:

- Check calibration of the device every two years.
- Never knowingly use a defective device.
- Immediately replace parts that are broken, worn, missing, incomplete, damaged or contaminated.
- Contact the nearest authorized service center should repair or replacement become necessary.

Further, the user of the device bears sole responsibility for any malfunction that results from improper use, faulty maintenance, improper repair, damage or alteration by anyone other than SunTech Medical or authorized service personnel.

While usage will have an impact, it is expected the monitor will be in service for 8 years. Typically, an electromechanical pump determines the lifetime of the monitor. Service and support, including relevant accessories, will be available up to 7 years following the last date this product is manufactured by SunTech Medical. For a list of service centers, see section 18, Technical Assistance.
Cleaning After Use

Oscar 2
The Oscar 2 monitor cannot be sterilized. DO NOT immerse the monitor in any fluid, or attempt to clean with any liquid detergents, cleaning agents, or solvents. You may use a soft, damp cloth to remove dirt and dust from the monitor. If the unit does become immersed in water, do not use; contact our service department.

Orbit ABPM Cuff & Accessories
Remove the bladder for cleaning. Dampen a soft cloth with a mild medical grade disinfectant and wipe the bladder, let it air dry. The fabric shell of the Orbit cuff may be machine-washed in cold water with a mild disinfectant. Line dry this cuff only - machine drying can cause damage to the fabric shell of the Orbit ABPM cuff.

The bladder needs to be inserted back into the cuff sleeve so the pneumatic hose portion of the bladder goes through the tube opening on the cuff. Please note that the pneumatic hose connection should face upward when using the Orbit ABPM cuff on either the right or the left arm.

⚠️ CAUTION: Do not machine wash bladder

Maintenance and Repairs After Use
Visually inspect cables, material, pneumatic hoses, and the monitor case for cracks, fraying, or kinks. DO NOT use the monitor or cuff if there are any signs of damage. Please contact our service department if any damage or defects are identified.

The Oscar 2 does not contain any user serviceable internal parts and should only be opened by an authorized service representative. To return for service, please send to your nearest SunTech office, listed above, care of Support and Service. Alternatively, please visit our website, www.SunTechMed.com, to request more information.

Calibration Verification Procedure
It is recommended that you check the accuracy of the Oscar 2 once every two years. If needed, an authorized service center may recalibrate the pressure transducers in the monitor. To verify calibration, the Oscar 2 must first be placed into the proper mode. Follow the steps below:

1. Remove and then replace one of the two “AA” batteries.
2. After the number of readings is displayed, press and hold down the Start/Stop key.
3. The unit will display the software version.
4. The unit will display the battery voltage.
5. You will then hear a click as the valves are closed.
6. You will now see “0 mmHg” displayed.

The monitor’s calibration can now be checked against a calibrated mercury column.

1. Place a t-tube (part #98-0030-00) between the hoses connecting the monitor and the cuff.
2. Wrap the cuff around a suitably sized can or bottle. This acts as the reservoir for the unit.
3. Attach the third end of the “T” tube into a calibrated mercury column, which gives you access to the bulb and a reference.
4. Using the bulb of the calibrated mercury column, inflate the cuff to 250 mmHg.
5. Once the pressure has stabilized at this level, the LCD should match the mercury column by ± 2.0 mmHg.
6. Check the unit against the column every 50 mmHg from 250 to 50 mmHg. The monitor should be within ± 2.0 mmHg. If it is not, the monitor needs to be returned to the service department for recalibration or repair.

NOTE: To return the Oscar 2 to its normal operating mode, remove and replace one of the batteries.
17. Limited Warranty

Oscar 2 Ambulatory Blood Pressure Monitoring System
SunTech Medical provides to the original purchaser the following limited warranty from the date of original invoice.

<table>
<thead>
<tr>
<th>Product</th>
<th>Warranty Period</th>
</tr>
</thead>
<tbody>
<tr>
<td>Serialized blood pressure monitor</td>
<td>24 months</td>
</tr>
<tr>
<td>SunTech ABPM Cuffs</td>
<td>6 months</td>
</tr>
<tr>
<td>Accessories (i.e. patient hoses, interface cables, etc.)</td>
<td>90 days</td>
</tr>
</tbody>
</table>

SunTech Medical, Inc. warrants each instrument to be free from defects in material and workmanship. Liability under this warranty covers servicing of the instruments when returned from the customer’s facility prepaid to the prospective factory depending on location. SunTech Medical will repair any component(s) or part(s) that it finds to be defective during the period of this limited warranty. Should a defect become apparent, the original purchaser should notify SunTech Medical of the suspected defect. The instrument should be carefully packaged and shipped prepaid to the appropriate service center listed in section 18, Technical Assistance.

The instrument will be repaired in the shortest possible time and returned prepaid by the same shipping method as received by the factory.

This limited warranty is void if the instrument has been damaged by accident, misuse, negligence, or serviced by any person not authorized by SunTech Medical.

This limited warranty contains the entire obligation of SunTech Medical and no other warranties expressed, implied, or statutory are given. No representative or employee of SunTech Medical is authorized to assume any further liability or grant any further warranties except as set herein.

18. Technical Assistance

For any questions, please reference the Help section within the software, this user manual, or our website. If these do not fully address your problem, please contact our service department.

USA, Canada, and Latin America
SunTech Medical, Inc.
507 Airport Blvd, #117
Morrisville, NC 27560-8200
Phone: 1-919-654-2300
1-800-421-8626
Fax: 1-919-654-2301

Europe, Middle East, and Africa
SunTech Medical, Ltd.
Oakfield Industrial Estate
Stanton Harcourt Road
Eynsham, Oxfordshire OX29 4TS
England
Phone: +44 (0) 1865-884-234
Fax: +44 (0) 1865-884-235

Asia-Pacific
SunTech Medical (Shenzhen) Co., Ltd.
105 HuanGuan South Road, Suite 15 2~3/F
DaHe Community Guanlan,
LongHua District, Shenzhen
GuangDong PRC 518110
Tel: +86-755-29588810
+86-755-29588986 (Sales)
+86-755-29588665 (Service)
Fax: +86-755-29588829
19. Radio Frequency Compliance Requirements

This device contains a transmitter module identified by FCC ID: XDULE40-D2 and Industry Canada: IC 8456A-LE4D2. It has been tested and found to comply with the limits for a Class B device. Changes or modifications not expressly approved by the manufacturer could void the user’s authority to operate the equipment. This device complies with Part 15 of the FCC Rules for the United States. Operation is subject to the following two conditions: (1) This device may not cause harmful interference, and (2) this device must accept any interference received, including interference that may cause undesired operation.

This equipment needs special precautions regarding EMC and needs to be installed and put into service according to the EMC information provided in this document. This equipment has been tested and found to comply with the limits for medical devices to IEC60601-1-2: 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.
- Consult the manufacturer or field service technician for help.
- Portable and mobile RF communications equipment can affect Medical Electrical Equipment.

WARNING: Use of accessories, transducers, and cables other than those specified may result in increased emissions or decreased immunity of the Oscar 2.

WARNING: The Oscar 2 should not be used adjacent to or stacked with other equipment. If adjacent or stacked use is necessary, the Oscar 2 should be observed to verify normal operation in the configuration in which it will be used.

20. Electromagnetic Compatibility System Requirements

<table>
<thead>
<tr>
<th>Guidance and manufacturer’s declaration – electromagnetic emissions</th>
</tr>
</thead>
<tbody>
<tr>
<td>The Oscar 2 is intended for use in the electromagnetic environment specified below. The customer or the user of the Oscar 2 should assure that it is used in such an environment.</td>
</tr>
<tr>
<td>Emissions</td>
</tr>
<tr>
<td>RF emissions</td>
</tr>
<tr>
<td>CISPR 11</td>
</tr>
</tbody>
</table>
### Harmonic emissions
IEC 61000-3-2

<table>
<thead>
<tr>
<th>N/A</th>
</tr>
</thead>
<tbody>
<tr>
<td>public low-voltage power supply network that supplies buildings used for domestic purposes.</td>
</tr>
</tbody>
</table>

### Voltage fluctuations/flicker emissions
IEC 61000-3-3

| N/A |

### Guidance and manufacturer’s declaration – electromagnetic immunity

The Oscar 2 is intended for use in the electromagnetic environment specified below. The customer or the user of the Oscar 2 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>Electrostatic discharge (ESD) IEC 61000-4-2</td>
<td>±6 kV contact ±8 kV air</td>
<td>±6 kV contact ±8 kV air</td>
<td>Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 40%.</td>
</tr>
<tr>
<td>Electrical fast transient/burst IEC 61000-4-4</td>
<td>N/A</td>
<td>N/A</td>
<td></td>
</tr>
<tr>
<td>Surge IEC 61000-4-5</td>
<td>N/A</td>
<td>N/A</td>
<td></td>
</tr>
<tr>
<td>Voltage dips, short interruptions and voltage variations on power supply input lines IEC 61000-4-11</td>
<td>N/A</td>
<td>N/A</td>
<td></td>
</tr>
<tr>
<td>Power frequency (50/60 Hz) magnetic field IEC 61000-4-8</td>
<td>3 A/m</td>
<td>3A/m</td>
<td>Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.</td>
</tr>
</tbody>
</table>

In the event of an error, the device will auto-recover within 5 seconds.

<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 IEC 61000-4-6</td>
<td>N/A</td>
<td>N/A</td>
<td>Portable and mobile RF communications equipment should be used no closer to any part of the Oscar 2, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter. Recommended separation distance</td>
</tr>
<tr>
<td>Radiated RF IEC 61000-4-3</td>
<td>3 V/m 80 MHz to 2.5 GHz</td>
<td>3 V/m</td>
<td>Portable and mobile RF communications equipment should be used no closer to any part of the Oscar 2, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter. Recommended separation distance</td>
</tr>
</tbody>
</table>

\[
d = [3.5/V1] \sqrt{P} \quad 80\text{MHz to 800MHz}
\]

\[
d = [3.5/E1] \sqrt{P} \quad 800\text{MHz to 2.5GHz}
\]

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, should be less than the compliance level in each frequency range. Interference may occur in the vicinity of equipment marked with the following symbol:

![Radio wave symbol]

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

a. Field strengths 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 Oscar 2 is used exceeds the applicable RF compliance level above, the Oscar 2 should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as re-orienting or relocating the Oscar 2.

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

### Recommended separation distances between portable and mobile RF communications equipment and the Oscar 2

The Oscar 2 is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the Oscar 2 can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the Oscar 2 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>150 kHz to 80 MHz</td>
<td>80 MHz to 800 MHz</td>
</tr>
<tr>
<td>d = [3.5/V1] \sqrt{P}</td>
<td>d = [3.5/E1] \sqrt{P}</td>
</tr>
<tr>
<td>0.01</td>
<td>0.12</td>
</tr>
<tr>
<td>0.1</td>
<td>0.38</td>
</tr>
<tr>
<td>1</td>
<td>1.2</td>
</tr>
<tr>
<td>10</td>
<td>3.8</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 meters (m) can be estimated using the equation applicable to the frequency of the transmitter, where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer.

**NOTE 1** At 80 MHz and 800 MHz, the separation distance for the higher frequency range applies.

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

---

### 21. Product Disposal

**Device**

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. Please dispose of materials according to local regulations for medical waste.

**Battery Disposal**

The SunTech Oscar 2 contains a small lithium ion battery on a Printed Circuit (PC) board that contain materials which may be hazardous to human health. The battery cannot be easily removed and therefore the Oscar 2 must be disposed of in an environmentally responsible way or returned to SunTech Medical. A prepaid return label can be obtained. Please see our website for more information about our environmental policy at [http://www.suntechmed.com/about-suntech/environmental-policy](http://www.suntechmed.com/about-suntech/environmental-policy).
Cuff
Do not return used cuffs. Used blood pressure cuffs may be contaminated medical waste and should be dealt with in accordance to your local regulations for medical waste.

22. References