BP+Vitals





Spot-check Vital Signs Device User Manual

Changes

This manual is identified as Part number: 80-0067-00. The most recent is available for download from the SunTech Medical website. Should you notice errors or omissions in this manual, please notify us at:

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Welcome to the SunTech CT40

Thank you for choosing the SunTech CT40 for accurate spot-checking of vital signs.

The SunTech CT40 is designed to be simple and efficient to use and the CT40 features:

- mobility for improved and compact efficiency in a cost-effective package
- automatic BP checks
- averaging of multiple BP readings
- optional use of stethoscope for traditional auscultatory BP measurement
- robust memory
- connection to EMR system
- multiple options for pulse oximetry and temperature measurement

SunTech CT40 Description and Operation

The CT40 vital signs device can perform automatic blood pressure, pulse oximetry and body temperature measurements for clinical professionals. For measuring blood pressure, a blood pressure cuff is placed around the patient's non-dominant upper arm. The cuff is inflated automatically, and blood pressure is determined by the oscillometric method-which senses pressure waves in the artery when occluded by pressure in the cuff. Measurement of the frequency of the pressure waves enables heart rate to also be measured. The pulse oximetry function non-invasively measures the patient's percent oxygen saturation of arterial hemoglobin using principles of plethysmography via a SpO₂ sensor placed on the patient's finger. Temperature can be measured using one of two different kinds of temperature probes. The oral/axillary/rectal temperature probe contains a thermistor that generates a voltage based on changes in temperature, and these voltages are recorded by the temperature circuitry. The touchless infrared temperature probe detects radiated infrared energy from the temporal artery. The CT40 is a portable device, approximately 205 x 190 x 140 mm in size and weighs approximately 1440 g without battery. Control buttons allow the user to stop/start a BP measurement, save a set of measurements to memory, change between BP modes, and return to the home screen. There is also a selection knob that is used to scroll and select different device options. The backlit LCD display shows the user device status and measurement information. The device uses a microprocessor with software, which is not accessible to the user. The unit is powered by a single rechargeable lithium-ion battery at the rear of the device. Two USB-A port connections can be used to connect, an optional printer, barcode scanner or Wi-Fi dongle. There is also RJ11 Ethernet port for network connectivity, and a mini-USB port used to connect the device to a PC or laptop for advanced device configuration.

Note: For purposes of this manual, the SunTech CT40 (Model 260) may be referred to as "the SunTech CT40," "the CT40" or "the device".

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1. Safety Considerations

Intended Use

The SunTech CT40 is a clinical grade, automated blood pressure measurement device with optional temperature and pulse oximetry modules for spot-check vital sign measurements in physician offices, long-term care facilities, and low-acuity areas in hospitals. The CT40 can be used in combination with a clinical IT network to transfer and store patient measurement data on an EMR system.

Indications for Use

The SunTech CT40 is intended for use as an aid or adjunct to diagnosis and treatment measuring systolic and diastolic blood pressures, heart rate, temperature and pulse oximetry (Sp02). The system is only for measurement, recording, and display. It makes no specific diagnosis.

The SunTech CT40 is intended to be used on adult and pediatric patients using appropriately sized SunTech Onepiece Durable (OPD) BP cuffs.

The device is intended to be used in outpatient departments, emergency treatment rooms, low-acuity areas of hospitals, community clinics, private clinics, and other medical institutions. It is not intended for helicopter transport, hospital ambulance or home use. It is not intended for near active HF SURGICAL EQUIPMENT and the RF shielded room of an ME SYSTEM for magnetic resonance imaging, where the intensity of EM DISTURBANCES is high.

The CT40 is intended for use by a qualified clinician on adult and pediatric patients. The CT40 will be operated by a nurse, mid-level practitioner, nurse's aide, clinician or physician. The operator is required to have some basic knowledge of the measurement of vital signs.

User Responsibility

Your SunTech CT40 is designed to perform in conformity with the description thereof contained in this operation manual and accompanying labels and inserts, when assembled, operated, maintained and repaired in accordance with the instructions provided.

Further, the user of this 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.

Use of SunTech CT40

Use only One-piece Durable (OPD) BP cuffs supplied by SunTech Medical.

Observe the patient carefully during the measurement. Ensure pressure compatibility to all patients. If any abnormality occurs, either in the unit or the patient, suspend the operation immediately and disconnect the BP cuff, SpO₂ sensor and thermometer (if applicable) from the patient.

Accuracy of any BP reading, or oxygen saturation measurement may be affected by the position of the patient, their physical condition and use outside of the operating instructions detailed in this guide. The interpretation of BP and oxygen saturation measurements should only be made by a physician.

Safety and effectiveness when used with pregnant women, children under 3 years of age and neonates have not been established.

Pulse Oximetry (SpO₂)

ChipOx SpO₂ Module: Use only pulse oximeter (SpO₂) sensors supplied by SunTech Medical or original Nellcor[®] pulse oximeter (SpO₂) sensors supplied by Covidien[®] (except for forehead reflectance sensors).

Masimo® SpO₂ Module: Use only original Masimo® pulse oximeter (SpO₂) sensors and cables.

Check the application site of the SpO₂ sensor frequently to confirm proper positioning of the sensor and to check the circulation and skin sensitivity of the patient.

Wireless Networking

When connecting this device to a wireless network, it is important to use only the hardware specified by SunTech Medical (see Accessories section for details). Unsupported USB accessories, including unsupported Wireless adapters, have been intentionally disabled and will not function with this device.

Possible Adverse Reactions

In the area of the BP cuff or SpO₂ sensor, allergic exanthema (symptomatic eruption) may result, including the formation of urticaria (allergic reaction including raised edematous patches of skin or mucous membrane and intense itching) caused by the fabric material of the cuff or sensor.

Following the application of the BP cuff, petechia formation (a minute reddish or purplish spot containing blood that appears in the skin) or Rumpel-Leede phenomenon (multiple petechia) may appear on the arm, which may lead to idiopathic-thrombocytopenia (spontaneous persistent decrease in the number of platelets, associated with hemorrhagic conditions) or phlebitis (inflammation of a vein).

In the event that the user or patient sustains any serious injury in relation to the device, the incident should be reported to both the manufacturer and the competent authority of the Member State in which the user and/or patient is established. To contact SunTech medical about an incident, please call or email at:

Tel: 800.421.8626 919.654.2300 Email: <u>CustomerSupport@SunTechMed.com</u>

Contraindications:

This system is contraindicated for use in the presence of a Magnetic Resonance Imaging (MRI) device.

Warnings and Cautions



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

WARNING: Pressurization of the cuff can temporarily cause loss of functionality of SpO₂ if simultaneously using device on the same limb.

WARNING: Not designed for neonates.

WARNING: Do not apply the BP cuff to a limb being used for IV infusions as the cuff inflation can temporarily block the infusion, causing harm to the patient.

WARNING: Check frequently by observing the limb that operation of the AUTOMATED SPHYGMOMANOMETER does not result in prolonged impairment of the circulation of the patient.

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: The CT40 is NOT defibrillator protected.

WARNING: Do not use in the presence of flammable anesthetics or other cuff substances in combination with air, oxygen-enriched environments, or nitrous oxide.

WARNING: Do not use the device if it has failed its diagnostics self-test, or if it displays a greater than zero pressure with no BP cuff attached or a value of oxygen saturation with no SpO₂ sensor attached.

WARNING: Do not use if device is dropped and/or damaged. Have a qualified service representative check the unit before using again.

WARNING: Do not remove unit covers. Doing so may cause electrical shock to the user. The device does not contain any user serviceable components.

WARNING: Do not immerse the device in any fluid, place fluids on top, or attempt to clean the unit with any liquid detergents, cleaning agents, or solvents. This may cause an electrical hazard. Refer to the cleaning section of this guide for instructions on cleaning. If any of these situations apply, please contact SunTech Medical.

WARNING: Remove power before servicing device. Failure to remove power could cause electrical shock or death.

WARNING: A pulse oximeter should NOT be used as an apnea device.

WARNING: Pulse rate measurement is based on the optical detection of a peripheral flow pulse and therefore may not detect certain arrhythmias. The pulse oximeter should not be used as a replacement or substitute for ECG based arrhythmia analysis.

WARNING: Do not use the device or any of its accessories during magnetic resonance imaging (MRI) scanning. Induced current could potentially cause burns.

WARNING: Tissue damage can be caused by incorrect application or use of an SpO₂ sensor, for example by wrapping the sensor too tightly. Inspect the sensor site as directed in the sensor Directions for Use to ensure skin integrity and correct positioning and adhesion of the sensor.

WARNING: Do not use high frequency surgical equipment with the CT40 as this may cause loss of stored data

WARNING: No modification of this equipment is allowed.

WARNING: Federal (U.S.) law restricts this device to sale by or on the order of a physician.



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: Check calibration of this device every two years.

CAUTION: Calibration should be done by biomedical technician or other authorized personnel.

CAUTION: Never knowingly use a defective device.

CAUTION: Immediately replace parts that are broken, worn, missing, incomplete, damaged or contaminated.

CAUTION: Contact the nearest SunTech approved service center should repair or replacement become necessary. A list of approved service centers appears in the guide or on our website at www.SunTechMed.com.

CAUTION: The reliability of the device depends upon conformance with the operation and service instructions, as detailed in this manual.

CAUTION: Only replace battery with same type and model number.

CAUTION: To avoid the risk of electrical shock, this equipment must be only connected to supply mains with protective earth.

CAUTION: Do not connect the device to equipment that does not meet EN60601-1. When the device is attached to a patient, the device's communication ports must only be connected to equipment that meets EN60601-1 standard.

CAUTION: Use only SunTech branded cuffs approved for use on the CT40 device.

CAUTION: Use only Masimo[®] oximetry sensors for SpO₂ measurements with the Masimo[®] SpO₂ module. Other oxygen transducers (sensors) may cause improper performance.

CAUTION: Do not machine wash the cuff bladder. Water could be trapped in the cuff and cause damage to the NIBP module and/or inaccurate BP reading.

CAUTION: Do not use damaged SpO_2 or temperature sensors. Do not use a SpO_2 sensor with exposed optical components.

CAUTION: Do not immerse the SpO_2 or temperature sensors in water, solvents, or cleaning solutions (the sensors and connectors are not waterproof). Do not sterilize by irradiation, steam, or ethylene oxide. See the cleaning instructions in the Sensor Directions for Use.

CAUTION: Do not use damaged patient cables. Do not immerse the patient cables in water, solvents, or cleaning solutions (the patient cable connectors are not waterproof). Do not sterilize by irradiation, steam, or ethylene oxide.

CAUTION: Do not position the device so that it is difficult to access and remove the power cord from the electrical supply. The AC power cord is the means of disconnection to the supply mains.

Icons, Symbols and Abbreviations

Icons and Symbols

The following icons and symbols are used in this guide, on the SunTech CT40 equipment and packaging.

Image	Description	Standard/Source
	Warning message	ISO 7010-W001
\triangle	Caution message	ISO 7000-0434A
	Manufacturer	ISO 7000-3082
\sim	Manufacture Date	ISO 7000-2497
~~	Country of Manufacture, Country of Origin	ISO 3166-1
EC REP	Authorized representative in the European Community	ISO 15223-1
MD	Medical Device	Medical Device Regulations
REF	Catalog Number	ISO 7000-2493
SN	Serial Number	ISO 7000-2498
LOT	Batch or Lot Code	ISO 7000-2492

Ţ	Fragile, handle with care	ISO 7000-0621
Ť	Keep dry	ISO 7000-0626
-20	Temperature limit	ISO 7000-0632
	Humidity limitation	ISO 7000-2620
in	Consult instructions for use	ISO 7000-1641
③	Refer to instruction manual/booklet	ISO 7000-M002
¥	Type B Applied Part	IEC 60417-5840
Ŕ	Type BF Applied Part. Part is isolated from earth ground.	IEC 60417-5334
×.	Indicates that separate collection for waste electric and electronic equipment (WEEE) is required.	IEC 60417-6414
CE 0413	CE Mark: Product meets the Medical Device Directive and is CE marked to indicate conformance	EU Directive
SpO ₂	SpO ₂ Sensor. Type BF Applied Part	IEC 60417-5333
€ ~ - - - - - - - - - - - - -	USB-A or USB micro	ISO 7000-3650
	Warning: Federal (U.S.A.) law restricts this device to sale by or on the order of a licensed health care practitioner.	FDA
(((•)))	Non-ionizing electromagnetic radiation	IEC 60417-5140
6	Indicates the arm circumference which is appropriate for the cuff	SunTech Design
	Cuff index marker, OPD	SunTech Design
	Artery marker indicating proper placement – Arrow and symbol should be placed over the brachial artery.	SunTech Design
	Cuff range indication	SunTech Design
	Device is not made with natural rubber latex	SunTech Design

X	Device is not made with PVC	SunTech Design
	Class II Equipment	IEC 60417-5172
IPX1	Protection against vertically falling drops of water	IEC 60529
	Expiration Date	ISO-7000-2607
\otimes	Single Use Only	ISO 7000-1051
#	Model number	IEC 60417-6050

Commonly Used Abbreviations

BP	Blood Pressure	
BPM	Beats Per Minute	
EMR	Electronic Medical Record system	
K-sounds	Korotkoff sounds	
MAP	Mean Arterial Pressure (Not available in the U.S.)	
DIA	Diastolic BP	
NIBP	Non-Invasive Blood Pressure	
OPD	One-piece Durable	
SpO ₂	Percent Oxygen Saturation of Arterial Blood (hemoglobin)	
SYS	Systolic BP	

2. Setting Up the SunTech CT40

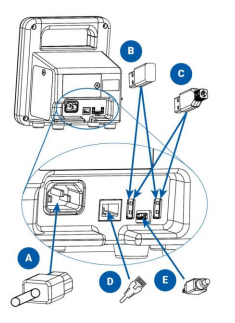
Unpacking the Device

As you unpack your SunTech CT40, check to make sure you have all the proper components.

Refer to the separate packing label stating which components you received based on the options you ordered with your device.

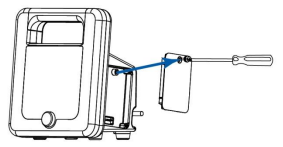
Rear Panel Configuration

SunTech CT40 connections on the back of the device:

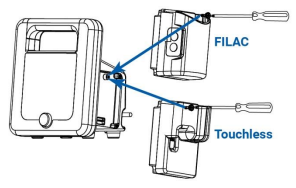


- A. Connect AC power cord
- B. Wi-Fi USB dongle (optional)
- C. Printer or barcode scanner USB cable (optional)
- D. Ethernet cable (optional)
- E. Mini USB cable to connect to PC or laptop

Side/Temperature Panel Configuration



If no temperature module will be added, then attach the right panel of the CT40, using the included Phillips-head screwdriver. This panel will be on the right side, when the screen is facing the user.



To add a temperature module, attach the module on the right side of the main unit. Fasten with the included Phillips-head screwdriver.

For more information regarding normal wear and replacement parts, reference Section 10, Taking Care of SunTech CT40.

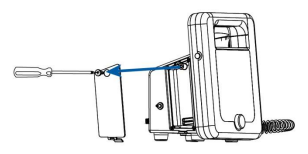
For FILAC[®]:

Temperature unit connector: Insert the probe/well-assembly into the top of the FILAC® module.

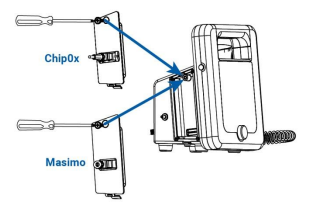
For Touchless:

Plug in the touchless thermometer cable connector into the already inserted module.

Side/SpO₂ Panel Configuration



Remove the left panel of the CT40, using a Phillips head screwdriver. This panel will be on the left side, when the screen is facing the user.



To add an SpO₂ module, attach the module on the left side of the main unit. Fasten with the included Phillips-head screwdriver. For more information regarding normal wear and replacement parts, reference Section 10, Taking Care of SunTech CT40.

For Masimo®:

Attach the Masimo[®] extension cable to the connector on the module, making sure that the connectors lock together. Then, attach the Masimo[®] SpO₂ sensor to the other end of the extension cable making sure that the retaining clip is in place.

For ChipOx:

Attach the ChipOx SpO₂ sensor to the connector on the module while the retention clip is held upwards. Once the sensor/cable connector is inserted, push the clip downward over the connector to hold it securely in the module. For easy insertion, ensure module is parallel to the device.

Install Batteries

Install the rechargeable battery into the battery bay. Please read the battery instructions and the label on its surface before use. Allow 8 to 12 hours for charging. All segments of the Battery Symbol will be lit when the device is fully charged.



Replace only with SunTech part number: 98-0900-00. For more information regarding how to order, see Section 11 for Accessories & Replacement Parts.

If not installed in the device, the battery shall be kept away from heat, fire or other high temperature environments. Keep the battery in a dry place stored at room temperature.

Do not disassemble, attempt to repair or use the battery for any other device or for any other purpose.

Do not place near any metal or use metal to shield the battery from physical damage as this may cause battery overheating and/or a fire risk.

Do not short across the contacts of the battery or attempt to discharge the battery by shorting as a risk of fire or explosion may result.

Do not expose or immerse the battery in water or attempt to clean with any cleaning agents. Only wipe battery with a damp cloth if necessary.

Wash the affected area if electrolyte spills on skin or clothes. Leaking electrolyte may cause discomfort to the skin. If it gets into the eyes, do not rob the eyes. Flush eyes immediately with water and seek medical attention.

Battery Disposal

In addition to the main battery, the SunTech CT40 device contains an additional non-serviceable lithium ion battery which is not removable by the user. Lithium ion batteries contain materials which may be hazardous to human health. Do NOT dispose of the battery or CT40 device in domestic waste! The removable rechargeable battery should be properly recycled 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/aboutsuntech/environmental-policy.

Attach Connections to the Device

After connecting the Power Cable, plug into an available AC power outlet to charge batteries.

Connect the BP cable, SpO₂ sensor cable, and temperature sensor (if option is included) to the device.

Connect the barcode scanner and/or printer to the device if these options are included.

Use the Power Button on the left side of the SunTech CT40 to turn it on.

NOTE: The blue LED around the Selection Knob will be lit whenever the device is powered on.

NOTE: To ensure optimal performance, plug printer into the CT40 device and charge printer battery fully (approx. 11 hours).

Mounting Options

The SunTech CT40 is designed to be used on a tabletop or mounted to a wall or mobile stand. Your device may be delivered with the appropriate mounting kit, depending on the configuration your facility ordered. Please see the separate mounting hardware instructions.

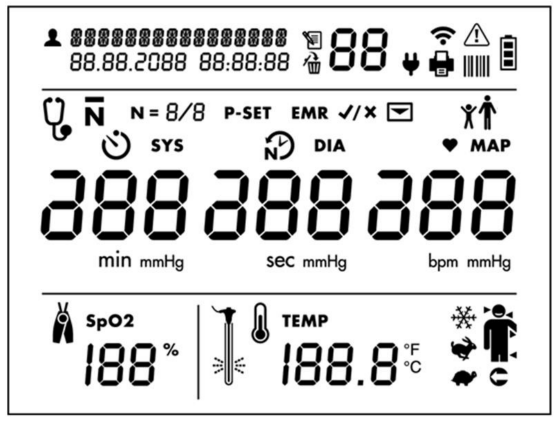
System Configuration

The SunTech CT40 can be used out of the box with the factory settings. Any changes desired to the factory settings can be made using the Advanced Configuration Application. The Advanced Configuration Application can be accessed when the device is connected to a Windows device. Advanced configuration, such as connecting to a network or EMR system, should be done by a qualified technician. See Advanced Configuration Application Guide (SunTech document 80-0072-00-MO) for details.

Advanced device configuration, such as choosing either BP or SpO₂ as the pulse rate source or activating MAP measurement (not available in the U.S.), is also done using the Advanced Configuration Application.

3. Getting to Know the SunTech CT40

Measurement Display



Patient Identification and Date/Time Stamp

L Patient ID

Memory Mode Information

Memory icon

Hemory Delete icon

Connections to Ancillary Devices and Networks

?: Wi-Fi radio on (segments will illuminate in series)

Wi-Fi radio connected (segments will be illuminated continuously)

➡ AC power connected

Printer connected and powered on

Barcode scanner connected and powered on

Error Alert Symbol

WARNING: User must take immediate action (see additional information in Section 12)

Battery Level Indicator

Battery symbol (All bars illuminated indicate a full charge)

Blood Pressure Measurement Mode Selection Icons

C: Auscultatory Sphygmomanometer Mode

N: Averaging mode

N =: Number of measurements taken in averaging mode

P-SET : Maximum pressure setting for Auscultatory Sphyg Mode

👏 : Time before first measurement in minutes (Illuminates when Averaging Mode is selected)

D: Time between measurements in seconds (Illuminates when Averaging Mode is selected)

Blood Pressure Results (DIA and SYS)

DIA and mmHg: Diastolic blood pressure

SYS and mmHg: Systolic blood pressure

EMR Transmission Icons

EMR \checkmark **X**: EMR icon prompts the user to confirm whether to send measurement data as message to EMR system. (Only shown if EMR connectivity has been set up via Configuration Application.)

: Message icon (Indicates if EMR messaging is successful)

If successful, the icon will flash on and off with check mark.

If not successful, it will flash on and off with X.

Adult or Child Patient Selection

Press and hold toggle button for 4 seconds, until selection knob starts flashing. Release toggle button and use the selection knob to select the icon needed. Selected icon will blink after 1 second. Press the selection knob to confirm the selection.

 $m{\Gamma}$: Adult BP Mode icon. This icon is illuminated when the Adult BP mode is selected.

X: Pediatric BP Mode icon. This icon is illuminated when the Pediatric BP mode is selected. The CT40 will remain in the selected patient mode until a new mode is selected.

Heart Rate and MAP Measurements

•: Measured in BPM (Beats per minute), is illuminated when a heart rate is shown in the heart rate display.

MAP: Measured in mmHg is illuminated when a MAP value is shown (only if MAP functionality is enabled). **NOTE:** This is a factory setting. MAP is not available in the U.S.

Pulse Oximetry Measurement Display

5p02 : Expressed in % of arterial oxyhemoglobin

Temperature Measurement Settings and Results Display



TEMP : Body temperature in either Fahrenheit or Celsius (user selectable)

🔆 : Cold Pre-heat mode

🕏 : Predictive mode

🖤 : Direct mode

T: Human thermometry mode. Temporal (arrow on left side of head), Oral (arrow on right side of head), Axillary (arrow under arm), or Rectal (arrow pointing to rectum).

lace : Object thermometry mode. Illuminates when object thermometry mode has been selected.

Apply temp probe cover

Control Panel

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Memory Button

Press this button to enter Memory Mode and to store, view or delete saved measurements.

Save Current Reading: A quick press gives the user an option to save the patients reading, suggested to be done after each reading.

Access Previous Reading: A long press (holding the button down for 4 seconds) gives the user the option to access the bank of prior readings. Please see Section 4 to read more about the saving and accessing saved readings.

Toggle Mode Button

Press this button to toggle between three different BP measurement modes: Single Measurement, Averaging Measurement and Auscultatory SphygMode.

A quick press gives the user an option to switch between BP measurements modes.

A long press (holding the button down for four seconds) gives the user the option to switch between pediatric or adult mode. More information on switching from adult to pediatric mode can be found in Section 5.

Selection Knob

This round knob rotates clockwise and counterclockwise and can be pressed in to select different options on the SunTech CT40. This button can be used to change parameters and to set the intended parameter.

Start/Stop Button

Press this button to start or stop a BP measurement.

Home Button

Pressing this button returns you to the single measurement mode.

Device Setup

Press the power button to start Initial Power-up. The display will gradually illuminate over a 3-second period. If this is the first time the device has been turned on since it left the factory or since the device was set to factory defaults, the user will be prompted to set the time and date in the following format: MM.DD.YYYY HH:MM:SS. The "MM" portion of the field will flash first, prompting you to set the month. Turn the Select Knob to select the appropriate month and press the Selection Knob. Then the next field, "DD" will flash, prompting you to select the day, and so on and so forth, until you have set "SS" for seconds.

After setting the date and time, the device is ready to use! It is possible to make additional configuration changes to the SunTech CT40 using the Advanced Configuration Application.

To Set the Time and Date

The time and date can be changed by simultaneously holding both the Power and Start/ Stop buttons for 6 seconds. Refer to the previous section "Device Setup" for instructions on setting the time and date using the selector knob.

The SunTech CT40 Vital Signs Device also allows for Automatic Time Synchronization via Network (NTP) settings. Refer to the advanced configurations manual for information regarding either of these changes.

Visual Events

When an error occurs, the Warning icon will blink, in addition to the relevant parameter experiencing the error.

Blood pressure error is displaced in the BP heart rate field.

Temperature error will displace in the lower right hand.

SpO₂ error will displace in lower left side of the screen.

Battery symbol will flash if error is related to battery.

Depending on the out-of-range value, the display will indicate whether the value is out-of-range (hi) or out-of-range (lo). If the out-of-range parameter is the systolic or diastolic pressure (or both), and MAP is enabled, MAP is not displayed for that measurement. **NOTE: MAP is not available in the U.S.**

See Section 12, "Status Messages and Events," for more details.

4. Good to Know Before You Begin

Power Modes

Initial Power-up occurs the first time the device is turned on after being received from the factory, after the device has been reset to factory defaults or after the battery has been changed. See "Device Setup" for information about Initial Power-up.

Nominal Power-up refers to every time the device is turned on after Initial Power-up. Simply press the power button and the display will illuminate.

NOTE: The blue LED around the Selection Knob will be lit whenever the device is powered on.

Power-down occurs when you press the power button when the device is powered up. During power-down, all LCD segments and icons illuminate for 1 second and then the entire display gradually darkens from normal brightness to black and then one beep sounds. At this point, the device enters **Sleep Mode**, a low-power state, and is considered off.

Automatic Power-down occurs when the device has not been used (i.e., no measurements taken, or buttons pressed) for 1 hour. The device will automatically execute the Power-down sequence and enter Sleep Mode.

Your SunTech CT40 offers flexibility to function in ways that best meet your work setting.

Documenting Measurements

Before you start taking measurements, it's important to document the patient ID per your facility's procedures. To do so, the barcode scanner must be used to enter the patient ID into the CT40.

With a Barcode Scanner

If your SunTech CT40 is connected to a barcode scanner, you can scan your patient's barcode ID and it will show at the top left of your screen. Then, the patient ID will remain attached to all measurements taken, saved in memory, and/or sent to your facility's EMR.

Without a Barcode Scanner

If you don't require a barcode scanner, all measurement results will be displayed on the main screen and can be saved to the device's memory, written down in a chart, printed out (with optional thermal printer) or captured in another way that meets your facility's documentation procedures.

Printing

The SunTech CT40 can be connected to an optional SunTech thermal printer to easily print out measurement results. See Section 11 for ordering information.

After connecting the optional CT40 thermal printer (see Section 2 and Section 3), ensure that the green LED on the top of the printer is lit and unblinking before printing.



To print the information currently displayed on the CT40 screen, press the print button on the top of the printer (see diagram above).

Saving Measurements

With EMR Connectivity

Technicians may have already connected the SunTech CT40 to your facility's EMR system. If so, you can easily send measurements directly to the EMR.

With the patient's current measurements displayed on the screen, press the Memory button

. The EMR

icon and Check Mark icon EMR 🗸 will begin flashing.

Press the Selection Knob to send the data to the EMR. If the measurements are delivered to the EMR successfully,

you will see the EMR Message icon 🗖 and the Check Mark icon 🗸 blink four times.

To store the displayed measurements in the device memory without sending to the EMR:

Press the Memory Button

While the EMR and Check Icons are flashing, turn the Selection Knob until the X icon *x* is displayed and then press the Selection Knob. The measurements will be saved to the device memory only.

Without EMR

The SunTech CT40 does not have to be connected to an EMR system to save measurements. You can save them to the device's own memory. With the patient's current measurements displayed on the screen:

Press the Memory button

All the measurement values on the screen, the Check icon ✓ and the Memory icon 🖻 will flash on and off 2 times.

The display will then return to a blank screen/shows null values, but the Check Mark icon will flash 2 more times, this verifies the measurements are successfully stored to the device's memory.

Note: The CT40 does not autosave. The user must save the reading to memory after every reading.

To view results stored in memory, press the Memory Button for more than four seconds. You will see the most recently saved measurements displayed on the screen and can turn the Selection Knob counterclockwise to see older measurements and then clockwise to see newer measurements.

TIP: When you send results to memory, make note of the "Memory Location" number to keep track of an individual patient's measurements. Then later, you can match the Memory Locator number with a patient's name if you pull up or print out results for documentation.

The memory will hold up to 99 records. At 100 readings the device will continue to save but will save by overwriting the oldest records.

5. Using SunTech CT40 for BP Measurement

The SunTech CT40 provides flexibility to allow you to measure BP using several different modes, depending on your facility's preferred procedure, the patient's condition or other care considerations.

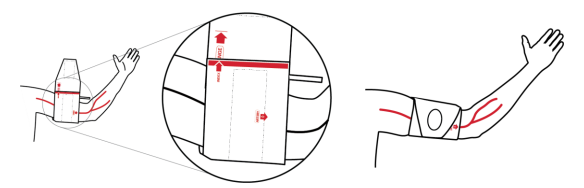
Step 1: Preparing the Patient

Patient Position

According to AHA guidelines, the patient should be seated for at least five minutes before taking a blood pressure reading. The patient should be seated with feet flat on the floor and back supported. The patient's upper arm should be fully supported and resting at heart level. Encourage the patient to relax, and to be still and quiet.

Select and position BP cuff

Selecting the wrong size cuff will produce an inaccurate blood pressure measurement. When wrapped around the patient's arm, the Cuff Index Line should fall within the Range Markers printed on the cuff. If not, select a larger or smaller cuff to ensure optimal BP measurement accuracy. The Artery Marker printed on the cuff must point to the brachial artery, located inside the patient's upper arm, between bicep and tricep. Wrap the cuff snugly around the patient's upper arm, so that the bottom edge of the cuff is approximately 1 inch above elbow joint.



Step 2: Select Between Adult Vs Pediatric Mode

The SunTech CT40 offers two different measurement modes for two very different types of patients: Adults and Pediatric patients (children between the ages of 3 and 12).

To switch between Adult and Pediatric modes:

T: Adult BP Mode icon X: Pediatric BP Mode icon

Hold the Toggle Mode button

down for 4 seconds until the adult and pediatric icons start blinking.

Turn the Selection Knob in either direction to select the desired icon

After making your selection, wait 2 seconds. The selected icon will be the one flashing.

Once you have selected the intended mode, press the Selection Knob to finalize this selection.

Note: Adult and Pediatric mode can be used in conjunction with any type of measurement mode.

Step 3: Select Measurement Mode

The SunTech CT40 offers three different ways to measure BP.

Automatic Single BP: You can take a single, automatic measurement (Single Measurement Mode).

Averaged BP: Automatically take and average up to 5 multiple BP measurements (Averaging Measurement Mode).

Note: This mode is not intended for unsupervised continuous surveillance of patients undergoing anaesthetic procedures as well as emergency care where there can be a clinical need for frequent readings.

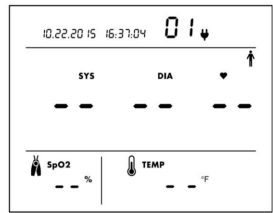
Auscultatory/ Sphyg mode. Allows you to verify an automated oscillometric BP reading with a manual auscultatory measurement (Auscultatory SphygMode).

Note: Use of Auscultatory SphygMode may be required to meet certain clinical trial protocols. It also can be used to verify a BP reading if the automated BP results diverge significantly from a patient's prior history, or if you as the clinician deem it necessary. Auscultatory SphygMode mimics a traditional sphygmomanometer and does not measure Systolic or Diastolic blood pressure.

Note: Averaging Measurement Mode can only be used for Automatic BP Measurements. It cannot be used to automatically average measurements taken in Auscultatory SphygMode.

Single Measurement Mode

This is the device's default mode. The SunTech CT40 will be in this mode each time it is powered up Pressing the Home button will revert to single measurement mode.

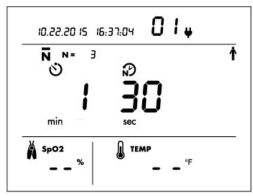


Example of center display when Single Measurement Mode is selected

Note: If you see the Auscultatory SphygMode Icon $\frac{1}{9}$ or the Averaging Mode Icon $\overline{\mathbf{N}}$, press the Mode Toggle Button

in succession until these icons no longer show to return to Single Measurement Mode.

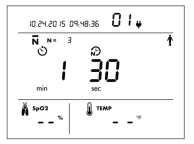
Averaging Measurement Mode



Select Mode

Set # of Measurements to Average

Press the Mode Toggle Button until you see the Averaging Measurement Mode Icon **N**. **N**. From single measurement mode, that will only be one button press. The Number of Measurements Icon **N** = will flash as will the Selection Knob. To keep the default of 3 measurements, press Knob. Or turn Knob to choose a different number and press Knob to select.

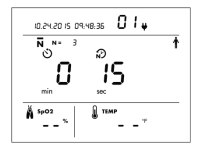


Example of Averaging Measurement Mode setup screen with default values.



You can select 2 to 5 measurements to average*. In this example, 4 measurements are set to be averaged. Set Time Between Measurements

Default values for Time Before First Measurement ightigginary and TimeBetween Measurements <math>
ightigginary willflash in sequence as the user isprompted to set each value. Tokeep defaults, press Knob. Orturn Knob to choose new valuesand press Knob to select.



On left is Time Before First Reading will start. Select 0-5 min. Default = 0. On right is Time Between Measurements. Select 15-120 sec. Default = 15 sec.

*Note: In averaging mode, you are unable to see individual readings. Averaged Results will be displayed at the end of all the readings. By default, all measurements taken (from 2 to 5) will be included in the averaging calculation. Using the Advanced Configuration Application, the device can be programmed to always throw out the first measurement taken and not include it in the averaging calculation.

**Note: All values selected during the Averaging Measurement Mode will be displayed the next time that this mode is used, so that the same customized averaging protocol can be set and used each time that Averaging Measurement Mode is used.

Auscultatory SphygMode



Example shows Auscultatory SphyqMode setup display. Three-digit number is the target cuff inflation pressure setting. The default is 160 mmHg.

until

Select Mode

Press the Mode Toggle Button

you see the Auscultatory SphygMode Icon

From single measurement mode, that will be two button presses.

Choose a Pressure Value

The pressure setting icon **P-SET** and Selection Knob will flash. To keep the default maximum cuff inflation setting of 160 mmHG, press the Knob. Or choose another pressure level between 100-280 mmHG and press the Knob to select. The device will remember the pressure setting and display the same pressure setting the next time SphygMode is used.

Step 4: How to Measure BP in Each Mode

When in the appropriate home screen (see above)

Single Measurement Mode

Ensure values are cleared from display except for SYS value representing current pressure in cuff.

Press Start/Stop Button



Inflate/deflate cycle proceeds automatically until results are displayed.

A long beep will sound to indicate measurement is complete.

Averaging Measurement Mode

When selecting this mode, the

N = will flash to allow the user to decide how many readings they would like averaged.

Select number of readings by rotating the selection knob and push the selection knob to set the number of readings.

The minute number will flash, and the user can select how many minutes until the first reading starts by rotating and pushing the selection knob.

The user can then select how often the readings will occur and select this by rotating the knob and pushing the knob to selection wanted.

Auscultatory SphygMode

Put stethoscope over artery.



Press Start/Stop

tο

inflate cuff automatically. Similar to an analog sphyg gauge, the display shows current pressure in cuff. When cuff reaches max pressure, deflation will begin automatically.

Listen for K-sounds while watching displayed pressure. Note SYS and DIA.



The timer will then count down the seconds until the first reading.

After all measurements are taken and averaged, results will display.

A long beep will sound to indicate measurement is complete.



ID.24.20 IS D9:48:36 82 + V N - 2 I 2 7 min mmHg 868 88 Sec mmHg bpm mmHg

Example of SYS, DIA and HR* after Exar BP measurement in Single all m Measurement Mode. aver

Example of SYS, DIA and HR* after all measurements have been averaged.



Example of SphygMode display with cuff pressure at zero. P-set value is default 160 mmHg.

*Note: Pulse rate source, whether BP or SpO₂, can be selected at any time using the Advanced Configuration Application. The default setting uses BP as HR source. If used outside of the United States, the device also can be configured to measure MAP using the Configuration Application. If set up to measure MAP, the MAP and HR measurements will alternate in 3-second intervals. **NOTE: MAP is not available in the U.S.**

Step 5: Record Results

When the measurement cycle is complete, values will be displayed for systolic and diastolic pressure, plus heart rate if BP is selected as the pulse rate source. * Manually record, print and/or digitally save your measurements according to your healthcare facility's preferred process. To send results to an EMR or to the device memory, follow the steps in Section 4.

Step 6: Prepare for New Patient

After all measurements have been recorded or captured, remove the cuff from the patient and clean it according to your facility's requirements. This disinfecting and storage step applies to BP cuffs as well as other optional accessories whose use will be covered in the following sections.

Pressing the Memory Button will clear all patient and measurement data from the display, in addition to saving the patient and measurement information in memory. It is recommended that the display is cleared before taking readings on a new patient.

6. Using SunTech CT40 for Heart Rate Measurement

There are two ways to measure heart rate with the SunTech CT40.

BP Heart Rate Measurement

With this method, HR is captured automatically during BP measurement.

SpO₂ Heart Rate Measurement

If you have the optional pulse oximetry module on your device, you can measure HR through the SpO_2 sensor. See next section (Section 7) for more on use of this option.

The SunTech CT40 uses BP as the default HR source. HR source can be selected using the Advanced Configuration Application.

7. Using SunTech CT40 for Pulse Oximetry

An optional SpO₂ module allows you to measure the oxygen saturation of arterial blood flow and display the reading on the SunTech CT40. There are two different types of optional SpO₂ modules: Masimo[®] SET[®] and ChipOx.

General Principle of Operation:

The pulse oximeter directs red and infrared light into a capillary bed and measures the change in light absorption during the pulsatile cycle. Red and infrared light-emitting diodes (LEDs) in oximetry sensors serve as the light sources, and a photodiode serves as the photo detector.

Traditional pulse oximetry assumes that all pulsations in the light absorbance signal are caused by oscillations in the arterial blood volume.

The traditional pulse oximeter calculates the ratio of pulsatile absorbance (AC) to the mean absorbance (DC) at each of two wavelengths (typically one visible, the other infrared). The oximeter then calculates the ratio of these two arterial absorbance signals. This value is used to find the saturation SpO_2 in a standard table built into the oximeter software. The values in the table are based upon human blood studies against a laboratory co-oximeter on healthy adult volunteers in induced hypoxia studies. The resulting value is reported as oxygen saturation level (%).



WARNING: Pulse oximeter probes and sensors are designed for use with specific pulse oximeter devices. Only use probes and extension/adapter cables that are specified for each SpO₂ module.

WARNING: The responsible organization and/or operator must verify the compatibility of the oximeter, oximeter probe, and extension/adapter cable before use. Otherwise, patient injury can result.

WARNING: Misapplication of a pulse oximeter probe with excessive pressure for prolonged periods can induce pressure injury.



CAUTION: Inaccurate measurements may be caused by:

Incorrect sensor application or use. Proper sensor uses, and placement is critical for good performance.

Significant levels of dysfunctional hemoglobin. (e.g., carboxyhemoglobin or methemoglobin)

Intravascular dyes such as indocyanine green or methylene blue.

Dyes or any substance containing dyes that affect light absorption may cause erroneous readings.

• Some nail polish colors (particularly dark shades) or artificial fingernails may reduce light transmission and affect pulse oximetry accuracy. Remove any nail polish or artificial fingernails before using the SpO₂ sensor.

Exposure to excessive illumination, such as surgical lamps (especially ones with a xenon light source), bilirubin lamps, fluorescent lights, infrared heating lamps, or direct sunlight (exposure to excessive illumination can be corrected by covering the sensor with a dark or opaque material)

Excessive patient movement.

• Abnormal pulse rhythm.

Placement of a sensor on an extremity with a blood pressure cuff, arterial catheter, or intravascular line. Always place the SpO2 sensor on the arm without a blood pressure cuff, arterial catheter, or intravascular line.

CAUTION: Pulse rate measurement is based on the optical detection of a peripheral flow pulse and therefore may not detect certain arrhythmias. The pulse oximeter should not be used as a replacement or substitute for ECG based arrhythmia analysis.

CAUTION: SpO₂ is empirically calibrated to functional arterial oxygen saturation in healthy adult volunteers with normal levels of carboxyhemoglobin (COHb) and methemoglobin (MetHb). A pulse oximeter cannot measure elevated levels of COHb or MetHb. Increases in either COHb or MetHb will affect the accuracy of the SpO₂ measurement.

CAUTION: COHb levels above normal tend to increase the level of SpO₂. The level of increase is approximately equal to the amount of COHb that is present.

CAUTION: High levels of COHb may occur with a seemingly normal SpO2. When elevated levels of COHb are suspected, laboratory analysis (CO-Oximetry) of a blood sample should be performed.

CAUTION: For increased MetHb, the SpO_2 may be decreased by levels of MetHb of up to approximately 10% to 15%. At higher levels of MetHb, the SpO_2 may tend to read in the low to mid 80s. When elevated levels of MetHb are suspected, laboratory analysis (CO-Oximetry) of a blood sample should be performed.

CAUTION: Venous congestion may cause under reading of actual arterial oxygen saturation. Therefore, assure proper venous outflow from monitored site. Sensor should not be below heart level (e.g. sensor on hand of a patient in a bed with arm dangling to the floor).

CAUTION: Venous pulsations may cause erroneous low readings (e.g. tricuspid value regurgitation).

CAUTION: The pulsations from intra-aortic balloon support can be additive to the pulse rate on the oximeter pulse rate display. Be sure to verify patient's pulse rate.

CAUTION: Elevated levels of Total Bilirubin may lead to inaccurate SpO₂, measurements.

CAUTION: With very low perfusion at the monitored site, the readings may read lower than core arterial oxygen saturation.

CAUTION: Do not immerse the sensor or patient cable in water or, solvents, or cleaning solutions (The sensors and connectors are not waterproof).

CAUTION: Loss of pulse signal can occur in any of the following situations:

The sensor is too tight.

There is excessive illumination from light sources such as a surgical lamp, a bilirubin lamp, or sunlight.

A blood pressure cuff is inflated on the same extremity as the one with a SpO₂ sensor attached.

The patient has hypotension, severe vasoconstriction, severe anemia, or hypothermia.

There is arterial occlusion proximal to the sensor.

The patient is in cardiac arrest or is in shock.

NOTE: See SP02 Sensor Specifications Table on page 45, the range of the peak wavelengths and maximum optical output power of the light emitted by the pulse oximeter probe, which can be especially useful to clinicians.

NOTE: This device does not include an alarm system that detects an SpO2 or pulse rate physiological alarm condition. This device is not designed for long-term monitoring and should only be used for spot-check measurements.

System Description:

The SunTech CT40 spot check device with pulse oximetry module consists of a base CT40 device for the display of Sp02 measurement data and other user information, either a ChipOx or Masimo SET Sp02 module which is attached to the CT40 base unit as described in Section 2, and the specific reusable pulse oximeter probe and/or cable that comes packaged with each module. Device setup is described in more detail in **Section 2**. Both the Masimo[®] SET and ChipOx Sp0₂ modules are calibrated to display functional oxygen saturation (Sp0₂).

Taking SpO₂ Measurements:

1. Insert a finger (preferably the index, middle or ring finger) into the SpO2 sensor until the end of the finger reaches the finger stop. Do not use the thumb.

2. The fingernail should face the side with the red light. Make sure that long fingernails do not interfere with proper finger position

3. An SpO₂ reading will be displayed after a few seconds. Remove the finger from the SpO₂ sensor, and the last measurement will be displayed and flashed.

4. If the sensor is not removed from the finger, SpO₂ readings will be automatically stopped after 10 minutes of continuous measurement, and the last measurement will be displayed and flashed.

5. To start a new measurement, re-apply the SpO₂ sensor to the finger as described in Step 1.

NOTE: If the signal quality from the sensor drops below 90%, no measurement will be displayed in the SpO₂ section

SpO2, and the percent symbol % will begin flashing. Check sensor for correct placement and usage.

Special Notes for Masimo® SET SpO₂ module:

NOTE: Possession or purchase of this device does not convey any express or implied license to use the device with unauthorized sensors or cables which would, alone or in combination with this device, fall within the scope of one or more of the patents relating to this device.

NOTE: If the accuracy of any measurement does not seem reasonable, first check the patient's vital signs by alternate means and the check the MS board pulse oximeter for proper functioning.

NOTE: A functional tester cannot be used to assess the accuracy of a Masimo pulse oximeter sensor or pulse oximeter. Using Clinical Dynamics SmartSat SpO2 simulator, the device can display the entire range of SpO2 values.

Masimo® Sensors:

Before use, carefully read the M-LNCS[™] DCI/DCIP sensor directions for use, which includes cleaning and disinfection instructions.

Masimo[®] M-LNCS[™] sensors are provided non-sterile.

Additional Masimo® Information:

In addition to the General Principles of Operation mentioned above, the Masimo[®] SET[®] MS board pulse oximeter uses the following principle:

The MS board decomposes the ratio of pulsatile absorbance (AC) to the mean absorbance (DC) at each of two wavelengths, 660 nm and 905 nm (S(660) and S(905)) into an arterial signal plus a noise component and calculates the ratio of the arterial signals without the noise:

S(660) = S1 + N1

S(905) = S2 + N2

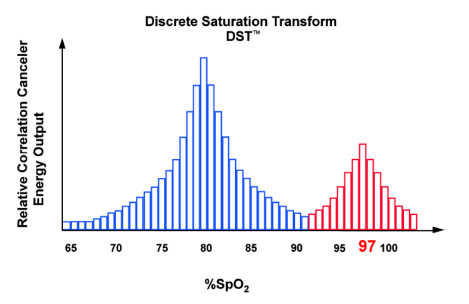
R = S1/S2

Again, R is the ratio of two arterial pulse-added absorbance signals and its value is used to find the saturation SpO_2 in an empirically derived equation into the oximeter's software. The values in the empirically derived equation are based upon human blood studies against a laboratory co-oximeter on healthy adult volunteers in induced hypoxia studies.

The above equations are combined and a noise reference (N') is determined:

N' = S(660) - S(905) x R

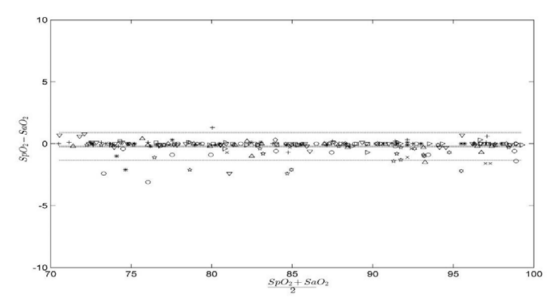
If there is no noise N' = 0: then S(660) = S(905) x R which is the same relationship for the traditional pulse oximeter. The equation for the noise reference is based on the value of R, the value being sought to determine the SpO₂. The MS board software sweeps through possible values of R that correspond to SpO₂ values between 1% and 100% and generates an N' value for each of these R-values. The S(660) and S(905) signals are processed with each possible N' noise reference through an adaptive correlation canceler (ACC) which yields an output power for each possible value of R (i.e., each possible SpO₂ from 1% to 100%). The result is a Discrete Saturation Transform (DST^M) plot of relative output power versus possible SpO₂ value as shown in the following figure where R corresponds to SpO₂ = 97%:



The DST plot has two peaks: the peak corresponding to the higher saturation is selected as the SpO_2 value. This entire sequence is repeated once every two seconds on the most recent four seconds of raw data. The MS board SpO_2 therefore corresponds to a running average of arterial hemoglobin saturation that is updated every two seconds.

Masimo[®] Graphical Plot:

Below is a graphical plot of discreet A_{RMS} values measured with Masimo[®] SET Oximetry in a clinical study using DCI/DCIP pulse oximetry sensors:



The tested device meets the stated accuracy specification of RMS Error = 2.00% for the range 70% to 100% SaO₂. The accuracy is not specified below 70% SaO₂. In addition, here are detailed measured A_{RMS} values over the specified discreet ranges:

Range	ARMS
90-100%	0.60%
80-90%	0.54%
70-80%	0.67%

Special Notes for ChipOx SpO₂ module:

NOTE: Use only Sp02 sensors from SunTech approved for use on the CT40 ChipOx Sp02 module.

NOTE: A functional tester cannot be used to assess the accuracy of a ChipOx pulse oximeter sensor or pulse oximeter. Using FLUKE Biomedical Index 2 SpO₂ simulator, the device can display the entire range of SpO₂ values.

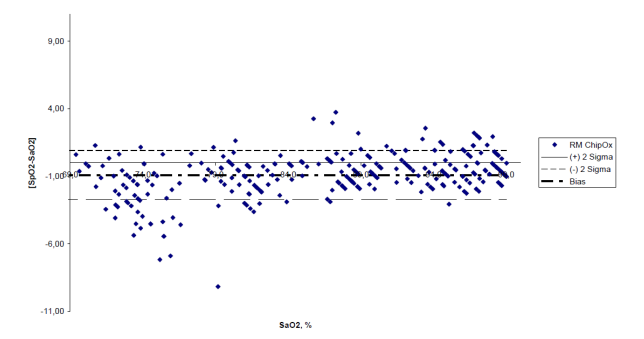
ChipOx Sensors:

Before using, carefully read the ChipOx sensor directions for use, which includes cleaning and disinfection instructions.

ChipOx sensors are provided non-sterile.

ChipOx Graphical Plot:

Below is a graphical plot of discreet A_{RMS} values measured with ChipOx Oximetry in a clinical study using ChipOx pulse oximetry sensors:



The tested device meets the stated accuracy specification of RMS Error = 1.89% for the range 70% to 100% SaO₂. The accuracy is not specified below 70% SaO₂. In addition, here are detailed measured A_{RMS} values over the specified discreet ranges:

Range	ARMS
90-100%	1.54%
80-90%	2.10%
70-80%	2.56%

8. Using SunTech CT40 for Temperature Measurement

The SunTech CT40 has several options for measuring a patient's body temperature.

Covidien® FILAC® 3000 Thermometry Module

Your device may come equipped with a Covidien[®] FILAC[®] 3000 digital thermometer capable of taking an oral, axillary (under arm) or rectal temperature reading. Your thermometer should come with a blue isolation chamber/probe unit for taking oral and axillary temperature or with a red isolation chamber/probe unit for taking rectal temperature.

NOTE: Always apply a new probe cover before taking temperature. Do not reuse probe cover. Use only probe covers designed for use with FILAC[®] 3000 probes. Using the incorrect probe cover can severely damage the probe and cause measurement inaccuracies.

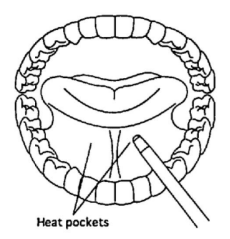
NOTE: To change between Celsius and Fahrenheit, press the °C/°F button on the FILAC® 3000 temperature module. Press again as needed.

NOTE: After removing the probe from the isolation chamber, the Install/Remove Probe Cover icon $\text{Probe}^{\text{remove}}$ will flash to remind you to apply a new probe cover. A new probe cover must be applied to take a measurement.

After you withdraw the probe from the isolation chamber and apply a probe cover, the thermometer begins working automatically. You will see the Human Thermometry Mode icon illuminated on your display with the appropriate arrow illuminated (i.e., pointing to the mouth on right side of head for oral temperature taking). Press the Site button on your thermometer unit to switch between Oral and Axillary mode.

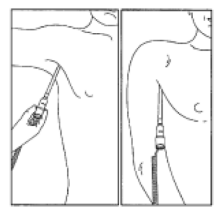
Oral Temperature Taking

Insert the probe tip under the tongue on one side or the other. Ask the patient to close his mouth. Hold the probe in place until there is a long beep and the temperature reading displays.



Axillary Temperature Taking

With the patient's arm uplifted, place the probe tip into the patient's armpit, directly on the skin. Ask the patient to lower his arm and hold still. Hold the probe perpendicular to the arm until there is a long beep and the temperature reading displays.



Rectal Temperature Taking

Apply lubricant to the probe cover and insert it gently into the patient's rectum only one-half inch to three-fourth inch (12 mm to 19 mm) for adults or one-fourth to one-half inch (6 mm to 13 mm) for children. Hold the probe still until there is a long beep and the temperature reading displays.

NOTE: If the temp probe is returned to the probe well before the "long beep" is heard, no temperature measurement will be displayed.

NOTE: Be sure to eject the used probe cover into an approved bio-hazard container before attempting to place the probe back in the isolation chamber.

Other FILAC® 3000 Temperature Settings

Your device's FILAC® 3000 thermometer operates in several different modes.

Quick Mode is an oral predictive measurement mode that provides a fast reading. Quick Mode allows you to rapidly identify patients with normal body temperatures. If the patient temperature is outside of the "normal" range, the FILAC[®] 3000 electronic thermometer will automatically switch into its standard predictive mode to provide a

more accurate reading. Quick Mode is indicated by a rabbit icon on the display 👻 . Quick Mode is not available when in Cold Mode or in Direct Mode.

The FILAC[®] 3000 thermometer normally operates in predictive mode to provide fast and accurate temperature measurements. However, in instances when no measurement site is detected, or the temperature does not stabilize, the thermometer will automatically switch to Direct Mode \clubsuit . Additionally, the FILAC[®] 3000 electronic thermometer will automatically switch into Direct Mode if the ambient temperature is greater than 35°C (95°F). If this happens, you will hear two short beeps and see the Direct Mode turtle icon appear \clubsuit . This icon will be continuously displayed whenever the thermometer is functioning in Direct Mode. The Direct Mode auto feature is always functional for both the Red or Blue isolation chamber/probe combinations.

Cold Mode can be selected if a patient's body temperature is expected to be lower than normal, such as when he is coming out of surgery. Cold Mode is activated by pressing the Site Selection button and °C/°F button

simultaneously on the temp module. When selected, the Cold Mode snowflake icon 🔆 is displayed, and the probe will preheat to 33°C (91°F). The accuracy and measurement time of Cold Mode measurements are equivalent to standard prediction measurements at the respective body sites.

Touchless Thermometry Module*

If your device comes with the optional touchless thermometry module, you can easily and quickly measure the patient's temperature using a touchless infrared temporal thermometer. To use, remove the thermometer from the holster on the side of the SunTech CT40. Touch the thermometer's START button to turn on the touchless thermometer.

On your SunTech CT40 display, you will see the Human Thermometry Mode icon selected with the arrow pointing

to the side of the head . Hold the thermometer probe 2 cm to 3 cm from the patient's temple and press the START button on the back of the thermometer. The reading is complete when you hear a beep, and the temperature measurement will appear on your device display. Place the thermometer back in the device holster.

NOTE: Be sure to wipe away any sweat from the area around the temporal artery and be sure to remove eyeglasses and push hair away from measurement site before initiating a measurement.

NOTE: The area just behind the ear lobe can be used as an alternative measurement site to the temporal artery.

NOTE: The SunTech Touchless Thermometer should not be used near sources of heat or cold, such as a heater or air conditioner vent. Also, do not use outdoors. Use this product in a room where the temperature is between 16° C and 40° C (61° F and 104° F).

NOTE: If patients are subject to a significant change in temperature from outside to inside, the patient should be allowed to acclimate for approximately 15 minutes before using the touchless thermometer.

NOTE: IR temperature readings provided by this device are closely aligned to core temperatures rather than an axillary or rectal reading. Take this into account when reviewing readings.

To change the measurement units from °C to °F or vice-versa, ensure that the thermometer is on by pressing the START button. Then press the MODE and MEM buttons on the thermometer simultaneously for 3-4 seconds.

The SunTech Touchless Thermometer can also be used to measure the temperature of an object, such as a vessel containing liquid, or food. To use Object Mode, ensure that the thermometer is turned on by pressing the START

button. Press the Mode Button briefly. The Object Mode icon \checkmark will then appear in the device display. Place the thermometer 2-3 cm from the desired measuring point on the object and press the START button. Hold thermometer in place until you hear a long beep. The temperature will be displayed on the screen.

NOTE: Remember that surface temperature and inner temperature of an object can vary greatly.

*Touchless Thermometry module is not available in all countries. Contact SunTech to find out if this feature is available in your area

9. Using SunTech CT40 EMR and Memory Functions

Section 4 "Good to Know Before You Begin" provides some basic information about the SunTech CT40's EMR and memory functions. Here are more details about how these functions work.

Barcode Scanner Setup

If your scanner was purchased from SunTech Medical Inc, they are provided, programmed for use with the CT40. If the scanner is not working with the CT40, scan the barcode below with the scanner plugged into any powered USB.



If your scanner was not purchased from SunTech Medical Inc, refer to the user manual of your scanner to ensure the scanner is programmed to provide a carriage return at the end of a reading.

Memory Mode

When you power up the unit, the memory identifier appears at the top of the display beside the Memory Icon E. This is the number that will be associated with the next set of measurement results that are sent to the device memory.



Press the Memory Button **Will** on the front of the device to save results to memory. The Memory Icon, check mark icon and all values that are displayed on the screen (at the time the button is pressed) will flash before the results are saved. The following values can be saved to memory.

Parameter Name	Field Type
Patient ID	Alphanumeric Text
Time Stamp	Numeric, HH:MM:SS
Date Stamp	Numeric, MM.DD.YYYY or DD.MM.YYYY
Memory Location	Numeric
Systolic BP	Numeric
Diastolic BP	Numeric
ВР Туре	Text: AVG or SM
Pulse Rate	Numeric
Pulse Rate Source	Text: BP or SpO ₂
Mean Arterial Pressure	Numeric
SpO ₂	Numeric
Temperature	Numeric
Temperature U/M	Text: C or F
Temperature Site	Text: ORL, AXL, RCT, FHD

NOTE: If a parameter has no data showing at the time the memory button is pressed, then all values for that parameter will be stored as 'null'.

To recall measurements from memory:

Press and hold the Memory Button for four seconds on the front of the device.

You will then be able to use the Selection knob to scroll through prior readings

Press the Memory Button or the Home Button to exit memory mode.

Clearing Measurements from Memory

To clear a single measurement from memory:

Press the Selection Knob while in memory mode. The Memory Delete icon $\stackrel{\checkmark}{10}$ and the Check icon $\stackrel{\checkmark}{10}$ will flash, along with the Memory Identifier number.

Ensure you are viewing the measurement you wish to delete.

Press the Selection Knob to delete that measurement. Or rotate the knob clockwise until the X icon X flashes and press the Selection Knob to cancel the deletion.

To clear all measurement values from memory:

While in Memory Mode, press and hold the Memory Button for less than three seconds, and while holding, then press the Selection Knob.

The Memory Delete icon and Check icon will flash in sync with the Selection Knob. All other values on the screen will be blank.

Press the Selection Knob again to delete all measurements or rotate the knob clockwise until the X icon X flashes and press the Selection Knob to cancel the deletion.

EMR Transmissions

Section 4 covers the basics of sending measurements to EMR. Here are more details.

Validating Patient ID with EMR

Scan the patient's barcode ID with your barcode scanner. If your SunTech CT40 is connected to an EMR, a verification query will automatically be sent to the EMR to confirm the validity of the patient ID. If the EMR

responds that the patient ID is valid, then the patient ID icon, field values and Check Icon 💙 will flash four times.

If the EMR responds that the Patient ID is invalid, then the Patient icon field, field values and X icon will flash six times and a message will appear in the Patient ID field indicating that the Patient ID entered is not valid.

NOTE: The Patient ID must be validated by the EMR before any vital signs measurements can be sent to the EMR.

Sending Measurements to EMR

Sending measurements to an EMR will work only if your SunTech CT40 has been configured to connect to an EMR.

After scanning a valid Patient ID, and taking vital signs measurements, press the Memory Button

EMR icon **EMR** and Check Mark Icon \checkmark will start flashing. Press the Selection Knob to send the data to the

EMR. If the measurements are delivered to the EMR successfully, you will see the EMR Message Icon 🖾 and the

Check Mark Icon ✓ flash four times.

If the EMR has a problem receiving the measurements, the EMR Message icon and X icon x will flash eight times and you will hear 4 short beeps. You will see a message indicating the nature of the problem, such as:

"Rejected" or "Rejected for an error." This message occurs when the device is connected to the network but the EMR rejects the message. A corresponding message will appear in the Patient ID field during a three-second

period when the Message Icon \square and X icon X flash on and off.

"No response (time out)." This message occurs when the device is connected to the network but the EMR is not responding. It also can occur if network connectivity is lost. If there is a network connection, the SunTech CT40 will continue to try to transmit the message every 10 seconds. A message in the Patient ID field will alert you that another attempt to send the message will be made in 10 seconds. If network connectivity is lost, you will see a corresponding message in the Patient ID field, and the device will continue trying to reestablish a connection and send the measurements at intervals set during EMR setup. If problems persist, you may wish to contact your facility's information technology department. As a backup, the patient's measurements are saved in the device memory when the memory button was pressed and can be retrieved using the procedure detailed in the beginning of Section 9.

Note: After the memory button is pressed, all values to be stored to memory will flash off and on twice, along with the check mark icon, memory icon and memory location identifier. After flashing, the display will revert to null values for the patient ID and all measurement data, and the check mark icon will flash off and on twice more.

10. Taking Care of SunTech CT40

Cleaning

CAUTION: The SunTech CT40 and patient hose is not sterilizable. Do not immerse the device or hose in any fluid or attempt to clean with any liquid detergents, cleaning agents, or solvents.

For the device, dampen a soft cloth with mild medical grade disinfectant and wipe the device to remove surface dust and dirt. Dry surface thoroughly before use.

For patient hose, wipe down with mild disinfectant (Clorox, Dispatch, Quaternary Disinfectant, Sani-Wipes, etc.) to remove surface dust and dirt. Dry thoroughly before use.

Cuffs

 \angle **CAUTION**: Do not machine wash the cuff bladder. Water could be trapped in the cuff and cause damage to the NIBP module and/or inaccurate BP reading.

Cleaning Manual Process

- 1. Use a fresh solution of STERIS® Prolystica® 2X Concentrate Neutral Detergent with warm tap water (50-60°C).
- Immerse the cuff in the solution and hand wash and wipe the shell with a lint-free cloth for 1 minute per side to remove debris, ensuring liquid does not enter tube connector. A soft bristle brush may be used to remove visible soil.
- 3. Rinse sleeve with warm water until all cleaning solution is removed, pat dry, and line dry.

Disinfection Process

NOTE: Before disinfecting, ensure that no liquid enters tubing by using a plug or taping off.

- 1. Spray cuff with Quaternary Ammonium until soaked.
- 2. Leave to soak for at least 10 minutes.
- 3. Rinse with distilled water, ensuring liquid does not enter tube connector, and line dry.

SpO₂ Sensors and Thermometers

CAUTION: Never immerse sensors, clips or thermometers in fluids. Do not pour or spray any liquids onto the sensor or thermometers. Caustic or abrasive cleaners will cause permanent damage. Do not open the case of the SpO₂ sensor finger clip sensor more than 45° or the case will be damaged.

Clean the SpO₂ sensor and thermometers with a soft cloth dampened with a mild medical grade disinfectant or isopropyl alcohol. Remove all tape residues. Allow the sensor and thermometer to dry thoroughly before reusing.

Preventative Maintenance

System Self Checks

The SunTech CT40 performs a range of system checks during normal operation. If the device detects a problem, it will display an error code.

Replaceable Parts

On a routine basis, inspect the device, cuffs, hoses, and sensor wires for cracks, fraying, or kinks. Immediately replace any damaged part. Below is a list of parts that may wear and require replacement with normal use.

- ChipOx SpO2 Sensor
- Masimo Sp02 Sensor
- Masimo SpO2 Patient Cable
- HuBDIC Temperature Sensor

Refer to the list of Accessories & Replacement Parts in Section 11 for purchasing replacements and below for proper product disposal. For more information on how to replace the part, refer back to Section 2, Setting up SunTech CT40.

Replacing and Disposal of the Battery

CAUTION: Fire, explosion and severe burn hazard. Replace only with SunTech part number: 98-0900-00

When the battery no longer charges, or it needs excessive recharging, it may need to be replaced. See the "Installing Battery" section above for proper installation and precautions to be taken when installing the rechargeable battery. Please dispose of the old battery per the instructions below.

Note: The expected lifetime of the Lithium-Ion Battery is approximately 2.7 years.

Battery Disposal

X

The SunTech CT40 device contains a lithium ion battery that contains materials which may be hazardous to human health. Do NOT dispose of battery in domestic waste! Instead, please dispose of in an environmentally responsible way, or return the battery 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

Product Disposal

X

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).

Cuff Disposal



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.

Routine Calibration

Have the SunTech CT40 blood pressure function checked every two years to verify the accuracy of the pressure transducers and indicators. Please have this done by a biomedical technician or SunTech Customer Service.

CAUTION: Calibration should be done by a biomedical technician or other person familiar with the SunTech CT40 device.

If a biomedical technician performs this function, they should contact SunTech Medical for instructions to access "Verify Calibration". Instructions are also available in the SunTech CT40 service manual (SunTech Part 80-0068-XX-SM).

For customers in the Americas:

SunTech Medical, Inc. Service Department 507 Airport Boulevard, Suite 117 Morrisville, NC 27560 USA Tel: 800.421.8626 919.654.2300 Fax: 919.654.2301

11. Accessories, Components, & Replacement Parts

Contact your SunTech Medical sales representative to purchase the following items:

Main Unit

Part Number	Description	Details
91-0028-16	10'/3.0m Patient BP Hose, Bayonet to Bayonet	Each
98-0600-41	OPD Child (12-19cm) BP Cuff, Bayonet	Box 5
98-0600-43	OPD Small Adult (17-25cm) BP Cuff, Bayonet	Box 5
98-0600-45	OPD Adult (23-33cm) BP Cuff, Bayonet	Box 5
98-0600-4A	OPD Adult Plus (28-40cm) BP Cuff, Bayonet	Box 5
98-0600-47	OPD Large Adult (31-40cm) BP Cuff, Bayonet	Box 5
98-0600-4C	OPD Large Adult Plus (40-55cm) BP Cuff, Bayonet	Box 5
98-0600-49	OPD Thigh (38-50cm) BP Cuff, Bayonet	Box 5
91-0003-00	AC Power Cord, Americas	Each
91-0003-05	AC Power Cord, Europe	Each
91-0003-06	AC Power Cord, UK	Each
91-0100-02	Mini-USB to USB Cable	Each

98-0900-00 Battery, 18650 Li-ion pack, 7.26V 6.6Ah, Rechargeable Ea

Masimo[®] SET SpO₂ Module

Part Number	Description	Details
(*)	Masimo® M-LNCS DCI Adult SpO2 Reusable Sensor, 2501	Each
(*)	Masimo® M-LNC 10'/3.0m Patient Cable, 2525	Each

(*) contact your local Masimo® distributor to purchase)

ChipOx (Nellcor[®] Compatible) SpO₂ Module

Part Number	Description	Details
52-0010-00	Reusable Adult SpO ₂ Finger Sensor, ChipOx	Each

Covidien® FILAC® 3000 Temperature Module

Part Number	Description	Details
52-0009-00	F3000 Oral/Axillary Temp Probe, 9'	Each
45-0006-00	F3000 Oral/Axillary Isolation Chamber, Blue	Each
52-0009-01	F3000 Rectal Temp Probe, 9'	Each
45-0006-01	F3000 Rectal Isolation Chamber, Red	Each
98-0131-01	F3000 Disposable Temp Probe Covers (25 boxes/tray, 20 covers/box)	1 tray
98-0130-01	F3000 Disposable Temp Probe Covers (25 boxes/tray, 20 covers/box)	10 trays

Touchless IR Temperature Module

Part Number	Description	Details
98-0412-00	Replacement IR Thermometer with cable	Each

Stands/Printer/Scanner

Part Number	Description	Details
46-0040-00	Deluxe Mobile Stand	Each
46-0040-01	Mobile Stand Printer Mount	Each
98-0256-00	CT40 Thermal Printer (w/USB, paper)	Each
45-0005-00	CT40/50 WIFI Dual Band USB Dongle	Each

71-0543-00	Thermal Printer Paper	Each
98-0257-03	Barcode Scanner, 1D	Each
98-0257-04	Barcode Scanner, 2D	Each

Alternate Functional Scanners

*These accessories are tested to work with the SunTech CT40, but are not sold or supported by SunTech Medical Inc. To work with the SunTech CT40, these barcode scanners must be programmed to provide a carriage return suffix at the end of a scan.

Model Number	Description	Details
DS4308HC	Zebra, 2D Medical Grade Barcode Scanner	Each
HS-1M	Jadak, 2D Medical Grade Barcode Scanner	Each
FG2100	Farsun, 1D Point of Sale Grade Barcode Scanner	Each
CM-880	Canmax, 1D Point of Sale Grade Barcode Scanner ¹	Each
1900HHD-5-INT	Honeywell, 2D Medical Grade Scanner	Each
1 Canmay has taken this r	product and of life and recommends CM-890 as a replacement	

1. Canmax has taken this product end of life and recommends CM-890 as a replacement.

12. Status Messages & Events

If the SunTech CT40 has a problem taking a measurement, you may see a code on the display screen.

The *icon* will blink with the respective measurement display result; (i.e blood pressure, SpO₂, Temperature), while displaying the code.

Below are common error codes. Take action as directed on the screen, or as suggested in the table below.

Status Messages

Code Displayed	Action User Should Take
	Blood Pressure:
1-2, 4, 87 & 88	Check the cuff is in the correct position.
	Check that the cuff is properly tightened, and tubing properly connected.
	Check that there is no excessive clothing between the arm and cuff.
	Check that the cuff applied is the correct size.

	The patient may have been moving too much.
	Take another BP reading.
3	No action possible. Values are outside the reportable range.
85 & 89	Check that the hose has no sharp bends and is not pinched.
	Check that the patient is not lying on the cuff.
	Check that the cuff is in the correct position.
	Take another BP reading.
86	If cancelled reading was unintended, allow cuff to deflate & restart reading.
	If cancelled reading was intended, no action needed.
90, 91, 97-107	Power off and repower on the CT40.
	If issue persists, call SMI Customer Service. Service Required
	Temperature:
505 & 506	Touchless:
	Make sure patient has been at rest for a few minutes. If patient has been in cold environment, make sure patient has acclimated to indoor temperature
	FILAC [®] :
	Make sure patient has not had any warm beverages recently (oral), and then retry measurement.
501-504, 507-514,	Remove and then replace the probe in the well.
516-517	Retry reading.
	Power off and repower on the CT40.
	Retry reading. If issue persists, call customer service.
520-524 & 526	Power off and repower on the CT40.
	Retry reading.
	If issue persists, replace the probe. Retry reading.
	If issue persists, replace module or call Customer Service.
530	Use in normal room temperature environment suitable to the device.
	SpO ₂ :
121-124, 133-136, 150-156	Remove sensor from the patient and ensure proper positioning. Attempt new measurement ensuring proper protocol as in section 7 of the User Manual.
	Unplug sensor and reconnect.
	If issue persists, power off and repower on the CT40.
	Retry reading. If issue persists, call Customer Service.

126-132	Power off and repower on the CT40.
137-149 & 157-192	If issue persists, replace module or call Customer Service
	Electronic Medical Records (EMR):
601-605	Check wired connection or wireless strength and proximity.
606-610, 612, 614	Call biomedical engineer or IT support to check EMR configuration.
611	Ensure proper barcode and Patient ID scanned.
613	Multiple patients have the same patient ID.
	CT40 System:
300-302	Power off and repower on the CT40.
341-344	Power off and repower on the CT40.
	Return for Service.
	Powering Device:
201	Run only from A/C power source.
	Have battery service performed at earliest convenience.
202	Power off and repower on the CT40.
	If repower attempt does not work, plug in power source and power off and repower on CT40 again.
	Retry reading. If issue persists, call Customer Service.
203-211	Clear the error and power off and repower on the CT40.
	Replace battery.
212-214	Call Customer Service.
215	Charge Battery.
	Barcode Scanner:
701-706	Check and ensure proper barcode and Patient ID was scanned
	Rescan Patient ID
	Disconnect / Reconnect cable from USB port on CT40.
	Replace Barcode Scanner
	Printing:
381	Close Printer Door.
382	Replace Paper.
386	Charge printer battery. Reset printer.

383- 387 & 388-390 Check cables. Reset Printer. Replace Printer.

Out of Range Measurements

The SunTech CT40 can measure BP that falls within the following ranges.

Systolic: Adult 40-260 mmHg, Pediatric 40-230 mmHg

Diastolic: Adult 20-200 mmHg, Pediatric 20-160 mmHg

Heart Rate: 30 - 220 BPM

The SunTech CT40 will display temperature and SpO₂ within the following ranges.

FILAC® Temperature: 30.1 to 42.9 °C (86.2 - 109.2 °F)

Touchless IR Temperature: 34 to 42.5 °C (93.2 - 108.5 °F)

SpO₂: 70 - 100%

If measurement is above or below specified ranges:

Single measurement mode: The highest or lowest possible threshold range will blink in the appropriate BP measurement field.

Averaging mode: 🔨 will blink in addition to the status code "3" in the blood pressure measurement display

Temperature: depending on the out or range value, the display will indicate whether the value is out of range (hi) or out of range (Lo).

SP02: results falling below the measurement range will display "Lo" in the results field.

Service Centers

For customers in the Americas, Asia, and the Pacific:



SunTech Medical, Inc. Service Department 507 Airport Boulevard, Suite 117 Morrisville, NC 27560 USA Tel: 800.421.8626 919.654.2300 Fax: 919.654.2301

13. Frequently Asked Questions

What is the expected battery life for the SunTech CT40?

When batteries are fully charged, the SunTech CT40 is designed to run on internal battery power for 120 measurement cycles (BP, temperature and SpO₂ measurements) before requiring recharge. When only the bottom segment of the battery symbol is illuminated, there is between 11 percent and 40 percent battery power remaining.

Will the device automatically power down?

When no measurements have been taken or buttons pressed for one hour, the device will automatically powerdown and enter the low-power state and is considered "off."

What is the significance of different beeps I hear during operation?

One long beep (approximately 3-second beep): Successful end of BP measurement.

Three long beeps: Severe hardware error.

One short beep (1-second beep): Power-up or Power-down complete.

Four short beeps: There has been an error.

If you abort a BP measurement by pressing the Start/Stop button, you will hear one short beep. Then after the pressure has been completely dumped from the cuff, you will hear one long beep.

Can pulse oximetry, temperature measurement, a barcode scanner or printer be added later too?

Yes, you can add vital signs modules and accessories to the SunTech CT40 at any time.

Can EMR connectivity be established at a later date if not enabled as part of initial configuration?

Yes, EMR settings can be accessed via the Advanced Configuration Application at any time.

Why is no heart rate data showing on the display?

If the device is configured to use the Sp02 sensor as the pulse rate source, and the sensor is not attached to the patient, there will not be heart rate data on the display. Only dashes will appear, or MAP results if MAP measurement is enabled. Note that MAP is not available in the United States.

How many measurements can be stored in the SunTech CT40 memory?

The SunTech CT40 can store 99 unique sets of measurements. After the 99th measurement has been saved, the next measurement will have the Memory Location Identifier label "01" and will overwrite the most recent measurement (measurement 99). After all results are cleared from the memory, the Memory Location Identifier also will return to "01."

14. Technical Information

EMC Statement

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: 2014. 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. When the device accuracy of any value displayed on the device is questionable, determine the patient's vital signs by alternative means. If it is determined that the device is malfunctioning, stop using the device and have the device checked by a qualified service technician before further use. If this equipment does cause harmful interference to other devices, which can be determined by turning the equipment off and on, the user is encouraged to try to correct the interference by one or more of the following measures:

Reorient or relocate the receiving device.

Increase the separation between the equipment.

Connect the equipment into an outlet on a circuit different from which the other device(s) are connected.

Consult the manufacturer or field service technician for help.

Portable and mobile RF communications equipment can affect Medical Electrical Equipment.

WARNING: Portable RF communications equipment (including peripherals such as antenna cables and external antennas) should be used no closer than 30 cm (12 inches) to any part of the CT40 system, including cables specified by the manufacturer. Otherwise, degradation of the performance of this equipment could result.

WARNING: Use of accessories, transducers, and cables other than those specified may result in increased emissions or decreased immunity of the CT40.

WARNING: The CT40 should not be used adjacent to or stacked with other equipment. If adjacent or stacked use is necessary, the CT40 should be observed to verify normal operation in the configuration in which it will be used.

Guidance and Manufacturer's Declaration – Electromagnetic Emissions			
The CT40 Device is intended for use in a Professional Healthcare Facility within the electromagnetic environment specified below. The customer or the user of the CT40 Device should assure that it is used in such an environment. This equipment has been tested and found to comply with the limits for medical devices to IEC60601-1-2: 2014.			
Emissions test	sions test Compliance Electromagnet Guidance		
RF emissions CISPR 11	Group 1	The CT40 Device uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.	
	Class B	The CT40 Device is suitable for use	

Guidance and Manufacturer's Declaration – Electromagnetic Emissions

Harmonic emissions IEC 61000-3-2	Class A	in all establishments other than
Voltage fluctuations/ flicker emissions IEC 61000-3-3	Complies	domestic and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.

Guidance and Manufacturer's Declaration – Electromagnetic Immunity

The CT40 Device is intended for use in a Professional Healthcare Facility within the electromagnetic environment specified below. It is not intended for helicopter transport, hospital ambulance or home use. It is not intended for near active HF SURGICAL EQUIPMENT and the RF shielded room of an ME SYSTEM for magnetic resonance imaging, where the intensity of EM DISTURBANCE is high. The customer or the user of the device should assure that it is used in such an environment. This equipment has been tested and found to comply with the limits for medical devices to IEC 60601-1-2: 2014.

If device should be subjected to Electrostatic Discharge (ESD) or Electromagnetic Interference from another device, the device should recover on it's own within 5 seconds. If device does not recover, turn off device, and wait 5 seconds. Turn on the device and verify that the device is operating normally. If device does not operate properly or fails to recover, contact SunTech Medical.

Immunity Test	Applies to	Compliance Level	Electromagnetic Environment- Guidance for Professional Healthcare Facility Environment
Electrostatic discharge (ESD) IEC 61000-4-2	All device input and output connections and cables.	± 2, 4, 6, 8kV contact ± 2, 4, 8, 15kV air discharge	Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30%. Users must eliminate static in their hands before use it.
Radiated RF EM fields IEC 61000-4-3	All device input and output connections and cables.	3V/m 80 MHz to 2700MHz 80% AM at 1kHz	Radiated electromagnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment
Radiated RF Wireless communication equipment IEC 61000-4-3	All device input and output connections and cables.	See Table A below	This device has been subjected to RF wireless communication bands from cell phones, and other communication devices
Electrical fast transient/burst IEC 61000- 4-3	All device input and output connections and cables.	± 2kV for power supply lines 100kHz repetition frequency	Mains power quality should be that of a typical commercial or hospital environment
Surge IEC 61000-4-5	All device input and output connections	± 0.5, 1, 2kV	(Professional Healthcare Facility)

	and cables.		
	All device input and output connections and cables.	± 0.5, 1kV	
Conducted Disturbances induced by RF fields IEC 61000-4-6	All device input and output connections and cables.	3V 0.15MHz – 80MHz 6V in ISM bands between 0,15 MHz and 80MHz 80% AM at 1kHz	Mains power quality should be that of a typical commercial or hospital environment. All handheld and patient coupled parts should be consistent with intended use.
	DC Input and all cables	(>3m)	be consistent with intended dae.
Power Frequency magnetic field IEC 61000-4-8	All device input and output connections and cables.	30 A/m 50 or 60 Hz	Power Frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.
NOTE: a) U _T is the a.c. mains b) E.g. 25/30 means 2	voltage prior to applic 25 periods at 50 Hz or 3		
		0% UT: 0.5 cycle a) At 0°, 45°, 90°, 135°, 180°, 225°, 270° and 315°	Mains power quality should be that of a typical commercial or hospital environment (Professional Healthcare Facility)
Voltage dips, short interruptions and voltage variations on power supply input lines IEC 61000-4-11	Device input (AC power)	0% Uτ: 1 cycle 70% Uτ: 25/30 cycles b) Single Phase: at 0°	If the user of the CT40 device requires continued operation during power mains interruption, it is recommended that the device
		0% UT: 250/300 cycles b)	be powered from an uninterruptible power supply or a battery.
Conducted RF IEC 61000- 4-6	All device input and output connections and cables.	3V 10V ISM bands 150kHz to 80MHz	Portable and mobile RF communications equipment should be used no closer to any part of the device including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter. Minimum separation distance for higher IMMUNITY TEST LEVELS shall be calculated using the following equation.
			Where P is the maximum outpur power rating of the transmitter ir

Guidance and Manufacturer's Declaration – Electro	omagnetic Immunity
	watts (W) according to the transmitter manufacturer and d is the recommended separation distance in meters (m) and E is the Immunity Test Level in V/m.
	$E = \frac{6}{d}\sqrt{P}$
	Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey, a should be less than the compliance level in each frequency range

Rated maximum output power of transmitter. Watts (W)	Separation distance according to frequency of transmitter meters (m)			
	150 kHz to 80 MHz $d = 1,2\sqrt{P}$	80 MHz to 800 MHz $d = 1,2\sqrt{P}$	800 MHz to 2,5 GHz $d = 2,3\sqrt{P}$	
0.01	0.12	0.12	0.23	
0.1	0.38	0.38	0.73	
1	1.2	1.2	2.3	
10	3.8	3.8	7.3	
100	12	12	23	

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

NOTE 1: at 80 MHz and 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.

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

Test Frequency (MHz)	Band a) (MHz)	Service b)	Modulation b)	Maximum Power (W)	Distance (m)	IMMUNITY TEST LEVEL (V/m)
358	380 - 390	TETRA 400	Pulse Modulation b) 18Hz	1,8	0.3	27
450	430 - 470	GMRS 460, FRS 460	FM c) 5 kHz deviation 1 kHz sine	2	0.3	28
710 745 780	704 - 787	LTE Band 13, 17	Pulse Modulation b) 217Hz	0,2	0.3	9
810 870 930	800 - 960	GSM 800/900, TETRA 800, iDEN 820, CDMA 850, LTE Band 5	Pulse Modulation b) 18Hz	2	0.3	28
1720 1845 1970	1700 - 1990	GSM 1800, CDMA 1900, GSM 1900, DECT, LTE Band 1, 3, 4, 25, UMTS	Pulse Modulation b) 217Hz	2	0.3	28
2450	2400 - 2570	Bluetooth, WLAN, 802.11 b/g/n, RFID 2450, LTE Band 7	Pulse Modulation b) 217Hz	2	0.3	28
5240 5500 5785	5100 - 5800	WLAN 802.11 a/n	Pulse Modulation b) 217Hz	2	0.3	9
NOTE If neces ME EQUIPMEI a) For s b) The o	ssary to achieve th NT or ME SYSTEM come services, only carrier shall be mo n alternative to FN	may be reduced the uplink freque dulated using a 5	to 1m. The 1m te encies are includ 0% duty cycle sq	est distance is pe ed Juare wave signa	ermitted by IEC 6	1000-4-3

c) As an alternative to FM modulation, 50% pulse modulation at 18 Hz may be used because while it does not represent actual modulation, it would be worst case.

Specifications, General

Conditions for Use:	Operating: 10°C (50°F) to 40°C (104°F) 15 – 90% RH non-condensing - 70 kPa - 106 kPa. Operating the device in an environment at maximum temperature can produce temperatures exceeding 41°C (41.6°C highest recorded) on a patient applied part. It is up to the operator to determine if this temperature is too high based upon the condition of a patient and, if so, to ensure the ambient temperature of the environment is 38°C or below.
Storage:	-20°C (-4°F) to 55°C (131°F) 15 – 90% RH non-condensing - 50 kPa - 106 kPa. Performance can be affected if, used or stored outside the specified temperature, humidity, or altitude listed in the ranges above.
Power:	Internal power supply. Input: 100-240 VAC @ 1.5A max, 50-60 Hz. Output +9VDC @ 5A IEC 320 type input connector.

Calibration:	The accuracy of cuff - pressure transducers/indicators should be verified bi-annually.
Safety Systems:	Independent hardware over-pressure circuit and redundant software overpressure algorithm to limit cuff pressure to less than 300 mmHg (+20/-10mmHg). Independent hardware timing circuit and redundant software timer algorithm to limit the duration of a blood pressure cycle to less than 180 seconds.
Dimensions:	Size (without thermometer option): 8" x 6.75" x 5.2")
Standards:	ANSI/AAMI ES60601-1:2005/(R)2012 and A1:2012, C1:2009/(R)2012 and A2:2010/(R)2012, EN 60601-1:2006/A1:2013, IEC 60601-1-2: 2014, IEC 80601-2-30: 2018, ISO 80601-2-61:2017, COR1:2018, ISO 15223-1:2016, ISO 10993-1, 2018, ISO 10993-5, 2009, EN ISO 10993-10, 2010, ISO 81060-2:2018, EN 50419: 2006, EN ISO 14971:2012, EN ISO 81060-1: 2012, IEC 60601-1-6: 2013, ISO 80601-2-56:2017
Classifications:	Equipment Classification: Class IIa per MDD, Class II (Electrical Shock), Continuous mode of operation, CE (413)
Ingress of liquid	Ingress Protection: IPX1 for BP and SpO ₂ modules. IPX0 for temperature modules (no protection of ingress to liquids).
Weight	0.6 lbs. (284) gm

Specifications, Blood Pressure Measurement

Measurement:	Oscillometric with step deflation	
Range:	Pressure: Diastolic: 20-200 mmHg (adult), 20-160 mmHg (pediatric). Systolic: 40-260 mmHg (adult), 40-230 mmHg (pediatric)	Heart Rate: 30-220 BPM (beats per minute)
Accuracy (BP):	Meets or exceeds ANSI/AAMI/ISO 81060-2:2018 stan (±5mmHg mean error with 8mmHg standard deviatior	-

Notes on Blood Pressure Data

Any blood pressure reading can be affected by the measurement site, the position of the patient, exercise, 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 pacemakers and common arrhythmias such as atrial or ventricular premature beats or atrial fibrillation, arterial sclerosis, poor perfusion, diabetes, age, pregnancy, pre-clampsia, renal diseases, patient motion, trembling, and shivering.

SP0₂ Sensor Specifications

Function	ChipOx®	Masimo [®] M-LNCS DCI
Saturation Normal (no motion)	± 2% (70-100%A _{rms})	± 2% (70-100%A _{rms})
Saturation Motion	N/A	± 3% (70-100%A _{rms})
Low Perfusion	N/A	SpO ₂ +/-2%

Pulse Heart rate	20-300 BPM: ±3 BPM	± 3% BPM
Pulse Heart rate w/motion	20-300 BPM: ±3 BPM	± 5% BPM
Wayalangth & Output Dawar	RED: 660nm @ 3.5-4.5mW	RED: 660nm @ ≤ 15mW
Wavelength & Output Power	IR: 905nm @ 3.5-4.5mW	IR: 905nm @ ≤ 15mW

For more detailed information, see the instructions for use included with each device.

Temperature Sensor Specifications

Function	Touchless IR Thermometer		Covidien® Electronic Thermometer
	Body Temperature:		30.1 to 42.9 °C (86.2 - 109.2 °F)
Temperature accuracy	36 to ~39°C	± 0.2°C	
	96.8 to 102.2 °F	± 0.2°C	Quick Mode (Oral): ± 0.3°C
	34.0 to ~35.9°C	± 0.3°C	Std. Mode (Axillary/Rectal): ± 0.1°C
	93.2 to ~96.6 °F,		Direct Mode: ± 0.1°C
	39.1 to ~42.5°C	± 0.3°C	
	102.4 to ~108.5		
Response Time	< 2 seconds		3-10 sec (Oral Quick Mode)
			6-10 (Oral Standard Mode)
			8-12 sec. (Axillary Mode) 10-14 sec. (Rectal Mode) 60-120 sec. (Direct Mode all sites)
Resolution	0.1°C		0.1°C

Wi-Fi Dongle Specifications

Function	Operating Band(s)	D-Link DWA-171
Frequency	Frequency Band	Max. Output Power
	5.15 – 5.25 GHz	200 mW
5 G	5.25 – 5.35 GHz	200 mW
	5.47 – 5.725 GHz	1 W
2.4 G	2.4 – 2.4835 GHz	100 mW
Standards	802.11ac/n/g/a/b	

Limited Warranty

SunTech Medical, Inc. provides to the original purchaser the following limited warranty from date of invoice.

Main unit, SpO ₂ modules, temp modules	Three years
OPD Blood Pressure Cuffs	Two years
Accessories & Components (SpO ₂ sensors, temp probes, BP hose, etc.)	90 days

SunTech Medical, Inc. warrants each instrument to be free from defects in material and workmanship. Liability under this warranty covers servicing of the instrument when returned from the customer's facility within the United States prepaid to the factory. SunTech Medical, Inc. 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 first notify SunTech Medical, Inc. of the suspected defect. The device should be carefully packaged and shipped prepaid to:



SunTech Medical, Inc. Service Department 507 Airport Boulevard, Suite 117 Morrisville, NC 27560 USA Tel: 800.421.8626 919.654.2300 Fax: 919.654.2301

The device 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 device has been damaged by accident, misuse, negligence, act of God or serviced by any person not authorized by SunTech Medical, Inc. This limited warranty contains the entire obligation of SunTech Medical, Inc. and no other warranties expressed, implied or statutory are given. No representative or employee of SunTech Medical, Inc. is authorized to assume any further liability or grant any further warranties except as herein.

Conflict Minerals

SunTech's current Conflict Minerals Statement can be accessed at:

https://www.suntechmed.com/support/additional-resources/1918-conflict-minerals-statement