DINAMAP® PRO
Series 100-400 Monitor
Operation Manual

CRITIKON
VITAL ANSWERS™
DINAMAP® PRO Monitor
Operation Manual

This manual is for DINAMAP® PRO Monitor Models 100, 200, 300, and 400, all with printers.

- PRO 100: BP and Pulse
- PRO 200: BP, Pulse, and Temp
- PRO 300: BP, Pulse, and SpO₂
- PRO 400: BP, Pulse, Temp, and SpO₂

The model of the Monitor determines which menu option buttons appear on the LCD. Please refer to applicable sections.

Reissues and Updates
Changes occurring between issues are addressed through Change Information Sheets, Addendums, and replacement pages. If a Change Information Sheet does not accompany this manual, it is correct as printed.

Errors and Omissions
If errors or omissions are found in this manual, please notify:
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Part No. 776995C

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Illustrations may show design models; production units may incorporate changes.
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Introduction

About the DINAMAP® PRO Monitor
DINAMAP® PRO Monitors provide noninvasive determination of systolic blood pressure, diastolic blood pressure, mean arterial pressure, pulse rate, temperature, and oxygen saturation. These portable AC- and DC-operated monitors are primarily intended for use in hospital acute care settings such as outpatient surgery, accident and emergency, labor and delivery, GI/endoscopy, and medical/surgical units.

The PRO Monitor comes in four different models: PRO 100, 200, 300, and 400, all with printers.
- PRO 100: BP and Pulse
- PRO 200: BP, Pulse, and Temp
- PRO 300: BP, Pulse, and SpO₂
- PRO 400: BP, Pulse, Temp, and SpO₂

All of the main operations of the PRO Monitor are easy to use. Please review the factory default settings and, where applicable, enter settings appropriate for your use. The “Using the Monitor” section of this manual explains how to use the Monitor in its most simple form, while the “Using the Menu System” section explains how to customize measurements by using the menu system.

Indications
The PRO Monitor is intended to monitor one patient at the bedside.

Contraindications
This device is not designed, sold, or intended for use except as indicated.

Federal law (U.S.A.) restricts this device to sale by or on the order of a clinician.

Warnings
- Do not use the PRO Monitor in the presence of magnetic resonance imaging (MRI) devices. There have been reports of sensors causing patient burns when operating in an MRI environment.
• Do not use the Monitor in the presence of flammable anesthetics.

• To help prevent unintended current return paths with the use of high frequency (HF) surgical equipment, ensure that the HF surgical neutral electrode is properly connected.

• To avoid personal injury, do not perform any servicing unless qualified to do so.

• WARNING: These Monitors should not be used on patients who are connected to cardiopulmonary bypass machines.

• If powering the Monitor from an external power adapter or converter, use only Critikon-approved power adapters and converters.

• The Monitor does not include any user-replaceable fuses. Refer servicing to qualified service personnel.

• To reduce the risk of electric shock, do not remove the cover or the back. Refer servicing to a qualified service person.

• If the accuracy of any determination reading is questionable, first check the patient’s vital signs by alternate means and then check the PRO Monitor for proper functioning.

Cautions

• Do not use replacement batteries other than the type supplied with the Monitor. Replacement batteries are available from Critikon. See Appendix D.

• The PRO Monitor is designed to conform to Electromagnetic Compatibility (EMC) standard IEC 601-1-2, 1993 and will operate accurately in conjunction with other medical equipment which also meets this requirement. To avoid interference problems affecting the Monitor, do not use the Monitor in the presence of equipment which does not conform to these specifications.
Introduction

- Place the PRO Monitor on a rigid, secure surface. Monitor must only be used with mounting hardware, poles, and stands recommended by Critikon. See Appendix D.

- The weight of the accessory basket contents should not exceed 6.6 lb (3 kg).

- Arrange the power cord, air hoses, and all cables carefully so they do not constitute a hazard.

- Verify calibration of BP parameter (temp and pulse oximeter do not require calibration). Ensure that the display is functioning properly before operating the PRO Monitor.

- Do not immerse the Monitor in water. If the Monitor is splashed with water or becomes wet, wipe it immediately with a dry cloth.

- Do not gas sterilize or autoclave.

- The PRO Monitor, when used with Critikon-approved applied parts and accessories, is protected against defibrillator damage.

Notes

- Waveforms may be distorted and readings inaccurate when electrosurgical cautery equipment is used while monitoring with the PRO Monitor.

- The electromagnetic compatibility profile of the PRO Monitor may change if accessories other than those specified for use with the PRO Monitor are used.

- Trend data are retained in the PRO Monitor when it is turned off, except when the default is overridden by selecting the Trend button under the Service menu.
**Product Compliance**

The DINAMAP® PRO Monitor is classified in the following categories for compliance with IEC 601-1:

- Class I, internally powered
- Transportable
- For continuous operation
- Not suitable for use in the presence of flammable anesthetics
- Not for use in the presence of an oxygen-enriched atmosphere (oxygen tent)
- Type BF applied parts
- IPX1, degree of protection against ingress of water
- Sterilization/Disinfection, see Appendix F

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**CE 0086**

This product conforms with the essential requirements of the Medical Device Directive. Accessories without the CE mark are not guaranteed to meet the Essential Requirements of the Medical Device Directive.
Getting Started

Unpacking the Monitor and Accessories
Before attempting to use the PRO Monitor, take a few minutes to become acquainted with the Monitor and its accessories. Unpack the items carefully, and check them against the contents checklist enclosed in one of the accessory boxes. This is also a good time to check for any damage or shortage. If there is a problem or shortage, contact Critikon.

It is recommended that all the packaging be retained, in case the Monitor must be returned for service in the future.

Power Sources
The PRO Monitor is designed to operate from either an internal lead-acid battery, AC mains or an IEC 601-1 compliant DC power source (see Appendix A). For replacement rechargeable batteries, please refer to the Service section of this manual.

The Monitor contains five fuses. Two AC line input fuses are mounted internally and are replaceable only by qualified service personnel. The remaining three fuses are auto-resetable and mounted within the Monitor. These fuses protect the low voltage DC input, the battery, and the +5 V output on the host port connector.

Powering the Monitor
Before the PRO Monitor is used for the first time, the battery should be charged in the Monitor for at least 8 hours.

Refer to the illustration of the rear panel connections. Looking at the rear of the PRO Monitor, remove the battery compartment cover. Insert the rechargeable battery into the compartment so that the battery terminals fit into the power clips at the bottom of the compartment. Then replace the cover. Insert the power cord plug into the mains external power socket (2) and plug into an AC outlet.

Refer to the illustration of the front panel controls and indicators. With mains or external DC power connected, the green external power indicator LED (14) will light to indicate that external power is being applied and that the battery is charging. If the battery is not inserted, the external
For continued safety, use only a power cord of listed type SJT, three-conductor, min. No. 18 AWG, terminated in a medical/hospital grade attachment plug, provided with the following cord tag: "Hospital Grade Plug." Grounding and 500 charge/discharge cycles. When it is necessary to replace the battery, refer to the “Compatibility Table and Reorder Codes” listed in Appendix D. To ensure full charge cycles, replace only with a recommended battery. If the Monitor is to be stored for some time, first charge the battery and then remove it and store it separately from the Monitor.

During battery-only operation, the yellow battery power indicator LED (17) will light. When the battery becomes discharged beyond the low battery threshold, the indicator will begin to flash, and the Monitor will sound warning beeps every 30 seconds. At this point, the Monitor should be connected to an AC outlet to recharge the battery. If the Monitor continues to be used without charging the battery, the message **WARNING: THE BATTERY IS TOO LOW FOR MONITOR TO FUNCTION. TURN MONITOR OFF** appears. The Monitor shuts down all functions until it is turned off and the battery is recharged or replaced. To run the Monitor on AC power, it must be powered off and then on again.

Battery charging will take place as long as the Monitor remains connected to an external AC power source. A battery that is fully discharged can be fully recharged in 1 hour 50 minutes when the Monitor is switched off or 8 hours if the Monitor is switched on.

**Notes**

- To prolong the life of the battery, keep the Monitor connected to an AC outlet whenever possible. NEVER allow the battery to become completely discharged. A fully charged battery will power the Monitor for approximately 2 hours and should survive between 200 and 500 charge/discharge cycles. When it is necessary to replace the battery, refer to the “Compatibility Table and Reorder Codes” listed in Appendix D. To ensure full charge cycles, replace only with a recommended battery. If the Monitor is to be stored for some time, first charge the battery and then remove it and store it separately from the Monitor.

- For continued safety, use only a power cord of listed type SJT, three-conductor, min. No. 18 AWG, terminated in a medical/hospital grade attachment plug, provided with the following cord tag: "Hospital Grade Plug." Grounding
Getting Started

integrity can only be maintained when equipment is connected to an equivalent receptacle marked "Hospital Grade."

- Where the integrity of the external earth conductor in the installation or its arrangement is in doubt, the Monitor must be operated from its internal battery.

General Caution

- Do not touch either the pin of the DC input connector (3) or the terminals within the battery compartment (1) and the patient at the same time.
Rear Panel Connections

1 Battery compartment cover: Retains and protects internal battery
2 Mains input: Used to connect to AC power supply
3 External power socket: To be used with approved Critikon AC-DC power converter ONLY
4 Inactive temperature cable storage: Inactive temperature probe cable attaches here (Models 200 and 400)
5 Pole clamp: Used to clamp monitor to pole or stand
6 Data interface connector: Host communications port (15 way D-type RS-232 serial port) for use only with equipment conforming to IEC 601-1, configured to comply with IEC 601-1-1


**Front Panel Controls and Indicators**

7 Systolic pressure display: 3-digit red LED indicates measured systolic BP in mmHg

8 Active temperature probe holster: Temperature probe that is being used stored here (Models 200 and 400)

9 Diastolic pressure display: 3-digit red LED indicates measured diastolic BP in mmHg

10 Temperature probe cover storage: Box of probe covers stored here (Models 200 and 400)

11 Inactive temperature probe holster: Extra temperature probe can be stored here (Models 200 and 400)

12 Temperature display: 4-digit red LED indicates measured temperature (Models 200 and 400)

13 °C °F display: Indicates whether temperature is being displayed in degrees Celsius or Fahrenheit (Models 200 and 400)
14 External power indicator: Green LED indicates external power status and battery charging status of monitor
15 Temperature probe connector: Temperature probe cable attaches here (Models 200 and 400)
16 ON/OFF switch: Controls on/off state of monitor; push for power on and push again for power off
17 Battery power indicator: Yellow LED indicates operation and charge status of internal battery
18 SpO₂ sensor connector: SpO₂ sensor extension cable attaches here (Models 300 and 400)
19 Mean arterial pressure display: 3-digit red LED indicates measured MAP in mmHg and shows instantaneous cuff pressure during BP determination
20 SpO₂ pulse indicator: Yellow LED in heart symbol flashes to indicate that real-time pulse rate measurements are being derived from SpO₂ signals (Models 300 and 400)
21 Rotor: Used to highlight and select items in LCD menus; if monitor is off, pressing rotor will switch monitor on
22 Pulse BPM display: 3-digit yellow LED shows pulse rate in beats per minute
23 SpO₂ display: 3-digit red LED indicates oxygen saturation in % (Models 300 and 400)
24 SpO₂ artifact indicator LED: Illuminates when motion artifact is detected (Models 300 and 400)
25 LCD (liquid crystal display): Displays all alarms, user interface messages, and configuration options
26 Alarm silence switch: Alternately mutes and enables audible alarms; when pushed once after alarm sounds (silence on), switch lights to indicate that audible alarms have been silenced for 2 minutes
27 AUTO BP key: Press to start Auto BP mode
28 Light sensor: Automatically measures ambient light to set LED display intensity
29 START/STOP BP key: Press to start or stop a BP, Auto, Stat, or Vitals determination
30 Cuff connector: BP cuff hose attaches here
Getting Started

Switching the Monitor On and Off

To switch the DINAMAP PRO Monitor on, push the power ON/OFF switch (16) or press the rotor (21).

As the Monitor powers up, it will run a short self-test routine, which will flash all the indicator lights and then beep the warning speaker. After a few seconds the system will be ready for operation, as indicated by the appearance of the main menu on the LCD (25).

To switch the Monitor off, push the power ON/OFF switch (16) again. This will terminate any measurements that may be in progress and automatically deflate the cuff.

When the Monitor is operating on the internal battery only, battery life is enhanced by the use of the sleep mode. However, the PRO Monitor will not enter sleep mode if an alarm is active. If no controls are used and no determinations are being made, the Monitor will enter sleep mode after a time which can be preset by the operator. All LED displays will be blanked except for a dash in the far-left systolic position, and any existing readings will be transferred to the LCD, which displays the message “Sleep Mode Active.” Moving the rotor or pressing a key will “wake up” the Monitor.

Liquid Crystal Display (LCD)

Menu Area
This area displays the name of the menu that has option buttons available for selection. Normal text in the menu area appears dark on a light background, while the text
selected buttons appears light on a dark background.

**Note:** Some menus have six option buttons. In these cases, there is no space available to display the menu title.

**Area 2**
This area displays data from one of three different sources.

- Source 1: SpO₂ plethysmograph (Models 300 and 400)
- Source 2: Last three BP readings
- Source 3: Error and warning messages

**Note:** Refer to “Display Button” in the “Using the Menu System” section for instructions on setting Area 2.

**Area 3**
This area displays the time, the time lapsed since the last Auto BP determination (if in Auto BP mode), the battery icon (if operating on battery power, the time and battery icon toggle), and the BP and Printer modes.

**Using the Printer**

**Installing the Paper**
Turn the PRO Monitor so that the side is facing you. While grasping the side of the Monitor, firmly press the notched indentations on the printer door to open it. The printer door will pop open. With the Monitor powered on, place the roll of paper into the compartment so that the end of the paper comes off the top, and thread it between the two printer plates. As the paper touches the plates, the paper will begin to auto-feed itself into the printer. Feeding the end of the paper strip through the slot in the door, firmly press the notched indentation on the side of the printer door to close it. Use the paper release lever to clear a paper jam or manually feed the paper.

**Note:** Make sure that the roll of paper is tightly wound.
Getting Started

Any time the paper is loaded, the printer automatically prints a test strip with the DINAMAP® PRO name on it. If no print is visible on the paper, check that the paper roll has been installed in the correct position (refer to diagram). To tear off the printout, use a slight sideways action to pull the paper sharply up across the serrated edge of the door.

Printer Alarms

If the Monitor is switched on with no paper installed or with the printer door open, the message “No Paper” will appear next to “PRNT” in Area 3 of the LCD. When new paper is installed and the printer door is closed, the message will change to “Manual” for Manual print or “Auto” for Auto print, depending on the status before the paper change.

If the paper runs out during a print request or if an attempt is made to print when no paper is installed, the message “Printer - No Paper” will appear in Area 2 of the LCD and an audible alarm will sound. In addition, the message “No Paper” will appear next to “PRNT” in Area 3 of the LCD. To clear the alarm, press the rotor. The message in Area 3 of the LCD will remain until new paper is installed and the printer door is closed. (See “Using the Menu System.”)

Installing new paper will cause the Critikon DINAMAP PRO header to be printed, thereby confirming that the paper is installed correctly and that the printer is operational. The message next to “PRNT” in Area 3 of the LCD will change to “Auto” or “Manual” to identify the operating mode of the printer. After power-off, the operating mode of the printer returns to the previous user-selected setting (Auto or Manual) unless specified otherwise in the Print button under the Service Button.

Storage

Store thermal paper in a cool, dry place. The printed strip (thermal paper recording) should not be

- exposed to direct sunlight,

- exposed to temperatures over 100 °F/38 °C or relative humidity over 80%, or

- placed in contact with adhesives, adhesive tapes, or plasticizers such as those found in all PVC page protectors.
Note: When in doubt about long-term storage conditions, store a photocopy of the thermal paper recording.

Cautions

- The paper is thermally activated; therefore, do not store it in a hot place as discoloration may result.

- Use only replacement paper rolls (58 mm) from Critikon.
Using the Monitor

Noninvasive Blood Pressure Determination

Description

The BP parameter is included in Models 100, 200, 300, and 400. Blood pressure is monitored noninvasively in the PRO Monitor by the oscillometric method, which measures the amplitude of the pressure oscillations within the blood pressure cuff. Further information about the oscillometric method is in Appendix C.

The PRO Monitor has four BP modes: 1. Manual, 2. Auto, 3. Stat, and 4. Vitals (UK: All Obs). The mode, which is selected by the user, is shown on the LCD (25). The BP measurements are automatic, and once the cycle is complete the LED displays (7, 9, 19, 22) show systolic pressure, diastolic pressure, mean arterial pressure, and pulse rate.

1. Manual BP determinations are started by pressing the START/STOP BP key (29). In the Manual mode, the blood pressure is determined one time.

2. Auto BP determinations are started by selecting the AUTO BP key (27) or the Auto button under the Set BP (UK: BP Mode) button in the Main menu.

When Auto mode is selected, a number at the right of the Auto button indicates the time interval between each reading. To change the time interval, choose the box around the number and turn the rotor until the desired interval is reached. The interval can be set between 1 and 120 minutes (1, 2, 3, 4, 5, 10, 15, 20, 25, 30, 45, 60, 90, and 120 minutes). Press the rotor to confirm the setting.

3. Stat determinations are started by selecting the Stat button under the Set BP button (UK: BP Mode) in the Main menu. In the Stat mode, the blood pressure is determined as many times as possible in 5 minutes.

4. Vitals (UK: All Obs) determinations are started by selecting the Vitals (UK: All Obs) button in the Main menu. (Refer to the “Using the Menu System” section.) Selection of this button initiates a BP determination.
while allowing SpO₂ and predictive temperature determinations to be monitored and recorded (depending on Monitor model). In the Vitals (UK: All Obs) mode, the blood pressure is determined one time.

Before each BP determination, the Monitor performs a test to ensure that the cuff pressure is below a specified level. The determination is delayed until this condition is met. During the delay, the BP values are displayed as zero.

The Monitor senses the type of hose being used and automatically uses adult/pediatric monitoring parameters or neonatal monitoring parameters, as appropriate.

Audible and visible alarms occur when a value for systolic pressure, diastolic pressure, mean arterial pressure, or pulse rate is outside the selected high or low limit.

Instructions for cleaning and disinfecting BP cuffs are in Appendix F.

**General Warnings**

- The PRO Monitor will not measure blood pressure effectively on patients who are experiencing seizures or tremors.

- Arrhythmias will increase the time required by the PRO Monitor to determine a blood pressure and may extend the time beyond the capabilities of the Monitor.

- In Manual mode, the PRO Monitor displays the results of the last blood pressure determination for 2 minutes or until another determination is completed. If a patient’s condition changes between one determination and the next, the Monitor will not detect the change or indicate an alarm condition.

- Devices that exert pressure on tissue have been associated with purpura, skin avulsion, compartmental syndrome, ischemia and/or neuropathy. To minimize these potential problems, especially when monitoring at frequent intervals or over extended periods of time, make sure the cuff is applied appropriately and...
Using the Monitor

examine the cuff site and the limb distal to the cuff regularly for signs of impeded blood flow.

- Do not apply external pressure against cuff while monitoring. Doing so may cause inaccurate blood pressure values.

- Use care when placing cuff on extremity used to monitor other patient parameters.

- The PRO Monitor is designed for use only with dual-tube cuffs.

- Use only accessories recommended by Critikon. Failure to use recommended accessories may result in inaccurate readings. See Appendix D.

- Blood pressure cuffs should be removed from the patient when the Monitor is powered off. If the extremity remains cuffed under these conditions or if the interval between blood pressure determinations is prolonged, the patient’s limb should be observed frequently and the cuff placement site should be rotated as needed.

General Cautions

- Accuracy of BP measurement depends on using a cuff of the proper size. It is essential to measure the circumference of the limb and to select the proper size cuff. The air hoses are color-coded according to size of the patient. The gray 12- or 24-foot hose (3.66 m or 7.3 m) is required on patients who require cuff sizes from infant through thigh cuffs. The teal (blue-green) 12-foot hose (3.66 m) is required for the neonatal cuff sizes #1 through #5.

- If it becomes necessary to move the cuff to another limb, make sure the appropriate size cuff is used.

- The pulse rate derived from a BP determination may differ from the heart rate derived from an EKG waveform because the PRO Monitor measures actual peripheral pulses, not electrical signals or contractions from the heart. Differences may occur because electrical signals at the heart occasionally fail
to produce a peripheral pulse or the patient may have poor peripheral perfusion. Also, if a patient’s beat-to-beat pulse amplitude varies significantly (e.g., because of pulsus alternans, atrial fibrillation, or the use of a rapid-cycling artificial ventilator), blood pressure and pulse rate readings can be erratic, and an alternate measuring method should be used for confirmation.

**General Notes**

- A patient’s vital signs may vary dramatically during the use of cardiovascular agents such as those that raise or lower blood pressure or those that increase or decrease heart rate.

- Because treatment protocols based on the patient’s blood pressure may rely on specific values and differing measurement methods, such as auscultatory, clinicians should note a possible variance from values obtained with the PRO Monitor in planning patient care management. The PRO Monitor values are based on the oscillometric method of noninvasive blood pressure measurement and correspond to comparisons with intra-aortic values within ANSI/AAMI Standards for accuracy (a mean difference of ±5 mmHg, and a standard deviation of ±8 mmHg).

- Several conditions may cause the BP parameter to calculate and display only the mean arterial pressure (MAP) without a systolic and diastolic reading. These conditions include very low systolic and amplitude fluctuations, so an accurate calculation for these values can’t be made (e.g., patient in shock); too small of a difference between systolic and MAP calculations in relationship to the difference between diastolic and MAP; or a leak has occurred in the PRO Monitor (1. Check all BP connections 2. Monitor may need calibration and leak testing). If only the MAP value is displayed, the systolic and diastolic will display dashes (—) and an alarm message “N99-BP FAILED” will be displayed.
Using the Monitor

Procedures
1. Connect the end of the air hose which has quick-release clips to the cuff connector (30) on the front of the Monitor. Make sure that the hose is not kinked or compressed.
   **Note:** To disconnect the hose from the Monitor, squeeze the quick-release clips together and pull the plug from the cuff connector (30).
2. Select the appropriate blood pressure measurement site. Because normative values are generally based on this site and as a matter of convenience, the upper arm is preferred. When upper arm size or shape, the patient’s clinical condition, or other factors prohibit use of the upper arm, the clinician must plan patient care accordingly, taking into account the patient’s cardiovascular status and the effect of an alternative site on blood pressure values, proper cuff size, and comfort. The figure shows the recommended sites for placing cuffs.
   **Warning:** Do not place the cuff on a limb being used for intravenous infusion or any area where circulation is compromised or has the potential to be compromised.

3. If patient is standing, sitting, or inclined, ensure that cuffed limb is supported to maintain cuff at level of patient’s heart. If cuff is not at heart level, the difference in systolic and diastolic values due to hydrostatic effect must be considered. Add 1.80 mmHg to values for every inch (2.54 cm) above heart level. Subtract 1.80 mmHg from values for every inch (2.54 cm) below heart level.
4. Select appropriate cuff size. Measure patient’s limb and select appropriately sized cuff according to size marked on cuff or cuff packaging. When cuff sizes overlap for a specified circumference, choose the larger size cuff.
   **Precaution:** Accuracy depends on use of proper size cuff.
5. Inspect cuff for damage. Replace cuff when aging, tearing, or weak closure is apparent. Do not inflate cuff when unwrapped.
   Precaution: Do not use cuff if structural integrity is suspect.

6. Connect the cuff to the air hose. Thread the cuff connectors onto the hose connectors until finger tight. Do not overtighten.
   Warning: It is mandatory that the appropriate hose and cuff combination be used. Any attempt to modify the hose will inhibit the Monitor from switching between the neonatal and adult measurement modes.
   Note: In normal use, each cuff will have its own hose, so it will not usually be necessary to disconnect them. If it is necessary to do so, carefully unscrew the cuff from the hose. Care should be taken in reconnecting the cuff to a hose, ensuring that threads of the cuff and hose are in alignment and no cross-threading occurs.

7. Inspect patient’s limb prior to application.
   Precaution: Do not apply cuff to areas where skin is not intact or tissue is injured.

8. Palpate artery and place cuff so that patient’s artery is aligned with cuff arrow marked “artery.”

9. Squeeze all air from cuff and confirm that connection is secure and unoccluded and that tubing is not kinked.

10. Wrap cuff snugly around the patient’s limb. Cuff index line must fall within the range markings. Ensure that hook and loop closures are properly engaged so that pressure is evenly distributed throughout cuff. If upper arm is used, place cuff as far proximally as possible.

11. Proper cuff wrapping should be snug, but should still allow space for a finger between patient and cuff. Cuff should not be so tight as to prevent venous return between determinations.
   Warning: Using a cuff that is too tight will cause venous congestion and discoloration of the limb, but using a cuff that is too loose may result in no readings and/or inaccurate readings.

Using the Monitor

Manual Mode

To start a determination, press the START/STOP BP key (29). A normal, uninterrupted Manual cycle takes about 40 seconds. The cuff pressure must drop below 5 mmHg (neonate) or 15 mmHg (adult) before another determination can be started. BP information will be displayed for 2 minutes on the LED unless another determination is started within that time frame. This applies to Manual and Vitals (UK: All Obs) modes. After power-off, the operating mode returns to the default setting of Manual. The default setting of Manual can be overridden to return to the previous user-selected setting (Auto or Manual) by selecting Set BP (UK: BP Mode) under the Service menu.

Note: The START/STOP BP key is an on-off switch; pressing it will stop any BP determination (Manual, Auto, Stat, or Vitals) that is in progress.

Auto Mode

Auto BP determinations are started by selecting the AUTO BP key (27) or the Auto button under the Set BP button (UK: BP Mode) in the Main menu.

Selecting the AUTO BP key (27) brings up the Set BP menu (UK: BP Mode) and automatically starts an Auto BP determination as long as the Monitor is in Manual BP mode. If the Monitor is already in Auto BP mode, selection of the AUTO BP key (27) brings up the Set BP menu (UK: BP Mode) without starting a new determination until the preset time interval has expired. Pressing the START/STOP BP key during a series of Auto BP determinations will cancel the determination in progress.

When Auto mode is selected, a number at the right of the Auto button indicates the time interval between each reading. To change the time interval, choose the box around the number and turn the rotor until the desired interval is reached. The interval can be set between 1 and 120 minutes (1, 2, 3, 4, 5, 10, 15, 20, 25, 30, 45, 60, 90, and
120 minutes). Press the rotor to confirm the setting. After power-off, the operating mode returns to the default setting of Manual. The default setting of Manual can be overridden to return to the previous user-selected setting (Auto or Manual) by selecting Set BP (UK: BP Mode) under the Service menu.

In the Auto mode, the pressure must be below 5 mmHg (neonate) or 15 mmHg (adult) for at least 30 seconds before another determination can be started. BP information will be displayed on the LED until the next determination is started. This applies to Auto mode only. **Note:** To cancel an Auto BP determination, select the Manual button in the Set BP menu (UK: BP Mode).

**Stat Mode**

Multiple BP readings can be taken at any time by selecting the Stat button under the Set BP button (UK: BP Mode) in the Main menu. Stat mode can also be accessed by pressing the AUTO BP key (27) and then selecting the Stat button when the Set BP menu (UK: BP Mode) appears.

If a Manual determination is not in progress, a 5-minute series of determinations will start. If a Manual determination is in progress, that determination will become the first in the series. A normal, uninterrupted Stat sequence will give the first set of systolic, diastolic, and mean arterial pressure values and pulse rate within 15 to 20 seconds. Selecting the Stat button during a series of Stat determinations will cancel the determination in progress and the rest of the series. BP information will be displayed on the LED until the determination has been canceled or completed. This applies to Stat mode only.

The series begins with cuff inflation to a pressure above the previous systolic pressure or, if no previous systolic value is stored, to approximately 160 mmHg for adult/pediatrics. The initial target pressure selection for neonates is 110 mmHg. Artifact rejection is relaxed in the Stat mode for adult/pediatric patients to allow for accelerated determinations. If a BP or Stat reading has been made previously, the first new systolic value will flash on the LED display (7) within a few seconds and will continue to flash until the end of the determination. At that point a short tone will sound and the updated systolic, diastolic, and mean
Using the Monitor

arterial pressures and pulse rate will appear on their LED displays (7, 9, 19, 22). The Monitor will begin another determination once the pressure is below 5 mmHg for 8 seconds (neonates) or 15 mmHg for 4 seconds (adults), unless the 5-minute period has ended or the determination has been canceled.

Note: Alarm limits are disabled while in Stat mode.

Predictive Temperature Determination

Description
The temperature parameter is included in Models 200 and 400. The PRO Monitor uses IVAC* technology and can be used with both oral and rectal temperature probes. Two modes of operation are available: predictive and monitor. In predictive mode, a final temperature is displayed with an audible tone. In monitor mode, the display is updated continually as the patient's temperature rises or falls.

Note: If the PRO is unable to complete a predictive determination, then it enters monitor mode. These temperature readings are not stored in trends and not reported via host comms.

During a temperature determination, the temperature display (12) provides a progress meter and probe ready indicator. In the far-left position, a single horizontal line indicates the probe is ready to start a determination after removal from the probe holster. In the far-right position of the temperature display, a “chase sequence” around the outside space indicates a predictive temperature determination is in progress. During monitor mode, the temperature readings flash constantly.

Temperature is shown on the temperature display in degrees Celsius or Fahrenheit, and the unit of measure is indicated by the °C °F display (13). The default, which is Celsius, can be changed in the Clinician Menu (please refer to the “Using the Menu System” section of this manual).

*IVAC is a trademark of Alaris Medical Systems
General Warning

- The performance of the Monitor may be degraded if it is operated outside of the environmental conditions specified in Appendix A.

General Cautions

- Be careful not to overextend the coiled cord of the temperature probe. Overextension can damage the probe coil connector interfaces.

- Accurate oral temperatures (blue) can only be obtained by placing the probe under the tongue in the right or left sublingual pocket. Temperatures in other locations in the mouth can vary by more than 2 °F or 1 °C.

- Accurate rectal temperatures can only be obtained by using the red temperature probe. Red and blue temperature probes are not interchangeable.

- Do not allow the tip of the predictive temperature probe to come into contact with a heat source (e.g., hands or fingers) prior to taking a temperature determination. If this occurs, allow 5 seconds for the probe tip to cool before proceeding.

- Use only IVAC* probes and probe covers. The size, shape, and thermal characteristics of the probe covers can affect the performance of the instrument. Inaccurate readings or retention problems may occur unless IVAC probes and probe covers are used.

Procedures

1. Connect the temperature probe cable to the temperature probe connector (15).
2. Remove the temperature probe from the probe holster. Place a protective temperature probe cover on the probe and insert the probe appropriately.
3. The determination begins automatically. Hold the temperature probe steady until the determination is complete. This takes approximately 60 seconds, during which time a pattern of lines on the temperature display (12) appears as a “chase sequence” to indicate progress.

*IVAC is a trademark of Alaris Medical Systems
Using the Monitor

When the determination is complete, an audible tone sounds and the temperature appears on the display.

4. Record the temperature and remove the probe. Discard the disposable cover by holding the probe as you would a syringe and pressing the button on the probe handle. Place the probe in the probe holster. Once you place the probe in the probe holster, the temperature values will be cleared.

Notes

- If the probe tip temperature is 94.0°F or higher (34.4°C) when taken out of the probe holster, the thermometer will not be able to perform a predictive measurement. Instead, the thermometer will automatically go into monitor mode. The temperature reading will then flash. A correct final temperature reading may require 3 minutes or longer. The Monitor will not beep at final temperature. It will continue to monitor the patient’s temperature until the probe is removed from the patient and returned to the probe holster.
- To cool the temperature probe down, wipe with alcohol.
- If there is a long delay from the time the probe is removed from the probe holster until it is inserted into the patient’s mouth, it is possible that the instrument will not display a final temperature. If this occurs, insert the probe into the probe holster, remove it again, and start a new measurement.
- If an alarm is actively sounding, an audible tone will not sound.
- If tissue contact is lost, the chase sequence on the temperature display (12) stops. If tissue contact is not made within 1 minute, the Monitor will alarm.
SpO₂

Description
The SpO₂ parameter is included in Models 300 and 400. To begin SpO₂ monitoring, simply place the SpO₂ sensor on the patient’s finger; monitoring begins automatically. Functional oxygen saturation (SpO₂) of arterial blood is noninvasively and continuously monitored in the PRO Monitor using pulse oximetry technology from NELLCOR*. Functional SpO₂ is the ratio of oxygenated hemoglobin to hemoglobin that is capable of transporting oxygen. This ratio, expressed as a percentage, is shown on the SpO₂ display (23), which is continually updated.

Heart rate derived from SpO₂ appears in the Pulse BPM display (22), and the SpO₂ pulse indicator (20) flashes synchronization with the real-time pulse rate measurements that are derived from the SpO₂ signal. A tone sounds at a rate corresponding to the pulse rate and at a pitch corresponding to the SpO₂ saturation level. The pitch is highest at 100% oxygen saturation, and it becomes lower as the saturation level falls. The Monitor can display a pulse amplitude bar and a plethysmographic waveform on the LCD (25). The pulse amplitude bar graph is proportional to the arterial blood flow. The artifact indicator LED (24) lights continuously when the Monitor detects motion sufficient enough to affect readings.

Audible and visible alarms occur when SpO₂ levels are outside the alarm limits. When a limit alarm occurs, a message appears in Area 2 of the LCD display.

If you select the Alarms button, the Alarms menu appears. This menu is used to adjust the violation limits for BP and SpO₂. Refer to “Alarms Button” in the “Using the Menu System” section.

*NELLCOR is a trademark of Mallinckrodt, Inc.
Using the Monitor

If you select the Suspend button, the SpO₂ alarm is suspended for 2 minutes and then the PRO returns to normal SpO₂ monitoring. A message informing the user that SpO₂ is suspending appears in Area 2 and dashes appear in the SpO₂ LED while the SpO₂ alarm suspend is counting down. Selecting Cancel will cancel the SpO₂ alarm suspension and return to monitoring SpO₂.

Low SpO₂ is suspending

2:00 Cancel

If the Monitor is unable to detect a pulse for 10 seconds during normal SpO₂ monitoring, the values in the LED flash, alternating patient values with dashes. The Monitor returns to normal SpO₂ reporting of values when several consecutive good pulse determinations are made.

General Warnings

- Do not use the SpO₂ function during magnetic resonance imaging (MRI). Adverse reactions include potential burns to patients as a result of contact with attachments heated by the MRI radio frequency pulse, potential degradation of the magnetic resonance image, and potential reduced accuracy of SpO₂ measurements. Always remove oximetry devices and attachments from the MRI environment before scanning a patient.

- The use of cardio-green and other intravascular dyes at certain concentrations may affect the accuracy of the SpO₂ measurement.

- The SpO₂ function is calibrated to read functional arterial oxygen saturation. Significant levels of dysfunctional hemoglobins such as carboxyhemoglobin or methemoglobin may affect the accuracy of the SpO₂ measurement.

General Cautions

- As with any clip-on sensor, pressure is exerted. The clinician should be cautious in using a clip-on sensor on
patients with compromised circulation (e.g., because of peripheral vascular disease or vasoconstricting medications).

- Do not perform any testing or maintenance on a sensor while it is being used to monitor a patient.

- Bright light sources (e.g., infrared heat lamps, bilirubin lights, direct sunlight, operating room lights) may interfere with the performance of the SpO₂ function. To prevent such interference, cover the sensor with opaque material.

**General Notes**

- A patient’s vital signs may vary dramatically during the use of cardiovascular agents such as those that raise or lower blood pressure or those that increase or decrease heart rate.

- The PRO Monitor is compatible only with NELLCOR sensors.

- Software development, software validation, and Risk and Hazard Analysis has been performed to a registered quality system.

**Procedures**

1. Select a sensor that is appropriate for the patient and the clinical situation.

   **Warning:** Do not use a damaged sensor or one with exposed electrical contacts.

   **Note:** Use only NELLCOR sensors, which are available from:

   **USA**
   Mallinckrodt, Inc.
   675 MacDonnell Blvd
   PO Box 5840
   St. Louis, MO 63134
   Phone: 1-800-NELLCOR (USA) Fax: 1-888-222-9799

   **UK**
   Nellcor
   10 Talisman Business Center
   London Road
   Bicester
   Oxfordshire OX6-OJX
   UK
   Phone: 44-189-632-2700
Using the Monitor

2. Following the directions for use supplied with the sensor, apply the sensor to the patient.

Warnings
Patient safety:

- If you fail to apply the sensor properly, the patient’s skin could be injured or the ability of the PRO Monitor to measure oxygen saturation could be compromised. For example, a clip-on sensor should never be taped shut. Taping the sensor could damage the patient’s skin or impair the venous return, thus causing venous pulsation and inaccurate measurement of oxygen saturation.

- Excessive pressure from the sensor may cause necrosis of the skin.

Monitor performance:

- When an SpO₂ sensor is on a limb that has a blood pressure cuff, the SpO₂ data will not be valid when the cuff is inflated. If SpO₂ readings are required during the entire blood pressure determination, attach the SpO₂ sensor to the limb opposite the one with the blood pressure cuff.

- Remove nail polish and artificial nails. Placing a sensor on a polished or an artificial nail may affect accuracy.

Cautions

Patient safety:

- Do not place any clip-on sensor in a patient’s mouth or on a patient’s nose or toe.

- Do not place a clip-on finger sensor on a patient’s thumb or across a child’s foot or hand.

- Observe the sensor site to assure adequate distal circulation.

Monitor performance:

- For best performance, place the sensor at heart level.

- Placing a sensor distal to an arterial line may interfere with adequate arterial pulsation and compromise the measurement of SpO₂.
- Place the sensor so that the LEDs and the photodiode are opposite each other.

3. Plug the SpO₂ sensor into the SpO₂ sensor extension cable. Then plug the SpO₂ sensor extension cable into the SpO₂ sensor connector (18).

4. Proceed with monitoring. SpO₂ determinations run continuously and can run simultaneously with other measurements.
Troubleshooting
This section discusses potential difficulties and suggestions for resolving them. If the difficulty persists, contact a qualified service person or your local Critikon representative.

The service manual, which is for use by qualified service personnel provides additional troubleshooting information.

PROBLEM: The pulse amplitude bar indicates a pulse, but no oxygen saturation or pulse rate values appear on the screen.
CAUSE:
- Excessive patient motion may be making it impossible for the SpO₂ function to find a pulse pattern.
- The sensor may be damaged.
- The patient’s perfusion may be too low to allow the SpO₂ function to measure saturation and pulse rate.

SOLUTION:
Check the patient.
- If possible, keep the patient still; check whether the SpO₂ sensor is applied securely and properly, and replace it if necessary; move the sensor to a new site; or use a disposable adhesive sensor that may tolerate more motion.
- Replace the sensor.

PROBLEM: The SpO₂ value or the pulse rate changes rapidly; the pulse amplitude bar is erratic.
CAUSE:
- Excessive patient motion may be making it impossible for the SpO₂ function to find a pulse pattern.
- An electrosurgical unit (ESU) may be interfering with performance.

SOLUTION:
Check the patient.
- If possible, keep the patient still; check whether the sensor is applied securely and properly, and replace it if necessary; move the sensor to a new site; use a sensor that tolerates more motion.
If an ESU is interfering:

- Move the SpO₂ cable as far from the ESU as possible.
- Plug the Monitor and the ESU into different AC circuits.
- Move the ESU ground pad as close to the surgical site as possible.
- The sensor may be damp or may need to be replaced with a new sensor.
- If the patient weighs less than 3 kg or more than 40 kg, apply an OXISENSOR N-25 oxygen transducer to an appropriate site. This sensor has added protection against electrosurgical interference.

PROBLEM: The oxygen saturation measurement does not correlate with the value calculated from a blood gas determination.

CAUSE:

- The SpO₂ calculation may not have correctly adjusted for the effects of pH; temperature; CO₂; fetal hemoglobin; or 2,3-DPG.
- Accuracy can be affected by incorrect sensor application or use; intravascular dyes; bright light; excessive patient movement; venous pulsations; electrosurgical interference; and placement of a sensor on an extremity that has a blood pressure cuff, arterial catheter, or intravascular line.

SOLUTION:

- Check that calculations have been corrected appropriately for the relevant variable. In general, calculated saturation values are not as reliable as direct laboratory hemoximeter measurements.
- If there is excessive light, cover the sensor with opaque material.
- Circulation distal to the sensor site should be checked routinely. The site must be inspected every 8 hours to ensure adhesion, skin integrity, and correct optical alignment. If skin integrity changes, move the sensor to another site.
PROBLEM: A valid SpO₂ signal was present but has disappeared.

CAUSE:

- A BP determination on the same limb is in progress.

SOLUTION:

- An alarm message (No signal) will appear on the screen, and the audible alarm will sound immediately.

PROBLEM: A bad signal error has occurred.

CAUSE:

- Weak or "noisy" signal.

SOLUTION:

- Check the patient.
  - If possible, keep the patient still; check whether the sensor is applied securely and properly, and replace it if necessary; move the sensor to a new site; or replace the sensor.
  - Change sensor type.
  - Consider increasing perfusion using heat.
  - If there is excessive light, cover the sensor with opaque material.

PROBLEM: A sensor error indicating a bad sensor has occurred.

CAUSE:

- The sensor or cable may be defective, or the cabling may be improperly connected.

SOLUTION:

- Check the patient.
  - If possible, keep the patient still; check whether the sensor/cable is applied securely and properly, and replace it if necessary.
  - Disconnect and reconnect the sensor.

- Try to keep the patient still, or change the sensor site to one with less motion.

- Observe all instructions, warnings, and cautions in this manual and in the directions for use of the sensor.
Using the Menu System

Introduction
The PRO Monitor is equipped with a liquid crystal display (25) and a rotor (21). Used together, these allow the operator to view and edit most of the Monitor's parameters and functions. When the Monitor is in use, a number of option buttons appear on the liquid crystal display (LCD). The model of the Monitor determines which menu option buttons appear on the LCD. The number of buttons and the specific options depend on the menu level. The rotor provides the means of choosing menu options and changing monitor settings.

Liquid Crystal Display
The LCD is divided into three areas, each of which has a distinct function.

Menu Area
This area displays the menu buttons that are available for selection. Normal text in the menu area appears dark on a light background, while the text of selected buttons appears light on a dark background.

Area 2
This area displays BP and SpO₂ data and error and warning messages. The Display mode menu is used to select the data to be displayed.

Area 3
This area displays the time, the time lapsed since the last Auto BP determination (if in Auto BP mode), the battery icon (if operating on battery power, the time and battery icon toggle), and the BP and printer modes.

Note: In cold ambient temperatures (below 50 °F / 10° C), updates on the LCD can be delayed by approximately 1 second. This delay on the LCD does not affect the performance of the Monitor.
The model of the Monitor determines which menu option buttons appear on the LCD.

Notes
- The model of the Monitor determines which menu option buttons appear on the LCD.
**Rotor**

Rotating the rotor causes option buttons to be highlighted (light text on a dark background). Turning the rotor produces a click. Turning it clockwise moves the highlighting clockwise over the available buttons, while turning it counterclockwise reverses the direction of the highlighting. Pressing the rotor selects the highlighted button and produces an audible tone.

Some menus (e.g., Alarms) contain values that can be changed by the operator. After the value is highlighted, the user selects it by pressing the rotor. Turning the rotor clockwise will cause the value to increase, and turning the rotor counterclockwise will cause the value to decrease. Pressing the rotor again will confirm the changed value.

**Menu Tree**

The menu tree on the previous page shows all possible choices available within the menu structure, from the top level downward.

**Main Menu**

This menu is the top level menu. It is displayed when the Monitor is first switched on and after the rotor has been inactive for 2 minutes, unless the Monitor is in sleep mode (Pwr Sav).

```
Vitals  More...
Set BP  Alarms
Trend  Print
```
Using the Menu System

Vitals Button (UK: All Obs)
Selection of this button initiates a BP determination while allowing SpO₂ and predictive temperature determinations to be monitored and recorded (depending on Monitor model). When the Vitals determination is complete, a single “warble” sounds and all patient data are displayed on the LEDs and held for 2 minutes or until cleared by the user. The LCD shows:

```
Values Held

Clear  Print
```

Note: If the printer is in “Auto” mode, the Print button does not appear as an option.

Notes

- If the Monitor is performing a Vitals determination, the Vitals button cannot be selected.

- If a BP determination is in progress, the Vitals button cannot be selected.

- A Vitals determination is canceled if the BP determination is canceled.

- A Vitals determination can be canceled by pressing either the AUTO BP or START/STOP key.

- During the 2 minute freeze period, SpO₂ monitoring and alarms are suspended.

Clear
Selection of this button halts measurements and returns the user to the Main menu.

Note: If the SpO₂ plethysmograph is displayed on the LCD, the waveform pauses for 2 minutes or until the Clear button is selected. SpO₂ values are also retained the same manner as the BP and Temperature values.

Print
Selection of this button causes the current data to be printed.
Notes

- The Print button appears only when Print is set to Manual mode.
- If the printer is in Auto print mode, the data will be printed automatically.

More... Button
Selection of this button displays the More... menu. The More... menu has six options (depending on model of Monitor), most of which have submenus. For this reason, instructions for the More... button are in a separate section.

Set BP Button (UK: BP Mode)
Selection of this button displays the Auto, Stat, and Manual BP menu.

Auto
Selection of this option starts an Auto BP determination. When Auto Mode is selected, a number at the right of the Auto button indicates the time interval between each reading. To change the time interval, choose the box around the number and turn the rotor until the desired interval is reached. The interval can be set between 1 and 120 minutes (1, 2, 3, 4, 5, 10, 15, 20, 25, 30, 45, 60, 90, and 120 minutes). Press the rotor to confirm the setting. After power-off, the operating mode returns to the default setting of Manual. The default setting of Manual can be overridden to return to the previous user-selected setting (Auto or Manual) by selecting Set BP under the Service menu. To cancel an Auto BP determination, select the Manual button in the Set BP (UK: BP Mode) menu.

Manual
Selection of this option starts a Manual BP determination. After power-off, the operating mode returns to the default setting of Manual. The default setting of Manual can be overridden to return to the previous user-selected setting (Auto or Manual) by selecting Set BP under the Service menu.
Using the Menu System

Tgt Pressure
Selection of this option allows the user to set the BP target inflation pressure. The initial target pressure can be set between 100 and 250 mmHg in 5 mmHg increments. The factory default is 160 mmHg for adults and 110 for neonates. (This is indicated by “AUTO” at the end of adjustable range.) When using a neonate blood pressure cuff, if the target pressure is set to greater than 140 mmHg under the Set BP or Clinical menu, the Monitor automatically defaults to a target pressure of 110 mmHg. If the target pressure is set between 100 and 140 mmHg, then that setting is the target pressure that will be used. When the target pressure is changed, the next determination will use the new target inflation value if no systolic is available. Initial target pressure is restored to the factory default setting after power-off. The target pressure can be adjusted permanently in the Clinician menu of the Service mode (refer to “Press” in the “Using the Menu System” section).

Stat
Selection of this option allows the user to start Stat determinations. When Stat is selected, blood pressure is determined as many times as possible in 5 minutes.

Note: Alarm limits are disabled while in Stat mode.

Main
Selection of this button returns the user to the Main menu.

Alarms Button
Selection of this button displays the Alarms menu. This menu is used to adjust the violation limits for BP, Pulse Rate, and SpO₂. The values and ranges for these parameters are not stored when the Monitor is turned off. The user may edit the limits, but they are restored to the default values each time the Monitor is switched on. To permanently change the alarm limits, refer to “Alarms” under “Service Button” in the “Using the Menu System” section.
Auto Selection of this button updates the alarm limits on the LCD relative to the current parameter values. Pressing this button will automatically cancel any limit violation alarm that becomes invalid as a result of a limit change. Alarm limits are updated as follows:

### ALARM VOLUME

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Range</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Systolic High</td>
<td>35 - 245</td>
<td>180</td>
</tr>
<tr>
<td>Systolic Low</td>
<td>30 - 240</td>
<td>30</td>
</tr>
<tr>
<td>Diastolic High</td>
<td>15 - 195</td>
<td>130</td>
</tr>
<tr>
<td>Diastolic Low</td>
<td>10 - 190</td>
<td>15</td>
</tr>
<tr>
<td>MAP High</td>
<td>20 - 215</td>
<td>140</td>
</tr>
<tr>
<td>MAP Low</td>
<td>15 - 210</td>
<td>50</td>
</tr>
<tr>
<td>Heart Rate High</td>
<td>35 - 250</td>
<td>160</td>
</tr>
<tr>
<td>Heart Rate Low</td>
<td>30 - 245</td>
<td>40</td>
</tr>
<tr>
<td>SpO₂ High</td>
<td>51 - 100</td>
<td>Off</td>
</tr>
<tr>
<td>SpO₂ Low</td>
<td>50 - 99</td>
<td></td>
</tr>
</tbody>
</table>

**Volume**

Selection of this button displays the alarm volume submenu. The volume range is from 1 to 10, with 10 being the loudest. The alarm volume is stored when the Monitor is turned off and restored to the user’s preference each time the Monitor is switched on. Selection of the Check button allows the current volume setting to be heard. Selection of the Main button returns the user to the Main menu.
Using the Menu System

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Label</th>
<th>High Limit</th>
<th>Low Limit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Systolic</td>
<td>SYS</td>
<td>+30</td>
<td>SYS -30</td>
</tr>
<tr>
<td>Diastolic</td>
<td>DIA</td>
<td>+30</td>
<td>DIA -30</td>
</tr>
<tr>
<td>MAP</td>
<td>MAP</td>
<td>+30</td>
<td>MAP -30</td>
</tr>
<tr>
<td>Heart Rate</td>
<td>BPM</td>
<td>+30</td>
<td>BPM -30</td>
</tr>
<tr>
<td>SpO2</td>
<td>SpO2</td>
<td>+5*</td>
<td>SpO2 -5</td>
</tr>
</tbody>
</table>

* If the reading plus the limit is greater than the valid range of measurement (e.g., SpO2 +5 is greater than 100%), the valid range of measurement becomes the limits.

Notes

- In no case will the updated alarm limits be set beyond the valid limits in the preceding table.
- If no values are available, the limits will remain unchanged.

Main
Selection of this button returns the user to the Main menu.

Trend Button
Selection of this button displays the Trend mode menu.

Display
Selection of this button allows the operator to view the trend data.

Note: If the trend data have been lost (e.g., if the clock settings have been changed), the message “Trend Empty” will appear instead of the Newer, Older, and Print page buttons.
Newer and Older. These buttons may be used to move forward and backward through the recorded data. If no information is available, these buttons will not appear.

Print page. Selection of this button causes the displayed information to be printed. If no information is available, this button will not appear.

Main. Selection of this button returns the user to the Main menu.

Clear
Selection of this button produces an advisory that the trend will be lost. Choosing Yes will erase the trend memory. Choosing No will retain the trend memory. This button disappears from the menu while printing and when Trend is empty.

Print All
Selection of this button prints all the historical data available. When selected, this button temporarily changes to Cancel until the history has completed printing. Once printing is complete, the Cancel button returns back to the Print All button. This button disappears from the menu when Trend is empty.

Main
Selection of this button returns the user to the Main menu.
Using the Menu System

Print Button
Selection of this button displays the Print menu.

<table>
<thead>
<tr>
<th>PRINT MENU</th>
</tr>
</thead>
<tbody>
<tr>
<td>Auto / Man</td>
</tr>
<tr>
<td>History</td>
</tr>
</tbody>
</table>

Auto/Man
Pressing this button toggles between Automatic and Manual Printing modes. The current mode is displayed on Area 3 of the LCD. The Automatic mode prints the readings after each determination. The Manual mode, which is the factory default mode, requires the user to press the Now button to print the readings.

Now
Selection of this button causes the last readings of the available parameters to be printed. If no readings are available, the message “No reading” is printed for that parameter. An error message appears if there is no paper in the printer.

History
Selection of this button causes the entire contents of the trend memory to be printed. When selected, this button temporarily changes to Cancel until the history has completed printing.

Main
Selection of this button returns the user to the Main menu.

More... Menu
This menu is used to set the various operating modes of the Monitor.
LEDs. Sleep mode is available only if the Monitor is operating from its battery. Sleep mode conserves power while the Monitor is not in use. Once the Monitor is in Sleep mode, the user can return it to normal operation by touching any button or the rotor.

**SpO₂ Button (Models 300 and 400)**
Selection of this button displays the SpO₂ mode menu, which is used to set the SpO₂ pulse tone volume.

```
Volume
5
Main
```

**Volume**
The pulse tone volume can be set in the range of Off to 9. The value Off should be selected if no pulse tone is desired. The volume setting is stored when the Monitor is turned off and is restored to the user’s preference each time the Monitor is switched on.

**Main**
Selection of this button returns the user to the Main menu.

**Config Button**
Selection of this button displays the Config mode menu, which allows the Power Save mode and time to be adjusted.

```
Pwr Sav  Time
Rotor  Main
```

**Pwr Sav (Sleep Mode)**
Selection of this button allows the operator to specify the time, in minutes, that elapses before the Monitor goes into "sleep" mode (LEDs blanked and LCD displaying values from LEDs). Sleep mode is available only if the Monitor is operating from its battery. Sleep mode conserves power while the Monitor is not in use. Once the Monitor is in Sleep mode, the user can return it to normal operation by touching any button or the rotor.
Using the Menu System

The Monitor enters Sleep mode only if the following are true:
- No alarm is active
- SpO₂ is not actively reporting patient statistics
- The keys and rotor have not been used for the preset time
- The Monitor is running from its battery
- No determinations are in progress
- The Monitor has been running from the battery for the entire preset time

The Monitor awakens from Sleep mode if any of the following occur:
- The rotor is turned or pressed
- Any of the keys are pressed
- An alarm condition is issued
- The battery supply level becomes discharged to a critical level
- A mains or suitable DC supply is connected
- An Auto BP or Temp determination starts
- An SpO₂ signal is detected
- A BP determination is started through the host comm

Time
Selection of this button allows the operator to change the internal time and date of the Monitor. The clock, which is maintained by an internal battery after power down, uses 24-hour format. The date is in the British format of dd/ mmm/yyyy; however, to avoid confusion the month number has been substituted with a three-letter abbreviation. Leap years are calculated automatically.
Sp02 data are available) and any error or warning messages that may appear. The Display mode setting is maintained when the Monitor is switched off and on.

NIBP is selected, Area 2 of the LCD will remain blank except for the pulse amplitude bar (if SpO2, nor 3 SpO2, or BP data. If neither 1 Main Selection of this button returns the user to the Main menu.

Display Button
Selection of this button displays the Display mode menu. This menu is used to specify whether Area 2 of the LCD will display SpO2 or BP data. If neither SpO2 nor 3 NIBP is selected, Area 2 of the LCD will remain blank except for the pulse amplitude bar (if SpO2 data are available) and any error or warning messages that may appear. The Display mode setting is maintained when the Monitor is switched off and on.
Using the Menu System

SpO₂ Pleth
When this option is checked and SpO₂ data are available, the plethysmograph waveform and the pulse amplitude bar will be displayed.

3 NIBP
When this option is checked, the last 3 NIBP readings will be displayed. If SpO₂ data is available, the pulse amplitude bar will also be displayed.

Main
Selection of this button returns the user to the Main menu.

Service Button
Selection of this button displays a keypad that allows the clinician to access some parts of the Service mode menu. To access the Clinician menu, use the rotor to select the numbers 1, 2, 3, 4 sequentially.

Notes

* SpO₂ is automatically disabled when entering Service mode.

* Service modes that affect the calibration or alignment of the instrument are not available to the user. These modes are described in the Service Manual.
Clinician Menu

Press. Selection of this button displays a panel for setting the default BP target inflation pressure. Adjusting the default target pressure will automatically update the current inflation target pressure and will be used for the next reading. The range of adjustment is 100 mmHg to 180 mmHg, and the setting is retained when the Monitor is turned off.

The initial target pressure can be set between 100 and 180 mmHg in 5 mmHg increments. The factory default is 160 mmHg for adults and 110 for neonates. This is indicated by the “AUTO” label at the end of the adjustable range. When the target pressure is changed, the next determination will use the new target inflation value if no systolic is available. When adjusted under the Clinician menu of the Service mode, the target pressure is adjusted permanently.

OK. Selection of this button returns the user to the Clinician menu.

Temp. Selection of this button displays the temperature submenu, which allows the user to choose the temperature label. When C (Celsius) is selected, the °C indicator lights. When F (Fahrenheit) is selected, the °F indicator lights.
Using the Menu System

C or F. Selection of this button toggles the temperature display between Celsius and Fahrenheit and produces an advisory that the trend will be lost. Choosing Yes will cause the Monitor to accept the new temperature label and erase the trend memory. Choosing No will cause the Monitor to retain the existing temperature label and the trend memory.

TREND WILL BE LOST

OK. Selection of this button returns the user to the Clinician menu.

Info. Selection of this button causes the most recent calibration date of the NIBP system to be displayed. Selection of OK returns the user to the Clinician menu.

Last Calibration
NIBP  14-Jul-1999

OK. Selection of this button returns the user to the Clinician menu.

More... Selection of this button displays the More... menu, which allows the user to permanently change default mode settings.
Trend. Selection of this button displays the message: **Automatically clear trend on power-up?** Selection of **Yes** overrides the default setting by clearing all trends on power-up and returns the Monitor to the More... menu. Selection of **No** retains the default setting by saving all trends after power-off and returns the Monitor to the More... menu. Selection of **Cancel** returns the user to the More... menu.

Print. Selection of this button displays the message: **Restore Print mode on power-up?** Selection of **Yes** restores the Print mode to the default setting (previous user-selected mode) after power-off and returns the Monitor to the More... menu. Selection of **No** restores the Print mode to the Manual mode after power-off and returns the Monitor to the More... menu. Selection of **Cancel** returns the Monitor to the More... menu.

Set BP. Selection of this button displays the message: **Restore BP mode on power-up?** Selection of **Yes** restores the BP mode to previous user-selected mode after power-off and returns the Monitor to the More... menu. Selection of **No** restores the BP mode to the default setting of Manual after power-off and returns the Monitor to the More... menu. Selection of **Cancel** returns the Monitor to the More... menu.

Alarms. Selection of this button displays the message: **Enter alarm configuration mode?** Selection of **No** returns the Monitor to the More... menu. Selection of **Yes** brings up the Alarms menu. Selection of **Reset** changes all alarm limits back to the factory defaults and returns the Monitor to the More... menu. Selection of **Save** permanently saves the user-selected alarm limits and returns the Monitor to the More... menu. Selection of **Cancel** returns the Monitor to the More... menu.
Using the Menu System

Main. Selection of this button returns the user to the Main menu.

Silence. Selection of this button will cause all alarms except the FAILSAFE alarm to be muted. A confirmation menu will appear in Area 2 of the LED. Selection of either Yes or No returns the user to the Clinician mode menu. If silence is confirmed, the Alarm Silence button (26) illuminates and alarms are permanently muted. If silence is not confirmed, the alarm will be audible.

Caution: Alarms will be muted until either the Monitor is switched off and on again or the Alarm Silence button (26) is pressed.

Confirm silence

Yes

No

Main. Selection of this button returns the user to the Main menu.
Error and Warning Messages
The error panel appears in Area 2 of the LCD and indicates the error and its code, if it has one. In this example, a limit violation alarm (which has no error code) has occurred. A list of alarm error messages and their codes is in Appendix B.

Alarm conditions are addressed in two ways: the Alarms button and OK button.

Alarms Button
Selection of this button takes the user to the Alarms menu, where the alarm limits can be adjusted. This button is available only when a parameter alarm limit has been violated.

OK Button
Selection of this button acknowledges the error. The Monitor clears the identified error and then returns the user to the Main menu.
Appendix A

**Technical Specifications**

**BP**

Cuff Pressure Range (Normal operating range)

<table>
<thead>
<tr>
<th></th>
<th>Adult</th>
<th>Neonate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Systolic</td>
<td>30 - 245</td>
<td>40 - 140</td>
</tr>
<tr>
<td>MAP</td>
<td>15 - 215</td>
<td>30 - 115</td>
</tr>
<tr>
<td>Diastolic</td>
<td>10 - 195</td>
<td>20 - 100</td>
</tr>
</tbody>
</table>

Default Target: Cuff Inflation

<table>
<thead>
<tr>
<th></th>
<th>Adult</th>
<th>Neonate</th>
</tr>
</thead>
<tbody>
<tr>
<td>160 ± 15</td>
<td>110 ± 15</td>
<td></td>
</tr>
</tbody>
</table>

Target Cuff Inflation: Adjustment Range (in 5 mmHg increments)

<table>
<thead>
<tr>
<th></th>
<th>Adult</th>
<th>Neonate</th>
</tr>
</thead>
<tbody>
<tr>
<td>100 to 250</td>
<td>100 to 140</td>
<td></td>
</tr>
</tbody>
</table>

Blood Pressure Measurement Range (mmHg)

<table>
<thead>
<tr>
<th></th>
<th>Adult</th>
<th>Neonate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Systolic</td>
<td>0 to 290</td>
<td>0 to 140</td>
</tr>
<tr>
<td>MAP</td>
<td>0 to 245</td>
<td>0 to 115</td>
</tr>
<tr>
<td>Diastolic</td>
<td>0 to 195</td>
<td>0 to 100</td>
</tr>
</tbody>
</table>

Blood Pressure Accuracy

Meets or exceeds ANSI/AAMI standard SP-10 (mean error ≤5 mmHg, standard deviation ≤8 mmHg)

Maximum Determination Time

<table>
<thead>
<tr>
<th></th>
<th>Adult</th>
<th>Neonate</th>
</tr>
</thead>
<tbody>
<tr>
<td>120 s</td>
<td></td>
<td>85 s</td>
</tr>
</tbody>
</table>

Overpressure Cutoff

<table>
<thead>
<tr>
<th></th>
<th>Adult</th>
<th>Neonate</th>
</tr>
</thead>
<tbody>
<tr>
<td>300 to 330</td>
<td>150 to 165</td>
<td></td>
</tr>
</tbody>
</table>

Pulse Rate Range

<table>
<thead>
<tr>
<th></th>
<th>Adult</th>
<th>Neonate</th>
</tr>
</thead>
<tbody>
<tr>
<td>30 to 200</td>
<td>30 to 200</td>
<td></td>
</tr>
</tbody>
</table>

Pulse Rate Accuracy

± 3.5%

**IVAC* Temperature**

Scale

<table>
<thead>
<tr>
<th></th>
<th>Fahrenheit (F)</th>
<th>Celsius (C)</th>
</tr>
</thead>
</table>

Range

Predictive mode

<table>
<thead>
<tr>
<th></th>
<th>Adult</th>
<th>Neonate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Max:</td>
<td>42.2° C; 108.0° F</td>
<td>31.6° C; 88.9° F</td>
</tr>
<tr>
<td>Min:</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*IVAC is a trademark of ALARIS Medical Systems
Monitor mode

<table>
<thead>
<tr>
<th>Functional Oxygen Saturation:</th>
<th>0 to 100%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Accuracy</td>
<td>0 to 69%: unspecified</td>
</tr>
</tbody>
</table>

**Adult Accuracy (70% - 100%)**

<table>
<thead>
<tr>
<th>NELLCOR Sensor</th>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>OXICLIQ-P pediatric sensor</td>
<td>2.5 digits</td>
</tr>
<tr>
<td>OXICLIQ-I infant sensor</td>
<td>2.5 digits</td>
</tr>
<tr>
<td>OXICLIQ-A adult sensor</td>
<td>2.5 digits</td>
</tr>
<tr>
<td>OXICLIQ-N neonatal/adult sensor</td>
<td>2.5 digits</td>
</tr>
<tr>
<td>OXIBAND pediatric/infant sensor</td>
<td>3.0 digits</td>
</tr>
<tr>
<td>OXIBAND adult/neonatal sensor</td>
<td>3.0 digits</td>
</tr>
<tr>
<td>DURA-Y ear clip</td>
<td>3.5 digits</td>
</tr>
<tr>
<td>REFLECTANCE sensor</td>
<td>3.5 digits</td>
</tr>
<tr>
<td>DURASENSOR adult</td>
<td>3.5 digits</td>
</tr>
<tr>
<td>PEDI-CHECK pediatric spot-check clip</td>
<td>3.5 digits</td>
</tr>
<tr>
<td>OXISENSOR II D-20 pediatric sensor</td>
<td>2.0 digits</td>
</tr>
<tr>
<td>OXISENSOR II D-25 adult sensor</td>
<td>2.0 digits</td>
</tr>
<tr>
<td>OXISENSOR II N-25 neonatal/adult sensor</td>
<td>2.0 digits</td>
</tr>
<tr>
<td>OXISENSOR II I-20 infant sensor</td>
<td>2.0 digits</td>
</tr>
<tr>
<td>OXISENSOR II D-25L adult sensor, long cable</td>
<td>2.0 digits</td>
</tr>
</tbody>
</table>

**Monitor mode accuracy**

<table>
<thead>
<tr>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>± 0.1°C</td>
</tr>
<tr>
<td>± 0.2°F  (when tested in a calibrated liquid bath; meets ASTM E1112, Table 1, in range specified)</td>
</tr>
</tbody>
</table>

**Predictive mode accuracy**

<table>
<thead>
<tr>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>± 1.0°F</td>
</tr>
<tr>
<td>± 0.6°C</td>
</tr>
</tbody>
</table>

**Determination time**

less than 60 seconds

**Use only IVAC probes and probe covers. The size, shape, and thermal characteristics of the probe covers can affect the performance of the instrument. Inaccurate readings or retention problems may occur unless IVAC probes and probe covers are used. Refer to Appendix D for reorder codes.**
### Neonatal Accuracy (70% - 100%)
When sensors are used on neonatal subjects as recommended, the specified accuracy range is increased by ±1 digit to account for the theoretical effect on oximeter measurements of fetal hemoglobin in neonatal blood (e.g., N-25 accuracy on neonates is ±3, rather than ±2).

**Note:** Refer to NELLCOR's sensor specifications.

<table>
<thead>
<tr>
<th>Specification</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Pulse Rate Range</strong></td>
<td>30 to 250 beats/min</td>
</tr>
<tr>
<td><strong>Accuracy</strong></td>
<td>±3 beats/min</td>
</tr>
<tr>
<td><strong>Audible Indicator</strong></td>
<td>Pitch changes with saturation; volume selectable from 0 (off) to 9</td>
</tr>
<tr>
<td><strong>Waveforms</strong></td>
<td>Pulse plethysmograph waveform on LCD gain compensated</td>
</tr>
<tr>
<td><strong>Sensor Connect/Disconnect From Patient</strong></td>
<td>Monitor will detect attachment or disconnection of sensor from patient within 15 s</td>
</tr>
<tr>
<td><strong>Pulse Detection</strong></td>
<td>Monitor will detect pulse or enter no signal state within 15 s of being attached to patient</td>
</tr>
<tr>
<td><strong>Loss of Pulse</strong></td>
<td>Monitor will detect loss of pulse from patient and enter no signal state within 10 s</td>
</tr>
</tbody>
</table>
| **Sensor Light Source**              | Infrared: 920 nm (nominal)  
Red: 660 nm (nominal) |
| **Power Dissipation**                | Infrared: 22.5 mW (max)  
Red: 30 mW (max) |
### Mechanical

**Dimensions**
- Height: 9.8 in (25.0 cm)
- Width: 9.8 in (24.8 cm)
- Depth: 6.9 in (17.5 cm)

**Weight, Including Battery**
- 7.8 lb (3.5 kg)

**Mountings**
- Self-supporting on rubber feet or pole mountable

**Portability**
- Carried by recessed handle or pole mounted

**Classification Information**
- Mode of operation: continuous
- Degree of protection against harmful ingress of water: Drip-proof IPX1

### Power Requirements

AC input voltage: 100-240 VAC, 50 / 60 Hz (nominal)
90 ~ 253 VAC, 47 ~ 63 Hz (range), 50VA.
Protection against electrical shock: Class 1
DC input voltage: 24 VDC (nominal), 12-30 VDC, 36VA, supplied from a source conforming to IEC 601-1.
AC input is protected by two internal fuses, replaceable by qualified service personnel only. DC input line is protected by an internal auto-resetting fuse.
Battery: 12 volt, 2.3 amp-hours protected by internal auto-resetting fuse.
Minimum operation time:
2 hrs (5 min cycle with adult cuff at 25 °C with power save mode
Appendix A

enabled) from full charge. Time for full recharge: 1 hr 50 min from full discharge when the Monitor is switched off and 8 hrs when the Monitor is switched on.

**Environmental**

- **Operating Temperature**: +5 °C to +40 °C (+41 °F to +104 °F)
- **Operating Atmospheric Pressure**: 700 hPa to 1060 hPa
- **Storage Temperature**: -20 °C to +50 °C (-4 °F to +122 °F)
- **Storage/Transportation Atmospheric Pressure**: 500 hPa to 1060 hPa
- **Humidity Range**: 0% to 95% noncondensing
- **Radio Frequency**: Complies with IEC Publication 601-1-2 (April 1993) Medical Electrical Equipment, Electromagnetic Compatibility Requirements and Tests and CISPR 11 (Group 1, Class A) for radiated and conducted emissions.

The DINAMAP® PRO Monitor is protected against vertically falling drops of water and conforms with the IEC 529 standard at level of IPX1. No harmful effects will come of vertically falling drops of water making contact with the Monitor.

**IPX1**
Appendix B

Alarm Codes
All alarm indications are accompanied by an audible signal unless Alarm Silence is selected.

A microprocessor system failure will generate a high-pitched audible alarm regardless of the setting of the Alarm Silence switch.

There are three categories of alarms: patient alarms, system alarms, and failsafe alarm.

Patient Alarms
Patient alarms include those alarms issued when the patient's systolic pressure, diastolic pressure, pulse rate, or oxygen saturation is outside the set limits. Whenever one of these conditions occurs, the associated display (SYSTOLIC, MAP, DIASTOLIC, PULSE, or SpO₂) will flash the most recent reading and an audible alarm will be issued.

Pressing the Alarm Silence switch (causing the integral LED to be lit) silences the audible alarm for 2 minutes, but the alarm display reading and SILENCE LED indicator will continue to flash at the same rate.

System Alarms
System alarms alert the operator to certain abnormal conditions or internal system failures. Pressing the rotor cancels the alarm information box which is displayed on the LCD. Codes for different procedural and system alarms are on the next page.

Failsafe Alarm
The failsafe alarm, which is the most powerful alarm of the PRO Monitor, indicates a serious failure of the Monitor. This alarm occurs immediately upon any failure of a self-test and indicates system failure. When the failsafe alarm occurs, the Monitor disables all features to ensure patient safety.
### Hierarchy of Alarms

Alarms in the DINAMAP® PRO Monitor are in three priority levels. They are:

<table>
<thead>
<tr>
<th>Alarm</th>
<th>Priority Level</th>
</tr>
</thead>
<tbody>
<tr>
<td>Failsafe</td>
<td>1</td>
</tr>
<tr>
<td>Patient and system</td>
<td>2 (High priority alarm)</td>
</tr>
<tr>
<td>Low battery</td>
<td>3</td>
</tr>
</tbody>
</table>

The Priority 1 alarm (i.e., Failsafe) will override any other alarm. Priority 2 alarms will override only the low battery alarm. The low battery alarm will not override any other alarm.
<table>
<thead>
<tr>
<th>Alarm Code</th>
<th>LED Display</th>
<th>LCD Display</th>
<th>Audible Tone and Volume</th>
<th>Effect of Alarm Silence Switch</th>
<th>Effect of Clear via SelectKnob</th>
<th>Probable Cause</th>
</tr>
</thead>
<tbody>
<tr>
<td>N99</td>
<td>No change</td>
<td>N99 - NIBP FAILED</td>
<td>High priority alarm. Volume adjustable</td>
<td>2 minutes silence</td>
<td>Clear</td>
<td>Unable to make an NIBP determination due to insufficient signal</td>
</tr>
<tr>
<td>N55</td>
<td>No change</td>
<td>N55 - TIMEOUT: PRESS</td>
<td>High priority alarm. Volume adjustable</td>
<td>2 minutes silence</td>
<td>Clear</td>
<td>One cuff pressure for &gt; 1 minute. Motion artifact</td>
</tr>
<tr>
<td>N44</td>
<td>No change</td>
<td>N44 - TIMEOUT: TOTAL</td>
<td>High priority alarm. Volume adjustable</td>
<td>2 minutes silence</td>
<td>Clear</td>
<td>Determination time &gt; 2 minutes. Motion artifact</td>
</tr>
<tr>
<td>N33</td>
<td>No change</td>
<td>N33 - TIMEOUT: INFL</td>
<td>High priority alarm. Volume adjustable</td>
<td>2 minutes silence</td>
<td>Clear</td>
<td>Inflation time &gt; 40 seconds or air leak detected</td>
</tr>
<tr>
<td>N00</td>
<td>No change</td>
<td>N00 - OVER PRESS</td>
<td>High priority alarm. Volume adjustable</td>
<td>2 minutes silence</td>
<td>Clear</td>
<td>Overpressure detected</td>
</tr>
</tbody>
</table>

**Procedural and Error Alarm Codes**
<table>
<thead>
<tr>
<th>Alarm Code</th>
<th>LED Display</th>
<th>LCD Description</th>
<th>Audible Tone and Volume</th>
<th>Effect of Alarm Silence Switch</th>
<th>Effect of Clear via SelectKnob</th>
<th>Probable Cause</th>
</tr>
</thead>
<tbody>
<tr>
<td>P55</td>
<td>No change</td>
<td>P55 - SpO₂ NO SIGNAL</td>
<td>High priority alarm. Volume adjustable</td>
<td>2 minutes silence</td>
<td>Clear</td>
<td>No or very low SpO₂ signal. Check or reposition sensor</td>
</tr>
<tr>
<td>P00</td>
<td>No change</td>
<td>P00 - NO SpO₂ SENSOR</td>
<td>High priority alarm. Volume adjustable</td>
<td>2 minutes silence</td>
<td>Clear</td>
<td>SpO₂ sensor not connected. No sensor code detected. Sensor failure</td>
</tr>
<tr>
<td>No Code</td>
<td>No change</td>
<td>SpO₂ PLACEMENT?</td>
<td>High priority alarm. Volume adjustable</td>
<td>2 minutes silence</td>
<td>Clear</td>
<td>SpO₂ signal weak or noisy. Sensor failure</td>
</tr>
<tr>
<td>No Code</td>
<td>Values zeroed</td>
<td>SpO₂ CABLE?</td>
<td>High priority alarm. Volume adjustable</td>
<td>2 minutes silence</td>
<td>Clear</td>
<td>SpO₂ sensor or cable possibly defective. Cable not connected properly</td>
</tr>
<tr>
<td>E33</td>
<td>No change</td>
<td>E33 - TEMP: FAIL</td>
<td>High priority alarm. Volume adjustable</td>
<td>2 minutes silence</td>
<td>Clear</td>
<td>Temperature probe not connected or inoperable</td>
</tr>
<tr>
<td>E00</td>
<td>No change</td>
<td>E00 - TEMP: FAIL</td>
<td>High Priority alarm. Volume adjustable</td>
<td>2 minutes silence</td>
<td>Clear</td>
<td>Predictive temperature determination &gt; 60 sec or attempting axillary temp</td>
</tr>
<tr>
<td>No Code</td>
<td>Blank</td>
<td>HIGH TEMP</td>
<td>High Priority alarm. Volume adjustable</td>
<td>2 minutes silence</td>
<td>Clear</td>
<td>Predictive temperature exceeds upper range</td>
</tr>
<tr>
<td>Alarm Code</td>
<td>LED Display</td>
<td>LCD Description</td>
<td>Audible Tone and Volume</td>
<td>Effect of Alarm Silence Switch</td>
<td>Effect of Clear via SelectKnob</td>
<td>Probable Cause</td>
</tr>
<tr>
<td>------------</td>
<td>-------------</td>
<td>-------------------------------</td>
<td>-------------------------</td>
<td>-------------------------------</td>
<td>-------------------------------</td>
<td>--------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>No Code</td>
<td>No Change</td>
<td>LOW BATTERY, Flashing battery icon</td>
<td>3 beeps every 10 seconds, adjustable volume</td>
<td>2 minutes silence</td>
<td>No effect</td>
<td>Replace or recharge battery. From onset of alarm. 5 NIBP measurements available. Beep rate increases linearly as battery discharges</td>
</tr>
<tr>
<td>No Code</td>
<td>Blank</td>
<td>LOW BATTERY - SYSTEM DISABLED</td>
<td>Steady tone, maximum volume</td>
<td>No effect</td>
<td>No effect</td>
<td>Replace or recharge battery. NIBP measurement disabled</td>
</tr>
<tr>
<td>No Code</td>
<td>No Change</td>
<td>PRINTER - NO PAPER</td>
<td>High Priority alarm. Volume adjustable</td>
<td>2 minutes silence</td>
<td>Clear</td>
<td>Paper ran out or printer door open</td>
</tr>
<tr>
<td>No Code</td>
<td>Values Posted</td>
<td>NIBP RANGE ERROR</td>
<td>High Priority alarm. Volume adjustable</td>
<td>2 minutes silence</td>
<td>Clear</td>
<td>NIBP algorithm returned value outside specified accuracy range</td>
</tr>
<tr>
<td>Other: N, P, E, I, S</td>
<td>Blank</td>
<td>Error code, description</td>
<td>Steady tone, maximum volume</td>
<td>No effect</td>
<td>No effect</td>
<td>Internal system fault</td>
</tr>
</tbody>
</table>

**Procedural and Error Alarm Codes (cont.)**
**Appendix C**

**Principles of Noninvasive Blood Pressure Determination**

The oscillometric method of determining BP is accomplished by a sensitive transducer which measures cuff pressure and minute pressure oscillations within the cuff. The first determination sequence initially pumps up to a cuff pressure of about 160 mmHg for adult/pediatric patients, or 110 mmHg for neonates depending on the initial target pressure preset. After inflating the cuff, the Monitor begins to deflate it and measures systolic pressure, mean pressure, and diastolic pressure. When the diastolic pressure has been determined, the Monitor finishes deflating the cuff and updates the systolic, diastolic, and MAP displays on the front panel.

The Monitor deflates the cuff one step each time it detects two pulsations of relatively equal amplitude. The time between deflation steps depends on the frequency of these matched pulses (pulse rate of the patient). However, if the Monitor is unable to find any pulse within several seconds, it will deflate to the next step. The process of finding two matched pulses at each step provides artifact rejection due to patient movement and greatly enhances the accuracy of the Monitor. The figure shows the BP determination sequence.
At each step the microprocessor stores cuff pressure, the matched pulse amplitude, and the time between successive pulses. The stepped deflation and matched pulse detection continues until diastolic pressure is determined or total cuff pressure falls below 7 mmHg. The Monitor then deflates the cuff (to zero detected pressure), analyzes the stored data, and updates the front panel displays.

The operating cycle is composed of four parts: inflation time, deflation time, evaluation time, and wait time. Wait time, which varies from mode to mode, is affected by the cycle time (Auto mode) or operator intervention (Manual mode). The figure shows the Basic Operating Cycle.

**BP Operating Cycle**

**Systolic Search**

If systolic pressure is not found, the Monitor can search at higher cuff pressures than the initial target pressure. If the determination is in a late stage, the Monitor will inflate the cuff to 70 mmHg above the initial target to get better data in the systolic region. If the determination is in an early stage, the Monitor will inflate the cuff to 50 mmHg above the initial target pressure. The maximum pressure allowed in systolic search is limited by the normal range for cuff pressures. In any operating mode, if a patient’s systolic pressure exceeds the inflation pressure of the Monitor, the Monitor will begin normal deflation sequence, detect the
Appendix C

absence of a systolic value, stop deflation, reinflate to a higher (than initial) inflation pressure (290 mmHg maximum), and resume normal deflation sequence. This additional inflation will occur only once per determination.

If a previous valid systolic pressure is displayed, and the new systolic pressure oscillations are compared with the previous valid determination and the Monitor “thinks” that the systolic was not obtained, the Monitor will inflate the cuff to a pressure of an additional 50 mmHg above the immediately preceding inflation. This additional inflation will occur only once per determination.

Do not use the auscultatory method to verify the accuracy of the Monitor. Because of differences in technique and technology, values may differ. The DINAMAP® Technology of the PRO Monitor compares BP values to an invasive arterial BP measurement technology. The auscultatory method uses audible sounds heard through a stethoscope and determines BP by the corresponding height of a column of mercury.

Invasive pressure monitoring directly measures the pressure exerted on a transducer and displays this pressure as a value. Noninvasive blood pressure monitoring is dependent on the flow of blood through the peripheral circulation.
# Appendix D

## Compatibility Table and Reorder Codes

<table>
<thead>
<tr>
<th>Description of Compatible Parts</th>
<th>Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>SOFT-CUF™, Infant</td>
<td>2500</td>
</tr>
<tr>
<td>SOFT-CUF™, Child</td>
<td>2501</td>
</tr>
<tr>
<td>SOFT-CUF™, Small Adult</td>
<td>2502</td>
</tr>
<tr>
<td>SOFT-CUF™, Adult</td>
<td>2503</td>
</tr>
<tr>
<td>SOFT-CUF™, Large Adult</td>
<td>2504</td>
</tr>
<tr>
<td>SOFT-CUF™, Thigh</td>
<td>2505</td>
</tr>
<tr>
<td>SOFT-CUF™, Neonatal Type 1</td>
<td>2521</td>
</tr>
<tr>
<td>SOFT-CUF™, Neonatal Type 2</td>
<td>2422</td>
</tr>
<tr>
<td>SOFT-CUF™, Neonatal Type 3</td>
<td>2523</td>
</tr>
<tr>
<td>SOFT-CUF™, Neonatal Type 4</td>
<td>2524</td>
</tr>
<tr>
<td>SOFT-CUF™, Neonatal Type 5</td>
<td>2525</td>
</tr>
<tr>
<td>DURA-CUF® Cuff, Infant</td>
<td>2783</td>
</tr>
<tr>
<td>DURA-CUF® Cuff, Child</td>
<td>2781</td>
</tr>
<tr>
<td>DURA-CUF® Cuff, Small Adult</td>
<td>2779</td>
</tr>
<tr>
<td>DURA-CUF® Cuff, Adult</td>
<td>2774</td>
</tr>
<tr>
<td>DURA-CUF® Cuff, Large Adult</td>
<td>2791</td>
</tr>
<tr>
<td>DURA-CUF® Cuff, Thigh</td>
<td>2796</td>
</tr>
<tr>
<td>DURA-CUF® Cuff, Assortment Cuff Pack</td>
<td>2699</td>
</tr>
<tr>
<td>DURA-CUF® Cuff, Child Pack</td>
<td>2697</td>
</tr>
<tr>
<td>CLASSIC-CUF™, Infant</td>
<td>2618</td>
</tr>
<tr>
<td>CLASSIC-CUF™, Child</td>
<td>2613</td>
</tr>
<tr>
<td>CLASSIC-CUF™, Small Adult</td>
<td>2608</td>
</tr>
<tr>
<td>CLASSIC-CUF™, Adult</td>
<td>2603</td>
</tr>
<tr>
<td>CLASSIC-CUF™, Large Adult</td>
<td>2643</td>
</tr>
<tr>
<td>CLASSIC-CUF™, Thigh</td>
<td>2648</td>
</tr>
<tr>
<td>CLASSIC-CUF™, Neonatal Type 1</td>
<td>2638</td>
</tr>
<tr>
<td>CLASSIC-CUF™, Neonatal Type 2</td>
<td>2633</td>
</tr>
<tr>
<td>CLASSIC-CUF™, Neonatal Type 3</td>
<td>2628</td>
</tr>
<tr>
<td>CLASSIC-CUF™, Neonatal Type 4</td>
<td>2623</td>
</tr>
<tr>
<td>CLASSIC-CUF™, Neonatal Type 5</td>
<td>2619</td>
</tr>
<tr>
<td>12 Foot (approx. 3.7 m) Long Adult / Pediatric Hose</td>
<td>107365</td>
</tr>
<tr>
<td>24 Foot (approx. 7.3 m) Long Adult / Pediatric Hose</td>
<td>107366</td>
</tr>
<tr>
<td>12 Foot (approx. 3.7 m) Long Neonatal Hose</td>
<td>107368</td>
</tr>
<tr>
<td>12 Foot (approx. 3.7 m) Long A/P Hose Quick Discon.</td>
<td>88847</td>
</tr>
<tr>
<td>IVAC** Oral Temperature Probe</td>
<td>088012</td>
</tr>
<tr>
<td>IVAC** Rectal Temperature Probe</td>
<td>088013</td>
</tr>
<tr>
<td>IVAC** Temperature Probe Covers</td>
<td>088015</td>
</tr>
<tr>
<td>DINAMAP® PRO Monitor Operation Manual</td>
<td>776995*</td>
</tr>
<tr>
<td>DINAMAP® PRO Monitor Service Manual</td>
<td>777105*</td>
</tr>
</tbody>
</table>

*PRO Monitor unique parts

**IVAC is a trademark of ALARIS Medical Systems
12 Volt Lead Acid Battery 633132
Accessory Pole/Basket 3210
Accessory Base 3211
Power Converter 621262*
Printer Paper (box of 10) 089100*
Power Cable 316579
NELCOR SpO₂ Extension Cable SCP10*
NELCOR Finger Sensor DS100A
BP Cal Kit 320246

*PRO Monitor unique parts
NELCOR is a trademark of Mallickrodt, Inc.

### NEONATAL CUFF TYPE

<table>
<thead>
<tr>
<th>LIMB CIRCUMFERENCE</th>
<th>REFERENCE NUMBER</th>
<th>USE WITH HOSE NO.</th>
</tr>
</thead>
<tbody>
<tr>
<td>#1 3 cm - 6 cm</td>
<td>2521/2638</td>
<td>107368 Teal Green, Neonatal, 12 ft (366 cm)</td>
</tr>
<tr>
<td>#2 4 cm - 8 cm</td>
<td>2422/2633</td>
<td></td>
</tr>
<tr>
<td>#3 6 cm - 11 cm</td>
<td>2523/2628</td>
<td></td>
</tr>
<tr>
<td>#4 7 cm - 13 cm</td>
<td>2524/2623</td>
<td></td>
</tr>
<tr>
<td>#5 8 cm - 15 cm</td>
<td>2525/2619</td>
<td></td>
</tr>
</tbody>
</table>

### ADULT/PEDIATRIC CUFF TYPE

<table>
<thead>
<tr>
<th>LIMB CIRCUMFERENCE</th>
<th>REFERENCE NUMBER</th>
<th>USE WITH HOSE NO.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Infant 8 cm - 13 cm</td>
<td>2703/2500/2618</td>
<td>107365 Adult 12 ft (366 cm) Air Hose Gray (made with Clippard screw connectors at cuff end)</td>
</tr>
<tr>
<td>Child 12 cm - 19 cm</td>
<td>2781/2501/2613</td>
<td>or 107366 Adult 24 ft (732 cm) Air Hose Gray (made with Clippard screw connectors at cuff end)</td>
</tr>
<tr>
<td>Small Adult 17 cm - 25 cm</td>
<td>2779/2502/2608/2607</td>
<td>or 88847 Long A/P</td>
</tr>
<tr>
<td>Adult 23 cm - 33 cm</td>
<td>2774/2503/2603/2602</td>
<td>12 ft (366 cm) Air Hose (quick disconnect)</td>
</tr>
<tr>
<td>Large Adult 31 cm - 40 cm</td>
<td>2791/2504/2643/2642</td>
<td></td>
</tr>
<tr>
<td>Thigh 38 cm - 50 cm</td>
<td>2796/2505/2648</td>
<td></td>
</tr>
</tbody>
</table>
Appendix F

Maintenance

Cleaning the Monitor

The Monitor and accessories are to be kept clean and used according to the instructions provided here and in the Service Manual.

The exterior of the Monitor may be wiped clean with a soft cloth slightly dampened with mild detergents. The Monitor and accessories should be inspected once yearly for wear and damage.

- Do not immerse unit.
- Do not clean with isopropyl alcohol or other solvents.
- Do not immerse hoses.

Cuff Cleaning and Disinfection

General

The cuff must be thoroughly cleaned with the specified detergent before reuse. The additional use of household bleach as described below provides at least intermediate-level disinfection.

- Apply cuff hose plugs before cleaning.
- The following cleansing procedure was repeated 20 times on DURA-CUF® Blood Pressure Cuffs and once on SOFT-CUF® Blood Pressure Cuffs without affecting the performance of the cuff.
- While this procedure is adequate for cleaning/disinfection, it may not remove all stains.
- Do not immerse hoses.
- Do not immerse cuffs without prior application of cuff hose caps.

Materials

- Enzymatic detergent such as ENZOL* enzymatic detergent (US) or Cidezyme* enzymatic detergent (UK)
- Distilled water
- 10% solution of household bleach (5.25% sodium hypochlorite) in distilled water

*Trademark
• Soft cloths and soft-bristled brushes
• Spray bottles

Procedure
1. Prepare the enzymatic detergent according to the manufacturer’s instructions and the 10% bleach solution, in separate spray bottles.
2. Spray the detergent liberally on device. If the material is dried on, allow the cuff to sit for 1 minute. For soil on the soft part of the closure or the cuff itself, wipe the material off with a soft cloth. For persistent contamination on the soft part of the closure, use a soft-bristled brush to loosen particles. Rinse with copious amounts of distilled water. Repeat until no visible contamination remains. For soil on the hook part of the closure, use a soft-bristled brush to remove the material, and rinse with copious amounts of distilled water. Repeat until no visible contamination remains.
3. Spray the 10% bleach solution on the affected area until the area is saturated. Allow the cuff to sit for 5 minutes.
4. Wipe away any excess solution and rinse the cuff again with distilled water. Allow 2 hours for drying.

The user has the responsibility to validate any deviations from the recommended method of cleaning and disinfection.

For additional information on infection control procedures, contact Critikon Technical Support.

Temperature Devices
Do not immerse predictive temperature probes. The probe may be cleaned with an alcohol solution. Use a cloth or sponge—just damp, not wet—and avoid getting any liquid into the interior of the probe.

SpO₂ Sensors
Adhesive sensors are sterile and for single use only. Reusable sensors should be cleaned before reuse with a 70% alcohol solution. Do not immerse the sensor completely in water, solvents, or cleaning solutions (because the connector is not waterproof). Do not sterilize the sensor by irradiation, steam, or ethylene oxide. If
Appendix F

disposable sensors or their packaging are damaged, they must be disposed of as advised in Appendix F.

Storage and Battery Care
If it becomes necessary to store the Monitor for an extended period of time, first fully charge then remove the battery. Then store the Monitor and the battery in the original packaging materials.

Batteries should always be fully charged before being placed in storage. Even after 6 months of storage, a fully charged battery can retain about 80% of its charge. A fully charged battery in good condition will provide sufficient power to operate a Monitor for approximately 2 hours, including temperature and BP measurements made at 5-minute intervals.

It is best to keep the battery charged as fully as practical and never store the Monitor with the battery in a discharged condition. When the battery will no longer hold a charge, remove and replace it with one of the same part number. Failure to replace the battery with the same Critikon part number may result in shorter battery life.

To charge the battery, insert the plug from either the AC mains power cord or the AC-DC power converter into an appropriate AC outlet. The battery will charge regardless of the position of any switches.

Battery charging will take place as long as the Monitor remains connected to an external AC power source. A battery that is fully discharged can be fully recharged in 1 hour 50 minutes when the Monitor is switched off or 8 hours if the Monitor is switched on.

Cautions
• To ensure that the battery will be ready for portable operation, keep the Monitor connected to a mains supply whenever possible.

• Repeated failure to fully charge the battery will result in a significant reduction in battery life.
- The expected lifetime of the battery largely depends on the way in which the Monitor is used. If the battery is allowed to completely discharge before being fully recharged, the battery should survive around 200 recharge cycles. If the battery is used in such a way that it never becomes more than one third discharged and is fully recharged whenever possible, it can survive up to 1200 cycles. This means that by thoughtful usage, the lifetime of the battery can be extended up to six times.

Replacement batteries may be obtained from Critikon.

**Note:** The replacement part number of the battery is 63313. Do not use other types.

**Fuses**
The Monitor contains five fuses. Two AC line input fuses are mounted internally and are replaceable only by qualified service personnel. The remaining three fuses are auto-resetable and mounted within the Monitor. These fuses protect the low voltage DC input, the battery, and the +5 V output on the host port connector.

**Calibration**
Calibration of the BP parameter should be checked at least once a year or when there is doubt about the validity of the readings.

**Leak Testing**
A leak test of the BP parameter should be performed at least once a year or when there is doubt about the validity of the pressure readings.

**Caution:** Refer calibration and leak testing to qualified service personnel. Full calibration details are available in the DINAMAP PRO Monitor Service Manual, available from Critikon.

**Disposal of Product Waste**
As you use the PRO Monitor, you will accumulate solid wastes that require proper disposal or recycling. These include batteries, patient applied parts, and packaging material.
Appendix F

Batteries
Caution: Do not incinerate batteries.
The sealed, rechargeable backup battery contains lead and can be recycled. The rechargeable memory battery is of the Nickel Metal Hydride form. Discharge this battery prior to disposal. Place the battery in packaging which electrically isolates its contents. Do not puncture or place the battery in a trash compactor. Do not incinerate the battery or expose it to fire or high temperatures. Dispose in accordance with regional body controlled guideline.

Patient Applied Parts
Certain patient applied parts, such as those with adhesive (disposable SpO₂ sensors), are intended for single use and should be disposed of properly as medical waste in accordance with regional body controlled guideline.

Other patient applied parts, such as blood pressure cuffs, should be cleaned according to instructions. Inspect reusable applied parts for wear, replace as necessary, and dispose of used product as medical waste in accordance with regional body controlled guideline.

Packaging Material
Retain original packaging materials for future use in storing or shipping the Monitor and accessories. This recommendation includes corrugated shippers and inserts.

Whenever possible recycle the packaging of accessories and patient applied parts.