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Foreword

The Panorama Operating Instructions are intended to provide the information required to operate the Panorama Patient Monitoring Network. For additional information and technical assistance, please contact an area or regional Representative.

A general understanding of the features and functions of the Panorama Patient Monitoring Network and its components is a prerequisite for the proper use of this equipment. Therefore, do not operate this equipment before reading these instructions thoroughly, including all appropriate warnings and cautions.

CAUTION: US Federal law restricts the sale of this device to, by or on the order of a physician.

NOTE: Figures in this manual are provided for reference purposes only. Screens will likely differ based on the monitoring device configuration, licenses available, parameters selected and patient configuration of the Panorama Patient Monitoring Network.

Warnings, Cautions, and Notes

Please read and adhere to all of the warnings and cautions listed throughout this manual.

A WARNING is provided to alert the user to potentially serious outcomes (death, injury or serious adverse events) to the patient or the user.

A CAUTION is provided to alert the user that special care should be taken for the safe and effective use of the device. They will include actions to be taken to avoid effects on patients or users that will not be potentially life threatening or result in serious injury, but about which the user should be aware.

A NOTE is provided when additional general information is available.
Warnings

WARNING: Only qualified and trained personnel or Service Personnel should attempt to service the equipment. Service is defined as any activity requiring the cover to be removed for internal adjustments, parts replacements, repairs or software upgrades of any kind to insure compatibility.

WARNING: To insure compatibility with the operating system and applications software, use only approved components to repair any part of the Panorama Patient Monitoring Network. Use of unauthorized software, devices, accessories, or cables may render the application unsuitable for medical patient monitoring. It may also result in increased electromagnetic emissions or decreased immunity of the system.

WARNING: Route cables neatly. Ensure cables are not in the way of patient or hospital personnel.

WARNING: Do not incinerate batteries, possible explosion may occur.

WARNING: Some of the equipment on the Panorama Patient Monitoring Network must utilize the hospital emergency power system. Failure to do so will result in loss of monitoring during extended periods of power failure. The back-up power time period, for the Panorama Patient Monitoring Network, is limited.

WARNING: The 18.1" flat panel may tip over, if the display head is inclined to an angle greater than 45° backward tilt. If the user elects to have the display head inclined to an angle greater than 45° backward tilt, the flat panel must be attached to a secure mounting surface via three screw locations on the bottom of the base.

WARNING: Do not put MPSO (Multiple Portable Socket Outlets, i.e., multiple outlet extension cords) used with the Panorama Patient Monitoring Network equipment on the floor. Connect only approved accessories and components to the same MPSO as the Panorama Patient Monitoring Network equipment. Do not overload MPSOs. Use only MPSOs that comply with the requirements of IEC 60601-1-1.

WARNING: Incorrect setting or silencing of patient alarms can jeopardize patient safety.

WARNING: Operator should check that the current alarm presets are appropriate prior to use on each patient.

WARNING: A hazard can exist if different alarm presets are used for the same or similar equipment in any single area.

WARNING: When the All Non-High Alarms Pause/Off sidebar button is enabled, the Panorama Central Station will not monitor any priority 2 and priority 3 physiological alarms, and will not display numeric physiological or technical alarms for the selected patient tile.
WARNING: When the All Alarms Pause/Off sidebar button is enabled, the Panorama Central Station will not monitor any lethal or non-lethal arrhythmias and will not display numeric physiological or technical alarms for the selected patient tile.

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

WARNING: The Telepack with Nonin SpO2 module should only be used with Nonin-manufactured PureLight® pulse oximeter sensors. Using other manufacturers’ sensors can result in inaccurate pulse oximeter performance.

WARNING: Do not use the Telepack with Nonin SpO2 module in an explosive atmosphere or in the presence of flammable anesthetics or gases.

WARNING: Loss of SpO2 monitoring when using the Telepack with Nonin SpO2 module can result if any objects hinder the pulse measurement. Ensure that no blood flow restrictions (e.g. blood pressure cuff) hinder pulse measurements.

WARNING: The Telepack with Nonin SpO2 module does not meet defibrillation-proof requirement per IEC 60601-1.

WARNING: Operation of the Telepack with Nonin SpO2 module below the minimum amplitude of 0.3% modulation may cause inaccurate SpO2 results.

WARNING: The Panorama Telepack is NOT intended for use in a magnetic resonance imaging (MRI) environment. The Telepack must be physically removed if a patient enters the MRI environment; induced currents may potentially cause burns.

WARNING: The Panorama Patient Monitoring Network, including all related system components, is NOT intended for use in a magnetic resonance imaging (MRI) environment.

WARNING: Do not use a damaged SpO2 sensor.

WARNING: Using unapproved accessories, sensors, and cables may result in increased emission and/or decreased immunity of this device.

WARNING: As with all medical equipment, carefully route cables and connections to reduce the possibility of entanglement or strangulation.

WARNING: PACEMAKER PATIENTS - Indication of the heart-rate may be adversely affected by cardiac pacemaker pulses or by cardiac arrhythmias. Keep pacemaker patients under close surveillance. Refer to “ECG Performance Requirements” on page 16-2 for disclosure of the pacer rejection capability.

WARNING: PACEMAKER PATIENTS - Rate meters may continue to count the pacemaker rate during occurrences of cardiac arrest or some arrhythmias. Do not rely entirely upon rate meter alarms. Keep pacemaker patients under close surveillance.
WARNING: PACEMAKER PATIENTS – On ventricular paced patients, episodes of Ventricular Tachycardia may not always be detected. Do not rely entirely upon the system's automated arrhythmia detection algorithm. Keep pacemaker patients under close surveillance.

WARNING: Changing the patient alarm limits at the WorkStation will automatically change the patient alarm limits at the Panorama Central Station AND the bedside monitor (if applicable) for the selected patient tile.

WARNING: Electrode polarization: some electrodes may be subject to large offset potentials due to polarization. Use only electrodes as recommended by Mindray. Recovery time after application of defibrillator pulses may be especially compromised. Squeeze bulb electrodes commonly used for diagnostic ECG recording may be particularly vulnerable to this effect. Electrodes of dissimilar metals should not be used unless the amplifier can handle polarization potentials as high as 1 volt (V).

WARNING: To reduce the hazard of burns in the high-frequency surgical neutral electrode connection, the electrodes should not be located between the surgical site and the electro-surgical unit return electrode.
Cautions

CAUTION: For proper operation do not obstruct the fan air holes.

CAUTION: For proper operation use only hospital grade power cords.

CAUTION: For proper operation, do not use the Panorama Patient Monitoring Network equipment with a frayed or damaged power cord.

CAUTION: For proper operation never place fluids on top of this equipment. In case of accidental spillage, wipe clean immediately and have the system serviced to ensure no hazard exists.

CAUTION: Do not block the speakers. Set the volume levels so that alarms can be heard at all times, as described in this Operation Manual.

CAUTION: Dispose of single use items in accordance with hospital policy.

CAUTION: The bedside device should be disconnected from the Panorama Central Station when changing the CO₂ units of measurement.

CAUTION: For proper monitoring of patient information, do not block the operators' clear view of the Panorama Patient Monitoring Network data display(s).

CAUTION: If passwords are changed from the factory defaults, users MUST ensure that they document new passwords in the space provided in this manual. For additional information refer to “Passwords Tab” on page 9-29. Failure to recall user defined passwords will render the system UNUSABLE.

CAUTION: The Panorama Telepack and Instrument Radio are wireless devices and may be subject to intermittent signal dropout. Panorama wireless devices are designed to perform with a maximum cumulative signal dropout of one minute in a 24-hour period under ideal conditions. A crowded wireless environment or interference from other medical or non-medical devices, either intentional (e.g., VOIP phones, pagers, asset tracking systems, etc.) or unintentional (e.g., AC motors, microwave ovens, paper shredders, etc.), may result in a significantly increased amount of signal dropout experienced by any one or multiple wireless device(s).

CAUTION: Operation of the wireless product requires the prior coordination with a frequency coordinator designated by the FCC for the Wireless Medical Telemetry Service.

CAUTION: Changes or modifications not expressly approved by Mindray DS USA, Inc. could void the user’s authority to operate the equipment.

CAUTION: The Telepack and the Telepack with Nonin SpO₂ module should not be used when the Electro Surgical Unit (ESU) is active.

CAUTION: To avoid potential system issues, deactivate the touch screen before cleaning.
CAUTION: DO NOT spray glass cleaner directly on a display as it could possibly leak inside a non-sealed unit and cause damage.

CAUTION: Follow your hospital protocol for handling of blood and body fluids.

CAUTION: The use of the Telepack may affect the functioning of other medical electrical equipment operating in the same frequency band.

CAUTION: Conductive parts of electrodes and associated connectors for applied parts and neutral electrode should not contact other conductive parts and earth.

CAUTION: The Telepack may not function properly, if it was subjected to extreme mechanical forces.

CAUTION: The Telepack may not function if the batteries installed in the device are corroded. Use only approved batteries (P/N 0146-00-0077-01) in the Telepack and observe proper polarity. Installing batteries with the incorrect polarity may cause the batteries to become excessively warm.

CAUTION: The Telepack should not be attached to the patient when the device is being programmed into the system.

CAUTION: Ensure that the ECG lead cable is plugged into the port firmly, the end cap of the battery is firmly secured and the serial port is also secured with the protective plug to prevent fluids from entering into the Telepack.

CAUTION: When disconnecting ECG leads from the patient, do not pull on the leadsets. This will decrease the life of the leadsets. Grasp the connectors on the leadset and pull gently.

CAUTION: Ensure that the lead wires are not inadvertently caught in the bed rails. If this happens, the insulation may get cut or the leadset may break.

CAUTION: Using more than one type of electrode on the same patient should be avoided because of variations in electrical resistance.

CAUTION: The LEDs on the Telepack may not be clearly visible under brightly lit conditions. The lighting should be adjusted to ensure that all LED indicators are clearly visible.

CAUTION: The user should monitor the Panorama Central Station for messages from the Telepack that indicate a low battery condition.

CAUTION: Periodically press the CHECK button on the Telepack to verify the battery condition.

CAUTION: Some pacemakers may contain a respiratory sensor that may produce artifact on an ECG waveform.

CAUTION: Prior to sterilization, lead sets must be removed from the Telepack. The Telepack and/or the Telepack with Nonin SpO₂ must not be sterilized.

CAUTION: Visually inspect the battery compartment for any foreign object prior to use.
CAUTION: Ensure that the batteries are removed when the Telepack is being stored.

CAUTION: Tissue damage or inaccurate SpO2 measurements may be caused by incorrect sensor application or use, such as wrapping too tightly, applying supplemental tape, failing to inspect the sensor site periodically, or failing to position appropriately. Carefully read the sensor directions for use, the Panorama Operating Instructions, and all precautionary information before use.

CAUTION: The accuracy of the SpO2 measurement may be affected if the total cable length (including extension cables) is greater than 3.28 yards (3 meters).

CAUTION: Inaccurate SpO2 measurements may be caused by:

- Incorrect sensor application or use
- Significant levels of dysfunctional hemoglobins, (e.g., carboxyhemoglobin or methemoglobin)
- Intra-vascular dyes such as indocyanine green or methylene blue
- Exposure to excessive illumination such as surgical lamps (especially ones with a xenon light source), bilirubin lamps, fluorescent lights, infrared heating lamps, or excessive ambient light. In such cases, cover the sensor site with opaque material.
- Excessive patient movement
- Venous pulsations
- Electro-surgical interference
- Placement of a sensor on an extremity that has a blood pressure cuff, arterial catheter, or intra-vascular line.
- Cold extremities due to reduced circulation.
- Nail polish or fungus

CAUTION: In certain situations in which perfusion and signal strength are low, such as in patients with thick or pigmented skin, inaccurately low SpO2 readings will result. Verification of oxygenation should be made, especially in patients with chronic lung disease, before instituting any therapy or intervention.

CAUTION: Many patients suffer from poor peripheral perfusion due to hypothermia, hypovolemia, severe vasoconstriction, reduced cardiac output, etc. These symptoms may cause a loss in vital sign readings.

CAUTION: Prolonged and continuous monitoring may increase the risk of skin erosion and pressure necrosis at the site of the sensor. Check the SpO2 sensor site frequently to ensure proper positioning, alignment and skin integrity at least every eight (8) hours; with the Adult and Pediatric re-usable finger sensor, check every four (4) hours; for patients of poor perfusion or with skin sensitive to light, check every 2 - 3 hours; more frequent examinations may be required for different patients. Change the sensor site if signs of circulatory compromise occur.
CAUTION: The Telepack with Nonin SpO₂ module has motion tolerant software that minimizes the likelihood of motion artifact being misinterpreted as good pulse quality. In some circumstances, however, the Telepack with Nonin SpO₂ module may still interpret motion as good pulse quality. This applies to all available outputs (i.e. SpO₂, HR, Pleth, PPG).

CAUTION: A functional tester cannot be used to assess the accuracy of the pulse oximeter probe or a pulse oximeter monitor.

CAUTION: The Telepack with Nonin SpO₂ module complies with International Standard IEC 60601-1-2:2007 for electromagnetic compatibility for medical electrical equipment and/or systems. This standard is designed to provide reasonable protection against harmful interference in a typical medical installation. However, because of the proliferation of radio-frequency transmitting equipment and other sources of electrical noise in healthcare and other environments, it is possible that high levels of such interference due to close proximity or strength of a source might disrupt the performance of this device. Medical electrical equipment needs special precautions regarding EMC, and all equipment must be installed and put into service according to the EMC information specified in this manual.

CAUTION: Use the designated cleaning plug (P/N 0003-00-0055) when cleaning and disinfecting the Nonin SpO₂ module. If fluids enter the device, it may affect the functionality of the module or render it unusable.

CAUTION: In compliance with the European Directive on Waste Electrical and Electronic Equipment (WEEE) 2002/96/EC, do not dispose of this product as unsorted municipal waste.

CAUTION: It is important to notify Service personnel when the system event message “System Service required - Contact Mindray or Authorized Mindray Distributor (300).” is displayed. This message indicates that while data was being written to or read from the system hard drive, an error was detected. Until service personnel correct this issue, patient data may be lost (for example, demographic information, alarm configurations, and historical patient data). Real time parameter monitoring and alarm checking will not be affected by this message. This message will be displayed until reset by service personnel.

CAUTION: The operator is responsible for checking the compatibility of the monitor, probe and cable before use.

CAUTION: DANGER OF EXPLOSION IF BATTERY IS INCORRECTLY REPLACED. REPLACE ONLY WITH THE SAME OR EQUIVALENT TYPE RECOMMENDED BY THE MANUFACTURER. DISPOSE OF USED BATTERIES ACCORDING TO THE MANUFACTURER’S INSTRUCTIONS.

ATTENTION: IL Y A DANGER D’EXPLOSION S’IL Y A REMPLACEMENT INCORRECT DE LA BATTERIE, REMPLACER UNIQUEMENT AVEC UNE BATTERIE DU MÊME TYPE OU D’UN TYPE ÉQUIVALENT RECOMMANDÉ PAR LE CONSTRUCTEUR. METTRE AU REBUT LES BATTERIES USAGÉES CONFORMÉMENT AUX INSTRUCTIONS DU FABRICANT.)
CAUTION: If the Panorama system experiences a communication loss for more than 60 seconds, all Telepack ST and/or arrhythmia alarms that were active at the time of communication loss will not produce central station alarm notifications once communication is restored. This is due to the automatic relearn function of the ST and/or arrhythmia algorithm after communication is restored.

CAUTION: Line Isolation Monitor transients may resemble actual cardiac waveforms, thus inhibiting heart rate alarms. Check lead wires for damage and ensure good skin contact prior to and during use. Always use fresh electrodes and follow proper skin preparation techniques.
Indications for Use

Panorama Patient Monitoring Network

The indications for use for the Panorama Patient Monitoring Network include:

- Viewing real time patient clinical and demographic data
- Graphical and numeric trending of clinical data
- Storing and printing of clinical and demographic data
- Setting independent alarm limits for data sent by the bedside monitor

The clinical data displayed by the Panorama Patient Monitoring Network is obtained from one or more compatible physiological monitors and includes: ECG (Lethal and Non-Lethal Arrhythmia Detection, ST Segment Analysis, QT Analysis and heart rate (HR)), Invasive Blood Pressure (IBP), Non-Invasive Blood Pressure (NIBP), Blood Oxygenation (SpO₂), Pulse Rate (PR), Respiration Rate (RESP), Respiration Gases (O₂, CO₂), Temperature (TEMP), Anesthetic Gases (AG), Cardiac Output (C.O.), Bispectral Index (BIS), and Hemodynamics.

The Panorama Patient Monitoring Network is intended for use in a fixed location in a healthcare facility setting as a central viewing station. The Panorama Patient Monitoring Network is not intended to be directly connected to the patient at any time or installed in a patient’s vicinity.

The Panorama Network includes the Panorama Telemetry System which acquires and monitors physiological data for ambulating patients within a defined coverage area. The system processes the physiological data to detect various ECG arrhythmia events and select physiological parameter limit violations. The Panorama Telemetry System is intended for installation in a hospital or clinical environment to provide clinicians with patient physiological data, while allowing for patient mobility.

The physiological parameters monitored through telemetry include ECG (Lethal and Non-Lethal Arrhythmia Detection, ST Segment Analysis, QT Analysis and heart rate (HR)), blood oxygenation (SpO₂), Pulse Rate (PR). Received data is calculated at the Panorama Central Station, and can be displayed, trended, stored and printed at the Panorama Central Station.

The Panorama Patient Monitoring Network is intended for use under the direct supervision of a licensed healthcare practitioner.

Unpacking Information

The Panorama Patient Monitoring Network must be installed by a certified Service Technician or authorized distributor. Please contact your authorized Service Representative for additional information.
Symbols and Descriptions

<table>
<thead>
<tr>
<th>SYMBOL</th>
<th>DESCRIPTION</th>
<th>SYMBOL</th>
<th>DESCRIPTION</th>
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<tbody>
<tr>
<td>!</td>
<td>Attention, Consult Accompanying Documents / Refer to Manual</td>
<td></td>
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<td>⚡</td>
<td>Alternating Current (AC)</td>
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<td>Telemetry Network Wireless</td>
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<td>⚪</td>
<td>Electrical Hazard</td>
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<td>Mark of compliance with U.S. and Canadian safety standards requirement.</td>
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<td>Not protected (ordinary)</td>
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<td></td>
</tr>
<tr>
<td>IPX 4</td>
<td>Protection against splashing water</td>
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Technical Support
For technical assistance please contact your area or regional Service Representative.

Additional information relating to the installation, servicing and repair of the Panorama Patient Monitoring Network is provided in the following documents:

Panorama WMTS Installation Guide (P/N 0070-00-0571)
Panorama Service Manual (P/N 0070-00-0634)
Panorama Printer Configuration Manual (P/N 0070-00-0561)
1.0 System Overview

This section provides a high level view of the authorized components of the Panorama Patient Monitoring Network.

- Licensing Key
- Supported Devices
- Antivirus
- Troubleshooting
1.1 Licensing Key

The Panorama Central Station uses a licensing key to regulate the operation of the system.

The licensing key controls the following settings on the Panorama Central Station:

- Number of monitoring channels; up to 24
- Number of ST licenses
- Number of arrhythmia licenses
- Number of displays
- Storage hours of full disclosure (1 hour (default), 24 hours, 48 hours, 72 hours)
- 12-lead data management using a View 12™ card
- Activates the communication between the Panorama Central Station(s), the ViewStation(s), and the WorkStation(s)
- Paging

For additional information regarding licensing keys, please have a system administrator contact a Service Representative.
1.2 **Supported Devices**

The Panorama Patient Monitoring Network consists of two local area networks (LAN): a Panorama Central Network (C-LAN) and a Panorama Patient Network (E-LAN). The C-LAN manages stored patient data and controls the data flow between systems. The E-LAN manages real-time data by controlling the flow of data that is transmitted to/from the bedside monitors, and the Panorama Wireless Network.

The Network may include some or all of the following devices:

- Panorama Central Station
- Panorama ViewStation
- Panorama WorkStation
- Panorama Telemetry Server
- Panorama Wireless Transceiver
- Panorama Gateway
- Panorama Web Viewer
- Passport 2 Bedside Monitor
- Passport V Bedside Monitor
- Spectrum Bedside Monitor
- Spectrum OR Bedside Monitor
- DPM 6 and DPM 7 Bedside Monitors
- Passport 12M/Passport 17M Bedside Monitors
- Passport 8/Passport 12 Bedside Monitors
- T1 Monitor
- V 12/V 21 Monitors
- Panorama Telepack
- Network Printer

### 1.2.1 Panorama Central Station

The Panorama Central Station is a device that monitors patients, collects and stores data, and performs alarm notification. Multiple Panorama Central Stations can be networked with other Panorama Central Stations using Ethernet via the C-LAN. Communication between the C-LAN and the Panorama Patient Network is managed through the Panorama Central Station. A maximum of two displays can be used with each Panorama Central Station.

### 1.2.2 Panorama ViewStation and WorkStation

The Panorama ViewStation provides the capability to display patient data (including digital data and waveforms) with assigned monitoring equipment, to be viewed at a remote location. Patient data is transmitted from a bedside or ambulatory monitoring device to a central station on the C-LAN and is then transmitted to the ViewStation and the WorkStation, if these remote workstations exist on the network. The central station, which transmits the patient data, is referred to as the **host** central station.

All the patient information that can be displayed at the host central station can be seen at the ViewStation and WorkStation once the patient is assigned to a patient tile at the remote workstation. Minor configuration changes, such as Display and Wave Gain settings, can be made directly from the ViewStation. However, changes to patient configuration settings such as alarm settings or patient demographic information must be performed at the host central station. More significant configuration changes can be made at the WorkStation, such as changing alarm limits, alarm responses, and altering patient demographic information.
The ViewStation and the WorkStation may use either a single or dual display. Single display setups can accommodate eight (8) or twelve (12) patient tiles. Dual display setups can accommodate up to eight (8) patient tiles per display, showing a maximum of twenty-four (24) patient tiles between the two displays.

1.2.3 Panorama Telemetry Server

The Panorama Telemetry Server is provided when the Panorama Wireless Network is required. It acts as a bridge to connect wireless monitoring devices, such as Telepacks or wireless bedside monitors, to the Panorama Central Station. The Panorama Telemetry Server receives data from the wireless monitoring devices and converts it into a protocol recognized by the Panorama Central Station. For wireless communication in the Wireless Medical Telemetry Service (WMTS) band, a Panorama Wireless Transceiver and radios are required, which transmit and/or receive data to/from remote devices using WMTS communication. For wireless communication in the Industrial Scientific Medical (ISM) band, access points and radios are required, which transmit and/or receive data to/from remote devices using ISM communication.

The Panorama Telemetry Server communicates with the following remote devices:

- Panorama Telepack-608 and Panorama Telepack-608 with Nonin SpO₂ module
- Panorama Instrument Radio-608

CAUTION: The Panorama Telepack and Instrument Radio are wireless devices and may be subject to intermittent signal dropout. Panorama wireless devices are designed to perform with a maximum cumulative signal dropout of one minute in a 24-hour period under ideal conditions. A crowded wireless environment or interference from other medical or non-medical devices, either intentional (e.g., VOIP phones, pagers, asset tracking systems, etc.) or unintentional (e.g., AC motors, microwave ovens, paper shredders, etc.), may result in a significantly increased amount of signal dropout experienced by any one or multiple wireless device(s).

1.2.4 Panorama Wireless Transceiver

NOTE: The Panorama Wireless Transceiver is used for WMTS installations only.

The Panorama Wireless Transceiver and its components serve as a bridge between the hardwired and the wireless infrastructures. A patient using instrument telemetry can transition from hardwired to wireless, and back, while being monitored from the same screen location at the central station. Communication for both hardwired and wireless monitoring is capable of supporting coverage over large areas, or on different floors of the hospital.

The Panorama Wireless Transceiver and its components control the local wireless network, and are responsible for:

- Providing protocol conversion for telemetry equipped Mindray products.
- Receiving control information and patient data from telemetry equipped Mindray products.
• Sending control information to telemetry equipped Mindray products.
• Acting as a conduit for moving data between the wireless network and the hardwired network via Panorama Telemetry Server.

The Panorama Wireless Transceiver device contains four LED indicator lights. The indicator lights function as follows:

<table>
<thead>
<tr>
<th>DEVICE LABEL</th>
<th>LED COLOR</th>
<th>DESCRIPTION</th>
</tr>
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<tbody>
<tr>
<td>Power</td>
<td>Green</td>
<td>Power is applied.</td>
</tr>
<tr>
<td>Fault</td>
<td>Red</td>
<td>There is a fault in the device configuration.</td>
</tr>
<tr>
<td>Configured</td>
<td>Yellow</td>
<td>The device is configured for use.</td>
</tr>
<tr>
<td>Status</td>
<td>Blue</td>
<td>The link between the Telemetry Server and the Wireless Transceiver is not functioning.</td>
</tr>
</tbody>
</table>

**CAUTION:** Operation of the wireless product requires the prior coordination with a frequency coordinator designated by the FCC for the Wireless Medical Telemetry Service.

1.2.5 Panorama Gateway

The Panorama Gateway enables hospitals to maintain a continuous and comprehensive history of a patient’s clinical information through the use of electronic medical records (EMR).

Bi-directional communication through the Panorama Gateway enables Admit data transfer between the C-LAN and the HIS/CIS.

**Additional Information**
For additional information regarding the Panorama Gateway, refer to “Communications with the Panorama Gateway” on page 5-2.

1.2.6 Panorama Web Viewer

The Panorama Web Viewer software application provides remote intranet access to any admitted patient on the Panorama Patient Monitoring Network.

**Additional Information**
For additional information regarding the Panorama Web Viewer application, refer to the Panorama Web Viewer Operating Instructions (P/N 0070-00-0681-XX).
1.2.7 Passport 2 Bedside Monitor

Passport 2 bedside monitors are used to monitor patient data and may use either a hardwired or a wireless (2.4 GHz or 608 MHz) configuration. The Passport 2 sends data to the Panorama Central Station for analysis and storage via an Ethernet connection or remotely via the Panorama Telemetry Server. The data collected by the Passport 2 is based on the type of lead wire set and sensors used.

If an alarm in the following table does not have an associated footnote, it may occur using any lead wire set.

Parameters/Waveforms

<table>
<thead>
<tr>
<th>PARAMETERS</th>
<th>SETTINGS</th>
<th>PANORAMA ALARMS</th>
</tr>
</thead>
<tbody>
<tr>
<td>ECG/Heart Rate</td>
<td>Leads I, II, III are available with a 3-wire lead set.</td>
<td>Low Heart Rate</td>
</tr>
<tr>
<td></td>
<td>Leads I, II, III, aVR, aVL, aVF, V are available with a 5-wire lead set.</td>
<td>High Heart Rate</td>
</tr>
<tr>
<td></td>
<td>Leads I, II, III, aVR, aVL, aVF, V1, V2, V3, V4, V5, V6 are available with a View 12™ card. (View 12 supported by Passport 2 hardwired and WMTS 608)</td>
<td></td>
</tr>
<tr>
<td>ST</td>
<td>ST analysis is in use at the bedside device.</td>
<td>ST Single</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ST Dual</td>
</tr>
<tr>
<td>Arrhythmia</td>
<td>Arrhythmia analysis is in use at the bedside device.</td>
<td>Asystole</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Bigeminy</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Brady2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Couplelet</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Irregular Heart Rate2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Pause1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High PVC</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Run</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Trigeminy</td>
</tr>
<tr>
<td></td>
<td></td>
<td>V-Fib</td>
</tr>
<tr>
<td></td>
<td></td>
<td>V-Rhythm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>V-Tach</td>
</tr>
<tr>
<td></td>
<td></td>
<td>PVC/min.</td>
</tr>
<tr>
<td>Respiration [Source: ECG, CO2 and Gas-module]</td>
<td>Respiration is in use at the bedside and is derived from a valid source.</td>
<td>Low Respiration</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High Respiration</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Apnea</td>
</tr>
<tr>
<td>SpO2</td>
<td>SpO2 sensor is in use at the bedside device.</td>
<td>Low SpO2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High SpO2</td>
</tr>
<tr>
<td>IBP1/IBP2 (Systolic, Diastolic, Mean)</td>
<td>IBPx option is in use at the bedside device.</td>
<td>Low IBPx Systolic</td>
</tr>
<tr>
<td></td>
<td>IBP parameter labels include IBP, Art, UA, LV, PA, CVP, ICP, LA and RA. The label used is based on the label selected at the bedside monitor.</td>
<td>High IBPx Systolic</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low IBPx Diastolic</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High IBPx Diastolic</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low IBPx Mean</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High IBPx Mean</td>
</tr>
</tbody>
</table>

**NOTE:**  
1. Alarm will only be detected when using a View 12™ card.  
2. Alarm will not be detected when using a View 12™ card.  
3. Waveform not supported when using a WMTS 608 bedside device.
Parameters/Waveforms (Continued)

<table>
<thead>
<tr>
<th>PARAMETERS</th>
<th>SETTINGS</th>
<th>PANORAMA ALARMS</th>
</tr>
</thead>
<tbody>
<tr>
<td>INSP/ET O$_2$</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>O$_2$ Inspired O$_2$ End Tidal</td>
</tr>
<tr>
<td>INSP/ET N$_2$O</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>N$_2$O Inspired N$_2$O End Tidal</td>
</tr>
<tr>
<td>INSP/ET Iso</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>Agent Iso Inspired Agent Iso End Tidal</td>
</tr>
<tr>
<td>INSP/ET Enf</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>Agent Enf Inspired Agent Enf End Tidal</td>
</tr>
<tr>
<td>INSP/ET Des</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>Agent Des Inspired Agent Des End Tidal</td>
</tr>
<tr>
<td>INSP/ET Sev</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>Agent Sev Inspired Agent Sev End Tidal</td>
</tr>
<tr>
<td>INSP/ET Hal</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>Agent Hal Inspired Agent Hal End Tidal</td>
</tr>
<tr>
<td>INSP/ET Agent</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>Agent Inspired Alarm Agent ET Alarm</td>
</tr>
<tr>
<td>NIBP (Systolic, Diastolic, Mean, Elapsed Time)</td>
<td>NIBP always displayed even if parameter not monitored.</td>
<td>Low NIBP Systolic High NIBP Systolic Low NIBP Diastolic High NIBP Diastolic Low NIBP Mean High NIBP Mean</td>
</tr>
<tr>
<td>Temp 1</td>
<td>Temperature sensor is in use at the bedside monitor.</td>
<td>Low Temperature 1 High Temperature 1</td>
</tr>
</tbody>
</table>

1 Alarm will only be detected when using a View 12™ card.
2 Alarm will not be detected when using a View 12™ card.
3 Waveform not supported when using a WMTS 608 bedside device.

### Additional Information

For additional information regarding Passport 2 bedside monitors, refer to the Passport 2 Operating Instructions (P/N 0070-00-0649-XX).

For information regarding the parameters that can be monitored by the Passport 2, refer to “Parameter Specifications” on page 12-1.
1.2.8 Passport V Bedside Monitor

Passport V bedside monitors are used to monitor patient data using a hardwired configuration. The Passport V sends data to the Panorama Central Station for analysis and storage via an Ethernet connection. The data collected by the Passport V is based on the type of lead wire set and sensors used.

If an alarm in the following table does not have an associated footnote, it may occur using any lead wire set.

Parameters/Waveforms

<table>
<thead>
<tr>
<th>PARAMETERS</th>
<th>SETTINGS</th>
<th>PANORAMA ALARMS</th>
</tr>
</thead>
<tbody>
<tr>
<td>ECG/Heart Rate</td>
<td>Leads I, II, III are available with a 3-wire lead set.</td>
<td>Low Heart Rate</td>
</tr>
<tr>
<td></td>
<td>Leads I, II, III, aVR, aVL, aVF, V are available with a 5-wire lead set.</td>
<td>High Heart Rate</td>
</tr>
<tr>
<td>ST</td>
<td>ST analysis is in use at the bedside device.</td>
<td>ST Single</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ST Dual</td>
</tr>
<tr>
<td>Arrhythmia</td>
<td>Arrhythmia analysis is in use at the bedside device.</td>
<td>Asystole</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Bigeminy</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Brady</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Couplelet</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Irregular Heart Rate</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High PVC</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Run</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Trigeminy</td>
</tr>
<tr>
<td></td>
<td></td>
<td>V-Fib</td>
</tr>
<tr>
<td></td>
<td></td>
<td>V-Rhythm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>V-Tach</td>
</tr>
<tr>
<td></td>
<td></td>
<td>PVC/min.</td>
</tr>
<tr>
<td>Respiration (Source: ECG, CO₂ and Gas Module)</td>
<td>Respiration is in use at the bedside and is derived from a valid source.</td>
<td>Low Respiration</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High Respiration</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Apnea</td>
</tr>
<tr>
<td>SpO₂</td>
<td>SpO₂ sensor is in use at the bedside device.</td>
<td>Low SpO₂</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High SpO₂</td>
</tr>
<tr>
<td>IBP1/IBP2 (Systolic, Diastolic, Mean)</td>
<td>IBPx option is in use at the bedside device. IBP parameter labels include IBP, Art, UA, LV, PA, CVP, ICP, LA and RA. The label used is based on the label selected at the bedside monitor.</td>
<td>Low IBPx Systolic</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High IBPx Systolic</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low IBPx Diastolic</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High IBPx Diastolic</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low IBPx Mean</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High IBPx Mean</td>
</tr>
<tr>
<td>NOTE: x = 1 or 2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>INSP/ET CO</td>
<td>Gas Module is in use at the bedside device.</td>
<td>High CO₂ Inspired</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low CO₂ End Tidal</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High CO₂ End Tidal</td>
</tr>
<tr>
<td>INSP/ET O₂</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>O₂ Inspired</td>
</tr>
<tr>
<td></td>
<td></td>
<td>O₂ End Tidal</td>
</tr>
<tr>
<td>INSP/ET N₂O</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>N₂O Inspired</td>
</tr>
<tr>
<td></td>
<td></td>
<td>N₂O End Tidal</td>
</tr>
<tr>
<td>INSP/ET Iso</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>Agent Iso Inspired</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Agent Iso End Tidal</td>
</tr>
<tr>
<td>INSP/ET Enf</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>Agent Enf Inspired</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Agent Enf End Tidal</td>
</tr>
</tbody>
</table>
Parameters/Waveforms (Continued)

<table>
<thead>
<tr>
<th>PARAMETERS</th>
<th>SETTINGS</th>
<th>PANORAMA ALARMS</th>
</tr>
</thead>
<tbody>
<tr>
<td>INSP/ET Des</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>Agent Des Inspired</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Agent Des End Tidal</td>
</tr>
<tr>
<td>INSP/ET Sev</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>Agent Sev Inspired</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Agent Sev End Tidal</td>
</tr>
<tr>
<td>INSP/ET Hal</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>Agent Hal Inspired</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Agent Hal End Tidal</td>
</tr>
<tr>
<td>INSP/ET Agent</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>Agent Inspired Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Agent ET Alarm</td>
</tr>
<tr>
<td>NIBP (Systolic,</td>
<td>NIBP always displayed even if parameter not</td>
<td>Low NIBP Systolic</td>
</tr>
<tr>
<td>Diastolic, Mean,</td>
<td>monitored.</td>
<td>High NIBP Systolic</td>
</tr>
<tr>
<td>Elapsed Time)</td>
<td></td>
<td>Low NIBP Diastolic</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High NIBP Diastolic</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low NIBP Mean</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High NIBP Mean</td>
</tr>
<tr>
<td>Temp 1</td>
<td>Temperature sensor is in use at the bedside</td>
<td>Low Temperature 1</td>
</tr>
<tr>
<td></td>
<td>monitor.</td>
<td>High Temperature 1</td>
</tr>
</tbody>
</table>

Additional Information

For additional information regarding Passport V bedside monitors, refer to the Passport V Operating Instructions (P/N 0070-00-0704-XX).

For information regarding the parameters that can be monitored by the Passport V, refer to “Parameter Specifications” on page 12-1.
1.2.9 Spectrum Bedside Monitor

Spectrum bedside monitors are used to monitor patient data and may use either a hardwired or a wireless (2.4 GHz or 608 MHz) configuration. Currently, the Spectrum sends data to the Panorama Central Station for viewing and storage via an Ethernet connection. The data collected by a Spectrum is dependent on the type of lead wire set and sensors used. If an alarm in the following table does not have an associated footnote, it may occur using any lead wire set.

**NOTE:** The CO/C1, Temp 2, and T Blood parameters are only available if they are being monitored by the Spectrum/Spectrum OR External Parameter Module (EPM). These parameters are not available if they are monitored by an Edwards Vigilance® monitor.

### Parameters/Waveforms

<table>
<thead>
<tr>
<th>PARAMETERS</th>
<th>SETTINGS</th>
<th>PANORAMA ALARMS</th>
</tr>
</thead>
<tbody>
<tr>
<td>ECG/Heart Rate</td>
<td>Leads I, II, III are available with a 3-wire lead set.</td>
<td>Low Heart Rate</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High Heart Rate</td>
</tr>
<tr>
<td></td>
<td>Leads I, II, III, aVR, aVL, aVF, V are available with a 5-wire lead set.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Leads I, II, III, aVR, aVL, aVF, V1, V2, V3, V4, V5, V6 are available</td>
<td></td>
</tr>
<tr>
<td></td>
<td>with a View 12™ card.</td>
<td></td>
</tr>
<tr>
<td>ST</td>
<td>ST analysis is in use at the bedside device.</td>
<td>ST Single</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ST Dual</td>
</tr>
<tr>
<td>Arrhythmia</td>
<td>Arrhythmia analysis is in use at the bedside device.</td>
<td>Asystole</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Bigeminy</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Brady2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Couple2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Irregular Heart Rate2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Pause1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High PVC</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Run</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Trigeminy</td>
</tr>
<tr>
<td></td>
<td></td>
<td>V-Fib</td>
</tr>
<tr>
<td></td>
<td></td>
<td>V-Rhythm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>V-Tach</td>
</tr>
<tr>
<td></td>
<td></td>
<td>PVC/min.</td>
</tr>
<tr>
<td>Respiration (Source:</td>
<td>Respiration is in use at the bedside and is derived from a valid source.</td>
<td>Low Respiration</td>
</tr>
<tr>
<td>ECG, CO2 and Gas-module)</td>
<td></td>
<td>High Respiration</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Apnea</td>
</tr>
<tr>
<td>SpO2</td>
<td>SpO2 sensor is in use at the bedside device.</td>
<td>Low SpO2</td>
</tr>
<tr>
<td>IBP1/IBP2/IBP3/IBP4</td>
<td>IBPx option is in use at the bedside device.</td>
<td>Low IBPx Systolic</td>
</tr>
<tr>
<td>(Systolic, Diastolic,</td>
<td></td>
<td>High IBPx Systolic</td>
</tr>
<tr>
<td>Mean)</td>
<td>IBP parameter labels include IBP, Art, UA, LV, PA, CVP, ICP, LA and RA.</td>
<td>Low IBPx Diastolic</td>
</tr>
<tr>
<td></td>
<td>The label used is based on the label selected at the bedside monitor.</td>
<td>High IBPx Diastolic</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low IBPx Mean</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High IBPx Mean</td>
</tr>
</tbody>
</table>

1. Alarm will only be detected when using a View 12™ card.
2. Alarm will not be detected when using a View 12™ card.
3. Waveform not supported when using a WMTS 608 bedside device.
## Parameters/Waveforms

<table>
<thead>
<tr>
<th>PARAMETERS</th>
<th>SETTINGS</th>
<th>PANORAMA ALARMS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>NOTE:</strong> $x = 1, 2, 3$ or $4$ (IBP3 and 4 are only available if the Spectrum/Spectrum OR External Parameter Module (EPM) is installed)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>INSP/ET CO$_2^3$</td>
<td>Gas Module is in use at the bedside device.</td>
<td>High CO$_2$ Inspired Low CO$_2$ End Tidal High CO$_2$ End Tidal</td>
</tr>
<tr>
<td>INSP/ET O$_2^3$</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>O$_2$ Inspired O$_2$ End Tidal</td>
</tr>
<tr>
<td>INSP/ET N$_2$O$^3$</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>N$_2$O Inspired N$_2$O End Tidal</td>
</tr>
<tr>
<td>INSP/ET Iso$^3$</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>Agent Iso Inspired Agent Iso End Tidal</td>
</tr>
<tr>
<td>INSP/ET Enf$^3$</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>Agent Enf Inspired Agent Enf End Tidal</td>
</tr>
<tr>
<td>INSP/ET Des$^3$</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>Agent Des Inspired Agent Des End Tidal</td>
</tr>
<tr>
<td>INSP/ET Sev$^3$</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>Agent Sev Inspired Agent Sev End Tidal</td>
</tr>
<tr>
<td>INSP/ET Hal$^3$</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>Agent Hal Inspired Agent Hal End Tidal</td>
</tr>
<tr>
<td>INSP/ET Agent$^3$</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>Agent Inspired Alarm Agent ET Alarm</td>
</tr>
<tr>
<td>NIBP (Systolic, Diastolic, Mean, Elapsed Time)</td>
<td>NIBP always displayed even if parameter not monitored.</td>
<td>Low NIBP Systolic High NIBP Systolic Low NIBP Diastolic High NIBP Diastolic Low NIBP Mean High NIBP Mean</td>
</tr>
<tr>
<td>Temp 1/Temp 2</td>
<td>Temperature sensor is in use at the bedside monitor.</td>
<td>Low Temperature 1 High Temperature 1 Low Temperature 2 High Temperature 2</td>
</tr>
<tr>
<td>Delta T</td>
<td>At least two temperature sensors in use at the bedside. Spectrum/Spectrum OR External Parameter Module (EPM) is installed.</td>
<td>Low Delta T Alarm High Delta T Alarm</td>
</tr>
<tr>
<td>Temp Blood</td>
<td>Spectrum/Spectrum OR External Parameter Module (EPM) is in use at the bedside.</td>
<td>Low T blood Alarm High T blood Alarm</td>
</tr>
<tr>
<td>Pulmonary Artery Wedge Pressure (PAWP)</td>
<td>One IBP must be labeled PA.</td>
<td>New PAWP Measurement</td>
</tr>
<tr>
<td>CO</td>
<td>Spectrum/Spectrum OR External Parameter Module (EPM) is in use at the bedside.</td>
<td>New CO Measurement</td>
</tr>
</tbody>
</table>

1. Alarm will only be detected when using a View 12™ card.
2. Alarm will not be detected when using a View 12™ card.
3. Waveform not supported when using a WMTS 608 bedside device.
Additional Information
For additional information regarding Spectrum bedside monitors, refer to the Spectrum Operating Instructions (P/N 0070-00-0648-XX).

1.2.10 Spectrum OR Bedside Monitor
Similar to the Spectrum, Spectrum OR bedside monitors are used to monitor patient data and send it to the Panorama Central Station for viewing and storage via an Ethernet connection. The Spectrum OR requires a hardwired configuration. The data collected by a Spectrum OR is dependent on the type of lead wire set and sensors used.

**NOTE:** The CO/CI, Temp 2, and T Blood parameters are only available if they are being monitored by the Spectrum/Spectrum OR External Parameter Module (EPM). These parameters are not available if they are monitored by an Edwards Vigilance® monitor.

### Parameters/Waveforms

<table>
<thead>
<tr>
<th>PARAMETERS</th>
<th>SETTINGS</th>
<th>PANORAMA ALARMS</th>
</tr>
</thead>
<tbody>
<tr>
<td>ECG/Heart Rate</td>
<td>Leads I, II, III are available with a 3-wire lead set.</td>
<td>Low Heart Rate</td>
</tr>
<tr>
<td></td>
<td>Leads I, II, III, aVR, aVL, aVF, V are available with a 5-wire lead set.</td>
<td>High Heart Rate</td>
</tr>
<tr>
<td>ST</td>
<td>ST analysis is in use at the bedside device.</td>
<td>ST Single</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ST Dual</td>
</tr>
<tr>
<td>Arrhythmia</td>
<td>Arrhythmia analysis is in use at the bedside device.</td>
<td>Asystole</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Bigeminy</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Brady^2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Couplet</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Irregular Heart Rate^2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Pause^1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High PVC</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Run</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Trigeminy</td>
</tr>
<tr>
<td></td>
<td></td>
<td>VFib</td>
</tr>
<tr>
<td></td>
<td></td>
<td>VRhythm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>VTach</td>
</tr>
<tr>
<td></td>
<td></td>
<td>PVC/min.</td>
</tr>
<tr>
<td>Respiration [Source: ECG, CO2 and Gas Module]</td>
<td>Respiration is in use at the bedside and is derived from a valid source.</td>
<td>Low Respiration</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High Respiration</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Apnea</td>
</tr>
<tr>
<td>SpO2</td>
<td>SpO2 sensor is in use at the bedside device.</td>
<td>Low SpO2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High SpO2</td>
</tr>
<tr>
<td>IBP1/IBP2/IBP3/IBP4 [Systolic, Diastolic, Mean]</td>
<td>IBPx option is in use at the bedside device.</td>
<td>Low IBPx Systolic</td>
</tr>
<tr>
<td></td>
<td>IBP parameter labels include IBP, Art, UA, LV, PA, CVP, ICP, LA and RA. The label used is based on the label selected at the bedside monitor.</td>
<td>High IBPx Systolic</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low IBPx Diastolic</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High IBPx Diastolic</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low IBPx Mean</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High IBPx Mean</td>
</tr>
<tr>
<td></td>
<td>x = 1, 2, 3 or 4 (IBP3 and 4 are only available if the Spectrum/Spectrum OR External Parameter Module (EPM) is installed)</td>
<td></td>
</tr>
<tr>
<td>INSP/ET CO2</td>
<td>Gas Module is in use at the bedside device.</td>
<td>High CO2 Inspired</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low CO2 End Tidal</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High CO2 End Tidal Rate</td>
</tr>
</tbody>
</table>
Parameters/Waveforms

<table>
<thead>
<tr>
<th>PARAMETERS</th>
<th>SETTINGS</th>
<th>PANORAMA ALARMS</th>
</tr>
</thead>
<tbody>
<tr>
<td>INSP/ET O₂</td>
<td>Gas Module is in use at the bedside device.</td>
<td>O₂ Inspired O₂ End Tidal</td>
</tr>
<tr>
<td>INSP/ET N₂O</td>
<td>Gas Module is in use at the bedside device.</td>
<td>N₂O Inspired N₂O End Tidal</td>
</tr>
<tr>
<td>INSP/ET Iso</td>
<td>Gas Module is in use at the bedside device.</td>
<td>Agent Iso Inspired Agent Iso End Tidal</td>
</tr>
<tr>
<td>INSP/ET Enf</td>
<td>Gas Module is in use at the bedside device.</td>
<td>Agent Enf Inspired Agent Enf End Tidal</td>
</tr>
<tr>
<td>INSP/ET Des</td>
<td>Gas Module is in use at the bedside device.</td>
<td>Agent Des Inspired Agent Des End Tidal</td>
</tr>
<tr>
<td>INSP/ET Sev</td>
<td>Gas Module is in use at the bedside device.</td>
<td>Agent Sev Inspired Agent Sev End Tidal</td>
</tr>
<tr>
<td>INSP/ET Hal</td>
<td>Gas Module is in use at the bedside device.</td>
<td>Agent Hal Inspired Agent Hal End Tidal</td>
</tr>
<tr>
<td>INSP/ET Agent</td>
<td>Gas Module is in use at the bedside device.</td>
<td>Agent Inspired Alarm Agent ET Alarm</td>
</tr>
<tr>
<td>NIBP (Systolic, Diastolic, Mean, Elapsed Time)</td>
<td>NIBP always displayed even if parameter not monitored.</td>
<td>Low NIBP Systolic High NIBP Systolic Low NIBP Diastolic High NIBP Diastolic Low NIBP Mean High NIBP Mean</td>
</tr>
<tr>
<td>Temp 1/Temp 2</td>
<td>Temperature sensor is in use at the bedside monitor.</td>
<td>Low Temperature 1 High Temperature 1 Low Temperature 2 High Temperature 2</td>
</tr>
<tr>
<td>Delta T</td>
<td>At least two temperature sensors in use at the bedside. Spectrum/Spectrum OR External Parameter Module (EPM) is installed.</td>
<td>Low Delta T Alarm High Delta T Alarm</td>
</tr>
<tr>
<td>Temp Blood</td>
<td>Spectrum/Spectrum OR External Parameter Module (EPM) is in use at the bedside.</td>
<td>Low T blood Alarm High T blood Alarm</td>
</tr>
<tr>
<td>Pulmonary Artery Wedge Pressure (PAWP)</td>
<td>One IBP must be labeled PA.</td>
<td>New PAWP Measurement</td>
</tr>
<tr>
<td>CO</td>
<td>Spectrum/Spectrum OR External Parameter Module (EPM) is in use at the bedside.</td>
<td>New CO Measurement</td>
</tr>
</tbody>
</table>

Additional Information

For additional information regarding Spectrum OR bedside monitors, refer to the Spectrum OR Operating Instructions (P/N 0070-00-0670-XX).

For information regarding the parameters that can be monitored by the Spectrum OR, refer to the “Parameter Specifications” on page 12-1.
### 1.2.11 DPM 6 and DPM 7 Bedside Monitors

DPM 6 and DPM 7 offer patient monitoring capabilities in a modular design with high resolution, touchscreen displays. Highlighted features include ETCO₂, 5-agent multi-gas analysis with auto-ID, extensive data storage and independent display support (DPM 7 only).

#### Parameters/Waveforms

<table>
<thead>
<tr>
<th>PARAMETERS</th>
<th>SETTINGS</th>
<th>PANORAMA ALARMS</th>
</tr>
</thead>
<tbody>
<tr>
<td>ECG/Heart Rate</td>
<td>Leads I, II, III are available with a 3-wire lead set.</td>
<td>Low Heart Rate</td>
</tr>
<tr>
<td></td>
<td>Leads I, II, III, aVR, aVL, aVF, V are available with a 5-wire lead set.</td>
<td>High Heart Rate</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Pacer Failure to Capture</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Pacer Failure to Sense</td>
</tr>
<tr>
<td>ST</td>
<td>ST analysis is in use at the bedside monitor.</td>
<td>ST</td>
</tr>
<tr>
<td>Arrhythmia</td>
<td>Arrhythmia analysis is in use at the bedside monitor.</td>
<td>Asystole</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Bigeminy</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Pause</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Couplet</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Irregular Heart Rate</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High PVC</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Run</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Trigeminy</td>
</tr>
<tr>
<td></td>
<td></td>
<td>V-Fib</td>
</tr>
<tr>
<td></td>
<td></td>
<td>V-Rhythm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>V-Tach</td>
</tr>
<tr>
<td></td>
<td></td>
<td>PVC/min.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Multifocal PVC</td>
</tr>
<tr>
<td></td>
<td></td>
<td>R-On-T PVC</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Extreme Bradycardia</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Extreme Tachycardia</td>
</tr>
<tr>
<td>Respiration</td>
<td>Respiration is in use at the bedside monitor and is derived from a valid</td>
<td>Low Respiration</td>
</tr>
<tr>
<td>[Source: ECG, CO₂ and Gas Module]</td>
<td>source.</td>
<td>High Respiration</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Apnea</td>
</tr>
<tr>
<td>SpO₂/Desat</td>
<td>SpO₂ sensor is in use at the bedside monitor.</td>
<td>Low SpO₂ (Desat)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High SpO₂</td>
</tr>
<tr>
<td>Ao, Art, Art2, BAP, CVP, FAP, ICP, LAP, IVP, PA, RAP, UAP, and UVP.</td>
<td>The label used is based on the label selected at the</td>
<td>Low Systolic</td>
</tr>
<tr>
<td></td>
<td></td>
<td>bedside monitor.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low Diastolic</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High Diastolic</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low Mean</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High Mean</td>
</tr>
<tr>
<td>INSP/ET CO₂</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>High CO₂ Inspired</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low CO₂ End Tidal</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High CO₂ End Tidal Rate</td>
</tr>
<tr>
<td>INSP/ET O₂</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>O₂ Inspired</td>
</tr>
<tr>
<td></td>
<td></td>
<td>O₂ End Tidal</td>
</tr>
<tr>
<td>INSP/ET N₂O</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>N₂O Inspired</td>
</tr>
<tr>
<td></td>
<td></td>
<td>N₂O End Tidal</td>
</tr>
<tr>
<td>INSP/ET Iso</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>Agent Iso Inspired</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Agent Iso End Tidal</td>
</tr>
<tr>
<td>INSP/ET Enf</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>Agent Enf Inspired</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Agent Enf End Tidal</td>
</tr>
<tr>
<td>INSP/ET Des</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>Agent Des Inspired</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Agent Des End Tidal</td>
</tr>
</tbody>
</table>
Parameters/Waveforms

<table>
<thead>
<tr>
<th>PARAMETERS</th>
<th>SETTINGS</th>
<th>PANORAMA ALARMS</th>
</tr>
</thead>
<tbody>
<tr>
<td>INSP/ET Hal</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>Agent Hal Inspired</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Agent Hal End Tidal</td>
</tr>
<tr>
<td>INSP/ET Agent</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>Agent Inspired Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Agent ET Alarm</td>
</tr>
<tr>
<td>NIBP (Systolic,</td>
<td>NIBP always displayed even if parameter not monitored.</td>
<td>Low NIBP Systolic</td>
</tr>
<tr>
<td>Diastolic, Mean,</td>
<td></td>
<td>High NIBP Systolic</td>
</tr>
<tr>
<td>Elapsed Time)</td>
<td></td>
<td>Low NIBP Diastolic</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High NIBP Diastolic</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low NIBP Mean</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High NIBP Mean</td>
</tr>
<tr>
<td>Temp 1/Temp 2</td>
<td>Temperature sensor is in use at the bedside monitor.</td>
<td>Low Temperature 1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High Temperature 1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low Temperature 2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High Temperature 2</td>
</tr>
<tr>
<td>Delta T</td>
<td>At least two temperature sensors in use at the bedside monitor.</td>
<td>High Delta T Alarm</td>
</tr>
<tr>
<td>C.O/C.I./T Blood</td>
<td>C.O. Module is in use at the bedside monitor.</td>
<td>High T Blood Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low T Blood Alarm</td>
</tr>
</tbody>
</table>

Panorama System Compatibility

The DPM6/7 user interface on the Panorama system varies from the user interface on the DPM6/7 monitors as follows:

- If CVP, ICP, UVP, LAP, or RAP are selected as labels, only the mean values will be displayed; the alarm limit sliders for Systole & Diastole will not appear.
- CVP on the bedside monitor will show a more precise value than at the Panorama system side (e.g. 6.8 mmHg vs. 7 mmHg).
- ST templates and ST reference values will not be available.
- ST Tab and Digital Data graphic user interface varies from the bedside.
- ST Single Alarms and ST Dual Alarms are not supported.
- ECG settings of the DPM6/7 monitors will not be available on the Panorama system.

Additional Information

For additional information regarding the DPM6 bedside monitor, refer to the DPM 6 Operating Instructions (P/N 046-000184-00).

For additional information regarding the DPM7 bedside monitor, refer to the DPM 7 Operating Instructions (P/N 046-000186-00).

For information regarding the parameters that can be monitored by the DPM 6 and DPM 7, refer to the “Parameter Specifications” on page 12-1.
Passport 12M/Passport 17M Bedside Monitors

Passport 12M and Passport 17M monitors offer patient monitoring capabilities in a modular design with high resolution, touchscreen displays. Highlighted features include ETCO₂, 5-agent multi-gas analysis with auto-ID, extensive data storage and independent display support (Passport 17M only).

Parameters/Waveforms

<table>
<thead>
<tr>
<th>PARAMETERS</th>
<th>SETTINGS</th>
<th>PANORAMA ALARMS</th>
</tr>
</thead>
<tbody>
<tr>
<td>ECG/Heart Rate</td>
<td>Leads I, II, III are available with a 3-wire lead set. Leads I, II, III, aVR, aVL, aVF, V are available with a 5-wire lead set. Lead I, II, III, aVR, aVL, aVF, V1-V6 are available with a 12-wire lead set.</td>
<td>Low Heart Rate Alarm High Heart Rate Alarm Pacer Failure to Capture Pacer Failure to Sense</td>
</tr>
<tr>
<td>ST</td>
<td>ST analysis is in use at the bedside monitor.</td>
<td>ST Alarm</td>
</tr>
<tr>
<td>Arrhythmia</td>
<td>Arrhythmia analysis is in use at the bedside monitor.</td>
<td>Asystole Bigeminy PauseCouplet Irregular Heart Rate High PVC Run Trigeminy V-Fib V-Rhythm V-Tach Multifocal PVC R-On-T PVC Extreme Bradycardia Extreme Tachycardia</td>
</tr>
<tr>
<td>Respiration [Source: ECG, CO₂ and Gas Module]</td>
<td>Respiration is in use at the bedside monitor and is derived from a valid source.</td>
<td>Low Respiration Rate Alarm High Respiration Rate Alarm Apnea</td>
</tr>
<tr>
<td>SpO₂</td>
<td>SpO₂ sensor is in use at the bedside monitor.</td>
<td>Low SpO₂ Alarm High SpO₂ Alarm</td>
</tr>
<tr>
<td>Art, UAP, LV, PA, CVP, ICP, LAP, RAP, Ao, BAP, FAP, UVP</td>
<td>The label used is based on the label selected at the bedside monitor.</td>
<td>Low Systolic Alarm High Systolic Alarm Low Diastolic Alarm High Diastolic Alarm Low Mean Alarm High Mean Alarm</td>
</tr>
<tr>
<td>CO₂</td>
<td>CO₂ Module is in use at the bedside monitor.</td>
<td>Low CO₂ ET Alarm High CO₂ ET Alarm High CO₂ Inspired Alarm</td>
</tr>
<tr>
<td>O₂</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>O₂ Inspired Alarm O₂ ET Alarm</td>
</tr>
<tr>
<td>N₂O</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>N₂O Inspired Alarm N₂O ET Alarm</td>
</tr>
</tbody>
</table>
Additional Information

For additional information regarding the Passport 12M/Passport 17M bedside monitors, refer to the Passport 12M/Passport 17M Operating Instructions (P/N 046-000184-00).
1.2.13 Passport 8/Passport 12 Bedside Monitors

Parameters/Waveforms

<table>
<thead>
<tr>
<th>PARAMETERS</th>
<th>SETTINGS</th>
<th>PANORAMA ALARMS</th>
</tr>
</thead>
<tbody>
<tr>
<td>ECG/Heart Rate</td>
<td>Leads I, II, III are available with a 3-wire lead set.</td>
<td>Low Heart Rate Alarm</td>
</tr>
<tr>
<td></td>
<td>Leads I, II, III, aVR, aVL, aVF, V are available with a 5-wire lead set.</td>
<td>High Heart Rate Alarm</td>
</tr>
<tr>
<td></td>
<td>Lead I, II, III, aVR, aVL, aVF, V1-V6 are available with a 12-wire lead set.(Passport 12 only)</td>
<td>Pacer Failure to Capture</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Pacer Failure to Sense</td>
</tr>
<tr>
<td>ST</td>
<td>ST analysis is in use at the bedside monitor.</td>
<td>ST Alarm</td>
</tr>
<tr>
<td>Arrhythmia</td>
<td>Arrhythmia analysis is in use at the bedside monitor.</td>
<td>Asystole</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Bigeminy</td>
</tr>
<tr>
<td></td>
<td></td>
<td>PauseCouplet</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Irregular Heart Rate</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High PVC</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Run</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Trigeminy</td>
</tr>
<tr>
<td></td>
<td></td>
<td>V-Fib</td>
</tr>
<tr>
<td></td>
<td></td>
<td>V-Rhythm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>V-Tach</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Multifocal PVC</td>
</tr>
<tr>
<td></td>
<td></td>
<td>R-On-T PVC</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Extreme Bradycardia</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Extreme Tachycardia</td>
</tr>
<tr>
<td>Respiration [Source: ECG, CO₂ and Gas Module]</td>
<td>Respiration is in use at the bedside monitor and is derived from a valid source.</td>
<td>Low Respiration Rate Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High Respiration Rate Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Apnea</td>
</tr>
<tr>
<td>SpO₂</td>
<td>SpO₂ sensor is in use at the bedside monitor.</td>
<td>Low SpO₂ Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High SpO₂ Alarm</td>
</tr>
<tr>
<td>Art, UAP, IV, PA, CVP, ICP, LAP, RAP, Ao, BAP, FAP, UVP</td>
<td>The label used is based on the label selected at the bedside monitor.</td>
<td>Low Systolic Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High Systolic Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low Diastolic Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High Diastolic Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low Mean Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High Mean Alarm</td>
</tr>
<tr>
<td>CO₂</td>
<td>CO₂ Module is in use at the bedside monitor.</td>
<td>Low CO₂ ET Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High CO₂ ET Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High CO₂ Inspired Alarm</td>
</tr>
<tr>
<td>O₂ (Passport 12 only)</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>O₂ Inspired Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>O₂ ET Alarm</td>
</tr>
<tr>
<td>N₂O (Passport 12 only)</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>N₂O Inspired Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>N₂O ET Alarm</td>
</tr>
</tbody>
</table>
1.2.14 T1 Monitor

Parameters/Waveforms

<table>
<thead>
<tr>
<th>PARAMETERS</th>
<th>SETTINGS</th>
<th>PANORAMA ALARMS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Iso, Enf, Des, Sev, Hal, Agent (Passport 12 only)</td>
<td>Gas Module is in use at the bedside monitor.</td>
<td>Agent Iso Inspired Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Agent Iso ET Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Agent Enf Inspired Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Agent Enf ET Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Agent Des Inspired Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Agent Des ET Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Agent Sev Inspired Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Agent Sev ET Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Agent Hal Inspired Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Agent Hal ET Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Agent ET Alarm</td>
</tr>
<tr>
<td>NIBP (Systolic, Diastolic, Mean)</td>
<td>NIBP always displayed even if parameter not monitored.</td>
<td>Low NIBP Systolic Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High NIBP Systolic Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low NIBP Diastolic Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High NIBP Diastolic Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low NIBP Mean Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High NIBP Mean Alarm</td>
</tr>
<tr>
<td>Temp 1/Temp 2</td>
<td>Temperature sensor is in use at the bedside monitor.</td>
<td>Low Temperature 1 Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High Temperature 1 Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low Temperature 2 Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High Temperature 2 Alarm</td>
</tr>
<tr>
<td>Delta T</td>
<td>At least two temperature sensors in use at the bedside monitor.</td>
<td>High Delta T Alarm</td>
</tr>
<tr>
<td>C.O/C.I./T Blood</td>
<td>C.O. Module is in use at the bedside monitor.</td>
<td>High T Blood Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low T Blood Alarm</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>PARAMETERS</th>
<th>SETTINGS</th>
<th>PANORAMA ALARMS</th>
</tr>
</thead>
<tbody>
<tr>
<td>ECG/Heart Rate</td>
<td>Leads I, II, III are available with a 3-wire lead set.</td>
<td>Low Heart Rate Alarm</td>
</tr>
<tr>
<td></td>
<td>Leads I, II, III, aVR, aVL, aVF, V are available with a 5-wire lead set.</td>
<td>High Heart Rate Alarm</td>
</tr>
<tr>
<td></td>
<td>Lead I, II, III, aVR, aVL, aVF, V1-V6 are available with a 12-wire lead set.</td>
<td>Pacer Failure to Capture</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Pacer Failure to Sense</td>
</tr>
<tr>
<td>ST</td>
<td>ST analysis is in use at the bedside monitor.</td>
<td>ST Alarm</td>
</tr>
<tr>
<td>Parameters</td>
<td>Settings</td>
<td>Panorama Alarms</td>
</tr>
<tr>
<td>---------------------</td>
<td>----------------------------------------------------------------------------</td>
<td>-------------------------------------------------------</td>
</tr>
<tr>
<td>Arrhythmia</td>
<td>Arrhythmia analysis is in use at the bedside monitor.</td>
<td>Asystole, Bigeminy, PauseCouplet,</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Irregular Heart Rate, High PVC,</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Run, Trigeminy, V-Fib, V-Rhythm, V-Tach,</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Multifocal PVC, R-On-T PVC, Extreme Bradycardia,</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Extreme Tachycardia</td>
</tr>
<tr>
<td>Respiration (Source:</td>
<td>Respiration is in use at the bedside monitor and is derived from a valid</td>
<td>Low Respiration Rate Alarm, High Respiration Rate</td>
</tr>
<tr>
<td>ECG, CO2 Module)</td>
<td>source.</td>
<td>Alarm, Apnea</td>
</tr>
<tr>
<td>SpO2</td>
<td>SpO2 sensor is in use at the bedside monitor.</td>
<td>Low SpO2 Alarm, High SpO2 Alarm</td>
</tr>
<tr>
<td>Art, UAP, LV, PA, CVP, ICP, LAP, RAP, Ao, BAP, FAP, UVP</td>
<td>The label used is based on the label selected at the bedside monitor.</td>
<td>Low Systolic Alarm, High Systolic Alarm, High Diastolic</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Alarm, Low Diastolic Alarm, High Diastolic Alarm,</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low Mean Alarm, High Mean Alarm</td>
</tr>
<tr>
<td>CO2</td>
<td>CO2 Module is in use at the bedside monitor.</td>
<td>Low CO2 ET Alarm, High CO2 ET Alarm, High CO2</td>
</tr>
<tr>
<td>NIBP (Systolic, Diastolic, Mean)</td>
<td>NIBP always displayed even if parameter not monitored.</td>
<td>Inspired Alarm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low NIBP Systolic Alarm, High NIBP Systolic Alarm,</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low NIBP Diastolic Alarm, High NIBP Diastolic Alarm,</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low NIBP Mean Alarm, High NIBP Mean Alarm</td>
</tr>
<tr>
<td>Temp 1/Temp 2</td>
<td>Temperature sensor is in use at the bedside monitor.</td>
<td>Low Temperature 1 Alarm, High Temperature 1 Alarm,</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low Temperature 2 Alarm, High Temperature 2 Alarm</td>
</tr>
<tr>
<td>Delta T</td>
<td>At least two temperature sensors in use at the bedside monitor.</td>
<td>High Delta T Alarm</td>
</tr>
</tbody>
</table>
1.2.15 V 12/V 21 Monitors

V 12/V 21 monitors offer patient monitoring capabilities in a modular design with high resolution, touchscreen displays. V 12/V 21 monitors are used to monitor patient data using a hardwired or a wireless configuration. V 12/V 21 monitors send data to the Panorama Central Station for analysis and storage via an Ethernet connection. The data collected by the V 12/V 21 monitors is based on the type of lead wire set and sensors used.

Parameters/Waveforms

<table>
<thead>
<tr>
<th>PARAMETERS</th>
<th>SETTINGS</th>
<th>PANORAMA ALARMS</th>
</tr>
</thead>
<tbody>
<tr>
<td>ECG/Heart Rate</td>
<td>Leads I, II, III are available with a 3-wire lead set.</td>
<td>Low Heart Rate, High Heart Rate, Extreme Bradycardia, Extreme Tachycardia, Pacer Failure to Capture Pacer Failure to Sense</td>
</tr>
<tr>
<td></td>
<td>Leads I, II, III, aVR, aVL, aVF, V are available with a 5-wire lead set.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Leads I, II, III, aVR, aVL, aVF, V, V2 are available with a 6-wire lead set.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>12 lead (I, II, III, aVR, aVL, aVF, V-6) requires a V 12/V 21 Patient Server (VPS).</td>
<td></td>
</tr>
<tr>
<td>ST Single / ST Dual</td>
<td>ST analysis is in use at the bedside monitor.</td>
<td>ST</td>
</tr>
<tr>
<td>Arrhythmia</td>
<td>Arrhythmia analysis is in use at the bedside monitor.</td>
<td>Asystole, Bigeminy, Couple, Irregular Heart Rate, Pause, Run, Trigeminy, VFib, VRhythm, VTach, PVC/min, Multifocal PVC, R-On-T PVC, Pause/Min Brady (SW 2.4.1.30 only), QTc</td>
</tr>
<tr>
<td>Respiration (Source: ECG, CO2 and Gas Module)</td>
<td>Respiration is in use at the bedside monitor and is derived from a valid source.</td>
<td>Low Respiration, High Respiration, Apnea</td>
</tr>
<tr>
<td>SpO2/Desat</td>
<td>SpO2 sensor is in use at the bedside monitor.</td>
<td>Low SpO2 (Desat), High SpO2</td>
</tr>
<tr>
<td>IBP1/IBP2/IBP3/IBP4/IBP5/IBP6/IBP7/IBP8 (Systolic, Diastolic, Mean)</td>
<td>IBPx option is in use at the bedside device. IBP parameter labels include IBP, ART1, ART2, UA, LV, PA, CVP, IAP, LA, RA. The label used is based on the label selected at the bedside monitor.</td>
<td>Low Systolic, High Systolic, Low Diastolic, High Diastolic, Low Mean, High Mean</td>
</tr>
<tr>
<td>NIBP (Systolic, Diastolic, Mean, Elapsed Time)</td>
<td>NIBP always displayed even if parameter not monitored.</td>
<td>Low NIBP Systolic, High NIBP Systolic, Low NIBP Diastolic, High NIBP Diastolic, Low NIBP Mean, High NIBP Mean</td>
</tr>
</tbody>
</table>
Parameters/Waveforms

<table>
<thead>
<tr>
<th>PARAMETERS</th>
<th>SETTINGS</th>
<th>PANORAMA ALARMS</th>
</tr>
</thead>
</table>
| Temp 1/Temp 2/Temp 3/TBlood | Temperature sensor is in use at the bedside monitor. | Low Temperature 1  
High Temperature 1  
Low Temperature 2  
High Temperature 2  
Low Temperature 3  
High Temperature 3 |
| CCO/CCI             | —                                     | High CCO Alarm  
Low CCO Alarm  
High CCI Alarm  
Low CCI Alarm |
| CCO STAT/CCI STAT   | —                                     | High CCO STAT Alarm  
Low CCO STAT Alarm  
High CCI STAT Alarm  
Low CCI STAT Alarm |
| SVR/SVRI            | —                                     | High SVR Alarm  
Low SVR Alarm  
High SVRI Alarm  
Low SVRI Alarm |
| SvO₂/ScvO₂          | —                                     | High SvO₂ Alarm  
Low SvO₂ Alarm  
High ScvO₂ Alarm  
Low ScvO₂ Alarm |
| Delta T             | At least two temperature sensors in use at the bedside monitor. | High Delta T Alarm |
| BIS/BIS L/BIS R     | —                                     | High BIS Alarm  
Low BIS Alarm  
High BIS L Alarm  
Low BIS L Alarm  
High BIS R Alarm  
Low BIS R Alarm |

Additional Information
For additional information regarding the V 12/V 21 bedside monitor, refer to the V 12/V 21 Operating Instructions (P/N 0070-00-0727-XX). For information regarding the parameters that can be monitored by the V 12/V 21, refer to the “Parameter Specifications” on page 12-1.
1.2.16 Panorama Telepack

The Panorama Telepack is a portable, wireless patient telemetry unit that can operate for an extended period of time on a disposable battery source. It can measure 3 or 5 Lead ECG and can also report on SpO2 data when the optional SpO2 module is attached. Telepack models are available for transmission at 608MHz (WMTS band), depending on the region of use. Unless otherwise specified, “Telepack” refers to the Telepack-608 (608 MHz) models.

The Telepack provides basic remote ECG monitoring of a patient, which includes heart rate, ST, PVC, and arrhythmia. The Telepack sends data to the Panorama Central Station for analysis and storage. The type of data that can be collected by a Telepack is dependent on the type of lead wire set and options configured for the connected patient.

Panorama Telepack-608

The WMTS Telepack utilizes the 608MHz wireless bands using a frequency division multiple access method. The Telepack has the capability to transmit to the Panorama Server through an antenna network installed throughout the specified coverage area.

Panorama Telepack with Nonin SpO2 Module

The Telepack may also monitor SpO2 if the Nonin SpO2 module is connected to the Telepack device. The module measures functional oxygen saturation of arterial hemoglobin (% SpO2) and pulse rate (bpm) for adult patients.

NOTE: The Panorama Central Station is used to program, configure, monitor and collect data for the Panorama Telepacks.

Parameters/Waveforms

<table>
<thead>
<tr>
<th>PARAMETERS</th>
<th>SETTINGS</th>
<th>PANORAMA ALARMS</th>
</tr>
</thead>
<tbody>
<tr>
<td>ECG/Heart Rate</td>
<td>Waveform leads I, II, III are available with a 3-wire lead set.</td>
<td>Low Heart Rate</td>
</tr>
<tr>
<td></td>
<td>Waveform leads I, II, III, aVR, aVL, aVF, V are available with a 5-wire lead set.</td>
<td>High Heart Rate</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Extreme Bradycardia</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Extreme Tachycardia</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Pacer Failure to Capture</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Pacer Failure to Sense</td>
</tr>
<tr>
<td>ST</td>
<td>ST analysis is enabled</td>
<td>ST Single</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ST Dual</td>
</tr>
</tbody>
</table>

* Alarm only available when the Telepack with Nonin SpO2 module is connected
Parameters/Waveforms

<table>
<thead>
<tr>
<th>PARAMETERS</th>
<th>SETTINGS</th>
<th>PANORAMA ALARMS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arrhythmia</td>
<td>Arrhythmia analysis is enabled</td>
<td>Asystole</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Bigeminy</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Couplet</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Irregular Heart Rate</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Pause</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Run</td>
</tr>
<tr>
<td>SpO2</td>
<td>Telepack with Nonin SpO2 Module</td>
<td>Low SpO2*</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High SpO2*</td>
</tr>
</tbody>
</table>

* Alarm only available when the Telepack with Nonin SpO2 module is connected

Additional Information

For information regarding how to program and configure a Panorama Telepack, refer to the “Wireless Tab” on page 9-63.

For additional information regarding Panorama Telepacks, refer to “Telepack (Optional)” on page 10-1.

For additional information regarding the Nonin SpO2 Module, refer to the “Telepack with Nonin SpO2 Module (Optional)” on page 10-10.

For information regarding the care and maintenance of a Panorama Telepack, refer to “User Maintenance” on page 11-1.

For information regarding the parameters monitored by a Panorama Telepack, refer to “Parameter Specifications” on page 12-1.
1.2.17  **Network Printer**

The Network Printer provides a vehicle for printing collected data from the Panorama Patient Monitoring Network. One Panorama Central Station is capable of supporting up to two printers.

The following network printers are approved for use with the Panorama Central Station:

- HP LaserJet 4050n
- HP LaserJet 4100n
- HP LaserJet 4200n
- HP LaserJet 4250n
- HP LaserJet 4350n
- HP LaserJet P4015n
- HP LaserJet M602
- HP LaserJet M605

**NOTE:** The HP LaserJet 4250n/4350n printer is intended for use with Panorama software version 8.2.5 or higher.

**Additional Information**

For information regarding the setup and configuration of a printer with the Panorama Central Station, refer to the Panorama Printer Configuration Manual (P/N 0070-00-0561).

1.3  **Wireless Network**

The Panorama Central Station supports wireless communication. The Panorama Central Station can communicate with Passport 2, Spectrum, Spectrum OR, T1, Passport 8/Passport 12, Passport 12M, DPM 6, Passport V and V 12/V 21 bedside monitors in a wireless way:

The wireless bedside monitor transmits waveforms and numerics of parameters (e.g. ECG, Resp, Temp, SpO2, CO2, NIBP), related alarms and alarm settings, patients' information, operating mode, trends and events, to the central station. The waveforms, numerics, alarms, alarm settings, patients' information, operating mode and historical data displayed on central station are consistent with the wireless bedside monitor.

The central station transmits alarm settings, parameter settings, patients' information and operating mode settings of the monitor to the wireless bedside monitor. The alarm settings, parameter settings, patients' information and operating mode settings of the wireless bedside monitor are consistent with central station.

1.3.1  **Requirements for Setting up the Wireless Network**

To set up the wireless network, the following requirements should be met:
The Cisco Aironet 1242 Access Point is the recommended wireless Access Point (AP). Performance cannot be guaranteed if any other wireless AP is used.

- The wireless AP should work at a rate no less than 802.11g.
- The channel bandwidth should be 20 MHz.
- WPA2-PSK security method is recommended for authentication and encryption.
- The throughput capacity of each device is as follows:
  - Passport 8 monitor is 729 kbps
  - Passport 12 monitor is 747 kbps
  - Passport 12M or DPM 6 monitor is 780 kbps
  - Passport V monitor is 720 kbps
  - V 12 or V 21 monitor is 480 kbps
  - T1 monitor is 758 kbps

The total throughput of the wireless devices connected to a wireless AP should be less than the effective transmitting capability of the wireless AP. Refer to the AP manufacturer for the AP effective transmitting capability.

- Where the monitor is located, the wireless AP signal strength should be no less than -65 dBm.
- Where the monitor is located, the signal strength of other Wi-Fi devices of the same channel should be no greater than -85 dBm.
- Where the monitor is located, the signal strength of other Wi-Fi devices of the adjacent channels should no greater than -50 dBm.
- The recommended distance between the monitor and other non-Wi-Fi wireless devices, including wireless devices at the frequency of 2.4GHz, cellular mobile communication networks, microwave ovens, interphones, cordless phones and electro-surgical units, is no less than 20 cm.

**CAUTION:** Always set the wireless network according to local wireless regulations.

**CAUTION:** Authentication and encryption other than WPA2-PSK may expose sensitive data and allow a malicious attack on the system.

**CAUTION:** Keep network authentication information, for example password, safe to protect the network from being accessed by unauthorized users.

**CAUTION:** Disconnecting from the network may result in data loss, including waveforms, numerics, alarm events, alarm settings, patient information, operating mode, or trends. Check the patient in case of network disconnection and solve the network problem as soon as possible.

**CAUTION:** Do not connect non-medical devices to the wireless network.
## 1.3.2 Wireless Network Related Problems

<table>
<thead>
<tr>
<th>SYMPTOMS</th>
<th>CORRECTION ACTION</th>
</tr>
</thead>
</table>
| Unable to connect the network.    | 1. Verify that the network is available.  
2. Check if the network type of the bedside monitor is correctly set. Take Passport 12M for an example, if LAN is used, set the network type to LAN and connect the Passport 12M to the network; if WLAN is used, set the network type to WLAN and connect the Passport 12M to the network.  
3. Check that the SSID and password of the monitor are consistent with those of the wireless AP.  
4. Check for IP address conflicts. If any, set the IP addresses correctly.  
5. Check if the Mindray recommended wireless AP is used. If not, verify the AP effective transmission rate meets the throughput requirements of the connected devices.  
6. Verify the AP channel bandwidth is 20 MHz.  
7. Verify where the monitor is located, the wireless AP signal strength is no less than -65 dBm.  
8. Verify where the monitor is located, the signal strength of other Wi-Fi devices at the same channel is no greater than -85 dBm.  
9. Verify where the monitor is located, the signal strength of other Wi-Fi devices of adjacent channels is no greater than -50 dBm.  
10. Verify that the recommended distance between the monitor and other non-Wi-Fi wireless devices, including wireless devices at the frequency of 2.4GHz, cellular mobile communication networks, microwave ovens, interphones, cordless phones and electro-surgical units, is no less than 20 cm. |

| The monitor is frequently offline or disconnects from the network. The transmission delay is too long. | 1. Check if the Mindray recommended wireless AP is used. If not, verify the AP effective transmission rate meets the throughput requirements of the connected devices.  
2. Verify the AP channel bandwidth is 20 MHz.  
3. Verify where the monitor is located, the wireless AP signal strength is no less than -65 dBm.  
4. Verify where the monitor is located, the signal strength of other Wi-Fi devices at the same channel is no greater than -85 dBm.  
5. Verify where the monitor is located, the signal strength of other Wi-Fi devices of adjacent channels is no greater than -50 dBm.  
6. Verify that the recommended distance between the monitor and other non-Wi-Fi wireless devices, including wireless devices at the frequency of 2.4GHz, cellular mobile communication networks, microwave ovens, interphones, cordless phones and electro-surgical units, is no less than 20 cm.  
7. Verify that no unauthorized devices are connected to the wireless AP. |
1.4 Antivirus

The Panorama Central Station, WorkStation and ViewStation are pre-installed with an anti-virus application McAfee S3 Control Solidifier for Windows. The Solidifier provides a real-time change monitoring and auditing of changes to software code and configuration on servers.

The Solidifier works in three modes:

- **Disabled mode**: After installation the default mode of the Solidifier is Disabled. You need to change the Solidifier mode to Enabled for getting it operational.

- **Enabled mode**: The Enabled mode is the operational mode of the Solidifier. The Solidifier is fully functional in this mode.

- **Update mode**: The Update Mode permits updates to a system and allows all update actions including addition, removal or modification of software on the system. It tracks every update action (change) and allows software maintenance.

Refer to Installing Panorama Central Station Software Instruction (P/N 0065-00-0409) for more information.
## 1.5 Troubleshooting

This section lists some of the potential issues that may occur with the monitor.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUES *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wandering ECG</td>
<td>Respiration artifact.</td>
<td>Try a different base line lead configuration.</td>
</tr>
<tr>
<td>Noisy ECG traces</td>
<td>Loose or dry electrodes.</td>
<td>Apply new electrodes.</td>
</tr>
<tr>
<td></td>
<td>Defective electrode wires.</td>
<td>Replace wires as necessary.</td>
</tr>
<tr>
<td></td>
<td>Patient cable or leads are routed too close to other electrical devices.</td>
<td>Eliminate 60Hz interference. Use ECG cable with built-in filter block.</td>
</tr>
<tr>
<td>Low Amplitude ECG</td>
<td>Electrode could be positioned over a bone or muscle mass.</td>
<td>Reposition electrodes. Select ECG SIZE button.</td>
</tr>
<tr>
<td>Excessive Electrosurgical Interference</td>
<td>Connections not tight and properly secured.</td>
<td>Ensure proper connection. (Electrode to lead, lead to cable, cable to monitor.)</td>
</tr>
<tr>
<td></td>
<td>Electrodes dry.</td>
<td>Reprep skin and apply fresh moist electrodes.</td>
</tr>
<tr>
<td></td>
<td>Cable or lead wires damaged.</td>
<td>Check with continuity tester.</td>
</tr>
<tr>
<td>Excessive alarms: heart rate, lead fault</td>
<td>Electrodes dry.</td>
<td>Reprep skin and apply fresh, moist electrodes.</td>
</tr>
<tr>
<td></td>
<td>Alarm limits set too close to patient's normal heart rate.</td>
<td>Readjust.</td>
</tr>
<tr>
<td></td>
<td>Rwave wrong size.</td>
<td>Must be twice the amplitude of other part of wave form.</td>
</tr>
<tr>
<td></td>
<td>Excessive patient movement or muscle tremor.</td>
<td>Reposition electrodes and secure with tape if necessary.</td>
</tr>
<tr>
<td>Low Amplitude ECG Signal</td>
<td>Gain set too low. (Set through SIZE button.)</td>
<td>Readjust as required.</td>
</tr>
<tr>
<td></td>
<td>Skin improperly prepared.</td>
<td>Abrade skin.</td>
</tr>
<tr>
<td></td>
<td>Possibly not patient's normal complex.</td>
<td>Check with 12 lead electrocardiogram.</td>
</tr>
<tr>
<td></td>
<td>Electrode could be positioned over a bone or muscle mass.</td>
<td>Reposition electrodes.</td>
</tr>
<tr>
<td>Waveforms Not Moving</td>
<td>FREEZE button may have been pressed.</td>
<td>Select the FREEZE button to unfreeze the trace.</td>
</tr>
<tr>
<td>Temperature Probes not Working</td>
<td>Poor contact from probes to body.</td>
<td>Check the body surface contact at the probe tip. Reposition or apply thermoconductive gel.</td>
</tr>
<tr>
<td>Display Appears to be Off</td>
<td>Mains power switch may not be on.</td>
<td>Check main power switch on side panel.</td>
</tr>
<tr>
<td></td>
<td>Unit may not be plugged into an AC outlet.</td>
<td>Check power cord (Is it plugged in?).</td>
</tr>
<tr>
<td></td>
<td>If used as a portable, battery pack may be drained.</td>
<td>If battery pack is drained, plug into an AC outlet to recharge the battery. A period of 16 hours is required for a full charge.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
<table>
<thead>
<tr>
<th>MESSAGE/ISSUES *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Disabled Alarm Tone</td>
<td>Alarm ACK button pressed.</td>
<td>Check for message.</td>
</tr>
<tr>
<td></td>
<td>Beep volume low.</td>
<td>Increase beep volume.</td>
</tr>
<tr>
<td>ECG Base Line With No Wave form</td>
<td>The Gain control is not set high enough. Set through SIZE button.</td>
<td>Readjust as required.</td>
</tr>
<tr>
<td></td>
<td>Lead wires and patient cable not fully inserted into proper receptacle.</td>
<td>Check insertion.</td>
</tr>
<tr>
<td></td>
<td>Cable or lead wires damaged.</td>
<td>Check with lead continuity tester.</td>
</tr>
<tr>
<td>Base Line Wander</td>
<td>Patient moving excessively.</td>
<td>Secure lead wires and cable to patient.</td>
</tr>
<tr>
<td></td>
<td>Patient's respiration.</td>
<td>Reposition electrodes</td>
</tr>
<tr>
<td></td>
<td>Electrodes dry.</td>
<td>Reprep skin and apply fresh moist electrodes.</td>
</tr>
<tr>
<td></td>
<td>Static build up around patient.</td>
<td>Check with hospital engineer.</td>
</tr>
<tr>
<td>Damped Invasive Wave form</td>
<td>Air bubbles in tubing.</td>
<td>Eliminate air from tubing.</td>
</tr>
<tr>
<td></td>
<td>Kinked catheter.</td>
<td>Slightly alter position of catheter.</td>
</tr>
<tr>
<td></td>
<td>Catheter against wall of blood vessel.</td>
<td>Check for leaks at connector.</td>
</tr>
<tr>
<td></td>
<td>Blood in tubing.</td>
<td>Pump pressure bag up to 300 mmHg.</td>
</tr>
<tr>
<td>No waveform for an invasive BP</td>
<td>Improper attachment of transducer to monitor.</td>
<td>Check transducer connection.</td>
</tr>
<tr>
<td></td>
<td>Faulty transducer.</td>
<td>Try a different transducer.</td>
</tr>
<tr>
<td>Recorder Report is Blank</td>
<td>Thermal paper may be installed incorrectly. (upside down)</td>
<td>Remove paper and reinstall with paper feeding off of the spool from the bottom.</td>
</tr>
<tr>
<td>Resp. Wave form Too Small</td>
<td>Patient breathing shallow or turned on side.</td>
<td>Change lead selection.</td>
</tr>
<tr>
<td>False Apnea Alarm</td>
<td>Apnea delay may be improperly set. Patient may be having frequent episodes of CVA.</td>
<td>Choose another apnea delay.</td>
</tr>
<tr>
<td></td>
<td>Scale set inappropriately.</td>
<td>Reposition electrodes to better detect respirations.</td>
</tr>
<tr>
<td>CHK Lead</td>
<td>Due to increased impedance.</td>
<td>Prep chest.</td>
</tr>
<tr>
<td></td>
<td>Chest hair under electrodes.</td>
<td>Change electrodes.</td>
</tr>
<tr>
<td></td>
<td>Dried electrode gel.</td>
<td>Replace electrode.</td>
</tr>
<tr>
<td></td>
<td>Electrode off.</td>
<td>Replace lead.</td>
</tr>
<tr>
<td></td>
<td>Lead off.</td>
<td>Replace lead wires.</td>
</tr>
<tr>
<td></td>
<td>Cracked lead wires.</td>
<td>Clean and abrade skin before applying electrodes.</td>
</tr>
<tr>
<td></td>
<td>Poor skin prep.</td>
<td></td>
</tr>
<tr>
<td>CVA</td>
<td>Can be caused by shallow breathing or an apnea event.</td>
<td>Check the patient</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Adjust scales or leads if necessary.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
<table>
<thead>
<tr>
<th>MESSAGE/ISSUES *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>No Resp. Wave form or</td>
<td>Patient not connected to a patient safety cable.</td>
<td>Turn respiration on (“OFF” will be displayed in Resp. window). Check</td>
</tr>
<tr>
<td>Rate Displayed</td>
<td></td>
<td>that proper patient cable is used.</td>
</tr>
<tr>
<td></td>
<td>Patient connected using Patient ESIS Choke/Cable.</td>
<td>Use 3-wire lead Patient Cable - non ESIS.</td>
</tr>
<tr>
<td>BAD CARD</td>
<td>View 12™ card is not the correct version or the card data has</td>
<td>Obtain replacement card.</td>
</tr>
<tr>
<td></td>
<td>become corrupt.</td>
<td></td>
</tr>
<tr>
<td>Artifact</td>
<td>View 12™ card is detecting muscle artifact or electrical interference</td>
<td>Check leads, follow skin preparation procedure. Check for electrical</td>
</tr>
<tr>
<td></td>
<td>from auxiliary devices.</td>
<td>interferences, replace wires as necessary.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
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2.0 Getting Started

This section provides an overview of the Panorama Central Station, the ViewStation and the WorkStation.

- The Panorama Central Station
- The ViewStation and the WorkStation
- The Main Screen
- Common Controls and Elements
- Quick Reference Guide
- Troubleshooting
2.1 The Panorama Central Station

The Panorama Central Station collects patient data from both wireless and hardwired devices and displays it based on its configured settings. Data is collected from the bedside monitors (i.e., Passport, Spectrum, DPM 6/DPM 7, Passport 12M/Passport 17M, Passport 8/Passport 12, T1 and V 12/V 21 monitors) and Panorama Telepacks (608 Mhz). The type of data that is collected and viewed is determined by the monitoring device and the selected parameters.

The Panorama Central Station uses either a single or dual display. Single display setups can accommodate eight (8) or 12 patient tiles. Dual display setups can accommodate up to eight (8) patient tiles per display, showing a maximum of 24 patient tiles between the two displays.

The full screen display (shown in FIGURE 2-1) includes patient tiles for viewing patient data and a menu bar for navigating the system.

NOTE: The Paging menu button will only be displayed if the Paging option has been installed.

FIGURE 2-1 Full Screen Display (Central Station with Paging Option Installed)
The Panorama Central Station uses a split-screen display (shown in FIGURE 2-2) to allow users to simultaneously enter patient information, and monitor patient data. The top half of the screen displays patient data, while the bottom half of the screen is converted to a workspace for entering and modifying patient information.

**FIGURE 2-2** Split-Screen Display (Central Station with Paging Option Installed)
2.2 The ViewStation and the WorkStation

The ViewStation and the WorkStation may display patient tiles from other Panorama Central Stations on the network from a remote location(s). The host central station’s Panorama Central Network (C-LAN) configuration settings, and the software that is installed at the remote location, determine the level of functionality available for each patient tile. A hospital area or department may also choose to restrict remote location access to its patient tiles by configuring its C-LAN configuration settings to off.

NOTE: C-LAN configuration settings are established by the System Administrator and are based on the needs of the individual hospital or hospital area.

The host central station C-LAN configuration settings available are:

<table>
<thead>
<tr>
<th>HOST CENTRAL STATION C-LAN NETWORK SETTING</th>
<th>REMOTE LOCATION PATIENT TILE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Full</td>
<td>Patient tiles from the host central station will have full interactive functionality when using a WorkStation or view-only functionality when using a ViewStation.</td>
</tr>
<tr>
<td>View Only</td>
<td>Patient tiles from the host central station will have view-only functionality regardless of the software installed at the remote location.</td>
</tr>
<tr>
<td>Off</td>
<td>Patient tiles from the host central station cannot be seen from any remote location.</td>
</tr>
</tbody>
</table>
FIGURE 2-3 Full Screen Display (ViewStation and WorkStation)

The full screen display (shown in FIGURE 2-3) includes patient tiles for viewing patient data and a menu bar for navigating the system.

**NOTE:** The Network and Audio Off/Audio Pause menu bar buttons are only displayed at the ViewStation and the WorkStation.

**NOTE:** The Audio Off/Audio Pause buttons share the same location within the menu bar. The button that is displayed is dependant on the pause duration setting in the System Setup. When the duration is set to Permanent, the Audio Off button is displayed. When the duration is set to another time period, the Audio Pause button is displayed. For the duration setting, refer to “Setting the System Alarms Options (Central Station)” on page 9-15.
The ViewStation and the WorkStation use a split-screen display (shown in FIGURE 2-4) to allow users to view patient information and monitor patient data. The top half of the screen displays patient data, while the bottom half of the screen is used to view patient data in more detail.

**FIGURE 2-4** Split-Screen Display (ViewStation)
2.3 The Main Screen

The Main Screen for the Panorama Central Station, the ViewStation, and the WorkStation is used to monitor patient data and to navigate the system.

The Main Screen includes the following elements:

- Patient Tiles (shown in FIGURE 2-1/FIGURE 2-2)
- Menu Bar (shown in FIGURE 2-1/FIGURE 2-2)
- Tabs (shown in FIGURE 2-2)

2.3.1 Patient Tiles

Patient tiles (shown in FIGURE 2-5) are the core elements of the Panorama system. Patient tiles display patient information and data that is obtained from an external monitoring device. One patient and one monitoring device are assigned to each patient tile. The number of active patient tiles displayed at the Panorama Central Station is dependent on the number of patient tiles that were purchased with the system.

Each patient tile is divided into the following sections:

- VIEW/Alarm ACK Button
- Care Group Indicator
- Digital Data Tile
- REC Button
- Patient Status Line
- Waveform Data Tile

FIGURE 2-5 Patient Tile

For information regarding admitting a patient to the Panorama Central Station, refer to “Auto-Admitting a Patient” on page 2-21. For information regarding the assignment of monitoring devices to a patient tile, refer to the “Equipment Setup Tab” on page 9-33.
VIEW/Alarm ACK Button

The VIEW/Alarm ACK (Alarm Acknowledge) buttons share the same location within the patient tile. The button that is displayed is dependant on the operating state of the patient tile. During normal operating conditions, the VIEW button (shown in FIGURE 2-5) is displayed in the patient tile. During an alarm condition, the Alarm ACK button is displayed in the patient tile.

NOTE: The Alarm ACK button is not available at the ViewStation or the WorkStation.

The VIEW button is available in a patient tile when no alarm conditions exist. When the VIEW button is selected, the full-screen display changes to the split-screen mode and provides access to the Bedside, Trends, Events, Disclosure and ST tabs.

<table>
<thead>
<tr>
<th>TAB</th>
<th>PURPOSE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bedside</td>
<td>Provides an expanded view of a patient’s digital and waveform data.</td>
</tr>
<tr>
<td>Trends</td>
<td>Provides list and graphic views of a patient’s historical digital data.</td>
</tr>
<tr>
<td>Events</td>
<td>Provides a list of the events that have occurred for a patient tile. If waveform data is associated with an event it may also be viewed in the Events tab.</td>
</tr>
<tr>
<td>Disclosure</td>
<td>Provides a view of a patient’s historical waveform data. Data may be displayed in a compressed or magnified view.</td>
</tr>
<tr>
<td>ST</td>
<td>Allows for the configuration and display of a patient’s ST data.</td>
</tr>
</tbody>
</table>

The Alarm ACK button is available in the patient tile when an alarm condition exists. The Alarm ACK button silences the audio portion of a patient alarm for the predetermined configured time.

Care Group Indicator

The Care Group indicator (shown in FIGURE 2-5) is a rectangular bar that represents the care group to which a patient tile is assigned. Colors can be assigned to care group indicators. A black indicator is displayed if a patient tile is not assigned to a care group.

For additional information regarding care groups, refer to the “Care Group Tab” on page 9-9.

NOTE: Colorized Care Group indicators are not available at the ViewStation or the WorkStation.
Digital Data Tile
The digital data tile (shown in FIGURE 2-5) may include the following patient information:

- Digital data measurements
  Examples of digital data parameters include: Heart Rate, Respiration, SpO₂, Temperature, NIBP, PVC, and ST. The parameters available to a patient tile are dependent on the selected monitoring device.
  
  Digital data is shown in the color assigned to the parameter.

- Alarms on/Alarms off icon
- ST analysis icon
- Arrhythmia analysis icon
- Heart Rate parameter lead

For information regarding supported devices and the parameters measured by each, refer to “Supported Devices” on page 1-3. For information regarding the parameters displayed in the digital data tile, refer to “Parameter Specifications” on page 12-2.

The following table lists the icons that may be shown in a digital data tile.

### Digital Data Icons

<table>
<thead>
<tr>
<th>ICON</th>
<th>MEANING</th>
</tr>
</thead>
<tbody>
<tr>
<td>![Icon]</td>
<td>All of the alarms for the selected patient are enabled.</td>
</tr>
<tr>
<td>![Icon]</td>
<td>At least one alarm for the selected parameter is disabled.</td>
</tr>
<tr>
<td>![Icon]</td>
<td>Arrhythmia analysis is active for the selected patient. This icon is only applicable to the Heart Rate tile.</td>
</tr>
<tr>
<td>![Icon]</td>
<td>ST analysis is active for the selected patient. This icon is only applicable to the Heart Rate tile.</td>
</tr>
<tr>
<td>![Icon]</td>
<td>Lead labels for the Heart Rate parameter include I, II, III, aVR, aVL, aVF, V1, V2, V3, V4, V5, and V6, depending on the lead set used. This icon is only applicable to the Heart Rate tile. For additional information, refer to “Parameter Specifications” on page 12-2.</td>
</tr>
</tbody>
</table>

During an alarm condition, the parameter’s digital data tile flashes in the color that is associated with its assigned alarm priority.

- The digital data flashes in red when a Priority 1 alarm occurs.
- The digital data tile flashes in yellow when a Priority 2 alarm occurs.
- The data is displayed in yellow (constant, not flashing) when a Priority 3 alarm occurs.

**REC Button**

The **REC** button (shown in FIGURE 2-5) prints an All Strips Report for the selected patient tile. Selecting the **REC** button sends a print request to the **Print Status** tab and prints an All Strips Report. The All Strips Report provides real-time monitoring data for the selected patient. For additional information, refer to the “All Strips Report” on page 8-2.
Patient Status Line

The Patient Status line (shown in FIGURE 2-5) displays patient demographic information, physiological event messages and technical event messages. Demographic information (Last Name, First Name, Bed, Patient ID) is only shown if it was provided. Demographic information is displayed as configured in the Unit Choices tab.

The following guidelines regulate the display of messages in the Patient Status line:

- A patient’s demographic information, if available, is displayed in white text.
- All Alarm Priority 1 physiological messages are displayed in red text.
- Alarm Priority 2/3 physiological messages are displayed in yellow text.
- If alarms have been paused or turned off, the messages are displayed.
- Technical event messages are displayed in white text.

Waveform Data Tile

The Waveform Data tile (shown in FIGURE 2-5) displays the waveform data that was selected for the patient tile. The Waveform Data portion of a patient tile is capable of displaying up to two of the following waveform data parameters: ECG, Pleth, Respiration, IBP, O₂, and Agent. The available parameters are dependant on the selected monitoring device. For information regarding supported devices, and parameters measured, refer to “Supported Devices” on page 1-3.

Waveforms display in a patient tile using the following guidelines:

- Each waveform tile includes its own baseline.
- Waveforms are shown in a patient’s configured color, wave gain, pacer enhancement and filter settings. If individual configurations are not set, the system default settings are used.

For information regarding the selection of patient waveform data, refer to the “Display Tab” on page 5-41. For information regarding the parameters that can be shown in the waveform tile, refer to “Parameter Specifications” on page 12-2.

2.3.2 Menu Bar

The Panorama Central Station, ViewStation, and the WorkStation menu bars (shown in FIGURE 2-6, FIGURE 2-7, and FIGURE 2-8) are used to view the system status, provide navigation for the application, and view the system time and date. All menu bars are fixed at the bottom of the screen.
Menu bar selections include:

- System Status Line
- Menu Buttons
- System Date/Time

**FIGURE 2-6** Panorama Central Station Menu Bar

**FIGURE 2-7** Panorama Central Station Menu Bar - Paging Option Enabled

**FIGURE 2-8** ViewStation and WorkStation Menu Bar

**System Status Line**

The System Status line (shown in FIGURE 2-7) displays messages regarding the status of the individual central station, ViewStation, or WorkStation. The System Status line is located above the menu buttons, in the menu bar. System status messages refer to the operation of the system. For a listing of System Status messages, refer to “System Event Messages and Error Codes” on page 13-26.

**Menu Buttons**

Menu buttons (shown in FIGURE 2-7) provide access to additional interactive screens. When a menu button is selected, a series of corresponding tabs is opened. The tabs that are displayed will vary depending on whether the patient tile is accessed from the central station, ViewStation, or from a WorkStation. For specific information related to a specific tab or button, refer to the appropriate chapter in this book or to the book index.
Table 2-1 provides a list of the tabs that are associated with each menu button.

### TABLE 2-1

<table>
<thead>
<tr>
<th>BUTTON</th>
<th>PURPOSE</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Patient Setup</strong></td>
<td>Provides access to the following tabs:</td>
</tr>
<tr>
<td></td>
<td>• The <strong>Demographics</strong> tab is used to enter and modify patient demographic information (i.e., Last name, First name, Patient ID, Bed, Height, Weight, etc.).</td>
</tr>
<tr>
<td></td>
<td>• The <strong>Transfer</strong> tab is used to move patient information from tile to tile within a Panorama Central Station or between central stations.</td>
</tr>
<tr>
<td></td>
<td>• The <strong>Discharge</strong> tab is used to remove a patient from centralized monitoring.</td>
</tr>
<tr>
<td></td>
<td>• The <strong>Display</strong> tab is used to select the parameter information displayed in a patient tile.</td>
</tr>
<tr>
<td></td>
<td>• The <strong>Standby</strong> tab is used to temporarily suspend monitoring while preserving patient settings and historical data.</td>
</tr>
<tr>
<td></td>
<td>• The <strong>Wave Gain</strong> tab is used to adjust the wave gain used for displayed patient waveforms.</td>
</tr>
<tr>
<td></td>
<td>• The <strong>Print Setup</strong> tab is used to select the parameters that are printed in a patient report.</td>
</tr>
<tr>
<td><strong>Patient Alarms</strong></td>
<td>Provides access to the following tabs:</td>
</tr>
<tr>
<td></td>
<td>• The <strong>Patient Alarm Setup</strong> tab is an optional, password-protected tab used to restrict access to the Patient Alarm Limits and Alarm Responses tabs.</td>
</tr>
<tr>
<td></td>
<td>• The <strong>Alarm Limits</strong> tab is used to set a patient alarm limit thresholds.</td>
</tr>
<tr>
<td></td>
<td>• The <strong>Alarm Responses</strong> tab is used to configure system behaviors during alarm conditions.</td>
</tr>
<tr>
<td><strong>Report</strong></td>
<td>Provides access to the following tabs:</td>
</tr>
<tr>
<td></td>
<td>• The <strong>Patient Reports</strong> tab is used to request patient-specific reports.</td>
</tr>
<tr>
<td></td>
<td>• The <strong>System Reports</strong> tab is used to request system reports.</td>
</tr>
<tr>
<td></td>
<td>• The <strong>Print Status</strong> tab is used to display the list of print requests that have been made and the status of each request.</td>
</tr>
</tbody>
</table>

* Disable ECG is available on version 8.7.1 and above.

** Indicates button is not available at the ViewStation or the WorkStation

*** Indicates button is only available at the ViewStation and the WorkStation
Table 2-1

<table>
<thead>
<tr>
<th>BUTTON</th>
<th>PURPOSE</th>
</tr>
</thead>
<tbody>
<tr>
<td>System Setup</td>
<td>Provides access to the following tabs:</td>
</tr>
<tr>
<td></td>
<td>• The Parameter Color tab is used to create the system default color and parameter assignment.</td>
</tr>
<tr>
<td></td>
<td>• The Print Setup tab is used to assign a system default printer for report printing.</td>
</tr>
<tr>
<td></td>
<td>• The Touch Screen tab is used to turn the touch screen on and off and recalibrate the touchscreen display.</td>
</tr>
<tr>
<td></td>
<td>• The Installation Setup tab is a password-protected tab used to restrict access to the following tabs:</td>
</tr>
<tr>
<td></td>
<td>• The Care Group tab is used to create the system level care group assignment.</td>
</tr>
<tr>
<td></td>
<td>• The Volume tab is used to adjust system default volume levels for physiological alarms.</td>
</tr>
<tr>
<td></td>
<td>• The System Alarms tab is used to create system default settings for alarms.</td>
</tr>
<tr>
<td></td>
<td>• The Passwords tab is used to create passwords that control access to the system.</td>
</tr>
<tr>
<td></td>
<td>• The Equipment Setup tab is used to enter information regarding external monitoring devices used with the Panorama Central Station.</td>
</tr>
<tr>
<td></td>
<td>• The More/Previous tab is used to navigate to the forward and backward between the sets of System Setup tabs.</td>
</tr>
<tr>
<td></td>
<td>• The Date/Time tab is used to select the system default date and time formats.</td>
</tr>
<tr>
<td></td>
<td>• The Unit Priorities tab is used to establish the system default order of parameters.</td>
</tr>
<tr>
<td></td>
<td>• The Unit Choices tab is used to adjust the system default settings for Discharge at Bedside, Trend Access Order, Signal Loss Delay, Paging Demographics, Paging Password Protection, Demographic Line Information, Disable ECG*, Height, Weight, Temperature, CO2, and Trend Display.</td>
</tr>
<tr>
<td></td>
<td>• The Wave Gain tab is used to adjust the system default wave gain setting for displaying waveforms.</td>
</tr>
<tr>
<td></td>
<td>• The Wireless tab is used to program wireless monitoring devices.</td>
</tr>
<tr>
<td>** Paging (Optional)</td>
<td>The Paging tab allows for the configuration of pager information for a Panorama Central Station that has the paging option enabled.</td>
</tr>
<tr>
<td>** Note: **</td>
<td>When using a dual display setup, the Paging menu bar button will be disabled if the Paging tab is active on the other display.</td>
</tr>
<tr>
<td>** Network**</td>
<td>Provides access to the following tab:</td>
</tr>
<tr>
<td></td>
<td>• The Network tab is used to access and select monitored patients on the Panorama Central Network (C-LAN) for viewing at a ViewStation or a WorkStation.</td>
</tr>
</tbody>
</table>

* Disable ECG is available on version 8.7.1 and above.  
** Indicates button is not available at the ViewStation or the WorkStation  
*** Indicates button is only available at the ViewStation and the WorkStation
The System Date/Time

The System Date/Time (shown in FIGURE 2-6, FIGURE 2-7, and FIGURE 2-8) portion of the menu bar displays the system’s date/time information. The system date/time is displayed in the format selected in the Date/Time tab. For additional information regarding the time and date formats available, refer to “Date/Time Tab” on page 9-41.

2.3.3 Tabs

Tabs (shown in FIGURE 2-9) are used to access different screens associated with the Panorama Central Station, ViewStation, and the WorkStation. Tabs are accessed using menu buttons, or the patient tile VIEW button. Each tab includes sidebar buttons for expanded functionality.

### TABLE 2-1

<table>
<thead>
<tr>
<th>BUTTON</th>
<th>PURPOSE</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Audio Off/Pause</strong></td>
<td>This button does not provide access to any tab. When selected, the Audio Off/Pause button silences the audio alarms for all patient tiles at a ViewStation or a WorkStation for the configured amount of time. The Audio Off/Pause button does not affect the audio or visual alarm indicators at the bedside monitor or at the host central station.</td>
</tr>
<tr>
<td>(Optional)</td>
<td></td>
</tr>
<tr>
<td><strong>Alarm Reset</strong></td>
<td>When patient is selected, the Alarm Reset button will have the following effects for the selected patient tile at a central station, a ViewStation or a WorkStation:</td>
</tr>
<tr>
<td></td>
<td>• Permanently silences all current audio alarms.</td>
</tr>
<tr>
<td></td>
<td>• Acknowledges all current alarms and latch alarms.</td>
</tr>
<tr>
<td></td>
<td>• Discontinues the audio off/pause state. The visual alarm is not affected, but the audio alarm is silenced. If a new alarm occurs, the alarm will sound.</td>
</tr>
<tr>
<td></td>
<td>• Discontinues the alarm off/pause state. The visual and audio alarms are turned on. If there is an active alarm, the visual and audio alarms are triggered.</td>
</tr>
<tr>
<td></td>
<td>The Alarm Reset button does not affect the audio or visual alarm indicators at the bedside monitor, or the visual alarm indicator at the host central station, ViewStation or WorkStation.</td>
</tr>
<tr>
<td><strong>Print</strong></td>
<td>The Print button provides an additional method to print patient information and reports.</td>
</tr>
<tr>
<td></td>
<td><strong>NOTE:</strong> The Print menu bar button will be disabled when in the Normal Screen mode.</td>
</tr>
</tbody>
</table>

* Disable ECG is available on version 8.7.1 and above.
** Indicates button is not available at the ViewStation or the WorkStation
*** Indicates button is only available at the ViewStation and the WorkStation
2.3.4 Sidebar Buttons
Sidebar buttons (shown in FIGURE 2-9) are used to perform an action (such as suspend alarms, enable arrhythmia, or return to the Normal Screen mode), add functionality to the tab (such as Skip to, Graphic, and Alarm Responses), or change the contents of the tab (such as Alarm Responses or Alarm Limits).

FIGURE 2-9 Tab Architecture
2.4 Common Controls and Elements

This section provides information regarding several types of controls and elements that are used throughout the system.

Common controls and elements include:

- Buttons
- Progress Bar
- Message Dialog Boxes
- Keyboard Dialog Box
- Keypad Dialog Box
- Weight Dial Dialog
- Height Dial Dialog

Buttons

Buttons provide additional options or perform an action when selected. For example:

- To perform an action. This type of button temporarily changes its appearance to indicate that it has been selected.

  FIGURE 2-10 illustrates the change in appearance for an Action button before and after it is selected.

  ![Figure 2-10 Action Buttons](image)

- To enable/disable a function. This type of button changes its appearance to indicate if the function is enabled or disabled.

  FIGURE 2-11 illustrates the change in appearance for a Function (or latched) button before and after it is selected. When one of these buttons is selected, its function will remain in effect until the button is released. In the first example shown in FIGURE 2-11, the alarms will be suspended until the button is unselected.

  ![Figure 2-11 Function and Check Box Buttons](image)
Progress Bar
The Panorama Central Station uses progress bars to indicate the progress of a lengthy operation.

![Progress Bar](FIGURE 2-12 Progress Bar)

Message Dialog Boxes
The Panorama Central Station uses message dialog boxes to verify an action (shown in FIGURE 2-13) and to provide information (shown in FIGURE 2-14).

A Question message dialog box contains Yes and No buttons.

- The Yes button confirms the action, closes the message box and enables all applicable sidebar buttons.
- The No button cancels the action, closes the dialog box and enables applicable controls and sidebar buttons.

![Message Dialog Box (Question)](FIGURE 2-13 Message Dialog Box (Question))

An Informational message dialog box contains an OK button.

The OK button closes the message box and enables all applicable controls and sidebar buttons.

![Message Dialog Box (Informational)](FIGURE 2-14 Message Dialog Box (Informational))

NOTE: All controls are disabled when a Message dialog box is displayed.
Common Controls and Elements

Getting Started

Keyboard Dialog Box
The Panorama Central Station uses a keyboard dialog box (shown in FIGURE 2-15) to enter and edit alphanumeric information. The buttons on the keyboard dialog box are similar to a standard computer keyboard.

**NOTE:** Tab controls are disabled when the keyboard dialog box is displayed.

![Keyboard Dialog Box](image)

The keyboard dialog box includes:

- The name of the selected field
- An Edit box to show the entered value
- A **Close** button to discard changes
- An **Enter** button to accept changes
- A Backspace («==) button to clear a single character in the edit box
- A **Caps Lock** button to enable or disable caps lock
  A Caps Lock indicator is shown when **Caps Lock** is enabled.

Keypad Dialog Box
The Panorama Central Station uses the keypad dialog box (shown in FIGURE 2-16) to enter and edit numeric information. The buttons in the keypad dialog box are similar to a standard calculator.

**NOTE:** Tab controls are disabled when the keypad dialog box is displayed.
Getting Started
Common Controls and Elements

The keypad dialog box includes:

- The name of the selected field
- An Edit box to show the entered value
- The acceptable range of values
- A **Clear** button to delete all of the numbers in the edit box
- A **CE** button to delete a single number from the edit box
- A **Cancel** button to discard changes
- An **OK** button to accept changes

**Weight Dial Dialog**

The Weight dial dialog can be used to quickly set the pounds and ounces (Neonate patient size only) for a patient’s weight.

**FIGURE 2-16 Keypad Dialog Box**

**FIGURE 2-17 Weight Dial Dialog (Spectrum bedside device, Neonate patient size)**
The Weight dial dialog includes:

- The name of the selected field
- A text box to show the selected value
- A dial control to select the weight
  - Turn in a clockwise direction to sequentially increase the numeric value
  - Turn in a counter-clockwise direction to sequentially decrease the numeric value
- Adjustment buttons to sequentially increase or decrease the numeric value
- A Done button to accept changes
- A Cancel button to discard changes

**Height Dial Dialog**

The Height dial dialog functions similarly to the Weight dial dialog. For additional information, refer to the Weight Dial Dialog.
2.5 Quick Reference Guide

The quick reference guide provides abbreviated procedural information for the operation of the Panorama Central Station.

For additional information, refer to the Panorama Quick Reference Guide (P/N 0002-08-7538).

Quick reference procedures include:

- Auto-Admitting a Patient
- Standby
- Discharge
- Display
- Patient Alarms (Limits and Responses)
- Viewing List Trends
- Viewing Events
- Viewing Disclosure Data

**NOTE:** Demographic information is bi-directionally shared between the bedside monitor and the Panorama Central Station.

2.5.1 Auto-Admitting a Patient

Auto-admit allows for the immediate monitoring of a patient at the Panorama Central Station.

To auto-admit a patient:

1. Verify that the monitoring device is in the Equipment List. For information on how to access the Equipment List, refer to the “Equipment Setup Tab” on page 9.33.
2. Connect/attach the monitoring device to the patient.
3. Turn on the monitoring device. The monitoring device begins monitoring the patient. The parameters that are monitored are dependent on the monitoring device.
4. Verify that the patient is being monitored at the central station.

**NOTE:** Auto-admit uses the system default settings for monitoring the patient.

2.5.2 Standby

Standby mode suspends the monitoring of a patient at the Panorama Central Station while retaining the patient’s historical data and setup priorities. It is important that a patient in this mode be alternatively monitored. The **Standby** tab is also available at the WorkStation.

To activate Standby mode at the central station or the WorkStation:

1. Select the Patient Setup menu button.
2. Choose a patient tile by selecting its digital data or waveform area.
3. Select the Standby tab.
   - The Panorama Standby Only button places the patient in Standby at the Panorama Central Station.
• The **Bedside and Panorama Standby** button places the patient in Standby at the Panorama Central Station and the bedside monitor.

**NOTE:** The Bedside and Panorama Standby button is not available when using a Telepack device.

4. Select a patient’s Standby location.
   a. Select a location from the patient location list
   b. Choose the **Select** button

To activate Standby mode at the bedside monitor:

• Select the **STANDBY** button on the front of the bedside monitor. This method places both the bedside monitor and the Panorama Central Station in **Standby** mode.

To remove a patient from the Standby mode at the Panorama Central Station or the WorkStation:

1. Select the **Patient Setup** menu button.
2. Choose a patient tile by selecting its digital data or waveform area.
3. Select the **Standby** tab.
4. Select the **Resume Monitoring** button. Monitoring is resumed at the Panorama Central Station.

To remove a patient from the Standby mode at the Panorama Central Station and the bedside monitor:

• Select the **STANDBY** button on the front panel of the bedside monitor. Monitoring resumes at the Panorama Central Station and the bedside monitor.

For additional information on Standby, refer to the "Standby Tab" on page 5-51.

2.5.3 Discharge

The discharge function discharges a patient from a patient tile and resets the Panorama Central Station for the next patient. This tab is available at the WorkStation.

To discharge a patient:

1. Select the **Patient Setup** menu button.
2. Choose a patient tile by selecting its digital data or waveform area.
3. Select the **Discharge** tab.
4. Select the **Bedside and Panorama Discharge** button.
5. A message dialog box is displayed.
   a. If at least two of the three required demographic fields (First Name, Last Name, Patient ID) are entered, the following message is displayed: Are you sure you want to discharge this patient? Selecting Yes will move the patient to the discharge list. The YES button discharges the patient. The NO button continues monitoring the patient.
   b. If at least two of the three required demographic fields are not completed, the following message is displayed: The patient’s data and submitted print jobs will be deleted because at least two of the following fields, First Name, Last Name or Patient ID have not been filled in. Do you still want to proceed with the discharge? The YES button discharges the patient. The NO button continues monitoring the patient.

To discharge a patient from a bedside monitor:

1. Select the DISCHARGE button on the front of the bedside monitor.

2. Select a discharge option.
   • The Discharge From Monitor option erases all patient information at the bedside monitor.
   • The Discharge From Central option erases all patient information at the Panorama Central Station.
   • The Discharge From Both option erases all patient information at the Panorama Central Station and at the bedside monitor.

   NOTE: When using the Discharge From Central and Discharge From Both options, if two of the three demographic fields (First Name, Last Name, and Patient ID) have been completed prior to the patient’s discharge, the patient is moved to the Discharge list at the Panorama Central Station and may be retrieved.

   NOTE: If a patient is discharged from the Panorama Central Station but not from the patient monitor, the Panorama Central Station tile will not be available for a new patient until a discharge has been performed at the patient monitor.

3. Verify the selection. The Are you sure? confirmation dialog box is displayed.
   • The Yes selection confirms the patient discharge.
   • The No selection discards the patient discharge.

For additional information regarding discharging a patient, refer to “Discharging a Patient from the Central Station and the Bedside Monitor” on page 5-37.

2.5.4 Display

The Display tab controls the order of patient data tiles and the waveforms that are shown on the Main Screen.

To change the patient tile display order at an individual Panorama Central Station, ViewStation, or WorkStation:

1. Select the Patient Setup menu button.

2. Choose a patient tile by selecting its digital data or waveform area.

3. Select the Display tab.
4. Select a digital parameter tile to change a digital parameter. Select a waveform tile to change a waveform.

5. Select a parameter from the Parameter list.

6. Choose the Select button to confirm the choice.

For additional information regarding parameter display, refer to the “Display Tab” on page 5-41.

2.5.5 Patient Alarms

Patient alarms provide for customized alarm limits and responses based on patient need. For additional information on patient alarms, refer to “Patient Alarm Setup Tab (Optional)” on page 6-2.

Alarm Limits

Alarm limits are the configured threshold settings used to monitor patient parameters. If a parameter does not fall within the configured threshold settings, an alarm occurs. When using a bedside monitor, patient alarm limit information will continuously be transmitted between the bedside monitor, the Panorama Central Station, ViewStation, and the WorkStation.

WARNING: Changing the patient alarm limits at the WorkStation will automatically change the patient alarm limits at the Panorama Central Station AND the bedside monitor (if applicable) for the selected patient tile.

To adjust alarm limits:

1. Select the Patient Alarms menu button.
2. Choose a patient tile by selecting its digital data or waveform area.
3. Select the Alarm Limits tab.
4. Select the parameter alarm limits to adjust. Use the slider bars to adjust the alarm limits.

To adjust alarm limits at the bedside monitor, refer to the bedside monitor’s Operating Instructions:

- Passport 2 (P/N 0070-00-0649-XX)
- Passport V (P/N 0070-00-0704-XX)
- Spectrum (P/N 0070-00-0648-XX)
- Spectrum OR (P/N 0070-00-0670-XX)
- DPM 6 (P/N 046-000184-00)
- DPM 7 (P/N 046-000186-00)
- V 12/V 21 (P/N 0070-00-0727-XX)
- Passport 12M/Passport 17M (P/N 046-005013-00)
- Passport 8/Passport 12 (P/N 046-004417-00)
- T1 monitor (P/N 046-002376-00)
NOTE: ST alarm limits for DPM 6/DPM 7, Passport 12M/Passport 17M, Passport 8/Passport 12 and T1 monitor must be adjusted at the bedside monitor.

For additional information on Alarm Limits, refer to “Alarm Limits Tab (Patient)” on page 6-4.

Alarm Responses
Alarm Responses determine the behavior of alarms when an alarm violation occurs at the Panorama Central Station.

To adjust alarm responses:
1. Select the Patient Alarms menu button.
2. Choose a patient tile by selecting its digital data or waveform area.
3. Select the Alarm Responses tab.
4. Select the parameter.
5. Adjust the alarm responses for the parameter.

For additional information on alarm responses, refer to “Alarm Responses Tab (Patient)” on page 6-12.

Muting Alarms
To mute a non-latched alarm (for example, IBP):
- Select the Alarm ACK button in the alarming patient tile to silence the alarm violation for the configured time period.

NOTE: The Alarm ACK button is not available at the ViewStation or the WorkStation.

For additional information on muting an alarm, refer to “VIEW/Alarm ACK Button” on page 2-8.

To mute a latched alarm (for example, lethal arrhythmias):
- Select the digital data area or waveform area of the alarming patient tile to acknowledge the alarm AFTER the alarm condition is resolved.
- Select the Alarm ACK button in the alarming patient tile to silence the alarm violation for the configured time period.

NOTE: The audio and visual indicators for a latched alarm will continue until the alarm is acknowledged, even if the alarm condition no longer exists.

NOTE: Selecting the digital data or waveform area of the alarming patient tile, at the host central station, also acknowledges the alarm at the ViewStation and the WorkStation.

For additional information about latched alarms, refer to “Latching” on page 3-3.
2.5.6 Viewing List Trends

The **Trends** tab displays all available digital data for a patient in a list format.

**NOTE:** To default to the List Trends when the Trends tab is selected, List must be selected in the Unit Choices tab Trend Access Order.

To view List Trends at the Panorama Central Station:

1. Select the **VIEW** button in the desired patient tile.
2. Select the **Trends** tab.
   - Select the **Print** menu bar button to print the trends list.

To adjust the trend display period:

1. Select the **Patient Setup** menu button.
2. Select the **Display** tab.
3. Select the **Trend Display** sidebar button.

For additional information regarding the Trends List, refer to “Trends Tab (List View)” on page 7-15.

2.5.7 Viewing Events

The **Events** tab displays all events and related waveform data for a patient.

To view events:

1. Select the **VIEW** button in the desired patient tile.
2. Select the **Events** tab.
   - Select the **Print** menu bar button to print the events list.
3. Select the **Events Filtering** sidebar button to change the filter used in the event list.

For additional information regarding events, refer to “Events Tab” on page 7-25.

4. To view the waveform data associated with an event, highlight the desired event and select the **View Waveform** sidebar button.
   - The event waveform can be printed by selecting the **Print** menu bar button.

2.5.8 Viewing Disclosure Data

The **Disclosure** tab displays all of the patient’s historical waveform data, and associated numeric information, for a selected time period.

To view disclosure data:

1. Select the **VIEW** button in the desired patient tile.
2. Select the **Disclosure** tab.
3. Select the **Display Choices** sidebar button to change the waveforms being viewed.
   a. Select the **Print** menu bar button or one of the menu bar **Print (x) Hour(s)** (**x**=number of hours to print) button to print the disclosure information.
   b. Select the **Skip To** sidebar button to view a specific time in a patient monitoring history.
4. For a magnified view of the waveform, highlight a waveform and select the **Zoom In** button.

   a. Select the **Print** menu bar button to print the magnified disclosure information.

   For additional information regarding disclosure, refer to the "Disclosure Tab" on page 7-42.
2.6 Troubleshooting

This section lists some of the potential issues that may occur when admitting a patient to the Panorama Central Station.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cannot admit patient/patient data not displayed at Central station.</td>
<td>Previous patient not discharged from system and/or monitoring device.</td>
<td>Discharge patient from system.</td>
</tr>
<tr>
<td>Monitoring device not assigned to Panorama Equipment List.</td>
<td>Assign device to Panorama Equipment List.</td>
<td></td>
</tr>
<tr>
<td>Panorama not installed as an option.</td>
<td>Access the Installation Menu, Options submenu at the bedside device. For additional information, refer to the operating instructions of the bedside monitor.</td>
<td></td>
</tr>
<tr>
<td>Network setting on monitoring device is incorrect.</td>
<td>Access the Installation Menu, System Information submenu at the bedside device. For additional information, refer to the operating instructions of the bedside monitor.</td>
<td></td>
</tr>
<tr>
<td>Telepack batteries may need to be replaced.</td>
<td>Test batteries, and if necessary, replace.</td>
<td></td>
</tr>
<tr>
<td>Telepack batteries improperly installed.</td>
<td>Reinstall batteries.</td>
<td></td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
3.0 *Alarm Behavior*

This section outlines the alarm types available in the system and the conditions that would trigger an alarm condition.

- General Alarm Behavior
- Physiological Alarms
- Technical Alarms
- System Alarms
3.1 General Alarm Behavior
The Panorama Central Station uses alarms to send notification when a patient’s physiological status changes, a patient has a specific technical issue, or when there is a system violation. When a patient is first admitted, the default system alarm settings are in effect. Individual patient alarm settings can be modified by using the Patient Alarms button at the central station or the WorkStation.

WARNING: Changing the patient alarm limits at the WorkStation will automatically change the patient alarm limits at the Panorama Central Station AND the bedside monitor (if applicable) for the selected patient tile.

By default, alarm settings send notification using:

• Visual indicators
• Audible indicators

The three alarm types used in this system include physiological, technical, and system.

3.2 Physiological Alarms
Physiological alarms indicate a violation of a monitored patient’s physiological alarm thresholds or arrhythmia analysis settings. For additional information regarding alarm thresholds and arrhythmia analysis settings, refer to “Alarms and Events” on page 13-1.

3.2.1 Alarm Behaviors
The visual and audio indicators associated with a physiological alarm are:

Visual Indicators
Visual indicators provide visual notification when an alarm threshold is violated. When a physiological alarm condition is triggered, the digital data portion of the patient tile flashes (if the alarm priority is set to 1 or 2), and a text message displays in the Patient Status line. The physiological alarm behavior varies depending on the parameter, and on the alarm priority associated with the parameter. This is discussed in further detail in the “Physiological Alarm Responses” on page 3-4.

Audio Indicators
Audio indicators provide audible notification when an alarm threshold is violated. Audio indicators may vary depending on the alarm severity, alarm type, system setup and configuration. Audio alarms are triggered by:

• The onset of an alarm condition, provided the alarm delay is not active.
• A configured alarm delay, if alarm delay is active.

Audio alarm options are configured in the System Alarms (System Alarm Options) tab. By default, audio alarms are enabled.

NOTE: Audio alarms will be triggered at the ViewStation and the WorkStation when there is an alarm occurring at the host central station.
Once the alarm condition is resolved and/or acknowledged, the audible alarm stops, unless the alarm is latched. For additional information regarding alarm limits, refer to “Alarm Limit Settings” on page 13-2. For additional information regarding alarm latching, refer to “Latching” on page 3-3.

3.2.2 Alarm Options
The alarm options associated with a parameter dictate how the system responds when an alarm occurs.

Bedside Alarm Tracking
Bedside Alarm Tracking provides an automatic bi-directional flow of alarm limit change and alarm delay information, between a patient bedside monitor and a Panorama Central Station.

Latching
Alarm latching denotes that the audio and visual indicators, associated with an alarm condition, will not automatically stop even when the patient alarm condition is resolved.

By factory settings, all alarm events and nurse call events are automatically latched. All, none, or lethal arrhythmia alarm only can be configured to be latched. Configuration of latched alarms is done in the System Alarms (System Alarm Options) tab.

The patient tile Alarm ACK button acknowledges the latched alarm, and temporarily silences the alarm condition for the configured time period. However, if that latched alarm is not acknowledged during the configured time period, the alarm indicators resume. Once a patient alarm condition is resolved, it must be acknowledged at the central station and at the bedside monitor. If it is not acknowledged, all audio and visual alarm indicators continue at the central station and at the bedside monitor, even if the alarm condition is resolved.

NOTE: For Passport V and DPM 6/DPM 7, Passport 12M/Passport 17M, Passport 8/Passport 12, T1 monitor, the alarm condition must be acknowledged first at the bedside, and then acknowledged at the Panorama Central Station.

To acknowledge a latched alarm condition at the central station, once the condition is resolved, perform one of the following to the affected patient tile:

- Select the digital data area
- Select the waveform area
- Select the Alarm ACK button

NOTE: Acknowledging a latched alarm at the host central station patient tile acknowledges the alarm at the associated ViewStation and WorkStation patient tile.

NOTE: Acknowledging a latched alarm at the bedside monitor does not acknowledge the alarm at the host central station, ViewStation, or WorkStation patient tile.
To acknowledge a latched alarm at the bedside monitor, refer to its respective Operating Instructions.

Once a latched alarm condition is acknowledged, the alarm indicators discontinue until another alarm condition occurs. Acknowledging the alarm at the central station will not discontinue the alarm at the bedside monitor, and acknowledging the alarm at the bedside monitor will not discontinue the alarm at the central station, ViewStation, or WorkStation.

**Apnea Latching**
The conditions for apnea latching follow standard latching procedures, however apnea latching is associated with an apnea alarm condition. This option can be enabled/disabled in the System Alarms (System Alarm Options) tab. By default, Apnea Latching is enabled.

**NOTE:** Respiratory apnea alarm condition events are detected by the bedside monitor.

**NOTE:** Acknowledging an apnea latched alarm at the central station patient tile also acknowledges the alarm at the associated ViewStation and WorkStation patient tile.

**NOTE:** Acknowledging an apnea latched alarm at the bedside monitor does not acknowledge the alarm at the host central station, ViewStation, or WorkStation patient tile.

**Alarm Delay**
Alarm Delay is the configurable time period required before an alarm is triggered. Alarm Delay will not effect the Apnea Alarm, ST Alarms, the non-lethal arrhythmia alarms, or the lethal arrhythmia alarms. When permitted, this option can be enabled/disabled in the System Alarms (System Alarm Options) tab. By default, an Alarm Delay time is not specified.

**ST Alarm Delay**
ST alarm delay postpones the trigger of the ST alarm for a configured time period. When permitted, this option can be configured in the System Alarms (System Alarm Options) tab.

### 3.2.3 Physiological Alarm Responses
The alarm responses associated with a physiological alarm are as follows.

### 3.2.3.1 Alarm Priorities
The Panorama Central Station supports three alarm priorities. When alarm priority is turned on, the choices available are Alarm Priority of 1, 2 or 3. Alarms may be turned off by using the Priority Off setting. Alarm priorities associated with lethal alarm parameters cannot be changed from their default value of Priority 1. However, all other parameters can be modified. The alarm priority used to configure a parameter defines the alarm severity.

When a parameter is assigned a Priority Off, there are no visual or audio alarm indicators associated with the alarm event, however **Print on Alarm** and **Save To Event** options are still available.
Alarm priorities are configured in the patient **Alarm Responses** and the **System Alarms** (System Alarm Responses) tabs.

For additional information about defining a parameters alarm priority, refer to “Configuring Alarm Responses” on page 6-14 and “Setting the System Alarms Responses” on page 9-27.

**Priority 1 Alarms**
Priority 1 alarms are the most severe alarm type. Lethal-arrhythmia alarms (Asystole, V-Tach, and V-Fib), Extreme Bradycardia and Extreme Tachycardia are automatically configured as Priority 1 alarms and cannot be modified. Priority 1 alarm conditions cause the applicable parameter sub-tile to flash in red, display a red text message in the Patient Status line, and use a Priority 1 alarm sound. A parameters’ alarm priority is configured in the **System Alarms** (System Alarm Responses) tab.

**Priority 2 Alarms**
Priority 2 alarms are less severe than Priority 1 alarms, and have different visual and audio indicators. Heart rate (HR), ST Single, ST Dual, SpO2 and QTc (only available with the Telepack and V 12/21 device) default to Priority 2 alarms. Priority 2 alarm conditions cause the applicable parameter sub-tile to flash in yellow (for a physiological alarm), display a yellow text message in the Patient Status line, and use a Priority 2 alarm sound (if alarm delay is not enabled). A parameters’ alarm priority is configured in the **System Alarms** (System Alarm Responses) tab.

**Priority 3 Alarms**
Priority 3 alarms are the least severe of the three alarm priorities. By default, temperature (T1/T2/T3/Delta T/T Blood), respiration, , IBP, NIBP, CCO/CCO STAT, CCI/CCI STAT, SvO2, SVR/SVRI, CO2, Apnea Delay, PVC, and BIS are Priority 3 alarms. Priority 3 alarm conditions cause the applicable parameter sub-tile to display in constant yellow, display a yellow text message in the Patient Status line, and use a Priority 3 alarm sound (if alarm delay is not enabled). A parameters’ alarm priority is configured in the **System Alarms** (System Alarm Responses) tab.

**Priority Off**
Alarm priority can be turned off for certain alarms. Parameters configured to use Priority Off do not have any visual or audio alarm indicators, however, the **Print on Alarm** and **Save To Event** alarm response options still function.

**3.2.3.2 Save To Event**
**Save To Event** enables the Panorama Central Station to save a 20 second snapshot of a specific alarm event so that it can later be retrieved and studied in more detail. The Save To Event option is disabled at the ViewStation.
3.2.3.3 Print on Alarm

Print on Alarm automatically generates a patient All Strips Report when a specified alarm is triggered. The patient tile Print on Alarm option is enabled at the ViewStation and functions in the same manner as the host central station. Changes made to the patient tile Print on Alarm option, at the ViewStation, will not affect the patient tile Print on Alarm option at the host central station. For additional information about the All Strips Report, refer to the “All Strips Report” on page 8-2.

3.3 Technical Alarms

Technical alarms are patient specific and are related to technical issues that occur with cables, leads, and parameter sensors. The ViewStation technical event alarm notification will follow those settings at the host central station.

3.3.1 Alarm Behaviors

The visual and audio indicators associated with a technical alarm are as follows:

Visual Indicators
Technical visual indicators display as white text messages in the Patient Status line.

Audio Indicators
Technical alarm audio indicators are based on Technical Events Sound settings in the System Alarms (System Alarm Options) tab, the event type and the Volume tab settings.

3.3.2 Alarm Options

There is one alarm option available for a technical alarm.

Technical Event Sound
The Technical Events Sound option enables/disables audio notification when a technical alarm is triggered. Some events, for example, a Communications Lost event, send an audio alarm notification even when the Technical Events Sound option is disabled.

3.3.3 Alarm Responses

Alarm responses are not available for technical alarms, with the exception of the Nurse Call technical event.

3.4 System Alarms

System alarm events are associated with system-wide issues at the Panorama Central Station.

3.4.1 Alarm Behaviors

The visual and audio indicators associated with a system alarm are as follows:

Visual Indicators
System alarm visual indicators display as white text messages in the System Status line.
Audio Indicators
System audio alarm indicators are based on the settings in the Volume tab.
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This section describes lead placement, arrhythmia, and ST analysis.

- Lead Placement Procedures
- Arrhythmia Algorithm
- ST Segment Analysis
- Arrhythmia Alarms
- Troubleshooting
4.1 Lead Placement Procedures

Site preparation and ECG electrode placement directly impact the quality of an ECG signal. Optimizing an ECG signal is imperative for accurate monitoring.

The processes involved in successful lead placement include:

- Skin Preparation
- Electrode Patches
- Lead Placement Methods: AHA and IEC

4.1.1 Skin Preparation

Proper skin preparation is essential to obtain accurate ECG data. Electrode sites should be clean, dry and should provide a smooth flat surface. Incidental electrical activity and inaccurate readings may occur due to incorrect skin preparation.

The following skin preparation is recommended for secure patch application:

1. Shave the chest hair in a 2-4 inch diameter of the electrode site.
2. Use a dry gauze pad to remove excess skin oils, skin cells and residue from the electrode sites. Never rub the skin until it is raw or bleeding.

**NOTE:** Prepare the electrode site with alcohol only if the skin is extremely greasy. If alcohol is used as a drying agent, always allow the skin to dry before placing the electrode patch on the skin.

4.1.2 Electrode Patches

**NOTE:** Store electrode patches at room temperature and open just prior to use.

**NOTE:** Avoid more than one type of electrode on a patient because of variations in electrical resistance.

**NOTE:** Avoid placing electrode patches directly over boney prominences or over any areas that move during activity such as shoulders or arms because muscle motion produces electrical activity. If an electrode patch is placed over a large muscle such as the pectorals, the monitor may detect this additional muscle activity, which could lead to false arrhythmia calls.

1. Peel the backing off of the electrode patch only when it is ready for use to prevent evaporation of the contact gel medium. Visually inspect the contact gel medium for moistness. If the gel medium is not moist, do not use the electrode patch. Dry electrode patches are not conductive.

**NOTE:** If using the snap type electrode wires, attach the electrode patch to the lead wire before placing patch on the patient.
2. Attach the electrode patch to the skin at the prepared site. Smooth the electrode patch down in a circular motion to ensure proper skin contact. If using soft gel electrodes, never push down directly over the contact gel medium as this may displace the gel and cause monitoring artifact. If using hard gel electrodes, it is recommended that during application, the center of the electrode should be slightly pressed onto the skin to ensure direct contact. Consult the electrode patch manufacturer’s instructions for specific use.

3. Secure the lead wires to the patient according to hospital practice. For additional information, refer to “Lead Placement Methods: AHA and IEC” on page 4-3. If using a Panorama Telepack, always secure the device to the patient according to hospital standard.

WARNING: As with all medical equipment, carefully route cables and connections to reduce the possibility of entanglement or strangulation.

NOTE: It is recommended that electrode patches be changed at least every 24-36 hours to maintain proper contact with the skin. Some patients may require electrodes to be changed more often. Electrode patches are disposable and should not be reused or reapplied. Try to avoid reusing the exact same electrode site during reapplication. If an electrode becomes wet with fluid, change the electrode patch.

4.1.3 Lead Placement Methods: AHA and IEC

The lead placement procedure that is utilized has a direct impact on the quality of an ECG waveform. The algorithm works best when a patient’s R wave is significantly larger than the P or T waves to avoid difficulty in identifying the appropriate waves. On some patients, electrode patch placement and/or the ECG lead viewed may need to be adjusted to obtain a more prominent R wave.

The following section outlines lead placement procedures for the American Heart Association (AHA) and the International Electro-Technical Commission (IEC).
3-wire Lead Set
A 3-wire lead set can monitor one of three ECG vectors (I, II, or III). The recommended 3-wire ECG lead placement is as follows.

- Place RA (white) electrode under right clavicle, mid-clavicular line within the rib cage frame.
- Place LA (black) electrode under left clavicle, mid-clavicular line within the rib cage frame.
- Place LL (red) electrode on the lower left abdomen within the rib cage frame.

5-wire Lead Set
A 5-wire lead set can monitor seven ECG vectors (I, II, III, aVR, aVL, aVF, and V) simultaneously. The recommended 5-wire ECG lead placement is as follows.

- Place RA (white) electrode under right clavicle, mid-clavicular line within the rib cage frame.
- Place LA (black) electrode under left clavicle, mid-clavicular line within the rib cage frame.
- Place LL (red) electrode on the lower left abdomen within the rib cage frame.
- Place RL (green) electrode on lower right abdomen within the rib cage frame.
- Place V (brown) chest lead in the proper positioning for desired lead V1-V6.
- Place N (black) electrode on lower right abdomen within the rib cage frame.
- Place C (white) chest lead in the proper positioning for the desired lead C1-C6.
6-wire Lead Set

The recommended 6-wire lead placement is as follows.

**FIGURE 4-5 6-wire Lead Placement (AHA)**
- Place the RA (white) electrode under the patient’s right clavicle, at the mid-clavicular line within the rib cage frame.
- Place the LA (black) electrode under the patient’s left clavicle, at the mid-clavicular line within the rib cage frame.
- Place the LL (red) electrode on the patient’s lower left abdomen within the rib cage frame.
- Place the RL (green) electrode on the patient’s lower right abdomen within the rib cage frame.
- Place the Vx (brown) electrode in one of the V-lead positions (V1 to V6) depicted in the following section.
- Place the Vy (yellow) electrode in one of the V-lead positions (V1 to V6) depicted in the following section.

**FIGURE 4-6 6-wire Lead Placement (IEC)**
- Place the R (red) electrode under the patient’s right clavicle, at the mid-clavicular line within the rib cage frame.
- Place the L (yellow) electrode under the patient’s left clavicle, at the mid-clavicular line within the rib cage frame.
- Place the F (green) electrode on the patient’s lower left abdomen within the rib cage frame.
- Place the N (black) electrode on the patient’s lower right abdomen within the rib cage frame.
- Place the Cx (white) electrode in one of the V-lead positions (V1 to V6) depicted in the following section.
- Place the Cy (white) electrode in one of the C-lead (C1 to C6) positions depicted in the following section.
View 12™ Card and View 12 Module
A View 12™ card or a View 12 Module utilizes a 10-wire ECG lead set that can monitor 12 ECG vectors (I, II, III, aVR, aVL, aVF, V1, V2, V3, V4, V5, and V6) simultaneously. The recommended lead placement for a View 12™ card is as follows.

- Place RA (white) electrode under the right clavicle, mid-clavicular line within the rib cage frame.
- Place LA (black) electrode under the left clavicle, mid-clavicular line within the rib cage frame.
- Place LL (red) electrode on the lower left abdomen within the rib cage frame.
- Place RL (green) electrode on lower right abdomen within the rib cage frame.
- Place V1 (brown) chest lead in the fourth intercostal space, right sternal border.
- Place V2 (brown) chest lead in the fourth intercostal space, left sternal border.
- Place V3 (brown) chest lead midway between V2 and V4 on a straight line.
- Place V4 (brown) chest lead in the fifth intercostal space, mid-clavicular line.
- Place V5 (brown) chest lead in the fifth intercostal space, anterior axillary line.
- Place V6 (brown) chest lead in the fifth intercostal space, mid-axillary line.

- Place R (red) electrode under the right clavicle, mid-clavicular line within the rib cage frame.
- Place L (yellow) electrode under the left clavicle, mid-clavicular line within the rib cage frame.
- Place F (green) electrode on the lower left abdomen within the rib cage frame.
- Place N (black) electrode on lower right abdomen within the rib cage frame.
- Place C1 (white) chest lead in the fourth intercostal space, right sternal border.
- Place C2 (white) chest lead in the fourth intercostal space, left sternal border.
- Place C3 (white) chest lead midway between C2 and C4 on a straight line.
- Place C4 (white) chest lead in the fifth intercostal space, mid-clavicular line.
- Place C5 (white) chest lead in the fifth intercostal space, anterior axillary line.
- Place C6 (white) chest lead in the fifth intercostal space, mid-axillary line.
Lead II Monitoring

The recommended lead placement for Lead II monitoring is as follows.

- Place RA (white) electrode under right clavicle, mid-clavicular line within the rib cage frame.
- Place LA (black) electrode under left clavicle, mid-clavicular line within the rib cage frame.
- Place LL (red) electrode on the lower left abdomen within the rib cage frame.

Select the ECG Lead II at the bedside monitor. Lead II is the direct electrical line between the RA (white) electrode and the LL (red) electrode.

- Place R (red) electrode under right clavicle, mid-clavicular line within the rib cage frame.
- Place L (yellow) electrode under left clavicle, mid-clavicular line within the rib cage frame.
- Place F (green) electrode on the lower left abdomen within the rib cage frame.

Select the ECG Lead II at the bedside monitor. Lead II is the direct electrical line between the R (red) electrode and the F (green) electrode.
Modified Chest Lead (MCL) Monitoring
The recommended lead placement for MCL monitoring is as follows.

- Place RA (white) electrode under left clavicle, mid-clavicular line within the rib cage frame.
- Place LA (black) electrode on the right sternal border, fourth intercostal in the rib cage frame.
- Place LL (red) electrode on the lower left abdomen within the rib cage frame.
- Place R (red) electrode under left clavicle, mid-clavicular line within the rib cage frame.
- Place L (yellow) electrode on the right sternal border, fourth intercostal in the rib cage frame.
- Place F (green) electrode on the lower left abdomen within the rib cage frame.

Select ECG Lead I for MCL 1 monitoring.
Lead I is the direct electrical line between the RA (white) electrode and the LA (black) electrode.

Select ECG Lead II for MCL 6 monitoring.
Lead II is the direct electrical line between the RA (white) electrode and the LL (red) electrode.

Select ECG Lead I for MCL 1 monitoring.
Lead I is the direct electrical line between the R (red) electrode and the L (yellow) electrode.

Select ECG Lead II for MCL 6 monitoring.
Lead II is the direct electrical line between the L (red) electrode and the F (green) electrode.
Neonatal Electrode Placement
When using a 3-wire lead set, ECG lead placement on a neonate is usually directed towards obtaining the best possible respiration data through the ECG thoracic impedance technique. Thoracic impedance is usually measured between the Right Arm and Left Arm electrode patches. These patches should be placed on the chest directly across from each other to optimize the measurement of the neonate’s chest movement.

The recommended lead placement for neonate monitoring is as follows.

- Place RA (white) electrode under patient left clavicle, mid-clavicular line within the rib cage frame.
- Place LA (black) electrode right sternal border, fourth intercostal space within the rib cage frame.
- Place LL (red) electrode on the patient’s lower left abdomen within the rib cage frame.

Monitoring in the Presence of a Transcutaneous Electrical Nerve Stimulator (TENS)
Since a TENS unit transmits electrical impulses, avoid placing ECG electrode patches near the TENS electrodes. ECG electrode patches may need to be repositioned and the ECG lead viewed may need to be adjusted until the optimum ECG tracing is obtained.
Monitoring a Pacemaker Patient

The recommended lead placement for a pacemaker patient is as follows.

A Pacemaker patient usually requires a different electrode patch placement configuration than a non-pacemaker patient.

Do not place an ECG electrode directly over the pacemaker generator. Place the electrode patches 3-5 inches away from the pacemaker generator area to avoid electrical interference. If the electrode patches are placed closer to pacemaker generator, the ECG will contain artifact, sometimes called “picket fence syndrome.” For example, if the pacemaker generator is located in the right subclavian area, relocate the Right Arm (white) electrode closer in towards the center of the chest.
WARNING: PACEMAKER PATIENTS - Indication of the heart-rate may be adversely affected by cardiac pacemaker pulses or by cardiac arrhythmias. Keep pacemaker patients under close surveillance. Refer to “ECG Performance Requirements” on page 16-2 for disclosure of the pacer rejection capability.

CAUTION: Some pacemakers may contain a respiratory sensor that may produce artifact on an ECG waveform.

4.2 Arrhythmia Algorithm

The Panorama Central Station uses an arrhythmia algorithm to monitor ECG waveform data. The arrhythmia analysis feature is intended to detect ventricular rhythms only. However, due to physiologic differences in patient populations, the arrhythmia algorithm may occasionally result in a false alarm or may not recognize certain arrhythmia patterns.

The algorithm creates ECG waveform templates based on a patient’s normal ECG data and uses them to analyze newly received data. The algorithm verifies that data is free from noise and artifact, and that it does not deviate from the patient’s normal ECG rhythms.

A normal ECG waveform typically includes consistent spacing between R waves, a sharp and well defined QRS complex, and an ECG baseline that is free of noise and artifact. The computerized arrhythmia algorithm works best when the patient’s R wave is significantly larger than the P wave and the T wave. If the R wave is not significantly larger than other lower voltage waves on the ECG tracing, the computer may have some difficulty in identifying the appropriate waves. On some patients, electrode patch placement and/or the viewed ECG lead may need to be adjusted in order to obtain a significant R wave.

![Sample Waveform](image)

FIGURE 4-19 Sample Waveform

Noise and Artifact

The presence of noise or artifact in an ECG waveform makes the accurate detection and classification of heart beats difficult. To best optimize performance, all leads should be free of noise. If artifact is present on any ECG lead/waveform, then the arrhythmia processing, alarm processing, and quality of the monitoring function may be adversely affected.
Some of the causes of ECG noise include poor skin preparation, improperly attached electrodes, dried electrode gel, defective lead wires, and patient movement. The algorithm uses several techniques to differentiate a patient’s QRS complexes from noise sources.

If noise levels are too high, the following will occur until the signal quality is re-established:

- Beat detection is suspended
- All rhythm calls are suspended
- An ECG Noise message is displayed in the Patient Status line at the Panorama Central Station when the noise continues beyond the configured noise delay

**Heart Rate Meter**

The average heart rate is calculated on the basis of the mean RR-interval of the last 16 beats, unless the heart rate calculated using the last 4 beats is less than or equal to 48, then this rate is used. After the start-up phase, the heart rate value is made available when 4 RR intervals are detected. All detected beats are used to compute the heart rate. A separate ventricular rate is used in the algorithm to determine rhythms like ventricular tachycardia and ventricular run.

**Filtering Pacer Signals**

In order to prevent pacer pulses from being mistaken for QRS complexes, they are removed from the ECG data sent to the arrhythmia algorithm for analysis. Pacer pulses are shown on the Panorama Central Station and bedside monitors as a long vertical line crossing through the waveform or a short vertical line above the waveform (depending on the selection in the Pacer Settings dialog) to indicate firing of the pacemaker impulse.

**ECG Amplitude**

The QRS detection threshold algorithm setting is adjustable from 0.16mV to 0.48 mV and defaults to 0.16mV. Changing the display gain (or wave gain) on the monitor does not affect the signal that is used by the algorithm for beat detection. For optimal performance, the leads selected for monitoring should have an amplitude of 0.5 to 1 mV or more.

**Learning**

A learning process is used to establish a normal beat template for a patient. The learn period is dependent on heart rate and dominant pattern. Learning should not be initiated during a primarily ventricular rhythm because an ectopic beat may be established as normal.

A learn should be initiated when beats are not being properly detected, or when they are being erroneously classified. However, if a signal is not strong enough, or lead data is extremely noisy, better signal quality must be established before a learn can be effective.
Beat Detection and Typing

The following tables describe the leads that are used for beat detection and beat typing.

**TABLE 4-1** Spectrum, Passport 2 and Passport V

<table>
<thead>
<tr>
<th>DESCRIPTION</th>
<th>3-WIRE LEAD SET</th>
<th>5-WIRE LEAD SET</th>
<th>VIEW 12™ CARD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Leads used for Beat Detection</td>
<td>Determined by viewed lead</td>
<td>II and V</td>
<td>V1 and V5</td>
</tr>
<tr>
<td>Leads used for Beat Typing</td>
<td>Determined by viewed lead</td>
<td>II, V, and I</td>
<td>V1, V5, and II</td>
</tr>
<tr>
<td>Leads used for V-Fib Detection</td>
<td>Determined by viewed lead</td>
<td>II and V</td>
<td>V1 and V5</td>
</tr>
</tbody>
</table>

**TABLE 4-2** V 12/V 21

<table>
<thead>
<tr>
<th>DESCRIPTION</th>
<th>3-WIRE LEAD SET</th>
<th>5-WIRE LEAD SET*</th>
<th>VIEW 12 MODULE*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Leads used for Beat Detection</td>
<td>Determined by viewed lead</td>
<td>II and V</td>
<td>II and V1</td>
</tr>
<tr>
<td>Leads used for Beat Typing</td>
<td>Determined by viewed lead</td>
<td>II, V, and I</td>
<td>II, V1, and I</td>
</tr>
<tr>
<td>Leads used for V-Fib Detection</td>
<td>Determined by viewed lead</td>
<td>II and V</td>
<td>II and V1</td>
</tr>
</tbody>
</table>

* Indicates that these leads are user configurable on the bedside.

**TABLE 4-3** Telepack

<table>
<thead>
<tr>
<th>DESCRIPTION</th>
<th>3-WIRE LEAD SET</th>
<th>5-WIRE LEAD SET*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Leads used for Beat Detection</td>
<td>Determined by viewed lead</td>
<td>II and V</td>
</tr>
<tr>
<td>Leads used for Beat Typing</td>
<td>Determined by viewed lead</td>
<td>II, V, and I</td>
</tr>
<tr>
<td>Leads used for V-Fib Detection</td>
<td>Determined by viewed lead</td>
<td>II and V</td>
</tr>
</tbody>
</table>

* Indicates that these leads are user configurable on the Panorama.

**NOTE:** See “ECG Analysis Leads (Telepack)” on page 5-17 for configuring the beat detection and typing leads of Telepack.

**TABLE 4-4** DPM 6/DPM 7, Passport 8/Passport 12, Passport 12M/Passport 17M and T1 monitor (Mortara Algorithm)

<table>
<thead>
<tr>
<th>DESCRIPTION</th>
<th>3-WIRE LEAD SET</th>
<th>5-WIRE LEAD SET</th>
<th>10-WIRE LEAD SET</th>
</tr>
</thead>
<tbody>
<tr>
<td>Leads used for Beat Detection</td>
<td>Determined by viewed lead</td>
<td>I and II</td>
<td>I and II</td>
</tr>
<tr>
<td>Leads used for Beat Typing</td>
<td>Determined by viewed lead</td>
<td>II, V, and I</td>
<td>II, V1, and I</td>
</tr>
<tr>
<td>Leads used for V-Fib Detection</td>
<td>Determined by viewed lead</td>
<td>I and II</td>
<td>I and II</td>
</tr>
</tbody>
</table>

**TABLE 4-5** DPM 6/DPM 7, Passport 8/Passport 12, Passport 12M/Passport 17M and T1 monitor (Mindray Algorithm)
When a beat is detected, the search for the next beat begins after a refractory period to avoid detecting T-waves as valid QRS complexes. The minimum QRS amplitude that can be detected is between 0.15 mV and 0.16 mV depending on the width of the QRS complex.

In the beat typing process, each new ECG beat is compared to reference templates that were previously stored in the system. A beat typing algorithm classifies the beats as Normal, Ventricular, Paced, etc.

- If an incoming beat matches a template that has already been classified, it is given the same label as the template. The template parameters are updated with the features from this new beat.

The real time ECG analysis library incorporates ventricular ectopic beat detection as a part of arrhythmia analysis.

- Parameters like compensatory pause, QRS width, QRS positive and negative areas, and R wave positive and negative amplitudes are used to differentiate ventricular ectopic beats from normals. This process uses multiple leads when available.

### 4.3 ST Segment Analysis

The ST segment of an ECG waveform (shown in FIGURE 4-19) represents the period from the end of ventricular de-polarization, to the beginning of ventricular re-polarization, or the end of the QRS complex (the J point) and the beginning of the T-wave. ST Segment analysis is used to monitor the adequacy of oxygen supply to and the viability of the heart muscle.

ST deviation is the vertical distance between the signal levels at the isoelectric (ISO) point and ST point.

The ISO point is located along the isoelectric segment between the end of the P wave and the start of the Q wave. The ISO point provides the baseline for this measurement.

The ST point is at a fixed distance from the J point which is the end of the QRS complex. The ST point can be configured to 40, 60, or 80 milliseconds past the J-point, independent of the heart rate. By default, the ST point is positioned as follows:

- at 80 milliseconds past the J point for heart rates less than or equal to 120 beats per minute
• at 60 milliseconds past the J point for heart rates greater than 120 beats per minute
  ST deviations are displayed in mm where 1 mm represents 0.1 mV. The reliability of ST
  measurements is lowered in the presence of atrial fibrillation, flutter, and erratic baseline
  changes.

Since ST deviation measurements are dependent on the correct locations of the ISO and ST
  points, selecting leads that contain the least amount of baseline flutter and artifact will
  improve measurement accuracy, but accurate ST deviation measurement is dependent on the
  correct location of the ISO and ST points.

ST segment deviations are measured on all available ECG leads.

The displayed ST template data is updated approximately every 10 seconds.

The ST algorithm has been tested for accuracy of the ST segment data. The significance of
  the ST segment changes need to be determined by a clinician.

Learning
The process of learning is used to establish the dominant beat morphology and templates as
  well as or a stable baseline for accurate ST analysis. To determine the templates for the
  dominant rhythm, the system evaluates the ECG signals from leads II, V, and I. It takes about
  16 beats to complete arrhythmia learn and 33 beats to complete the learning process for ST
  segment analysis. To establish an accurate baseline, it is recommended that learning be
  initiated when the patient is in stable condition, not moving, and has an ECG rhythm that is
  free of artifact. Learning should not be initiated during a primarily ventricular rhythm or other
  ECG rhythm irregularity because an ectopic beat may be established as normal.

Learning is automatically initiated by the system when:

• Arrhythmia analysis is turned on
• ST analysis is turned on
• Patient monitoring resumes and patient is removed from the Standby mode
• A 3-wire lead set is in use and the current ECG lead is changed
• Communication is reestablished after a Communications Lost event
• The leads have been off for 60 seconds or more before being reattached to the patient

ECG Filters
The ST segment of an ECG waveform contains low amplitude signals with low frequency
  content. To preserve low frequency signal content, the high pass filter is set to 0.05 Hz when
  ST analysis is turned on.

4.4 Arrhythmia Alarms
Arrhythmia alarms are activated based on the patterns in the patient ECG waveform rhythms.
When using a 5-lead wire set, signals from leads II and V are used for beat detection.
Information from these two leads, as well as from lead I is used to determine beat
  morphology (normal, ventricular ectopic etc). When using a 3-lead wire set, beat detection
  as well as beat morphology are based on information from the viewed lead.
The following lethal and non-lethal arrhythmia alarms are detected by the arrhythmia algorithm. For additional information regarding alarm behavior at the Panorama Central Station, refer to “Alarm Behavior” on page 3-1.

NOTE: Cardiac arrhythmia alarm condition events are detected and processed by the bedside monitor. When using a Telepack device, the detection and processing of the event is done by the Panorama Central Station.

NOTE: Arrhythmia alarms are not available for the Neonate patient size.

4.4.1 Lethal Arrhythmia Alarms

A lethal arrhythmia is an arrhythmia that can be life threatening to a patient if left untreated. Ventricular Tachycardia (V-Tach), Ventricular Fibrillation (V-Fib), and Asystole alarms are classified as lethal arrhythmia alarms. These alarms automatically default to priority 1.

NOTE: Lethal arrhythmia alarms may be configured to have a different alarm sound than other physiological alarms. The sounds used are determined during system setup and configuration.

NOTE: Lethal arrhythmia alarms can be set to latched alarms and if set to latched must be acknowledged. To acknowledge the alarm, select the digital data area or the waveform area of the affected patient tile.

NOTE: When a V-Tach or V-Fib alarm occurs, but its alarm delay has not expired, and is replaced by an Asystole alarm, the Asystole alarm will be paged. This will occur if Paging has been installed on the central station; the Asystole alarm paging delay timer has expired; and the alarm condition still exists when the paging delay timer has expired.

Asystole Alarm

An Asystole alarm is activated at the Panorama Central Station when no QRS complexes are detected for a period greater than the asystole delay period selected by the user. When using a View 12™ card, the asystole delay can be set to be between three (3) and eight (8) seconds. When using a 3 or 5-wire lead set, the asystole delay can be set to be is between three (3) and ten (10) seconds.

The Asystole alarm is a priority 1 alarm event that:

- Produces a priority 1 visual and audio alarm indication once the condition is identified.
- Causes a red Asystole text message to display in the Patient Status line.
- Causes the Heart Rate tile to flash red

Ventricular-Fibrillation (V-Fib) Alarm

A V-Fib alarm is activated at the Panorama Central Station when a fibrillatory waveform (P, QRS or T waves can no longer be identified) is detected. V-Fib is defined as “irregular, disorganized electrical activity of the heart”. The ECG signal for V-Fib continuously changes shape, interval, and amplitude. The V-Fib detection algorithm runs in parallel to the beat detection algorithm and continuously examines the incoming data.
The V-Fib alarm is a priority 1 alarm event that:

- Produces a priority 1 visual and audio alarm indication once the condition is identified.
- Causes a red **V-Fib** text message to display in the Patient Status line.
- Causes the Heart Rate tile to flash red.

**Ventricular Tachycardia (V-Tach) Alarm**

A **V-Tach** alarm is activated at the Panorama Central Station when the configured number of consecutive PVC's at a rate equal to or greater than the V-Tach threshold is detected.

The range for V-Tach rate is 100 to 180 beats/min. and the range for V-Tach threshold is 3-15 beats (as shown in FIGURE 4-20):

A V-Tach alarm is a priority 1 alarm event that:

- Produces the priority 1 visual and audio alarm indication once the condition is identified.
- Causes a red **V-Tach** text message to display in the Patient Status line.
- Causes the Heart Rate tile to flash red.

### 4.4.2 Non-Lethal Arrhythmia Alarms

A Non-Lethal Arrhythmia is an arrhythmia that is most likely not life threatening to a patient. Bigeminy, Bradycardia, Couple, Irregular Heart Rate, Multifocal PVC, Pause, PVC, Run, Trigeminy, R-On-T PVC, and Ventricular Rhythm (V-Rhythm) alarms are non-lethal arrhythmia alarms, and depending on the alarm, default to priority 2 or 3 (except for Bradycardia).

#### Bigeminy Alarm

The **Bigeminy** alarm is activated at the Panorama Central Station when three or more consecutive cycles of one PVC coupled to one normal beat are detected.

The Bigeminy alarm has priority settings of 2, 3, and OFF. When the setting is:

- Set to **2**, the Panorama Central Station produces a priority 2 sound once the condition is identified.
- Set to **3**, the Panorama Central Station produces a priority 3 sound once the condition is identified.
- Set to **OFF**, the Panorama Central Station does not produce any audio alarm indications for the Bigeminy alarm condition.
- The Bigeminy alarm causes a yellow **Bigeminy** text message to display in the Patient Status line.

#### Bradycardia (Brady) Alarm

The **Brady** alarm is activated at the Panorama Central Station when the heart rate falls to a value 10% lower than the user selected value for low heart rate alarm.

**NOTE:**  The Bradycardia alarm is available when using Passport 2, Spectrum, Spectrum OR and Passport V monitor.

**NOTE:**  The Brady alarm will not be suspended when the All Non-High Alarms Pause sidebar button is selected.
NOTE: Brady alarms may be configured to have a different alarm sound than other non-lethal physiological alarms. The sounds used are determined during system setup and configuration.

The Brady alarm has priority settings of 1 and OFF. When the setting is:

- Set to 1, the Panorama Central Station produces a priority 1 sound once the condition is identified.
- Set to OFF, the Panorama Central Station does not produce any audio alarm indications for the Bradycardia alarm condition.
- The Bradycardia alarm causes a Brady text message to display in the Patient Status line. The color of the message is dependant on the alarm priority configured.

Couplet Alarm

The Couplet alarm is activated at the Panorama Central Station when two consecutive PVCs are detected between normal beats.

The Couplet alarm has priority settings of 2, 3, and OFF. When the setting is:

- Set to 2, the Panorama Central Station produces a priority 2 sound once the condition is identified.
- Set to 3, the Panorama Central Station produces a priority 3 sound once the condition is identified.
- Set to OFF, the Panorama Central Station does not produce any audio alarm indications for the Couplet alarm condition.
- The Couplet alarm causes a yellow Couplet text message to display in the Patient Status line.

Irregular Heart Rate Alarm

The Irregular Heart Rate alarm is activated at the Panorama Central Station when the measured variation in the R-R interval over a period of time exceeds a preset limit established by the arrhythmia algorithm.

NOTE: The Irregular Heart Rate alarm is not available when using a View 12™ card.

The Irregular Heart Rate alarm has priority settings of 2, 3, and OFF. When the setting is:

- Set to 2, the Panorama Central Station produces a priority 2 sound once the condition is identified.
- Set to 3, the Panorama Central Station produces a priority 3 sound once the condition is identified.
- Set to OFF, the Panorama Central Station does not produce any audio alarm indications for the Irregular Heart Rate alarm condition.
- The Irregular Heart Rate alarm causes a yellow Irregular HR text message to display in the Patient Status line.

Multifocal PVC Alarm

The Multifocal PVC alarm is activated at the Panorama Central Station if during the last 15 beats, two or more premature ventricular beats with different morphologies are detected.
The Multifocal PVC alarm has priority settings of 2, 3, and OFF. When the setting is:

- Set to 2, the Panorama Central Station produces a priority 2 sound once the condition is identified.
- Set to 3, the Panorama Central Station produces a priority 3 sound once the condition is identified.
- Set to OFF, the Panorama Central Station does not produce any audio alarm indicators for the Multifocal PVC alarm condition.
- The Multifocal PVC alarm causes a yellow Multifocal PVC text message to display in the Patient Status line.

**Pause Alarm**

The Pause alarm is activated at the Panorama Central Station when no QRS complexes are detected during an interval that is greater than user defined times (1.2 to 2.0) the average R-R interval (user defined percent (120% to 200%) of the average R-R interval) and when the previous beat is not a PVC.

**NOTE:** The Pause alarm is only available when using View 12™ card, V 12/V 21, DPM 6/DPM 7, Passport 8/Passport 12, Passport 12M/Passport 17M, T1 device or Telepack.

The Pause alarm has priority settings of 1, 2, 3, and OFF. When the setting is:

- Set to 1, the Panorama Central Station produces a priority 1 sound once the condition is identified.
- Set to 2, the Panorama Central Station produces a priority 2 sound once the condition is identified.
- Set to 3, the Panorama Central Station produces a priority 3 sound once the condition is identified.
- Set to OFF, the Panorama Central Station does not produce any audio alarm indication for the Pause alarm condition.
- The Pause alarm causes a Pause text message to display in the Patient Status line. The color of the message is dependant on the configured alarm priority.

**PVC/minute Alarm**

The High PVC alarm is activated at the Panorama Central Station when the number of PVCs detected per minute exceeds the configured threshold. The PVC rate alarm can be set to Off, or 1 to 30 PVCs per minute.

**NOTE:** The PVC/min counter will be set to invalid and display (---) when Asystole, V-Tach, V-Fib, or V-Rhythm is detected when using a Spectrum, Passport 2 or a Telepack.

The High PVC alarm has priority settings of 2, 3, and OFF. When the setting is:

- Set to 2, the Panorama Central Station produces a priority 2 sound once the condition is identified.
- Set to 3, the Panorama Central Station produces a priority 3 sound once the condition is identified.
- Set to OFF, the Panorama Central Station does not produce any audio alarm indication for the High PVC alarm condition.
• The High PVC alarm causes a yellow **High PVC Alarm** text message to display in the Patient Status line.

**Run Alarm**

The **Run** alarm is activated on the Panorama Central Station when a user selected number of consecutive PVCs are detected and these occur at a rate that exceeds the user defined VTach rate (100 to 180). The number of consecutive PVCs that constitute a Run is one less than user defined threshold for VTach (3 to 15). In other words, a Run is a non sustained VTach. If the number of consecutive PVCs for VTach is set at 3 or more, a Run alarm will not be called. For additional information about VTach, refer to the definition for "Ventricular Tachycardia (VTach) Alarm" on page 4-17.

**FIGURE 4-20**

The Run alarm has priority settings of **2**, **3**, and **OFF**. When the setting is:

• **Set to 2**, the Panorama Central Station produces a priority 2 sound once the condition is identified.
• **Set to 3**, the Panorama Central Station produces a priority 3 sound once the condition is identified.
• **Set to OFF**, the Panorama Central Station does not produce any audio alarm indication for the Run alarm condition.
• The Run alarm causes a yellow **Run** text message to display in the Patient Status line.

**R-On-T PVC**

The **R-On-T PVC** alarm is activated at the Panorama Central Station when the following occurs:
- A beat is detected as a PVC and the previous beat is not a PVC
- And current BTB heart rate is > 120 bpm
- And current RR-interval is < previous R-R interval/2

The R-on-T PVC alarm has priority settings of 1, 2, and OFF. When the setting is:

- Set to 1, the Panorama Central Station produces a priority 1 sound once the condition is identified.
- Set to 2, the Panorama Central Station produces a priority 2 sound once the condition is identified.
- Set to OFF, the Panorama Central Station does not produce any audio alarm indicators for the R-On-T PVC alarm condition.

The R-On-T PVC alarm causes a yellow Run-On-PVC text message to display in the Patient Status line.

**Trigeminy Alarm**

The Trigeminy alarm is activated at the Panorama Central Station when three or more consecutive cycles of one PVC coupled to two normal beats are detected. This rhythm could also cause an Irregular HR alarm.

The Trigeminy alarm has priority settings of 2, 3 and OFF. When the setting is:

- Set to 2, the Panorama Central Station produces a priority 2 sound once the condition is identified.
- Set to 3, the Panorama Central Station produces a priority 3 sound once the condition is identified.
- Set to OFF, the Panorama Central Station does not produce any audio alarm indication for the Trigeminy alarm condition.

- The Trigeminy alarm causes a yellow Trigeminy text message to display in the Patient Status line.

**Ventricular Rhythm (V-Rhythm) Alarm**

The V-Rhythm alarm is activated at the Panorama Central Station when the number of consecutive PVCs is equal to or greater than three and the ventricular rate is less than the configured V-Tach threshold (as shown in FIGURE 4-20).

The V-Rhythm alarm has priority settings of 2, 3, and OFF. When the setting is:

- Set to 2, the Panorama Central Station produces a priority 2 sound once the condition is identified.
- Set to 3, the Panorama Central Station produces a priority 3 sound once the condition is identified.
- Set to OFF, the Panorama Central Station does not produce any audio alarm indication for the V-Rhythm alarm condition.

- The V-Rhythm alarm causes a yellow V-Rhythm text message to display in the Patient Status line.
Pauses/Minute Alarm

The **Pauses/Minute** alarm is activated at the Panorama Central Station when the number of pauses detected per minute exceeds the configured threshold. The Pauses/minute limit can be set to 1 to 15 Pauses per minute.

**NOTE:** The Pause/min alarm is only available when using a V 12/ V 21 and Telepack device.

The Pauses/Minute alarm has priority settings of 1, 2, 3, and OFF. When the setting is:

- Set to 1, the Panorama Central Station produces a priority 1 sound once the condition is identified.
- Set to 2, the Panorama Central Station produces a priority 2 sound once the condition is identified.
- Set to 3, the Panorama Central Station produces a priority 3 sound once the condition is identified.
- Set to OFF, the Panorama Central Station does not produce any audio alarm indication for the Pause alarm condition.
- The Pauses/Minute alarm causes a Pauses/Minute Alarm text message to display in the Patient Status line. The color of the message is dependant on the configured alarm priority.

QTc Alarm

The **High QTc** alarm is activated when the current QTc value is equal to or higher than the user selected value for the high QTc alarm limit.

**NOTE:** The High QTc Alarm is only available when using a V12/ V 21 or a Telepack device.

The QTc alarm has priority settings of 2 and 3. When the setting is:

- Set to 2, the Panorama Central Station produces a priority 2 sound once the condition is identified.
- Set to 3, the Panorama Central Station produces a priority 3 sound once the condition is identified.
- The High QTc alarm causes a High QTc Alarm text message to display in the Patient Status line.

4.5 Extreme Bradycardia/Extreme Tachycardia

The **Extreme Bradycardia** alarm is activated on the Panorama system when the configured alarm limit is reached or less. The Extreme Bradycardia alarm limit must be less than the Low Heart Rate alarm limit.

The **Extreme Tachycardia** alarm is activated on the Panorama system when the configured alarm limit is reached or greater. The Extreme Tachycardia alarm limit must be greater than the High Heart Rate alarm limit.

The Extreme Bradycardia or Extreme Tachycardia alarm is Priority 1 alarm event that:
• Produces a priority 1 visual and audio alarm indication once the condition is identified.
• Causes a red Extreme Bradycardia or Extreme Tachycardia text message to display in the Patient Status line.
• Causes the Heart Rate tile to flash red.

**NOTE:** The Extreme Bradycardia/Extreme Tachycardia alarm is available when using a DPM 6/DPM 7, Passport 12M/Passport 17M, Passport 8/Passport 12, T1 monitor, V 12/V 21 monitor, or Telepack device.
4.6 Troubleshooting

This section lists some of the potential messages and issues that may occur. For additional assistance troubleshooting arrhythmia analysis, please contact a Mindray Clinical Educator at 1-800-288-2121 (U.S and Canada only) or 201-995-8000 (outside U.S. and Canada).

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Noisy ECG traces</td>
<td>Loose or dry electrodes. Defective electrode wires. Patient cable or leads are routed too close to other electrical devices.</td>
<td>Apply fresh, moist electrodes. Replace wires as necessary. Eliminate 60 Hz interference.</td>
</tr>
<tr>
<td>Excessive Electro-surgical Interference</td>
<td>Wrong ECG cable used.</td>
<td>Use ECG cable with internal filter block. <strong>NOTE:</strong> Respiration monitoring via the ECG electrodes will not be available when using this cable.</td>
</tr>
<tr>
<td></td>
<td>Muscle noise. Inadequate skin preparation prior to application of electrode, tremors, tense subject, and/or poor electrode placement. Patient may have a pacemaker that contains a respiratory sensor.</td>
<td>Repeat skin preparation and electrode location procedures. Apply fresh, moist electrodes. Avoid areas of the torso that are very muscular. Reposition electrodes to reduce ECG artifact.</td>
</tr>
<tr>
<td>Intermittent Signal</td>
<td>Connections not tight and/or properly secured. Electrodes dry or loose. Cable or lead wires damaged.</td>
<td>Ensure proper connection. (cable to monitor, cable to lead, lead to electrode). Repeat skin preparation and apply fresh, moist electrodes. Check with continuity tester.</td>
</tr>
<tr>
<td>Excessive alarms: heart rate, lead fault</td>
<td>Electrodes dry Alarm limits set too close to patient’s normal heart rate. R-wave wrong size Excessive patient movement or muscle tremor.</td>
<td>Repeat skin preparation and apply fresh, moist electrodes. Adjust alarm limits. Must be twice the amplitude of the other waves, like the P and T waves. Reposition electrodes and secure with tape, if necessary.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
### Arrhythmia and ST Analysis Troubleshooting

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low Amplitude ECG Signal</td>
<td>Electrode could be positioned over a bone or muscle mass. Electrodes dry/old Skin improperly prepared This could be the patient's normal QRS complex. Gain set too low.</td>
<td>Move ECG patches away from the bone or muscle mass. Apply fresh, moist electrodes. Abrade skin. Verify with 12-lead electro-cardiogram. Readjust the ECG wave gain as required. Refer to the Wave Gain Tab [System] for instructions on adjusting the wave gain setting. NOTE: Changing the ECG wave gain only affects the signal displayed on the monitor. The signal analyzed by the arrhythmia algorithm and, consequently, beat detection and classification will not be affected.</td>
</tr>
<tr>
<td>No ECG Waveform</td>
<td>Gain set too low.</td>
<td>Readjust the ECG wave gain as required. Refer to the Wave Gain Tab [System] for instructions on changing the wave gain setting. NOTE: Changing the ECG wave gain only affects the signal displayed on the monitor. The signal analyzed by the arrhythmia algorithm and, consequently, beat detection and classification will not be affected.</td>
</tr>
<tr>
<td>Base Line Wander</td>
<td>Patient moving excessively. Patient's respiration Electrodes dry or loose Static build up around patient.</td>
<td>Secure lead wires and cable to patient. Reposition electrodes Repeat skin preparation and apply fresh, moist electrodes. Check with biomedical engineer.</td>
</tr>
</tbody>
</table>

---

* Messages are shown in all bold text.*
This page intentionally left blank.
5.0 *Patient Setup Functions*

This section describes patient setup at the Panorama Central Station.

- Demographics Tab
- Transfer Tab
- Discharge Tab
- Display Tab
- Standby Tab
- Wave Gain Tab (Patient)
- Print Setup Tab (Patient)
5.1 Demographics Tab

The **Demographics** tab (shown in FIGURE 5-1) allows for the entry and modification of patient demographic information (such as name, age, and weight) and patient settings (such as care group, pacers, arrhythmia, and ST). The **Demographics** tab is only accessible when the selected patient tile is assigned to a monitoring device. The default settings for patient demographics are defined by the **Unit Choices** tab, which is described in the “Unit Choices Tab” on page 9-50.

The **Demographics** tab is available at the ViewStation (shown in FIGURE 5-3) but is for viewing purposes only. When a patient tile is assigned at the ViewStation, it will always display the demographic information of the associated host central station patient tile, provided the patient tile is in an active state.

The **Demographics** tab is available at the WorkStation and may be used in the same manner as it is at the central station. Similar to the central station, demographic changes made to a patient tile at the WorkStation will be sent to the central station, View Only Workstation, and bedside monitor (if applicable).

Automatic updating of the demographic information at the remote workstation will also occur when communication is interrupted, for example:

- The monitoring device is placed in the Standby mode at the central station
- The Telepack is not communicating with its host central station (Communications Lost)
- The patient monitor is not communicating with its host central station (Comm Loss - Check Bedside Monitor)
- Communication between the host central station and the ViewStation and/or the WorkStation is lost (Central Communications Lost)

**NOTE:** The host central station(s) C-LAN configuration settings and the software installed at the remote location dictates the functionality available for each patient tile. For additional information, refer to “The ViewStation and the Workstation” on page 2-4.

5.1.1 Communications with the Panorama Gateway

If the central station is communicating with an Electronic Medical Records (EMR) system through the Panorama Gateway, entry of demographic information into the **Demographics** tab will differ slightly from the data entry instructions that follow. When using an EMR system, any changes to patient demographics made at the central station and/or bedside monitor will not be sent to the EMR system.

For the central station to be recognized by the EMR system, specific demographics content, referred to as the “Patient Key”, must be entered in the **Demographics** tab. Each facility has its own unique “Patient Key.” The “Patient Key” consists of the required Patient ID # and, optionally, one or more of the following Patient Menu demographics fields:

- First Name
- Last Name
Once the correct “Patient Key” has been entered, communication with the EMR system must be verified. Since the Panorama Gateway can be purchased with Admit/Discharge/Transfer (ADT) messaging only, Results messaging only or ADT messaging with Results messaging, this verification differs as follows:

**ADT Messaging**
For a Panorama Gateway that has ADT messaging, verify that communication with the EMR system has been established as follows:

- If demographics information corresponding to the following fields was not part of the “Patient Key” but was entered into the EMR system, these fields should auto-populate:
  - Patient ID #
  - First name
  - Last name
  - Date of Birth
  - Gender
- If the fields do not auto-populate, verify that the “Patient Key” was correctly entered and that the patient demographics were entered in the EMR system. If the fields still do not auto-populate after verification, contact the EMR administrator.

**Results Messaging**
For a Panorama Gateway that has Results messaging, verify that communication with the EMR system has been established by checking the charting system to ensure that patient’s vital signs data has been uploaded. If the charting system is not displaying this information, contact the EMR administrator.

### 5.1.2 Accessing the Demographics Tab

1. From the menu bar, select the **Patient Setup** menu button. The **Demographics** tab is displayed.
2. Choose a patient tile by selecting its digital or waveform area. The **Demographics** tab is enabled.

![Demographics Tab](image)
5.1.3 Entering Data in the Demographics Tab

This section outlines entering and modifying data in the Demographics tab.

**NOTE:** Patient demographic information cannot be edited at the ViewStation.

Select the Patient to be Set Up/Modified

Choose a patient tile by selecting its digital data or waveform area.

Device Label Field

The Device Label field displays the identification label for the device assigned to the selected patient tile.

The device label cannot be entered or modified in the Demographics tab. Device Label data is entered and modified via the “Equipment Setup Tab” as described on page 9-33.

Bed Field

The Bed field is used to display and enter the label data for bed assignment. The Bed field can only be entered, or modified, at the bedside monitor.
NOTE: When using a WMTS 608 bedside monitor, 15 characters may be entered at the bedside monitor but only the first five (5) characters will be displayed in the Bed field.

Entering a value in the Bed Field (Telepack)

1. From the Demographics tab, select the Bed field. The keyboard dialog box is displayed.
2. Enter a bed value.
   - The Backspace («») button clears a single character in the edit box.
   - The Caps Lock button enables/disables the caps lock function. A Caps Lock indicator displays when Caps Lock is enabled.
3. Accept or reject the text entered in the keyboard entry.
   - The Close button discards the changes.
   - The Enter button accepts the changes.

Device ID Field

The Device ID field displays the identification number for the device assigned to the selected patient tile.

The device ID cannot be entered or modified in the Demographics tab. Device ID data is entered and modified via the “Equipment Setup Tab” as described on page 9-33.

Last Name Field

The Last Name field is used to enter the last name of a patient. A patient's last name can be entered at the bedside monitor or the Panorama Central Station. Information entered at either the bedside, or at the central station, is automatically transmitted bi-directionally between the devices.

NOTE: If the central station is communicating with an EMR system via the Panorama Gateway, this field may auto-populate once the “Patient Key” has been correctly entered.

1. Select the Last Name field. The keyboard dialog box is displayed.
2. Enter the patient's last name (maximum of 15 characters).
   - The Backspace («») button clears a single character in the edit box.
   - The Caps Lock button enables/disables the caps lock function. A Caps Lock indicator displays when Caps Lock is enabled.
3. Accept or reject the text entered in the keyboard dialog box.
   - The Close button discards the changes.
   - The Enter button accepts the changes.

First Name Field

The First Name field is used to enter the first name of a patient. A patient's first name can be entered at the bedside monitor or the Panorama Central Station. Information entered at either the bedside, or at the central station, is automatically transmitted bi-directionally between the devices.
NOTE: If the central station is communicating with an EMR system via the Panorama Gateway, this field may auto-populate once the “Patient Key” has been correctly entered.

1. From the Demographics tab, select the First Name field. The keyboard dialog box is displayed.
2. Enter the patient’s first name (maximum of 15 characters).
   • The Backspace (<==) button clears a single character in the edit box.
   • The Caps Lock button enables/disables the caps lock function. A Caps Lock indicator displays when Caps Lock is enabled.
3. Accept or reject the text entered in the keyboard dialog box.
   • The Close button discards the changes.
   • The Enter button accepts the changes.

For additional information, refer to “Simultaneous Patient Demographic Tab Editing” on page 5-22.

Patient ID Field
The Patient ID field is used to enter the ID of a patient. A patient's ID can be entered at the bedside monitor or the Panorama Central Station. Information entered at either the bedside, or at the central station, is automatically transmitted bi-directionally between the devices.

1. From the Demographics tab, select the Patient ID field. The keyboard dialog is displayed.
2. Enter the patient’s ID.
   • The Backspace (<==) button clears a single character in the edit box.
   • The Caps Lock button enables/disables the caps lock function. A Caps Lock indicator displays when Caps Lock is enabled. Accept or reject the text entered in the keyboard dialog box.

For additional information, refer to “Simultaneous Patient Demographic Tab Editing” on page 5-22.

Height Field
The Height field is used to enter the height of a patient. A patient’s height can be entered at the bedside monitor or the Panorama Central Station. Information entered at either the bedside, or at the central station, is automatically transmitted bi-directionally between the devices.

1. From the Demographics tab, select the Height field. The Height dialog box (shown in FIGURE 5-4) is displayed.
2. Rotate the dial control or use the arrow buttons to select a value in the **Height** dialog box. Refer to the following value ranges.

**TABLE 5-1** All Spectrum, Passport 2 and Passport V (All Patients)

<table>
<thead>
<tr>
<th>CONTROLS</th>
<th>INCHES (in)</th>
<th>CENTIMETERS (cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Range</td>
<td>8.0 to 120.0</td>
<td>20.0 to 305.0</td>
</tr>
<tr>
<td>Dial Control</td>
<td>Increases number of inches when turned in a clockwise direction</td>
<td>Increases number of centimeters when turned in a clockwise direction</td>
</tr>
<tr>
<td></td>
<td>Decreases number of inches when turned in a counter-clockwise direction</td>
<td>Decreases number of centimeters when turned in a counter-clockwise direction</td>
</tr>
<tr>
<td>Single arrow</td>
<td>Increases/decreases inches by 0.5 in until 20.0 in is reached</td>
<td>Increases/decreases cm by 1.0 cm until 120.0 cm is reached</td>
</tr>
<tr>
<td></td>
<td>Increases/decreases inches by 1.0 in until 120.0 in is reached</td>
<td></td>
</tr>
<tr>
<td>Double arrow</td>
<td>Increases/decreases inches by 5.0 in until 20.0 in is reached</td>
<td>Increases/decreases cm by 10.0 cm until 305.0 cm is reached</td>
</tr>
<tr>
<td></td>
<td>Increases/decreases inches by 10.0 in until 120.0 in is reached</td>
<td></td>
</tr>
</tbody>
</table>

**TABLE 5-2** DPM 6/DPM 7, Passport 12M/Passport 17M, Passport 8/Passport 12 and T1 (All Patients)

<table>
<thead>
<tr>
<th>CONTROLS</th>
<th>INCHES (in)</th>
<th>CENTIMETERS (cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Range</td>
<td>8.0 to 118.0</td>
<td>20.0 to 300.0</td>
</tr>
<tr>
<td>Dial Control</td>
<td>Increases number of inches when turned in a clockwise direction</td>
<td>Increases number of centimeters when turned in a clockwise direction</td>
</tr>
<tr>
<td></td>
<td>Decreases number of inches when turned in a counter-clockwise direction</td>
<td>Decreases number of centimeters when turned in a counter-clockwise direction</td>
</tr>
<tr>
<td>Single arrow</td>
<td>Increases/decreases inches by 1.0 in</td>
<td>Increases/decreases cm by 1.0 cm</td>
</tr>
<tr>
<td>Double arrow</td>
<td>Increases/decreases inches by 10.0</td>
<td>Increases/decreases cm by 10.0</td>
</tr>
</tbody>
</table>
TABLE 5-3 V 12/V 21 (All Patients)

<table>
<thead>
<tr>
<th>CONTROLS</th>
<th>INCHES (in)</th>
<th>CENTIMETERS (cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Range</td>
<td>8.0 to 108.0 in</td>
<td>20.0 to 275.0 cm</td>
</tr>
<tr>
<td>Dial Control</td>
<td>Increases number of inches when turned in a clockwise direction</td>
<td>Increases number of centimeters when turned in a clockwise direction</td>
</tr>
<tr>
<td></td>
<td>Decreases number of inches when turned in a counter-clockwise direction</td>
<td>Decreases number of centimeters when turned in a counter-clockwise direction</td>
</tr>
<tr>
<td>Single arrow</td>
<td>Increases/decreases inches by 1.0 in</td>
<td>Increases/decreases cm by 1.0 cm</td>
</tr>
<tr>
<td>Double arrow</td>
<td>Increases/decreases inches by 10.0</td>
<td>Increases/decreases cm by 10.0</td>
</tr>
</tbody>
</table>

3. Accept or reject the value in the **Height** dialog box.
   - The **Done** button accepts the selection.
   - The **Cancel** button discards the selection, closes the dialog box, and returns to the **Demographics** tab.

For additional information, refer to “Simultaneous Patient Demographic Tab Editing” on page 5-22.

**Gender Field**

The **Gender** field is used to enter the gender of a patient. A patient's gender can be entered at the bedside monitor or the Panorama Central Station. Information entered at either the bedside, or at the central station, is automatically transmitted bi-directionally between the devices.

**NOTE:** If the central station is communicating with an EMR system via the Panorama Gateway, this field may auto-populate once the “Patient Key” has been correctly entered.

Select the **Gender** button until the desired gender selection is displayed.

**Weight Field**

The **Weight** field is used to enter the weight of a patient. A patient's weight can be entered at the bedside monitor or the Panorama Central Station. Information entered at either the bedside, or at the central station, is automatically transmitted bi-directionally between the devices.

For additional information regarding the Weight dialog box, refer to the “Weight Dial Dialog” on page 2-19.
1. From the Demographics tab, select the Weight field. The Weight dialog box (shown in FIGURE 5-5 and FIGURE 5-6) is displayed.

2. Rotate the dial control or use the arrow buttons to select a value in the Weight dialog box. Refer to the following value ranges.

3. Rotate the dial control or use the arrow buttons to select a value in the Weight dialog box. Refer to the following value ranges.

### TABLE 5-4 Spectrum, V 12/V 21 (Adult and Pediatric patient size)

<table>
<thead>
<tr>
<th>CONTROLS</th>
<th>POUNDS (lbs)</th>
<th>KILOGRAMS (kgs)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Range</td>
<td>1 lb to 1100 lbs</td>
<td>1 kg to 500 kg</td>
</tr>
<tr>
<td>Pound/Kilogram Dial</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Control</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Increases number of pounds when turned in a clockwise direction</td>
<td>Increases number of kilograms when turned in a clockwise direction</td>
</tr>
<tr>
<td></td>
<td>Decreases number of pounds when turned in a counter-clockwise direction</td>
<td>Decreases number of kilograms when turned in a counter-clockwise direction</td>
</tr>
<tr>
<td>Single arrow</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Increases/decreases by 1 lb each time button is selected</td>
<td>Increases/decreases by 1 kg each time button is selected</td>
</tr>
<tr>
<td>Double arrow</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Increases/decreases by 10 lbs each time button is selected</td>
<td>Increases/decreases by 10 kgs each time button is selected</td>
</tr>
</tbody>
</table>

### TABLE 5-5 Spectrum, V 12/V 21 (Neonate patient size)

<table>
<thead>
<tr>
<th>CONTROLS</th>
<th>POUNDS (lbs)</th>
<th>KILOGRAMS (kgs)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Range</td>
<td>Refer to TABLE 5-4</td>
<td>10 kg to 10000 kg</td>
</tr>
<tr>
<td>Single arrow</td>
<td>Refer to TABLE 5-4</td>
<td>Increases/decreases by 10 kgs each time button is selected</td>
</tr>
<tr>
<td>Double arrow</td>
<td>Refer to TABLE 5-4</td>
<td>Increases/decreases by 100 kgs each time button is selected</td>
</tr>
</tbody>
</table>
### TABLE 5-6  Spectrum, V 12/V 21 (Neonate patient size)

<table>
<thead>
<tr>
<th>CONTROLS</th>
<th>POUNDS (lbs)</th>
<th>KILOGRAMS (kgs)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Range</td>
<td>0 lb to 22 lbs 0 oz to 15 oz</td>
<td>Refer to TABLE 5-5</td>
</tr>
<tr>
<td>Pound Dial Control</td>
<td>Increases number of pounds when turned in a clockwise direction</td>
<td>Refer to TABLE 5-5</td>
</tr>
<tr>
<td></td>
<td>Decreases number of pounds when turned in a counter-clockwise direction</td>
<td></td>
</tr>
<tr>
<td>Ounces Dial Control</td>
<td>Increases number of ounces when turned in a clockwise direction</td>
<td>Refer to TABLE 5-5</td>
</tr>
<tr>
<td></td>
<td>Decreases number of ounces when turned in a counter-clockwise direction</td>
<td></td>
</tr>
</tbody>
</table>

### TABLE 5-7  Passport 2 and Passport V (All patient sizes) and Telepack (Adult patient size)

<table>
<thead>
<tr>
<th>CONTROLS</th>
<th>POUNDS (lbs)</th>
<th>KILOGRAMS (kgs)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Range</td>
<td>0.1 to 1100.0 lbs 0.1 to 500.0 kg</td>
<td></td>
</tr>
<tr>
<td>Pound/Kilogram Dial</td>
<td>Increases number of pounds when turned in a clockwise direction Increases number of kilograms when turned in a clockwise direction</td>
<td></td>
</tr>
<tr>
<td>Control</td>
<td>Decreases number of pounds when turned in a counter-clockwise direction Decreases number of kilograms when turned in a counter-clockwise direction</td>
<td></td>
</tr>
<tr>
<td>Single arrow</td>
<td>0.1 (Increases/decreases by 0.1 lb each time button is selected until 10.0 lbs is reached) 0.1 (Increases/decreases by 0.1 kg each time button is selected until 10.0 kg is reached)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>0.5 (Increases/decreases by 0.5 lbs each time button is selected once 10.0 lbs is reached) 0.5 (Increases/decreases by 0.5 kgs each time button is selected until 100.0 kgs is reached)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1.0 (Increases/decreases by 1.0 lbs each time button is selected until 1100.0 lbs is reached) 1.0 (Increases/decreases by 1.0 kgs each time button is selected until 500.0 kgs is reached)</td>
<td></td>
</tr>
<tr>
<td>Double arrow</td>
<td>0.1 (Increases/decreases by 1.0 lb each time button is selected until 10.0 is reached) 1.0 (Increases/decreases by 1.0 kg each time button is selected until 10.0 kg is reached)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>5.0 (Increases/decreases by 5.0 lbs each time button is selected once 100.0 lbs is reached) 5.0 (Increases/decreases by 5.0 kgs each time button is selected until 100.0 kgs is reached)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>10.0 (Increases/decreases by 10.0 lbs each time button is selected until 1100.0 lbs is reached) 10.0 (Increases/decreases by 10.0 kgs each time button is selected until 500.0 kgs is reached)</td>
<td></td>
</tr>
</tbody>
</table>
4. Accept or reject the selection made in the **Weight** dialog box.

   - The **Done** button accepts the selection.
   - The **Cancel** button discards the selection, closes the dialog box, and returns to the **Demographics** tab.

For additional information, refer to “Simultaneous Patient Demographic Tab Editing” on page 5-22.

### Patient Size Field

The **Patient Size** field is used to identify the size of the patient in the selected patient tile. The options for patient size include Adult, Pediatric and Neonate. The default patient size is Adult.

The value for the **Patient Size** field must be selected at the bedside monitor.

**NOTE:** Verify monitoring settings when the Patient Size is changed.

### Date Of Birth Field

The **Date Of Birth** field is used to enter the patient’s date of birth. A patient’s date of birth can be entered at the bedside monitor or the Panorama Central Station. Information entered at either the bedside, or at the central station, is automatically transmitted bi-directionally between the devices.

**NOTE:** If the central station is communicating with an EMR system via the Panorama Gateway, this field may auto-populate once the “Patient Key” has been correctly entered.

To enter a date of birth:

1. From the **Demographics** tab, select the **Date Of Birth** field. The **Date Of Birth** dialog box (shown in FIGURE 5-7) is displayed.
2. Rotate the dial controls to select the month, day and year values. Refer to the following value ranges.

<table>
<thead>
<tr>
<th>TIME FRAME</th>
<th>RANGE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Month</td>
<td>1 to 12</td>
</tr>
<tr>
<td>Day</td>
<td>Dependant on the month selected</td>
</tr>
<tr>
<td>Year</td>
<td>1891 to current year</td>
</tr>
</tbody>
</table>

3. Accept or reject the selection made in the Date Of Birth dialog box.
   - The Done button accepts the selections.
   - The Cancel button discards the selections, closes the dialog box and returns to the Demographics tab.

For additional information, refer to “Simultaneous Patient Demographic Tab Editing” on page 5-22.

Doctor Field
The Doctor field is used to enter the name of the patient’s doctor.

1. From the Demographics tab, select the Doctor field. The keyboard dialog box is displayed.
2. Enter the patient’s doctor (maximum of 15 characters).
   - The Backspace (<<) button clears a single character in the edit box.
   - The Caps Lock button enables/disables the caps lock function. A Caps Lock indicator displays when Caps Lock is enabled.
3. Accept or reject the text entered in the keyboard dialog box.
   - The Close button discards the changes.
   - The Enter button accepts the changes.

Comments Field
The Comments field is used to enter notes for the selected patient tile.

1. From the Demographics tab, select the Comments field. The keyboard dialog box is displayed.
2. Enter a note (maximum of 31 characters).
   - The Backspace (==) button clears a single character in the edit box.
   - The Caps Lock button enables/disables the caps lock function. A Caps Lock indicator displays when Caps Lock is enabled.
3. Accept or reject the text entered in the keyboard dialog box.
   - The Close button discards the changes.
   - The Enter button accepts the changes.

5.1.3.1 Pacer Configuration
Select the Pacer sidebar button to adjust a patient’s Pacer Enhancement, Pacer Filter settings, Pacer Indicator, and Pacing Rate (V 12/V 21 and Telepack only). The default Pacer Settings are determined in the system setup Unit Choices tab. For additional information regarding unit choices, refer to “Unit Choices Tab” on page 9-50 and to “Monitoring a Pacemaker Patient” on page 4-10.

- Pacer Enhancement visually marks pacemaker-induced ECG complexes with a colored vertical spike. It can be enabled or disabled in the Pacer Setting dialog box (shown in FIGURE 5-8 and FIGURE 5-9).

NOTE: A yellow vertical spike will be displayed when a paced beat has been detected and the ECG waveform color is not set to yellow.
A green vertical spike will be displayed when a paced beat has been detected and the ECG waveform color is set to yellow.

- Pacer Filtering adjusts the time period during which pacemaker-induced ECG complexes are filtered for ECG sampling. The preset options for the Pacer Filter are 0 to 80 ms (in increments of 20). The Pacer Filter is selected in the Pacer Setting dialog box (shown in FIGURE 5-8 and FIGURE 5-9).

- Pacer Indicator is a configurable long vertical spike through the waveform or a short vertical spike above the waveform.

- Pacing Rate is available only when V 12/V 21 or Telepack device is connected. The range available is 40-100 bpm. The factory default setting is empty. The Pacing Rate is configured in Pacer Setting dialog box (shown in FIGURE 5-9).

NOTE: When the Pacing Rate field is empty, the Pacer Failure to Sensor and Pacer Failure to Capture alarms are unavailable in the Alarm Responses tab and Patient Alarms Report for Telepack and V 12/V 21.
When the Pacing Rate field is empty, the Pacer Failure to Sensor and the Pacer Failure to Capture alarms on the V 12/V 21 are disabled.

NOTE: The Pacing Rate setting is disabled at the ViewStation or the WorkStation.
To enter Pacer Settings:

NOTE: For additional information regarding lead placement procedures for pacemaker patients, refer to “Monitoring a Pacemaker Patient” on page 4-10.

1. From the Demographics tab, select the Pacer sidebar button. The Pacer Setting dialog box is displayed.
2. Select the Pacer Enhancement button to enable or disable this setting for the patient.
   - A checkmark indicates that Pacer Enhancement is enabled.
   - An empty box indicates that Pacer Enhancement is disabled.
3. Select the desired Pacer Indicator by placing a check mark in the box next to the icon indicating the desired Pacer spike.
4. Select the Pacer Filter button until the desired setting is displayed. The selections for Pacer Filter range from 0 to 80 ms (in increments of 20).
5. Select the Pacing Rate text box. A keypad dialog box is displayed. Then enter a desired value from 40 to 100 bpm.
6. Accept or reject the selections made in the Pacer Setting dialog box.
   - The Done button accepts the selections.
   - The Cancel button discards the selections, closes the dialog box and returns to the Demographics tab.
5.1.3.2 Arrhythmia Setup

Select the **Arrhythmia** sidebar button to:

- Enable or disable arrhythmia analysis (for Telepack device only).
- Adjust the system default V-Tach Rate, Asystole Delay, and V-Tach Thresholds.
- Adjust the system default Pause Threshold RR (for V 12/V 21 and Telepack only).
- Adjust the system Default ECG Noise Delay, Primary Analysis Lead, Secondary Analysis Lead, and Beat Classification Lead (for Telepack only).

**NOTE:** The Arrhythmia sidebar button is not available at the ViewStation or the WorkStation.

**FIGURE 5-10** Arrhythmia Menu Dialog Box (Telepack, 5-wire lead set)

**FIGURE 5-11** Arrhythmia Menu Dialog Box (Passport 2, Passport V and Spectrum devices)

**FIGURE 5-12** Arrhythmia Menu Dialog Box (V 12/V 21)

1. Select the All Arrhythmia On/Off check box button (shown in FIGURE 5-10) to enable or disable arrhythmia analysis for the system.
• A checkmark in the All Arrhythmia On/Off box indicates that arrhythmia analysis is enabled.
• An empty box indicates that arrhythmia analysis is disabled.

NOTE: Arrhythmia analysis can be enabled for Telepack devices if an Arrhythmia license is available. When arrhythmia analysis is enabled, a license is automatically assigned to a patient.

NOTE: The All Arrhythmia On/Off check box button is only displayed and enabled when using a Telepack device. This check box is not available when using a bedside monitor (shown in FIGURE 5-11 and FIGURE 5-12). Arrhythmia analysis must be enabled or disabled at the bedside monitor.

2. Select the QT Monitoring On/Off check box button (shown in FIGURE 5-10) to enable or disable QT monitoring for the system.
   • A checkmark in the QT Monitoring On/Off box indicates that QT monitoring is enabled.
   • An empty box indicates that QT monitoring is disabled.

3. Select the QTc Formula button to change the QTc calculation formula. The available options are Hodges, Bazett, Fridericia and Framingham. The default factory setting is Hodges.

NOTE: The QT Monitoring On/Off check box and QTc Formula button are only displayed and enabled when using a Telepack device (shown in FIGURE 5-10).

NOTE: The QT Monitoring On/Off check box and QTc Formula button are displayed when using the V 12/V 21. The settings can only be adjusted on the V 12/21 bedside monitor. When the bedside monitor turns off QT monitoring, the QTc Formula displays Unknown (shown in FIGURE 5-12).

4. Rotate the dial control to change the V-Tach Rate. The range available is 100-180 bpm, in 5 bpm increments.

5. Select the Asystole Delay button to change the asystole delay time for arrhythmia processing. When using a 3 or 5-wire lead set, the range available is 3-10 seconds. When using a View 12™ card, the range available is 3-8 seconds.

6. Select the V-Tach Threshold button to change the V-Tach threshold settings. The range available is 3-15 beats.

7. Select the Pause Threshold button to change the Pause threshold settings. The range available is 1.2-2.0 RR%.

NOTE: Select the ECG Noise Delay button to change the noise delay setting. The range available is 3-30 seconds. The ECG Noise Delay button is only displayed and enabled when using a Telepack device. This button is not available when using a bedside monitor (shown in FIGURE 5-11 and FIGURE 5-12). ECG Noise Delay must be enabled or disabled at the bedside monitor.

8. Accept or reject the selections made in the dialog box.
   • Select the Done button to accept the selections.
   • Select the Cancel button to discard the selections, close the dialog box and return to the Unit Choices tab.
For additional information, refer to “Simultaneous Patient Demographic Tab Editing” on page 5-22.

5.1.3.3 ECG Analysis Leads (Telepack)
The Telepack uses information from two leads to detect beats and to compute heart rate (HR). These two leads are referred to as the primary and secondary leads. In addition, information from an additional lead (analysis lead) is made use of to classify the beats (normal, abnormal etc). The user can select any of the available leads (depending on whether a 3 or 5 lead cable is used) as primary, secondary or analysis leads.

For best results, the following guidelines should be used:

- the QRS complex should be tall, narrow and preferably either completely above or below the baseline. If at all possible, avoid selecting a lead where the QRS complex is biphasic.
- the P-waves and T-waves should be small compared to the QRS. They should be less than about 0.2 mV.

To select a lead as primary, secondary, or beat lead:

1. Select **Primary Analysis Lead** button (shown in FIGURE 5-10) to change the primary analysis lead setting. The lead available is I, II, III, aVR, aVL, aVF and V.
2. Select **Secondary Analysis Lead** button (shown in FIGURE 5-10) to change the secondary analysis lead setting. The lead available is I, II, III, aVR, aVL, aVF and V.
3. Select **Beat Classification Lead** button (shown in FIGURE 5-10) to change the beat classification lead setting. The lead available is I, II, III, aVR, aVL, aVF and V.

**NOTE:** The Primary Analysis Lead button, Secondary Analysis Lead button and Beat Classification Lead button are displayed and enabled only when using a Telepack device with 5-wire lead set.

**NOTE:** The Primary Analysis Lead button and Beat Classification Lead button are displayed but disabled when using a Telepack with 3-wire lead set.

5.1.3.4 Configuring Care Groups
Select the **Care Group** sidebar button to assign a specific care group to a patient tile. Care groups allow user-defined categorization of patient tiles. Care groups may represent a care-giver, hospital department, or a special condition. Each care group button has a color indicator that, when assigned to a patient, appears to the right of the **VIEW** button in the patient tile. The system default settings for Care Groups are defined in the System Care Group tab. For additional information, refer to the “Care Group Tab” on page 9-9.

The Care Group tab and the associated colorized indicators are not available at the ViewStation or the WorkStation.

To select a care group:

1. From the **Demographics** tab, select the **Care Group** sidebar button. The Care Group dialog box (shown in FIGURE 5-13) is displayed.
2. Select a **Care Group** button.

3. Accept or reject the **Care Group** selection.
   - The **Done** button accepts the care group selection.
   - The **Cancel** button discards the selection, closes the dialog box, and returns to the **Demographics** tab.

5.1.3.5 **ST Settings (Telepack)**

Select the **ST Settings** sidebar button to establish the type of ECG monitoring filter that will be used for the Telepack device at the central station or the WorkStation. There are two types of monitoring available:

**NOTE:** The **ST Settings** sidebar button is not available at the **ViewStation**.

- The **Monitor** filter establishes the frequency response range for non-ST ECG monitoring.
- The **ST** filter establishes the frequency response range for ST diagnostic monitoring.

To enter settings for a Telepack:

1. From the **Demographics** tab, select the **ST Settings** sidebar button. The **ST Settings** dialog box (shown in FIGURE 5-14) is displayed.

**FIGURE 5-13** Care Group Dialog Box

**FIGURE 5-14** ST Settings Dialog Box

2. Select the **ST** (optional) check box to enable or disable ST analysis for the selected patient.
   - A checkmark in the ST box indicates that ST analysis is enabled.
• An empty box indicates that ST analysis is disabled.
• When the ST check box is empty, the Monitor filter is automatically enabled.

NOTE: If ST licenses are available, an ST license is automatically assigned to the patient when ST is enabled.

NOTE: When using a 3-lead set, ST is performed on the lead being viewed.

3. Select the Monitor or ST filter mode.

NOTE: One filter box must always be populated. When a filter mode is selected, the other mode is automatically disabled.

• A checkmark in the Monitor filter box indicates that the Monitor filter is enabled and ST analysis is disabled.
• A checkmark in the ST filter box indicates that the ST analysis is enabled and the Monitor filter is disabled.

4. Accept or reject the selections.

• The Done button accepts the selections.
• The Cancel button discards the selections, closes the dialog box and returns to the Demographics tab.

5.1.3.6 Minimum QRS Threshold (Telepack)

Select the Minimum QRS Threshold sidebar button (shown in FIGURE 5-15) to adjust the QRS threshold.

FIGURE 5-15 Demographics Tab (Telepack)

The minimum detection threshold used in the automatic mode is about 0.16 mV. In case the P waves are very tall, one might consider moving the minimum QRS detection threshold up to be above the level of the P waves using the manual mode so events like ventricular standstill are not missed.
In the manual mode, two horizontal lines (one below and one above the baseline) appear on the screen (shown in FIGURE 5-17). These represent the current minimum detection threshold on the positive and negative sides of the baseline so manual adjustment for both positive and negative going QRS’s can be made. Using the commands provided (the arrows beside the sidebar buttons), move the minimum detection threshold up or down to ensure it is above the level of the P waves but below the peak of the R-wave. Since the P-wave height could vary a little from beat to beat, do not set the horizontal line representing the minimum detection threshold at or barely above the level of the peak of the P wave. Ensure that it is at least one or two millimeters above the peak of the P wave but below the peak of the R-wave.

1. From the Demographics tab, select the Minimum QRS Threshold sidebar button (shown in FIGURE 5-15).

**NOTE:** The Minimum QRS Threshold sidebar button is not available at the ViewStation or the WorkStation.

2. Wait for the waveform to draw, and then select the Adjust sidebar button (shown in FIGURE 5-16) to manually adjust the QRS threshold.

![FIGURE 5-16 Minimum QRS Threshold - Adjust Sidebar Button](image)

3. Select the arrows beside the sidebar buttons to move the waveform QRS threshold line up or down. The threshold setting range is between 0.16 mV to 0.48 mV. If needed, select the Resume sidebar button to reset the adjustment.

![FIGURE 5-17 Minimum QRS Threshold - Resume Sidebar Button](image)
4. To use the default threshold setting, select the Default sidebar button after selecting the Adjust sidebar button.

5. Select the Accept or Cancel sidebar button.
   - The Accept sidebar button accepts the configured settings and returns to the Demographics tab.
   - The Cancel sidebar button discards the configured settings and returns to the Demographics tab.

5.1.3.7 Retrieving Discharged Patients

Select the Retrieve Discharged Pt. sidebar button to retrieve a patient from the discharge list. A patient is only added to the discharge list if at least two of the three demographic fields (First Name, Last Name, Patient ID) were entered prior to discharge.

NOTE: The Retrieve Discharged Pt. sidebar button is not available at the ViewStation or the WorkStation.

The Retrieve Discharged Patient Setting view (shown in FIGURE 5-18) provides a list of the discharged patients that can be retrieved. The list has four columns labeled First Name, Last Name, Patient ID, and Discharge Time. Each row in the list contains at least two of the three demographic indicators and the discharge time for a patient. The list displays the 15 most recently discharged patients. When the next patient is added to the list, the oldest patient data on the list is removed and permanently deleted from the system.

- A discharged patient can be retrieved into a patient tile that is not actively monitoring a patient.
- A discharged patient can be retrieved into any patient tile that has an assigned device.
- A discharged patient can be retrieved into a tile assigned a different monitoring device than was previously assigned to that patient.

Retrieving a discharged patient into a patient tile:

- Allows for active monitoring of the retrieved patient
- Restores Trend, Event, and Disclosure records
- Restores Wave Gain and Trend Display settings
- Restores patient’s alarm limit settings to reflect those at the bedside monitor
- Does not restore patient’s alarm limit settings when using a Telepack device

NOTE: If the destination device and the source device are different, the source device’s sensor display setup will be reset to the system default settings of the destination device.

To retrieve a discharged patient:

1. Select the Retrieve Discharged Pt. sidebar button in the Demographics tab. The Retrieve Discharged Patient Setting view (shown in FIGURE 5-18) is displayed.
FIGURE 5-18 Retrieve Discharged Patient Setting View

2. Select the patient to be retrieved.
3. Accept or reject the patient retrieval in the Retrieve Discharged Patient Setting view.
   - The **Done** button accepts the patient retrieval.
   - The **Cancel** button discards the patient retrieval, closes the view, and returns to the Demographics tab.

5.1.3.8 Simultaneous Patient Demographic Tab Editing

**Bedside Menu**
If using a bedside device with the Panorama Central Station, patient demographic changes cannot be made at the bedside monitor, central station, and the WorkStation simultaneously. The bedside device will take precedence, followed by the central station, ending with the WorkStation. When an attempt is made to make changes simultaneously, the message **These settings are currently being edited remotely.** will be displayed at the central station or WorkStation, depending on where the change was initiated. If there are multiple WorkStations on the network, the first WorkStation that initiates the change will take precedence over any other WorkStation attempting to implement the same change.

These are the scenarios in which this message will be displayed:

**Patient Menu (Passport 2, Passport V, Spectrum, Spectrum OR, DPM 6/DPM 7, Passport 12M/Passport 17M, Passport 8/Passport 12, T1 and V 12/V 21 only)**

- The Patient Menu is active at the bedside monitor and then the Demographics tab is opened at the central station or the WorkStation
- The Demographics tab is active at the central station or WorkStation and then the Patient Menu is opened at the bedside monitor

**Arrhythmia Menu (Passport 2, Passport V, Spectrum, Spectrum OR, and V 12/V 21 only)**

- The Arrhythmia Menu is active at the bedside monitor and then the Arrhythmia Menu dialog is opened at the central station
- The Arrhythmia Menu dialog is active at the central station and then the Arrhythmia Menu is opened at the bedside monitor
CO Setup Menu (applies to the Spectrum, Spectrum OR, DPM 6/DPM 7, Passport 12M/Passport 17M, Passport 8/Passport 12 and V 12/V 21 only)

- The CO Setup Menu is active at the bedside monitor and then the Demographics tab is opened at the central station or the WorkStation
- The Demographics tab is active at the central station or the WorkStation and then the CO Setup Menu is opened at the bedside monitor

Hemodynamic Calculations Menu (applies to the Spectrum, Spectrum OR, and V 12/V 21 only)

- The Hemodynamic Calculations Menu is active at the bedside monitor and then the Demographics tab is opened at the central station or the WorkStation
- The Demographics tab is active at the central station or the WorkStation and then the Hemodynamic Calculations Menu is opened at the bedside monitor

Drug Calculator (applies to the Spectrum, Spectrum OR, and V 12/V 21 only)

- The Drug Calculator is active at the bedside monitor and then the Demographics tab is opened at the central station or the WorkStation
- The Demographics tab is active at the central station or the WorkStation and then the Drug Calculator is opened at the bedside monitor

Central Station or WorkStation

The message These settings are currently being edited remotely. will be displayed at the WorkStation when the WorkStation Demographics tab is active and then the Demographics tab is opened at the central station. In this scenario, the central station will take precedence, followed by the WorkStation.

This message will also be displayed at the WorkStation when the WorkStation patient Alarm Limits or Alarm Responses tab is active and then the Alarm Limits or Alarm Responses tab is opened at the central station. In this scenario, the central station will take precedence, followed by the WorkStation.

Acknowledging the Message

To close the These settings are currently being edited remotely. message dialog, select the OK button. Acknowledging this message will convert the Demographics tab at the central station or WorkStation to a read-only state and editing will not be permitted. Once the menu at the bedside device or central station is closed or times out (timeout applies to bedside device only), the Demographics tab will become active again and editing can resume.

Unsaved central station or WorkStation changes made in the First Name keyboard dialog, Date Of Birth dialog, Height dialog box, Patient ID keyboard dialog, Last Name keyboard dialog, Weight dialog box, or Arrhythmia Menu (applies to central station only) dialog box will be discarded.

For additional information regarding the bedside monitor menus, refer to the bedside monitor’s Operating Instructions:
- Passport 2 (P/N 0070-00-0649-XX)
- Passport V (P/N 0070-00-0704-XX)
- Spectrum (P/N 0070-00-0648-XX)
- Spectrum OR (P/N 0070-00-0670-XX)
- DPM 6 (P/N 046-000184-00)
- DPM 7 (P/N 046-000186-00)
- V 12/V 21 (P/N 0070-00-0727-XX)
- Passport 12M/Passport 17M (P/N 046-005013-00)
- Passport 8/Passport 12 (P/N 046-004417-00)
- T1 (P/N 046-002376-00)

**NOTE:** While the Demographics tab is in a read-only state because the Patient Menu is open at the bedside, the Pacer, Arrhythmia, and Care Group sidebar buttons will be enabled.

### 5.1.4 Sidebar Buttons

The following sidebar buttons are used to expand the functionality of this tab.

**Pacer**
This sidebar button was previously described on page 5-13.

**Arrhythmia**
This sidebar button was previously described on page 5-15.

**Minimum QRS Threshold**
This sidebar button was previously described on page 5-19.

**Care Group**
This sidebar button was previously described on page 5-17.

**Retrieve Discharged Pt.**
This sidebar button was previously described on page 5-21.

**More (Telepack)**
Select the *More* sidebar button to see additional sidebar buttons while using a Telepack and viewing the *Demographics* tab.

**Normal Screen**
Select the *Normal Screen* sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.
### Troubleshooting

This section lists some of the potential messages and issues that may occur while using the **Demographics** tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Can not admit Patient to this tile. No Device attached to this tile</td>
<td>There is no device assigned to the selected tile.</td>
<td>Select the <strong>OK</strong> button to close the dialog box. Select a patient tile that has a device attached to it or attach a device to the selected patient tile.</td>
</tr>
<tr>
<td>The <strong>Arrhythmia</strong> sidebar button is disabled.</td>
<td>No equipment is assigned to the patient tile.</td>
<td>Select a different patient tile.</td>
</tr>
<tr>
<td>The <strong>Pacer</strong>, <strong>Care Group</strong>, and the <strong>Retrieve Discharged Pt.</strong> sidebar buttons are disabled.</td>
<td>The selected patient is in the <strong>Standby</strong> mode.</td>
<td>Remove the selected patient tile from <strong>Standby</strong> mode.</td>
</tr>
<tr>
<td>The filter mode cannot be Monitor when ST is on</td>
<td>Monitor mode cannot be selected while ST is enabled.</td>
<td>Select the <strong>OK</strong> button to close the dialog box. Turn off ST select the correct filter mode.</td>
</tr>
<tr>
<td>Turning on ST will change your ECG filtering to ST mode</td>
<td>ST mode was selected while in Monitor mode.</td>
<td>Select the <strong>OK</strong> button to close the dialog box. Select the correct filter mode.</td>
</tr>
<tr>
<td>Retrieve Discharged Patient Failed: The destination tile is currently occupied.</td>
<td>A patient cannot be retrieved into an occupied tile.</td>
<td>Select the <strong>OK</strong> button to close the dialog box. Select a vacant patient tile before attempting to retrieve a patient.</td>
</tr>
<tr>
<td>No demographic data is displayed at ViewStation or WorkStation for the selected patient tile.</td>
<td>The host central station is in a Central Communications Lost mode.</td>
<td>Resolve the communication error at the host central station.</td>
</tr>
<tr>
<td>Patient tile does not contain a care group indicator</td>
<td>If viewing patient tile from the central station, a care group has not been indicated.</td>
<td>Assign the patient tile to a care group.</td>
</tr>
<tr>
<td>Patient tile is being viewed from a ViewStation or a WorkStation, which does not support this feature.</td>
<td></td>
<td>To discern if the patient is assigned to a care group, view the patient tile from the host central station.</td>
</tr>
</tbody>
</table>

*Messages are shown in all bold text.*
<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>These settings are currently being edited remotely.</td>
<td>The Care Group dialog box was closed while the bedside device's Patient Menu is active.</td>
<td>Select the <strong>OK</strong> button to close the message box and enable the sidebar button.</td>
</tr>
<tr>
<td>These settings are currently being edited remotely.</td>
<td>The <strong>Retrieve Discharged Patient Settings</strong> tab was closed while the bedside device's Patient Menu is active.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>The Arrhythmia Menu dialog box was closed while the bedside device's Patient Menu is active.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>The Pacer Settings dialog box was closed while the bedside device's Patient Menu is active.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>The CO Setup Menu is opened at the bedside monitor while the <strong>Demographics</strong> tab is active at the central station or WorkStation.</td>
<td>Use the bedside monitor to make the arrhythmia changes.</td>
</tr>
<tr>
<td></td>
<td>The <strong>Demographics</strong> tab is active at the central station or WorkStation and then the CO Setup Menu is opened at the bedside monitor.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>The Hemodynamic Calculations Menu is opened at the bedside monitor while the <strong>Demographics</strong> tab is active at the central station or WorkStation.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>The <strong>Demographics</strong> tab is active at the central station or WorkStation and then the Hemodynamic Calculations Menu is opened at the bedside monitor.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>The Arrhythmia Menu at the central station was closed and all changes were discarded.</td>
<td>Close the Arrhythmia Menu at the bedside monitor and then reenter the Arrhythmia Menu dialog box at the central station or WorkStation.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
5.2 Transfer Tab

The Transfer tab (shown in FIGURE 5-19) enables the transfer of a patient’s data from one device to another. Three types of transfers can be performed through the Transfer tab:

- Room To Room Transfer Without Device
- Room To Room Transfer With Device
- Central To Central Transfer Without Device

The Transfer tab is not available at the ViewStation or the WorkStation. Patient transfers can only be performed at a host central station.

5.2.1 Accessing the Transfer Tab

1. From the menu bar, select the Patient Setup menu button. The tabs associated with patient setup are displayed.
2. Choose a patient tile by selecting its digital tile or waveform area.
3. Select the Transfer tab. The Transfer tab is enabled.

![FIGURE 5-19 Transfer Tab](image)

5.2.2 Room To Room Transfer Without Device

A Room To Room Transfer Without Device is used to transfer patient data stored at the Panorama Central Station from one patient tile (the source) to another (the destination). Historical patient data is preserved upon successful completion of the transfer, with the exception of historical paging data. If Paging is enabled at the central station, a successful Room to Room Transfer Without Device shall delete any rows in the Failed Pages tab for the transferred patient.

For information regarding changes that occur at the destination tile when the transfer is complete, refer to “Effects of the Transfer” on page 5-28.

The Room To Room Without Device button is enabled under the following conditions:

- When a patient tile actively monitoring a patient is selected as the source
- When a patient tile actively monitoring a patient, but placed in Standby mode, is selected as the source
When a patient tile actively monitoring a patient, but experiencing a Communications Lost or a Comm Loss - Check Bedside Monitor event, is selected as the source.

To perform a Room To Room Transfer Without Device:

1. Select the patient tile to be transferred. Choose a patient tile by selecting its digital data or waveform area.

2. Select the Room To Room Without Device button. The following prompt is displayed: Please touch tile to which you wish to transfer the patient.

3. Select a destination tile that has a device assigned. The message All of the patient’s data will be deleted in the destination tile. Do you still want to continue with the transfer? is displayed.

   • The Yes button initiates the transfer. As the transfer proceeds, there is no indication in the Transfer tab.

   • If the destination tile is in Standby mode when the transfer is complete, the digital data labels and the demographic information will be displayed in the patient tile, but the digital data values and waveforms will not be displayed until the device is returned to the normal monitoring mode.

   • If the destination tile is in a Communications Lost or Comm Loss - Check Bedside Monitor state when the transfer is complete, the tile is blank and all buttons on the Transfer tab are unavailable. The tile remains blank until communication has been re-established.

When the transfer is complete, the source tile is blank and the destination tile remains blank until the patient is connected to the monitoring device.

   • If the destination tile is in a Waiting for Patient state when the transfer is complete, the tile will display the Waiting for Patient message until the destination device is activated.

   NOTE: When the transfer is complete, a Room Transfer event and an Admit event are recorded in the Event database.

   NOTE: After a Room To Room Transfer Without Device, the patient’s Paging settings will remain the same.

   • The No button terminates the transfer and returns to the Transfer tab.

   NOTE: When transferring patients out of tiles hosted by Passport V, DPM 6/7, Passport 12M/Passport 17M, Passport 8/Passport 12 or T1 monitors, a new patient will be automatically admitted to the Panorama for that source tile following the completion of the transfer. The new patient will immediately be in an active monitoring state and the Panorama will reflect the data being gathered at the bedside.

5.2.2.1 Effects of the Transfer

The following is a list of the changes that occur at the destination tile when a Room To Room Transfer Without Device is complete:

Display
The transferred patient’s parameter display changes only when the destination device does not support the source device parameters.

The transferred patient’s parameter display waveform 1 changes to ECG II when the destination device does not support the source device’s waveform 1 parameter and has a 5 lead ECG cable.

The transferred patient’s parameter display waveform 1 changes to the active lead when the destination device does not support the source device’s waveform 1 parameter and has a 3 lead ECG cable.

The transferred patient’s parameter display waveform 2 changes to OFF when the destination device does not support the source device’s waveform 2 parameter.

If the source device is a bedside monitor, the units for CO2 and temperature will change to that of the destination device.

If the destination device and the source device are different, the source devices sensor display setup will be reset to the system default settings of the destination device.

The care group is set to the system default at the destination device.

Demographics

For all transfers to a bedside monitor, the bed ID and patient size change to those of the destination device. The remaining demographic data only changes when the destination device has experienced a Communications Lost or Comm Loss - Check Bedside Monitor event. When communication is re-established, the remaining demographic data is taken from the destination device.

For all transfers to a Panorama Telepack, the bed ID is cleared and the patient size changes to Adult (the Telepack default). The remaining demographic data is preserved.

Alarms

The transferred patient’s Alarm Limits, Alarm Delay, and ST Alarm Delay change to those of the destination device.

All alarms for the transferred patient become enabled, discontinuing all Alarm Suspended settings.

5.2.3 Room To Room Transfer With Device

This is used to swap the patient’s database and equipment from one patient tile (the source) to another (the destination). Historical patient data is preserved upon successful completion of the transfer, with the exception of historical paging data. If Paging is enabled at the central station, a successful Room to Room Transfer With Device shall delete any rows in the Failed Pages tab for the transferred patient.

The Room To Room With Device button is disabled when a patient tile is not selected.

To perform a Room To Room Transfer With Device:

1. Select the patient tile to be transferred. Choose a patient tile by selecting its digital data or waveform area.

2. Select the Room To Room With Device button. The following prompt is displayed: Please select the tile you wish to transfer the device and the patient.
3. Select a destination tile. The message **Are you sure you want to proceed with the transfer?** is displayed.

- The **Yes** button initiates the transfer. As the transfer proceeds, there is no indication in the **Transfer** tab.

  If the destination tile is in **Standby** mode when the transfer is complete, the digital data labels and the demographic information will be displayed in the patient tile, but the digital data values and waveforms will not be displayed until the device is returned to the normal monitoring mode.

  If the destination tile is in a **Communications Lost** or **Comm Loss - Check Bedside Monitor** state when the transfer is complete, the tile is blank and all buttons on the **Transfer** tab are unavailable. The tile remains blank until communication has been re-established.

  When the transfer is complete, the source tile is blank and the destination tile remains blank until the patient is connected to the monitoring device.

  If the destination tile is in a **Waiting for Patient** state when the transfer is complete, the tile will display the **Waiting for Patient** message until the destination device is activated.

**NOTE:** When the transfer is complete, a Room Transfer event and an Admit event are recorded in the Event database.

**NOTE:** After a Room To Room Transfer With Device, the patient’s Paging settings will remain the same.

**NOTE:** The Telepack device involved in the transfer will relearn ST and arrhythmia.

- The **No** button terminates the transfer and returns to the **Transfer** tab.

### 5.2.3.1 Effects of the Transfer

The following is a list of the changes that occur at the destination tile when a **Room To Room Transfer With Device** is complete:

**Display**

- The source patient tile and destination patient tile swap places.

**Demographics**

- For all transfers, the transferred patient’s demographic data is preserved from the source tile.

**Alarms**

- Discontinue the transferred patient’s Alarm Off/Alarm Pause settings.

**NOTE:** When the transfer is complete, equipment corresponding to the source tile and destination tile swaps places in the Equipment Setup tab and Equipment Report.
5.2.4 Central To Central Transfer Without Device

A **Central To Central Transfer Without Device** is used to transfer patient data stored at one Panorama Central Station to a different Panorama Central Station. Historical patient data is preserved upon successful completion of the transfer, with the exception of historical failed page data. If Paging is enabled at the central station, a successful Central To Central Transfer Without Device will delete any rows in the **Failed Pages** tab for the transferred patient.

The **Central To Central Without Device** button is enabled under the following conditions:

- When a patient tile actively monitoring a patient is selected as the source
- When a patient tile actively monitoring a patient, but placed in **Standby** mode, is selected as the source
- When a patient tile actively monitoring a patient, but experiencing a **Communications Lost** or a **Comm Loss - Check Bedside Monitor** event, is selected as the source

For information regarding changes that occur at the destination tile when the transfer is complete, refer to “Effects of the Transfer” on page 5-34.

5.2.4.1 Source Panorama

1. A prerequisite for the transfer is that certain patient demographic information must exist. In the patient tile to be transferred, ensure that the appropriate information has been provided in at least two of the three demographic fields (Last Name, First Name, and Patient ID).

2. Ensure that the device name for the Source Panorama is properly entered into the Equipment List at the Destination Panorama, as described in the “Equipment Setup Tab” on page 9-33.

3. Ensure that the device name for the Destination Panorama is properly entered into the Equipment List at the Source Panorama, as described in the “Equipment Setup Tab” on page 9-33.

4. Select the patient tile to be transferred. Choose a patient tile by selecting its digital data or waveform area.

5. Select the **Central To Central Without Device** button. The **Select Destination Panorama** list box (shown in FIGURE 5-20) is displayed.

![FIGURE 5-20 Select Destination Panorama List Box](image-url)
6. Select the Destination Panorama device from the Select Destination Panorama list box.

   • Select the Done button. The message Are you sure you want to transfer the selected patient? is displayed with the following two choices:

   • Select the Yes button to initiate the first part of the transfer process and return to the Transfer tab. In the Transfer tab, the message Central Transfer in progress is displayed next to the Central To Central Without Device button, which has been disabled.

   • Select the No button to terminate the transfer and return to the Transfer tab.

   • Select the Cancel button to terminate the transfer and return to the Transfer tab.

If the transfer process is initiated, it will continue as described in following section, “Destination Panorama”. As the transfer process continues, the procedure will refer to messages and a progress bar that will display at the Source Panorama.

5.2.4.2 Destination Panorama

When a Central To Central Transfer Without Device has been requested from another Panorama Central Station, the message Transfer In Request from [the Source Panorama name]. Accept/Reject in Transfer Dialog. is displayed in the system status line of the Destination Panorama.

1. At the Destination Panorama, select a tile that has a monitoring device attached.

2. From the menu bar, select the Patient Setup menu button. The tabs associated with Patient Setup are displayed.

3. Select the Transfer tab. The Transfer tab (shown in FIGURE 5-19) is displayed.

4. Select the Accept/Reject button on the Transfer tab. The Transferred In Patient view (shown in FIGURE 5-21) is displayed.

![FIGURE 5-21 Transferred In Patient View](image-url)
Select the **Accept** button. The message **All of the patient's data will be deleted in the destination tile. Do you still want to continue with the transfer?** is displayed with the following two choices:

- The **Yes** button begins the transfer and returns to the **Transfer** tab. While the transfer is in progress, the following conditions exist:
  
  In the **Transfer** tab of the Destination Panorama, the message **Incoming Transfer In progress** is displayed next to the **Accept/Reject Patient** button, which has been disabled. A progress bar displays below the **Accept/Reject Patient** button.

  In the **Transfer** tab of the Source Panorama, a progress bar displays below the **Central To Central Without Device** button.

  If the destination tile is in **Standby** mode when the transfer is complete, the digital data displays but the waveform does not display. Upon returning to normal monitoring mode, the waveform is displayed.

  If the destination tile is in a **Communications Lost** or a **Comm Loss - Check Bedside Monitor** state when the transfer is complete, the tile is blank and all buttons on the **Transfer** tab are unavailable. The tile remains blank until communication has been re-established.

  If the destination tile is in a **Waiting for Patient** state when the transfer is complete, the tile will display the Waiting for Patient message until the destination device is activated.

  When the transfer is complete, the message **Transfer In Complete** is displayed on the system status line of the Destination Panorama and the message **Transfer Out Complete** is displayed on the system status line of the Source Panorama.

  When the transfer is complete, the source tile is blank and the destination tile remains blank until the patient is connected to the monitoring device.

**NOTE:** When the transfer is complete, a Central Transfer event and an Admit event are recorded in the event database.

- The **No** button returns to the **Transfer** tab without terminating the transfer.

- The **Reject** button terminates the transfer and returns to the **Transfer** tab. The message **Transfer Out rejected by [the Destination Panorama name] Central.** is displayed in the system status line of the Source Panorama.

- The **Cancel** button returns to the **Transfer** tab without terminating the transfer.

**NOTE:** When transferring patients out of tiles hosted by Passport V, DPM 6/DPM 7, Passport 12M/Passport 17M, Passport 8/Passport 12, or T1 monitors, a new patient will be automatically admitted to the Panorama for that source tile following the completion of the transfer. The new patient will immediately be in an active monitoring state and the Panorama will reflect the data being gathered at the bedside.
NOTE: A Central To Central Transfer Without Device can fail if the network connection is lost or if either Panorama loses power during the process. If a Central To Central Transfer Without Device fails while in progress, the patient information from the source tile is saved to a Failed Transfer List at the Source Panorama.

The Failed Transfer List stores a maximum of five failed transfers. If the maximum is reached, the Central To Central Without Device button is disabled. A Central To Central Transfer Without Device cannot be initiated until at least one of the failed transfers is either recovered, deleted or transferred using one of the sidebar buttons in the Transfer tab. For additional information, refer to “Sidebar Buttons” on page 5-35.

5.2.4.3 Effects of the Transfer

The following is a list of the changes that occur at the destination tile when a Central To Central Transfer Without Device is complete:

Display

- The transferred patient’s parameter display changes, but only when the destination device does not support the source device parameters.
- The transferred patient’s parameter display waveform 1 changes to ECG II when the destination device does not support the source device’s waveform 1 parameter and has a 5-lead ECG cable.
- The transferred patient’s parameter display waveform 1 changes to the active lead when the destination device does not support the source device’s waveform 1 parameter and has a 3-lead ECG cable.
- The transferred patient’s parameter display waveform 2 changes to OFF when the destination device does not support the source device’s waveform 2 parameter.
- If the source device is a bedside monitor, the units for CO₂ and temperature will change to that of the destination device.
- If the destination device and the source device are different, the source devices sensor display setup will be reset to the system default settings of the destination device.
- The care group is set to the system default at the destination device.

Demographics

- For all transfers to a bedside monitor, the bed ID and patient size change to those of the destination device. The remaining demographic data only changes when the destination device has experienced a Communications Lost or a Comm Loss - Check Bedside Monitor event. When communication is re-established, the remaining demographic data is taken from the destination device.
- For all transfers to a Panorama Telepack, the bed ID is cleared and the patient size changes to Adult (the Telepack default). The remaining demographic data is preserved.

Alarms

- The transferred patient’s Alarm Limits, Alarm Delay and ST Alarm Delay change to those of the destination device.
5.2.5 Sidebar Buttons

The following sidebar buttons are used to expand the functionality of this tab.

Recover Patient

When a Central To Central Transfer Without Device has failed, the Recover Patient sidebar button is used to retrieve patient information that was collected in the Failed Transfer List.

The list box in the Recover Failed Transfer Patients view (shown in FIGURE 5-22) has three columns labeled First Name, Last Name and Patient ID. The list contains at least two of these three demographic indicators.

1. Select a destination tile on the Source Panorama to which the patient information is to be recovered. Ensure that the destination tile has a device assigned.

2. Select the Recover Patient sidebar button. The Recover Failed Transfer Patients view is displayed.

3. Select the patient whose information is to be recovered from the list.
   - Select the Recover button. The message All of the patient's data will be deleted in the destination tile. Do you still want to continue with the transfer? is displayed with the following two choices:
     - The Yes button deletes all of the patient’s data that may have been previously stored for the destination tile, returns to the Transfer tab and the recovery process begins. When the recovery is complete, the patient information is removed from the Failed Transfer List and the destination tile displays the recovered patient information, except in the following situations:
       - If the destination tile is in Standby mode when the recovery is complete, the digital data displays but the waveform does not display. Upon returning to normal monitoring mode, the waveform is displayed.
       - If the destination tile is in a Communications Lost or a Comm Loss - Check Bedside Monitor state when the recovery is complete, the tile is blank and all buttons on the Transfer tab are unavailable. The tile remains blank until communication has been re-established. Once communication is re-established, the destination tile will populate with the demographic information available at the destination bedside monitor.
• The **No** button terminates the recovery process and returns to the **Transfer** tab.
• Select the **Cancel** button to return to the **Transfer** tab.

**Delete Failed Patient**

When the Failed Transfer List has reached its maximum of five failed transfers, the **Delete Failed Patient** sidebar button is used to delete patient information from the list.

The list box in the **Delete Failed Transfer Patients** view (shown in FIGURE 5-23) has three columns labeled First Name, Last Name and Patient ID. The list will contain at least two of these three demographic indicators.

1. Select the **Delete Failed Patient** sidebar button. The **Delete Failed Transfer Patients** view is displayed.

   ![FIGURE 5-23 Delete Failed Transfer Patients View](image)

2. Select the patient whose information is to be deleted from the list.
   • Select the **Delete** button. The message **Are you sure that you want to delete this failed transfer?** is displayed with the following choices:

   **NOTE:** If the **Yes** button is selected, all of the patient’s data will be unrecoverable. Once the patient is deleted from the list, the patient will not be displayed in the Retrieve Discharged List and therefore, cannot be retrieved.

   • Selecting the **Yes** button when the Delete Failed Transfer Patient list is empty deletes all of the selected patients data from the Failed Transfer List and then returns to the **Transfer** tab.

   • Selecting the **Yes** button when the Delete Failed Transfer Patient list contains more patients will delete all of the selected patients data from the Failed Transfer List and then keep the Delete Failed Transfer Patients list open.

   • When the deletion is complete, the patient information is removed from the Delete Failed Transfer Patient list.

   • The **No** button terminates the deletion process and returns to the **Transfer** tab.

   • The **Cancel** button returns to the **Transfer** tab.
Transfer From Failed List

The Transfer From Failed List sidebar button is used to resume a transfer of patient information that was collected in the Failed Transfer List when a Central To Central Transfer Without Device failed. The patient data and settings collected in the Failed Transfer List at one Panorama Central Station are transferred to a room in an area monitored by a different Panorama Central Station. Historical patient data is preserved upon successful completion of the transfer.

The list box in the Transfer Failed Transfer Patients view (shown in FIGURE 5-24) has three columns labeled First Name, Last Name and Patient ID. The list contains at least two of these three demographic indicators.

1. Select the Transfer From Failed List sidebar button. The Transfer Failed Transfer Patients view is displayed.

   ![FIGURE 5-24 Transfer Failed Transfer Patients View](image)

2. Select the patient whose information is to be transferred from the list.
   - The Transfer button displays the Select Destination Panorama list box (shown in FIGURE 5-20) displays. Go to step 6 on page 5-32 and continue the transfer process. When a Destination Panorama is selected, the patient information is removed from the Failed Transfer List.
   - The Cancel button returns to the Transfer tab.

Normal Screen

Select the Normal Screen sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.
### Troubleshooting

This section lists some of the potential messages and issues that may occur while using the Transfer tab.

#### Room To Room Transfer

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patient tile shown with demographic text and digital data labels, but no waveforms and/or digital values are displayed</td>
<td>The selected tile is still in the Bedside and Panorama Standby mode.</td>
<td>Return the monitoring device to its active monitoring state.</td>
</tr>
<tr>
<td>The Room To Room Without Device button is disabled.</td>
<td>The Transfer Failed list is full. The list stores a maximum of 5 failed transfers.</td>
<td>Delete a patient from this list before transferring.</td>
</tr>
<tr>
<td></td>
<td>A patient tile is not selected on the Main Screen.</td>
<td>Select the patient tile that is intended to be transferred.</td>
</tr>
<tr>
<td></td>
<td>The selected patient tile has a device assigned but is not monitoring a patient.</td>
<td>Select the patient tile that is intended to be transferred.</td>
</tr>
<tr>
<td></td>
<td>The selected patient tile has no device assigned.</td>
<td>Select the patient tile that is intended to be transferred.</td>
</tr>
<tr>
<td>The Failed Patient List is full. Delete a patient from this list before transferring.</td>
<td>The Transfer Failed list is full. The list stores a maximum of 5 failed transfers.</td>
<td>Delete a patient from this list before transferring.</td>
</tr>
<tr>
<td>Unable to select a patient tile on the Panorama Display.</td>
<td>A message box is currently being displayed.</td>
<td>Close the message box or wait for it to time out and close automatically.</td>
</tr>
<tr>
<td>The destination and source tile cannot be the same. Please try again.</td>
<td>The original source tile was selected when the prompt “Please touch tile to which you wish to transfer the patient.” was displayed.</td>
<td>Restart the transfer process and select an appropriate destination tile when prompted. Do not select the original source tile.</td>
</tr>
<tr>
<td>The destination tile must have a device assigned to perform the transfer.</td>
<td>The destination tile does not have a device assigned.</td>
<td>Restart the transfer process and select a destination tile that has a device assigned when prompted.</td>
</tr>
<tr>
<td>This Tab is in use by another display. Can not be updated from more than one display at the same time.</td>
<td>An attempt was made to open the Transfer tab on one display when it is already open on the other display.</td>
<td>Select the OK button and use the open Transfer tab.</td>
</tr>
<tr>
<td>At the destination tile, the digital data is displayed but the waveform is not displayed</td>
<td>The destination tile was in Standby mode when the transfer completed.</td>
<td>Return to normal monitoring mode.</td>
</tr>
<tr>
<td>Room to Room Transfer Without device failed</td>
<td>Another tab was selected during the transfer process.</td>
<td>Restart the transfer process and do not select another tab until the transfer process is complete.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
### Room To Room Transfer (Continued)

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>The destination tile is blank and all buttons on the Transfer tab are unavailable.</strong></td>
<td>The destination tile was in a <strong>Communications Lost</strong> or a <strong>Comm Loss - Check Bedside Monitor</strong> state when the transfer completed.</td>
<td>Re-establish communication.</td>
</tr>
</tbody>
</table>

**Unable to perform operation on tile: Central to Central Transfer Out in progress**

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>A Central to Central transfer Out has been initiated at the source central station but not been accepted or rejected at the destination central station for the selected tile.</strong></td>
<td>Accept or reject the transfer at the destination central station, and then initiate Room to Room Transfer again.</td>
<td></td>
</tr>
</tbody>
</table>

**Unable to perform operation on tile: Central to Central Transfer In in progress.**

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>A Central to Central transfer In is in progress for the selected tile, which is actively monitoring a patient, and a Room to Room transfer is attempted into the same selected tile.</strong></td>
<td>Wait until the Central to Central transfer In is complete, then attempt transfer again.</td>
<td></td>
</tr>
</tbody>
</table>

**Unable to perform operation on tile: Central to Central Transfer In in progress.**

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>A Central to Central transfer In has been initiated at the source central station but not accepted or rejected at the destination central station for the selected tile, and another Central to Central transfer is attempted for another patient into the same selected tile.</strong></td>
<td>Wait until the Central to Central transfer In is complete, then attempt transfer again.</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>The Central to Central Without Device button in the Transfer Tab was selected when a Central to Central transfer In is in progress for the selected tile, and the selected tile is actively monitoring a patient.</strong></td>
<td>Wait until the Central to Central transfer In is complete, then attempt transfer again.</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
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<tbody>
<tr>
<td><strong>A Central to Central transfer In is in progress for the selected tile, which has an assigned device that is not actively monitoring a patient, and a Room to Room transfer for another patient is initiated into the same selected tile.</strong></td>
<td>Wait until the Central to Central transfer In is complete, then attempt transfer again.</td>
<td></td>
</tr>
</tbody>
</table>

**The Waiting for Patient message is displayed in the heart rate digital tile.**

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>The destination device has not been activated.</strong></td>
<td>Activate the destination device.</td>
<td></td>
</tr>
</tbody>
</table>

* *Messages are shown in all bold text.*
Central To Central Transfer

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>The Central To Central Without Device button is disabled.</td>
<td>The Transfer Failed list is full. The list stores a maximum of 5 failed transfers.</td>
<td>Delete a patient from this list before attempting to transfer.</td>
</tr>
<tr>
<td></td>
<td>A patient tile is not selected on the Main Screen.</td>
<td>Select the patient tile that is intended to be transferred.</td>
</tr>
<tr>
<td></td>
<td>The selected patient tile has a device assigned but is not monitoring a patient.</td>
<td>Select the patient tile that is intended to be transferred.</td>
</tr>
<tr>
<td></td>
<td>The selected patient tile has no device assigned.</td>
<td>Select the patient tile that is intended to be transferred.</td>
</tr>
<tr>
<td>The Failed Patient List is full. Delete a patient from this list before transferring.</td>
<td>The Transfer Failed list is full. The list stores a maximum of 5 failed transfers.</td>
<td>Delete a patient from this list before transferring.</td>
</tr>
<tr>
<td>Available central list is empty.</td>
<td>There are no central stations found on the network.</td>
<td>Contact the System Administrator for assistance.</td>
</tr>
<tr>
<td>The patient cannot be transferred because at least two of the following fields: First Name, Last Name or Patient ID has not been entered.</td>
<td>The appropriate information has not been entered into at least two of the three demographic fields (Last Name, First Name and Patient ID) on the Demographics tab.</td>
<td>Enter the appropriate information into at least two of the three demographic fields (Last Name, First Name, and Patient ID) on the Demographics tab.</td>
</tr>
<tr>
<td>Selecting the Done button on the Select Destination Panorama list box terminates the transfer and returns to the Transfer tab.</td>
<td>A Destination Panorama device was not selected from the list box before selecting the Done button.</td>
<td>Repeat the transfer procedure and ensure that a Destination Panorama device is selected from the Select Destination Panorama list box before selecting the Done button.</td>
</tr>
<tr>
<td>Transfer Out Rejected. Timeout at [the Destination Panorama name] Central.</td>
<td>The Accept/Reject Patient button on the Transfer tab at the Destination Panorama was not selected within 10 minutes of initiating the Central To Central Transfer Without Device.</td>
<td>Repeat the transfer procedure and ensure that the Accept/Reject Patient button on the Transfer tab at the Destination Panorama is selected within 10 minutes of initiating the transfer.</td>
</tr>
<tr>
<td>Transfer In Request from [the Source Panorama name] cancelled due to user timeout.</td>
<td>At the Destination Panorama, either the Accept or Reject button on the Transferred In Patient view of the Transfer tab was not selected within 10 minutes of initiating the Central To Central Transfer Without Device.</td>
<td>Repeat the transfer procedure and ensure that either the Accept or Reject button on the Transferred In Patient view of the Transfer tab is selected within 10 minutes of initiating the Central To Central Transfer Without Device.</td>
</tr>
<tr>
<td>Transfer In Request from [the Source Panorama name] rejected: Transfer In already in progress</td>
<td>A transfer in was initiated and accepted at the destination central station, and then another transfer in to the same destination central station is attempted.</td>
<td>Wait until the first transfer in is complete, then repeat transfer procedure.</td>
</tr>
<tr>
<td></td>
<td>The Accept/Reject Patient button is disabled.</td>
<td>A transfer request has not been received from any other Panorama Central Station.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
## Central To Central Transfer (Continued)

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>The destination tile must have a device assigned to perform the transfer.</td>
<td>The destination tile does not have a device assigned.</td>
<td>Select the OK button on the message box and then select a destination tile that has a device assigned.</td>
</tr>
<tr>
<td>Please touch tile to which you wish to transfer the patient.</td>
<td>A destination tile has not been selected at the Destination Panorama.</td>
<td>Select a destination tile that has a device assigned.</td>
</tr>
<tr>
<td>Failed Transfer Out: System Error. (Source Panorama)</td>
<td>The selected Destination Panorama device was not properly defined in the equipment list.</td>
<td>Repeat the transfer procedure and ensure that the desired Destination Panorama device is properly defined in the Equipment List. Determine the cause of the interruption and correct.</td>
</tr>
<tr>
<td>Failed Transfer In: System Error. (Destination Panorama)</td>
<td>The selected Destination Panorama device was not properly defined in the equipment list.</td>
<td>Repeat the transfer procedure and ensure that the desired Destination Panorama device is properly defined in the Equipment List. Determine the cause of the interruption and correct.</td>
</tr>
<tr>
<td>This Tab is in use by another display. Can not be updated from more than one display at the same time.</td>
<td>An attempt was made to open the Transfer tab on one display when it is already open on the other display.</td>
<td>Select the OK button and use the open the Transfer tab.</td>
</tr>
<tr>
<td>At the destination tile, the digital data is displayed but the waveform is not displayed</td>
<td>The destination tile was in Standby mode when the transfer completed.</td>
<td>Return to normal monitoring mode.</td>
</tr>
<tr>
<td>Patient ID number truncated.</td>
<td>More than 10 characters were entered in the Patient ID field at the WMTS 608 bedside device.</td>
<td>Do not enter more than 10 characters when using a WMTS 608 bedside device because only 10 characters will be transferred to the central station.</td>
</tr>
<tr>
<td>The destination tile is blank and all buttons on the Transfer tab are unavailable.</td>
<td>The destination tile was in a Communications Lost or a Comm Loss - Check Bedside Monitor state when the transfer completed.</td>
<td>Re-establish communication.</td>
</tr>
<tr>
<td>Waiting for Patient message is displayed in the heart rate digital tile.</td>
<td>The destination device has not been activated.</td>
<td>Activate the destination device.</td>
</tr>
<tr>
<td>Available monitor list is empty.</td>
<td>The Central To Central Transfer Without Device button was selected but there are no other Panorama devices on the network.</td>
<td>Try and transfer to another central station on a different network. If unsuccessful, contact the System Administrator for assistance.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
Recovering Failed Transfer Patient Data

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>The <strong>Recover Failed Transfer</strong> sidebar button is disabled.</td>
<td>The Failed Transfer List is empty.</td>
<td>No solution required.</td>
</tr>
<tr>
<td>Selecting the <strong>Recover</strong> button has no effect.</td>
<td>A patient has not been selected from the Failed Transfer List.</td>
<td>Select the patient whose information is to be recovered from the list and then select the <strong>Recover</strong> button.</td>
</tr>
<tr>
<td>Select the tile to place the recovered patient in</td>
<td>A destination tile on the Source Panorama was not selected before pressing the <strong>Recover</strong> button.</td>
<td>Select a destination tile on the Source Panorama.</td>
</tr>
<tr>
<td>The <strong>destination tile must have a device assigned to perform the transfer</strong></td>
<td>The destination tile does not have a device assigned.</td>
<td>Select the <strong>OK</strong> button to return to the Recover Failed Transfer Patients view. Select a destination tile that has a device assigned and continue the transfer procedure.</td>
</tr>
<tr>
<td>The <strong>Waiting for Patient</strong> message is displayed in the heart rate digital tile.</td>
<td>The destination device has not been activated.</td>
<td>Activate the destination device.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.

Deleting Failed Transfer Patient Data

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>The <strong>Delete Failed Transfer</strong> sidebar button is disabled.</td>
<td>The Failed Transfer List is empty.</td>
<td>No solution required.</td>
</tr>
<tr>
<td>Selecting the <strong>Delete</strong> button has no effect.</td>
<td>A patient has not been selected from the Failed Transfer List.</td>
<td>Select the patient whose information is to be deleted from the list and then select the <strong>Delete</strong> button.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.

Transferring Failed Transfer Patient Data

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>The <strong>Transfer From Failed List</strong> sidebar button is disabled.</td>
<td>The Failed Transfer List is empty.</td>
<td>No solution required.</td>
</tr>
<tr>
<td>Selecting the <strong>Transfer</strong> button has no effect.</td>
<td>A patient has not been selected from the Failed Transfer List.</td>
<td>Select the patient whose information is to be transferred from the list and then select the <strong>Transfer</strong> button.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
5.3 Discharge Tab

The **Discharge** tab (shown in FIGURE 5-25) is used to discharge a patient from a patient tile. If the required demographic information was previously entered for the patient to be discharged, the patient will be moved to the list of retrievable patients. For additional information regarding retrieving a patient, refer to “Retrieving Discharged Patients” on page 5-21.

The **Discharge** tab is not available at the ViewStation. Patient discharge can be performed at the host central station, the WorkStation, or the bedside monitor. Any change in the patient tile’s state will be communicated and changed at the host central station and bedside monitor, if applicable.

**NOTE:** The host central station(s) C-LAN configuration settings and the software installed at the remote location dictates the functionality available for each patient tile. For additional information, refer to “The ViewStation and the WorkStation” on page 2-4.

5.3.1 Accessing the Discharge Tab

1. From the menu bar, select the **Patient Setup** menu button. The tabs associated with patient setup are displayed.
2. Choose a patient tile by selecting its digital tile or waveform area.
3. Select the **Discharge** tab. The **Discharge** tab is enabled.

![Discharge Tab](image)

**FIGURE 5-25** Discharge Tab

5.3.2 Discharging a Patient from the Central Station and the Bedside Monitor

**NOTE:** Discharge from the bedside monitor and the central station or WorkStation can only be performed when the central station **Discharge at Bedside** button in the Unit Choices tab is set to **ON**. For additional information regarding the Unit Choices tab, refer to the “Unit Choices Tab” on page 9-50.

1. Select the tile of the patient to be discharged. Choose a patient tile by selecting its digital data or waveform area.
2. Select the **Bedside and Panorama Discharge** button (shown in FIGURE 5-25). One of two messages will be displayed:

- If the patient tile contains all the required demographic information (at least two of the three demographic fields, First Name, Last Name, Patient ID must be populated), the **Are you sure you want to discharge this patient? Selecting Yes will move the patient to the discharge list** dialog box is displayed.
  - The Yes button discharges the selected patient from the central station and the bedside monitor. It also moves the patient to the central station discharge list.
  - The No button cancels the discharge

- If the patient tile does not contain all the required demographic information (at least two of the three demographic fields, First Name, Last Name, Patient ID must be populated) the message **The patient's data and submitted print jobs will be deleted because at least two of the following fields: First Name, Last Name or Patient ID has not been filled in. Do you still want to proceed with the discharge?** is displayed.
  - The Yes button completes the discharge deleting all patient data and unprocessed print jobs
  - The No button cancels the discharge

5.3.3 Discharging a Patient from the Central Station

**NOTE:** Discharge from the central station is only available when the bedside monitor is in a Comm Loss - Check Bedside Monitor state.

1. Select the tile of the patient to be discharged. Choose a patient tile by selecting its digital data or waveform area.

2. Select the **Discharge Patient** button. The **Local Discharge Only! Patient will not be discharged at bedside. Do you still want to proceed with discharge?** message is displayed.

- If the Yes button is selected and at least two of the three demographic fields, First Name, Last Name, Patient ID are populated, the **Are you sure you want to discharge this patient? Selecting Yes will move the patient to the discharge list** message box will be displayed.
  - The Yes button completes the discharge and moves the patient to the central station discharge list
  - The No button cancels the discharge

- If the Yes button is selected and the patient tile does not contain all the required demographic information (at least two of the three demographic fields, First Name, Last Name, Patient ID must be populated) the message **The patient's data and submitted print jobs will be deleted because at least two of the following fields: First Name, Last Name or Patient ID has not been filled in. Do you still want to proceed with the discharge?** is displayed.
  - The Yes button completes the discharge deleting all patient data and unprocessed print jobs
  - The No button cancels the discharge
• The No button cancels the discharge

5.3.4 Discharging a Patient from a Telepack Device
The same messages described in “Discharging a Patient from the Central Station and the Bedside Monitor” on page 5-43 will be displayed, with one exception. If the Yes button is selected after the message Are you sure you want to discharge this patient? Selecting Yes will move the patient to the discharge list is displayed, the Telepack will be discharged from the central station. Discharging a Telepack device can only be done from the central station. Once discharged, all the patient information will be cleared.

5.3.5 Sidebar Button - Normal Screen
Select the Normal Screen sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

5.3.6 Troubleshooting
This section lists some of the potential messages and issues that may occur while using the Discharge tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>No patient Selected!</td>
<td>This message appears in the demographic field when either a patient tile has not been selected or an empty tile was selected.</td>
<td>Select an active patient tile.</td>
</tr>
</tbody>
</table>

The patient’s data and submitted print jobs will be deleted because at least two of the following fields: First Name, Last Name or Patient ID has not been filled in. Do you still want to proceed with the discharge?

The selected patient does not have at least two of the three demographic fields completed. The demographic fields include First Name, Last Name, and ID.

Select the No button to close the dialog, then enter the required demographic information.

The patient’s data and submitted print jobs will be deleted because at least two of the following fields: First Name, Last Name or Patient ID has not been filled in. Do you still want to proceed with the discharge? dialog box is open and was automatically closed without any user interaction.

The bedside monitor has gone into a Comm Loss state. Check Bedside Monitor state.

Restore the connection between the Panorama Central Station and the monitoring device. If unable to restore the connection, select the Discharge Patient button to discharge the equipment from the patient tile.

* Messages are shown in all bold text.
### Message/Issue *Reason* *Solution*

**Unable to perform operation on tile: Central to Central Transfer Out in progress.**
A Central to Central transfer Out has been initiated at the source central station but not accepted or rejected at the destination central station for the selected tile, and the **Discharge Patient** button is pressed in the **Discharge** tab at the source central station.
Wait until the Central to Central transfer is complete, then discharge the patient.

**Unable to perform operation on tile: Central to Central Transfer In in progress.**
A Central to Central transfer In is in progress for the selected tile and the selected tile is actively monitoring a patient, and the **Discharge Patient** button is pressed in the **Discharge** tab at the destination central station.
Wait until the Central to Central transfer is complete, then discharge the patient.

The **Discharge Patient** button in the **Discharge** tab is disabled.
The **Discharge** at Bedside setting in the **Unit Choices** tab is set to **OFF**.
If a patient discharge is necessary, it must be performed from the bedside monitor.

The **Discharge** tab needs to be refreshed.
Select another tab and then return to the **Discharge** tab.

No patient is selected.
Select a patient tile.

The **Discharge Patient** button in the **Discharge** tab is enabled.
The bedside monitor has gone into a Comm Loss - Check Bedside Monitor state.
Verify the connection between the Panorama Central Station and the monitoring device.

The **Discharge** at Bedside setting in the **Unit Choices** tab is set to **OFF**.
No solution required.

**Local Discharge Only! Patient will not be discharged at bedside. Do you still want to proceed with discharge?**
The bedside monitor is in a Comm Loss - Check Bedside Monitor state.
Verify the connection between the Panorama Central Station and the monitoring device.

A Telepack device is in use.
No solution required.

**Local Discharge Only! Patient will not be discharged at bedside. Do you still want to proceed with discharge?**
The bedside monitor is no longer in a Comm Loss - Check Bedside Monitor state.
If the **Discharge at Bedside** button in the **Unit Choices** tab is set to ON, proceed with the discharge by selecting the **Discharge Patient** button. If it is set to OFF, the discharge must be performed from the bedside.

*Messages are shown in all bold text.*
5.4 Display Tab

The Display tab (shown in FIGURE 5-26) is used to select the digital and waveform parameters that will appear in the selected patient tile. The default settings for patient tiles are defined by the Unit Choices tab, which is described in the “Unit Choices Tab” on page 9-50. The layout and functionality of the Display tab will change based on the type of monitoring device assigned to the selected patient tile.

The Display tab is available at the ViewStation and the WorkStation (shown in FIGURE 5-27), and functions in the same manner as it does at the host central station. When a patient tile is assigned at the ViewStation or the WorkStation, it will initially use the display settings of the associated patient tile at the host central station, provided the patient tile is in an active state.

After this initial transfer, changes made to the host central station, the ViewStation and/or the WorkStation patient tile Display tab will be independent of each other unless communication is interrupted. Automatic updating of the display settings at the remote workstation patient tile will occur if any of the following occur:

• The monitoring device is placed in the Standby mode at the central station
• The Telepack is not communicating with its host central station (Communications Lost)
• The patient monitor is not communicating with its host central station (Comm Loss - Check Bedside Monitor)
• Communication between the host central station and the ViewStation and/or the WorkStation is lost (Central Communications Lost)

NOTE: The host central station(s) C-LAN configuration settings and the software installed at the remote location dictates the functionality available for each patient tile. For additional information, refer to “The ViewStation and the WorkStation” on page 2-4.

5.4.1 Accessing the Display Tab

1. From the menu bar, select the Patient Setup menu button. The tabs associated with patient setup are displayed.
2. Choose a patient tile by selecting its digital tile or waveform area.
3. Select the Display tab. The Display tab is enabled.

FIGURE 5-26 Display Tab
5.4.2 Components of the Display Tab

Digital Parameter Data
The Panorama Central Station is capable of showing three to five digital data parameters in each patient tile.

NOTE: The Parameter list box is not displayed in the Display tab until a parameter tile is selected.
NOTE: When using a WMTS 608 wireless bedside device with a View 12™ card, all 12 leads will be available in the Parameter list, but only the waveforms for leads I, II, III, aVR, aVL, aVF, and V1 will be displayed in the patient tile.

Digital data tiles display a minimum of three and a maximum of five parameters depending on the parameters selected.

A digital data tile consists of three equal-size tiles. The Heart Rate (HR) parameter is always shown in the first tile. The remaining two tiles can each be divided into two sub-tile areas. If the associated device is a Telepack, the Battery Percentage parameter (shown in FIGURE 5-30) will occupy one sub-tile.

![Battery Percentage Digital Data Tile (Telepack) Image]

FIGURE 5-30 Battery Percentage Digital Data Tile (Telepack)

Parameters occupy 1 or 2 sub-tile areas. The number of digital data parameters that can be displayed is dependent on the type of parameters selected. For example, the NIBP parameter uses two data tiles while the Respiration parameter uses one data tile.

![NIBP and Respiration Digital Data Tiles Image]

FIGURE 5-31 NIBP and Respiration Digital Data Tiles

Table 5-4 shows the number of sub-tiles that each parameter occupies. For a list of the parameters listed by monitoring device, refer to “Supported Devices” on page 1-3. For additional information regarding the setup of a patient tile, refer to the “Display Tab” on page 5-47.

**TABLE 5-9**

Number of Sub-tiles Occupied by a Parameter

<table>
<thead>
<tr>
<th>PARAMETERS</th>
<th># OF SUB-TILES OCCUPIED</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heart Rate</td>
<td>2</td>
</tr>
<tr>
<td>NIBP</td>
<td>2</td>
</tr>
<tr>
<td>Respiration</td>
<td>1</td>
</tr>
<tr>
<td>SpO₂</td>
<td>1</td>
</tr>
<tr>
<td>IBP</td>
<td>2</td>
</tr>
<tr>
<td>Temperature</td>
<td>1</td>
</tr>
<tr>
<td>CO₂</td>
<td>2</td>
</tr>
<tr>
<td>O₂</td>
<td>2</td>
</tr>
<tr>
<td>N₂O₂</td>
<td>2</td>
</tr>
<tr>
<td>Agent</td>
<td>2</td>
</tr>
</tbody>
</table>
NOTE: Digital data is shown in the color configured for the parameter. For additional information, refer to the “Parameter Color Tab” on page 9-2.

Heart Rate Tile

The Heart Rate tile (shown in FIGURE 5-32) shows the heart rate measurement, the lead through which the measurement is acquired and icons for Arrhythmia/ST analysis.

FIGURE 5-32 Heart Rate Digital Data Tile

The Heart Rate tile follows additional digital data rules.

For example, the Heart Rate tile:

• Always occupies the first digital data tile
• Displays an ECG lead label for the lead acquiring the data
• Displays an \[ \text{A} \] icon if arrhythmia analysis is enabled
• Displays an \[ \text{S} \] icon if ST analysis is enabled
• Displays the standby location for a patient in Standby mode

Tile Display Rearrangement

Depending on the number of parameters selected, a blank tile may be displayed in the Display tab. A blank tile will not be displayed on the Main Screen because the system will automatically populate the empty sub-tiles based on the type of monitoring device:

<table>
<thead>
<tr>
<th>PARAMETERS</th>
<th># OF SUB-TILES OCCUPIED</th>
</tr>
</thead>
<tbody>
<tr>
<td>PVC/min</td>
<td>1</td>
</tr>
<tr>
<td>Pause/min</td>
<td>1</td>
</tr>
<tr>
<td>ST</td>
<td>1</td>
</tr>
<tr>
<td>CO/CI</td>
<td>2</td>
</tr>
<tr>
<td>CCO/CCI</td>
<td>2</td>
</tr>
<tr>
<td>CCO STAT/CCI STAT</td>
<td>2</td>
</tr>
<tr>
<td>SVR</td>
<td>1</td>
</tr>
<tr>
<td>SVRI</td>
<td>1</td>
</tr>
<tr>
<td>SvO₂</td>
<td>1</td>
</tr>
<tr>
<td>ScvO₂</td>
<td>1</td>
</tr>
<tr>
<td>PAWP</td>
<td>1</td>
</tr>
<tr>
<td>BIS</td>
<td>1</td>
</tr>
<tr>
<td>QT</td>
<td>2</td>
</tr>
<tr>
<td>Battery Percentage</td>
<td>1</td>
</tr>
</tbody>
</table>
• **Telepack**: the parameters in the Main Screen tile will automatically rearrange to remove the blank tiles.

• **Bedside**: the parameters in the Main Screen tile will automatically rearrange to compress user selections with the remaining digital data tiles populating based on the next parameter available in the **Unit Priorities** tab.

### Waveform Parameter Data

The patient tile is capable of showing up to two waveforms for a patient (shown in FIGURE 5-33).

**NOTE:** The initial central station wave default settings come from the System Wave Default settings, if settings were configured.

![FIGURE 5-33 Display Tab - Waveform Data](image)

The first waveform is always an ECG waveform. The second waveform can be any other waveform parameter available for the associated device or, if desired, it can be turned off. Each waveform has its own baseline.

**NOTE:** Waveforms display in the color that was selected for the parameter. For additional information, refer to the "Parameter Color Tab" on page 9-2.
The following table shows the available waveform parameters.

<table>
<thead>
<tr>
<th>WAVEFORM PARAMETER</th>
<th>AVAILABLE WAVEFORMS</th>
</tr>
</thead>
<tbody>
<tr>
<td>ECG Lead</td>
<td>I, II, III, aVR, aVL, aVF, V, V1, V2, V3, V4, V5, V6</td>
</tr>
<tr>
<td></td>
<td>• The ECG waveforms available are dependent on the lead set used.</td>
</tr>
<tr>
<td></td>
<td>• ECG waveforms use the Pacer Enhancement and Filter Settings that were selected in the <strong>Demographics</strong> tab for the selected patient tile.</td>
</tr>
<tr>
<td>Waveform 2</td>
<td>I, II, III, aVR, aVL, aVF, V, V1, V2, V3, V4, V5, V6, Resp, SpO₂, IBP1, IBP2, IBP3, IBP4, CO₂, O₂, Agent (Iso, Enf, Sev, Hal, Des)</td>
</tr>
<tr>
<td></td>
<td>• The ECG waveforms available are dependent on the lead set used. The waveform chosen for the ECG Lead waveform will not be included in the parameter list because it cannot be used in both waveform positions.</td>
</tr>
<tr>
<td></td>
<td>• The remaining parameters available for the second waveform are dependent on the monitoring device associated with the patient tile.</td>
</tr>
</tbody>
</table>

### 5.4.3 Configuring the Display Tab

This section describes how to configure parameter tiles in the **Display** tab.

ViewStation and WorkStation patient tiles will initially display the same display settings of the associated patient tile at the host central station. For additional information regarding the transfer of display settings from the host central station, refer to the “**Trends Tab**” on page 7-14.

#### Selecting a Patient’s Parameter Data

1. Choose a patient tile by selecting its digital data or waveform area.
2. Select the parameter tile to be modified. The Parameter list is displayed.
3. From the Parameter list, select the parameter to be viewed or turn the parameter OFF.

**NOTE:** Digital parameters and the first waveform parameter cannot be turned off.

**NOTE:** The OFF option is only available in the Parameter list when Waveform 2 is selected in the Waveform Display area. When OFF is selected, the message “Are you sure you want to turn off the parameter” is displayed.

- Digital Parameters include Heart Rate, Respiration, IBP, NIBP, Temp, ST, PVC/Min, Pause/Min, SpO₂, N₂O, CO₂, O₂, Cardiac Output (CO), Cardiac Index (CI), Continuous Cardiac Output (CCO), Continuous Cardiac Index (CCI), System Vascular Resistance (SVR), System Vascular Resistance Index (SVRI), SvO₂, ScvO₂, T Blood, Delta T, Pulmonary Artery Wedge Pressure (PAWP) (not supported in DPM6/7, Passport 12M/Passport 17M, Passport 8/Passport 12, or T1 monitor), BIS (for V 12/ V 21 only), QT (for V 12/V 21 and Telepack only), Battery Percentage (for a Telepack device only), and Agent.

**NOTE:** The CO/CI, Temp 2, and T Blood parameters are only available if they are being monitored by the DPM6/DPM 7, Passport 12M/Passport 17M, Passport 8/Passport 12, V 12/V 21, Spectrum/Spectrum OR External Parameter Module (EPM). These parameters are not available if they are monitored by an Edwards Vigilance® monitor.
Patient Setup Functions

Display Tab

NOTE: The CCO/CCI, SVR, SVRI, CCO STAT/CCI STAT, SvO₂, ScvO₂, Temp 3 and BIS parameters are only available if they are being monitored by the V Series.

- Waveform Parameters include I, II, III, aVR, aVL, aVF, V, V1, V2, V3, V4, V5, and V6.

4. Choose the **Select** button to highlight the selected parameter. The parameter is shown in the selected patient tile.

NOTE: A digital parameter can only be shown in one parameter tile at any given time.

5.4.3.1 Configuring a Trend Display Interval

Select the **Trend Display** sidebar button to configure the interval at which a patient’s trend data will be displayed at the Panorama Central Station, ViewStation, and the WorkStation. Regardless of which Trend Display interval is selected, NIBP, CO/CI, and PAWP trend records will be displayed.

When a patient tile is assigned at the ViewStation or the WorkStation, it will initially display the Trend Display interval of the associated host central station patient tile, provided the patient tile is communicating with the central station. The **Trend Display** sidebar button is available at the ViewStation and the WorkStation and it may be used in the same manner as it is at the central station. For additional information regarding the transfer of trend interval settings from the host central station, refer to the “Trends Tab” on page 7-14.

The default setting for a Trend Display interval is determined by the **Unit Choices** tab. For additional information regarding the **Unit Choices** tab, refer to “Unit Choices Tab” on page 9-50.

NOTE: Trends data is automatically stored in one minute intervals. The Trend Display interval selected does not affect how trend information is stored in the database, just how it is displayed.


To select the Trend Display interval:

1. From the **Display** tab, select the **Trend Display** sidebar button. The Trend Display dialog box (shown in FIGURE 5-35) is displayed.

![FIGURE 5-35 Trend Display Dialog Box](image)
2. Select the Interval button until the desired interval period is displayed. Interval periods include 1 minute, 5 minutes, 10 minutes, 15 minutes, 30 minutes, 1 hour, and 2 hours.

3. Select the Done or the Cancel button.
   - The Done button accepts the selection.
   - The Cancel button discards the selection, closes the dialog box, and returns to the Display tab.

With the exception of NIBP, CO/CI, and PAWP, selecting the Done button will change the Trend Display interval in the Trends tab based on the standard clock time intervals. Refer to the following table:

<table>
<thead>
<tr>
<th>INTERVAL</th>
<th>TREND RECORDS DISPLAYED IN TRENDS TAB</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 minute</td>
<td>All trend records will be displayed.</td>
</tr>
<tr>
<td>5 minute</td>
<td>All trend records having a timestamp of 0, 5, 10, 20, 25, 30, 35, 40, 45, 50, and 55 minutes will be displayed.</td>
</tr>
<tr>
<td>10 minute</td>
<td>All trend records having a timestamp of 0, 10, 20, 30, 40, and 50 minutes will be displayed.</td>
</tr>
<tr>
<td>15 minute</td>
<td>All trend records having a timestamp of 0, 15, 30, and 45 minutes will be displayed.</td>
</tr>
<tr>
<td>30 minutes</td>
<td>All trend records having a timestamp of 0, and 30 minutes will be displayed.</td>
</tr>
<tr>
<td>1 hour</td>
<td>All trend records having a timestamp of 0 minutes (for example, 12:00, 1:00, 2:00) will be displayed.</td>
</tr>
<tr>
<td>2 hour</td>
<td>All trend records having a timestamp of 0 minutes for even hours only (for example, 12:00, 2:00, 4:00) will be displayed.</td>
</tr>
</tbody>
</table>
5.4.4 Sidebar Buttons
The following sidebar buttons are used to expand the functionality of this tab.

Restore Previous Settings
Select the **Restore Previous Settings** sidebar button to return patient display settings to those in use when the tab was opened.

Trend Display
This sidebar button was previously described on page 5-53.

Disable ECG
Disable ECG is available on version 8.7.1 and above. See page 9-53 for setting the system default for displaying the **Disable ECG** sidebar button.

If the system has been configured, the **Disable ECG** sidebar button is displayed when a Telepack device with an SpO₂ module is connected. Select the **Disable ECG** sidebar button to disable the display of ECG waveform(s) for the patient(s) if the current Heart Rate source is not ECG.

The Disable ECG sidebar button is disabled if the patient is in Standby mode, during communications loss, or if the Telepack does not have an SpO₂ module connected.

**NOTE:** If the Disable ECG sidebar button is latched and the Heart Rate source changes to ECG, the Disable ECG sidebar button will be unlatched and the ECG waveform(s) will be re-enabled.

Normal Screen
Select the **Normal Screen** sidebar button to return the display to the full screen mode, and close all tabs, message boxes and dialog boxes.

5.4.5 Troubleshooting
This section lists some of the potential messages and issues that may occur while using the **Display** tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>No Patient Selected!</td>
<td>This message is displayed in the demographic field when either a patient tile has not been selected or an empty tile was selected.</td>
<td>Select an active patient tile.</td>
</tr>
<tr>
<td>V2, V3, V4, V5, or V6 was selected from the Parameter list but the waveform is not displayed in patient tile.</td>
<td>A WMTS 608 wireless bedside device with a View 12 card is in currently in use. When using this device type, the V2-V6 waveforms will not be displayed.</td>
<td>Select a different waveform from the Parameter list. Change to a hardwired or 2.4 GHz wireless device.</td>
</tr>
<tr>
<td>The parameter tiles in the <strong>Display</strong> tab are disabled.</td>
<td>A valid patient tile is not selected.</td>
<td>Select an active patient tile.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
5.5 Standby Tab

The Standby tab (shown in FIGURE 5-36) is used to place a patient in Standby mode at the Panorama Central Station, the WorkStation or the bedside monitor. Standby mode temporarily suspends all patient monitoring, alarm processing, and trending activity.

The Standby tab is available at the WorkStation and it may be used in the same manner as it is at the central station. Any change in the patient tile’s state will be communicated and changed at the host central station and bedside monitor, if applicable.

NOTE: Patients placed in the Standby mode should be closely monitored.

NOTE: The host central station(s) C-LAN configuration settings and the software installed at the remote location dictates the functionality available for each patient tile. For additional information, refer to “The ViewStation and the WorkStation” on page 2-4.

5.5.1 Accessing the Standby Tab

1. From the menu bar, select the Patient Setup menu button. The tabs associated with patient setup are displayed.

2. Choose a patient tile by selecting its digital tile or waveform area.

3. Select the Standby tab. The Standby tab is enabled.

![FIGURE 5-36 Standby Tab](image)

5.5.2 Placing a Patient in Standby Mode

This section outlines placing a patient in the Standby mode from the Panorama Central Station.

1. Choose a patient tile by selecting its digital data or waveform area.

2. From the menu bar, select the Patient Setup menu button. The tabs associated with patient setup are displayed.

3. Select the Standby tab. The Standby tab is displayed.

4. Select the type of Standby mode to be used for the patient.

   a. Select the Panorama Standby Only/Standby button to discontinue patient monitoring at central station but continue monitoring the patient at the bedside.
NOTE: If using a Telepack device, the Panorama Standby Only button will be labeled Standby.

b. Select the Bedside and Panorama Standby button to discontinue monitoring the patient at the central station and at the bedside monitor.

The Patient Location list box is displayed (shown in FIGURE 5-37).

![FIGURE 5-37 Standby Tab with Patient Location List displayed](image)

5. Select a patient location from the Patient Location list box, which includes:
   - Standby
   - Cath Lab
   - X-ray
   - Therapy
   - Bath
   - ER
   - In Transit
   - OR

6. Choose the Select button to confirm the Standby location selected. The text for the standby location is shown in the patient tile, and all monitoring, alarm processing and trend storage is suspended.

5.5.3 Removing a Patient from Standby Mode

To remove a patient from Standby mode:

1. From the menu bar, select the Patient Setup menu button. The tabs associated with patient setup are displayed.

2. Select the Standby tab. The Standby tab is displayed.
   - If the patient is in the Panorama Standby Only mode, select the Resume Monitoring button.

   The message Are you sure that you want to return the patient from Panorama standby is displayed. Select the YES button to return the patient from standby mode and resume monitoring or the NO button to keep the patient in Standby mode.
   - If the patient is in the Bedside and Panorama Standby mode, select the STANDBY button on the bedside monitor.

5.5.4 Sidebar Button - Normal Screen

Select the Normal Screen sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.
### 5.5.5 Troubleshooting

This section lists some of the potential messages and issues that may occur while using the **Standby** tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>No Patient Selected!</td>
<td>This message is displayed in the demographic field when either a patient tile has not been selected or an empty tile was selected.</td>
<td>Select an active patient tile.</td>
</tr>
<tr>
<td>A patient in bedside standby cannot be directly placed into Panorama only standby</td>
<td>This message is displayed when the selected patient is already in <strong>Standby</strong> mode at the bedside monitor or was just released from Standby mode at the bedside monitor.</td>
<td>Select the OK button to close the dialog box. Resume monitoring at the bedside monitor before trying to change standby locations. Wait a moment and try to put the patient in <strong>Standby</strong> mode again.</td>
</tr>
<tr>
<td>Patient in auto standby may only be released from bedside</td>
<td>This message is displayed when the <strong>Bedside and Panorama Standby</strong> button is selected for a patient that is already in <strong>Standby</strong> mode at the Panorama Central Station and the Bedside monitor.</td>
<td>Select the OK button to close the dialog box. Release the patient from <strong>Standby</strong> mode at the bedside monitor before trying to change the patient's <strong>Standby</strong> mode.</td>
</tr>
<tr>
<td>The <strong>Standby</strong> tab for the selected patient tile is disabled at the WorkStation.</td>
<td>The central station from where the patient tile was networked is configured as View Only.</td>
<td>No solution required.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
5.6 Wave Gain Tab (Patient)

The **Wave Gain** tab (shown in FIGURE 5-38) is used to change the wave gain settings for the selected patient’s available waveforms. The layout, functionality, and buttons available in this tab will change based on the type of monitoring device assigned to the selected patient tile.

**NOTE:** Changing the ECG wave gain does not affect the arrhythmia algorithm.

The **Wave Gain** tab is available at the ViewStation and the WorkStation, and functions in the same manner as it does at the host central station. When a patient tile is assigned at the ViewStation or the WorkStation, it will initially display the wave gain settings of the associated host central station patient tile, provided the patient tile is communicating with the central station.

After this initial transfer, changes made to the host central station, the ViewStation and/or the WorkStation patient tile **Wave Gain** tab will be independent of each other unless communication is interrupted. Automatic updating of the wave gain settings at the remote workstation patient tile will occur if any of the following occur:

- The monitoring device is placed in the Standby mode at the central station
- The Telepack is not communicating with its host central station (Communications Lost)
- The patient monitor is not communicating with its host central station (Comm Loss - Check Bedside Monitor)
- Communication between the host central station and the ViewStation and/or the WorkStation is lost (Central Communications Lost)

The default host central station wave gain settings are defined by the System **Wave Gain** tab, which is described in the “Wave Gain Tab (System)” on page 9-61.

5.6.1 Accessing the Wave Gain Tab (Patient)

1. From the menu bar, select the **Patient Setup** menu button. The tabs associated with patient setup are displayed.
2. Choose a patient tile by selecting its digital tile or waveform area.
3. Select the **Wave Gain** tab. The **Wave Gain** tab is enabled.

![FIGURE 5-38 Wave Gain Tab](image)
5.6.2 Changing a Patient's Wave Gain Settings

This section outlines entering and modifying a patient's wave gain settings using the **Wave Gain** tab.

ViewStation and WorkStation patient tiles will initially display the same wave gain settings of the associated patient tile at the host central station. For additional information regarding the transfer of wave gain settings from the host central station, refer to "Wave Gain Tab (Patient)" on page 5-59.

1. Choose a patient tile by selecting its digital data or waveform area.
2. In the **Wave Gain** tab, select the **ECG mm/mV** button until the desired wave gain displays. The settings available for the ECG wave gain include 2.5, 5.0, 10.0, 20.0, and 30.0 mm/mV.
3. Select the **IBPx mmHg** buttons until the desired wave gain displays. The settings available for the IBPx wave gain include 0-20, 0-40, 0-80, 60-140, 0-160, 0-225, 0-320, and (-10) -10 mmHg.

**NOTE:** If the IBPx button(s) have been labeled at the bedside monitor, the same label(s) will be displayed on the individual wave gain button(s).

4. Select the **Resp Gain** button until the desired wave gain is displayed. The settings available for the resp gain wave gain include 1, 2, 3, 4, and 5.
5. Select the **CO2 (%)**, **kPa**, **mmHG**) button until the desired wave gain is displayed.
   - The settings available for the CO2 % wave gain include 5.0, 7.5, and 10.0 %.
   - The settings available for the CO2 kPa wave gain include 5.0, 7.5, and 10.0 kPa.
   - The settings available for the CO2 mmHg wave gain include 40, 60, and 100 mmHg.
6. Select the **O2 %** button until the desired wave gain is displayed. The settings available for the O2 wave gain include 18 - 30, 18 - 60, and 18 - 100 %.
7. Select the **AGENT %** button until the desired wave gain is displayed. The settings available for the agent wave gains include 0 - 1.0, 0 - 2.5, 0 - 5.0, 0 - 10.0, 0 - 15.0, and 0 - 20.0 %.
8. Select the **Done** button to accept the wave gain settings shown in the Wave Gain tab. A patient’s wave gain settings will not be changed unless the **Done** button is selected.
5.6.3 Sidebar Buttons
The following sidebar buttons are used to expand the functionality of this tab.

Restore Previous Settings
Select the **Restore Previous Settings** sidebar button to return patient display settings to those in use when the tab was opened.

Normal Screen
Select the **Normal Screen** sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

5.6.4 Troubleshooting
This section lists some of the potential messages and issues that may occur while using the patient Wave Gain tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>No Patient Selected!</td>
<td>This message is displayed in the demographic field when either a patient tile has not been selected or an empty tile was selected.</td>
<td>Select an active patient tile.</td>
</tr>
<tr>
<td>The patient’s wave gain settings did not change</td>
<td>The <strong>Done</strong> button was not selected after the patient’s wave gain settings were changed.</td>
<td>Adjust the patient’s wave gain settings and select the <strong>Done</strong> button when the settings are completed.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
5.7 Print Setup Tab (Patient)

The **Print Setup** tab (shown in FIGURE 5-40) is used to select the waveforms that will be printed when an All Strips, Event Zoom-In, or Full Disclosure Zoom-In report is generated for the selected patient tile. The waveforms included in this tab will change based on the parameters in use and the type of monitoring device that is assigned to the selected patient tile. The default settings for the waveform portion of the Print Setup tab are defined by the system **Print Setup** tab, which is described in the “Print Setup Tab (System)” on page 9-45.

The **Print Setup** tab is available at the ViewStation and the WorkStation, and functions in the same manner as it does at the host central station. When a patient tile is assigned at the ViewStation or the WorkStation, it will initially display the print setup configuration settings of the associated host central station patient tile, provided the patient tile is communicating with the central station.

After this initial transfer, changes made to the host central station, the ViewStation and/or the WorkStation patient tile **Print Setup** tab will be independent of each other unless communication is interrupted. Automatic updating of the print setup configuration settings at the remote workstation patient tile will occur if any of the following occur:

- The monitoring device is placed in the Standby mode at the central station
- The Telepack is not communicating with its host central station (Communications Lost)
- The patient monitor is not communicating with its host central station (Comm Loss - Check Bedside Monitor)
- Communication between the host central station and the ViewStation and/or the WorkStation is lost (Central Communications Lost)

5.7.1 Accessing the Print Setup Tab

1. From the menu bar, select the **Patient Setup** menu button. The tabs associated with patient setup are displayed.

2. Choose a patient tile by selecting its digital tile or waveform area.

3. Select the **Print Setup** tab. The **Print Setup** tab is enabled.

![FIGURE 5-40 Patient Print Setup Tab (Central Station)]
5.7.2 Changing a Patient's Print Setup

This section outlines entering and modifying a patient's print setup via the Print Setup tab.

Customizing a Patient’s Print Setup

1. Choose a patient tile by selecting its digital data or waveform area.

2. In the Print Setup tab, select a check box button to enable/disable the printing of a parameter. The patient's print setup settings are instantly changed.

   The Parameters available in the Print Setup tab are dependent on the lead set and/or monitoring device connected to the patient.

   • A check mark in a parameter check box indicates that the parameter is selected and will print the patient's All Strip Report, Event Zoom-In, and Full Disclosure Zoom-In Reports.

   • An empty parameter check box indicates that the parameter is not selected and will not print the patient’s All Strip Report, Event Zoom-In, and Full Disclosure Zoom-In Reports.

5.7.3 Sidebar Buttons

The following sidebar buttons are used to expand the functionality of this tab.

Restore Previous Settings

Select the Restore Previous Settings sidebar button to return patient display settings to those in use when the tab was opened.
Normal Screen
Select the **Normal Screen** sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

5.7.4 Troubleshooting
This section lists some of the potential messages and issues that may occur while using the patient **Print Setup** tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>No Patient Selected!</strong></td>
<td>This message appears in the demographic field when either a patient tile has not been selected or an empty tile was selected.</td>
<td>Select an active patient tile.</td>
</tr>
<tr>
<td>A waveform does not print</td>
<td>The waveform is not selected in the patient's <strong>Print Setup</strong> tab.</td>
<td>Select the waveform in the patient's <strong>Print Setup</strong> tab.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
6.0 Patient Alarm Functions

This section describes the configuration of patient alarms at the Panorama Central Station.

- Patient Alarm Setup Tab (Optional)
- Alarm Limits Tab (Patient)
- Alarm Responses Tab (Patient)
6.1 Patient Alarm Setup Tab (Optional)

The Patient Alarm Setup tab (shown in FIGURE 6-1) provides password protected access to the patient alarm configuration. When the correct password is entered, the Alarm Limits and the Alarm Responses tabs are opened.

The Patient Alarm Setup tab is only shown if the Password Protection option is set to ON in the System Alarms tab. If Password Protection is OFF, the Patient Alarms menu button provides direct access to the Alarm Limits and the Alarm Responses tabs. For additional information, refer to the “System Alarms Tab” on page 9-14.

The Patient Alarm Setup tab is available at the central station and the WorkStation.

6.1.1 Accessing the Patient Alarm Setup Tab

From the menu bar, select the Patient Alarms menu button. The Patient Alarm Setup tab is displayed.

![FIGURE 6-1 Patient Alarm Setup Tab](image)

6.1.2 Entering a Password

To enter a password:

1. Select the Password text box in the Patient Alarm Setup tab. The keyboard dialog box is displayed.

2. Use the keyboard dialog box to enter the Alarm password that was created in the Passwords tab (maximum 15 characters). For security purposes asterisks (*) are shown for each typed character. For additional information regarding passwords, refer to the “Passwords Tab” on page 9-29 or contact your System Administrator.

**NOTE:** Passwords are case sensitive.

- The Backspace (<==) button clears a single character in the edit box
- The Caps Lock button enables/disables the caps lock function. A Caps Lock indicator is shown when Caps Lock is enabled.
- The Close button discards the changes
- The Enter button accepts changes
3. The **Alarm Limits** (shown in FIGURE 6-2) and the **Alarm Responses** (shown in FIGURE 6-6) tabs are shown after the correct password is entered. For additional information about alarm limits and alarm responses, refer to “Alarm Limits Tab (Patient)” on page 6-4 and “Alarm Responses Tab (Patient)” on page 6-13.

6.1.3 Sidebar Buttons
The following sidebar button is used to expand the functionality of this tab.

**Normal Screen**
Select the **Normal Screen** sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

6.1.4 Troubleshooting
This section lists some of the potential messages and issues that may occur while using the **Patient Alarm Setup** tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Incorrect password. Please try again.</td>
<td>The wrong password was entered in the Password text box.</td>
<td>Select the OK button to close the dialog box. Enter the correct password in the text box.</td>
</tr>
<tr>
<td></td>
<td>The password was entered in the wrong case.</td>
<td>Re-enter the password turning on/off the keyboard dialog Caps Lock key.</td>
</tr>
<tr>
<td>Incorrect password. This is your last chance. Please try again</td>
<td>The wrong password was entered two consecutive times in the Password text box.</td>
<td>Select the OK button to close the dialog box. Enter the correct password in the text box.</td>
</tr>
<tr>
<td></td>
<td>The password was entered in the wrong case.</td>
<td></td>
</tr>
<tr>
<td>The Patient Alarm Setup tab closes and the central station is returned to a full screen monitoring display.</td>
<td>The wrong password was entered three consecutive times in the Password text box.</td>
<td>Reopen the Patient Alarm Setup tab and enter the correct password in the text box.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
6.2 Alarm Limits Tab (Patient)

The patient Alarm Limits tab (shown in FIGURE 6-2) adjusts patient alarm limit settings. The alarm limits available will change according to the type of monitoring device assigned to the selected patient tile. The fields discussed in this section may not apply to all users. For additional information regarding alarm limit settings, refer to “Alarms and Events” on page 13-1.

The patient Alarm Limits tab is available at the ViewStation (shown in FIGURE 6-3) but is for viewing purposes only. When a patient tile is assigned at the ViewStation, it will always display the alarm limit settings of the associated host central station patient tile.

The patient Alarm Limits tab is available at the WorkStation and functions in the same manner as it does at the host central station but contains fewer sidebar buttons. Similar to the central station, alarm limit changes made at the WorkStation will be sent to the central station, View Only Workstation, and bedside monitor (if applicable).

Automatic updating of the alarm limit settings at the remote workstation will also occur when:

- The monitoring device is placed in the Standby mode at the central station
- The Telepack is not communicating with its host central station (Communications Lost)
- The patient monitor is not communicating with its host central station (Comm Loss - Check Bedside Monitor)
- Communication between the host central station and the ViewStation and/or the WorkStation is lost (Central Communications Lost)

WARNING: Changing the patient alarm limits at the WorkStation will automatically change the patient alarm limits at the Panorama Central Station AND the bedside monitor (if applicable) for the selected patient tile.

NOTE: The host central station(s) C-LAN configuration settings and the software installed at the remote location dictates the functionality available for each patient tile. For additional information, refer to “The ViewStation and the WorkStation” on page 2-4.

6.2.1 Accessing the Alarm Limits Tab

From the menu bar, select the Patient Alarms button. If the Patient Alarm Setup tab is displayed, enter the password in the Password text box. The Alarm Limits and Alarm Responses tabs are displayed.
6.2.2 Configuring Patient Alarm Limits

The following procedure is used for modifying patient alarm limits in the **Alarm Limits** tab. The order in which the alarm slider controls are shown in the **Alarm Limits** tab is selected in the “Unit Priorities Tab” on page 9-47.

ViewStation patient tiles will always display the same patient alarm limit settings of the associated patient tile at the host central station. For additional information regarding the transfer of alarm limit settings from the host central station, refer to “Alarm Limits Tab (Patient)” on page 6-4.

**NOTE:** The alarm slider controls in the patient Alarm Limits tab are disabled at the ViewStation.

1. Choose a patient tile by selecting its digital data or waveform area.
2. Use the scroll bar or the arrow buttons to locate the parameter to be modified.
   - Double arrow buttons display the next/previous page of parameter slider controls (until the first/last page is reached).
   - Single arrow buttons display the next/previous slider control (until the last control is shown).

3. Set the **High Alarm Limit** button for the selected parameter. This button is either labeled **OFF** or contains an alarm limit value. A high alarm limit is set using the up/down arrow buttons on the parameter’s vertical scroll bar or by moving the alarm’s slider control.

4. Set the **Low Alarm Limit** button for the selected parameter. This button is either labeled **OFF** or contains an alarm limit value. A low alarm limit is set using the up/down arrow buttons on the parameter’s vertical scroll bar or by moving the alarm’s slider control.

Alarm limit adjustments made at the Panorama Central Station are automatically sent to the monitoring device.

For additional information regarding the default alarm limit ranges, refer to “Alarms and Events” on page 13-1.

**Auto Set Alarms Button**

The **Auto Set Alarms** button automatically recalculates alarm limit settings to the percent indicated (shown in Table 6-1) above and below the current measured value.

**NOTE:** The Auto Set Alarms button does not affect the SpO₂, ST, PVC/min, QTc, Pause/min, CCO, CCI, SvO₂, SVR, SVRI, BIS, all arrhythmias or Apnea Delay settings.

**NOTE:** The Auto Set Alarms button is not available at the ViewStation.

**TABLE 6-1**  Auto Set Percent Indicator

<table>
<thead>
<tr>
<th>ALARM</th>
<th>PERCENT INDICATOR (APPROXIMATE)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heart Rate</td>
<td>30%</td>
</tr>
<tr>
<td>Respiration</td>
<td>30%</td>
</tr>
<tr>
<td>IBP Systolic</td>
<td>20%</td>
</tr>
<tr>
<td>IBP Diastolic</td>
<td>20%</td>
</tr>
<tr>
<td>IBP Mean</td>
<td>20%</td>
</tr>
<tr>
<td>NIBP Systolic</td>
<td>20%</td>
</tr>
<tr>
<td>NIBP Diastolic</td>
<td>20%</td>
</tr>
<tr>
<td>NIBP Mean</td>
<td>20%</td>
</tr>
<tr>
<td>ET CO₂ (mmHg)</td>
<td>20%</td>
</tr>
<tr>
<td>INSP CO₂ (mmHg)</td>
<td>20%</td>
</tr>
<tr>
<td>ET CO₂ (%)</td>
<td>20%</td>
</tr>
<tr>
<td>INSP CO₂ (%)</td>
<td>20%</td>
</tr>
<tr>
<td>ET CO₂ (kPa)</td>
<td>20%</td>
</tr>
<tr>
<td>INSP CO₂ (kPa)</td>
<td>20%</td>
</tr>
<tr>
<td>Temp (°F)</td>
<td>3%</td>
</tr>
</tbody>
</table>
6.2.3 Simultaneous Alarm Limit Editing

If using a bedside device with the Panorama Central Station, alarm limit and alarm response configuration changes cannot be made at the bedside monitor, central station, and WorkStation simultaneously. The bedside device will take precedence, followed by the central station, ending with the WorkStation. When an attempt is made to make changes simultaneously, the message **These settings are currently being edited remotely.** will be displayed at the central station or WorkStation, depending on where the change was initiated. If there are multiple WorkStations on the network, the first WorkStation that initiates the change will take precedence over any other WorkStation attempting to implement the same change.

These are the scenarios in which this message will be displayed:

- The Alarm Limits Menu is active at the bedside monitor and then the patient **Alarm Limits** tab is opened at the central station or the WorkStation
- The **Alarm Limits** tab is active at the central station or the WorkStation and then the Alarm Limits Menu is opened at the bedside monitor

In both of these scenarios, unsaved changes made in the **Alarm Limits** tab will be discarded.

To close this message dialog, select the **OK** button. Acknowledging this message will convert the **Alarm Limits** tab at the central station to a read-only state and editing will not be permitted. Once the Alarm Limits Menu is closed at the bedside device, the **Alarm Limits** tab will become active again and editing can resume.

For additional information regarding the bedside monitor menus, refer to the bedside monitor’s Operating Instructions:

---

**TABLE 6-1** Auto Set Percent Indicator (continued)

<table>
<thead>
<tr>
<th>ALARM</th>
<th>PERCENT INDICATOR (APPROXIMATE)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temp (°C)</td>
<td>3%</td>
</tr>
<tr>
<td>Delta Temp (°F)</td>
<td>25%</td>
</tr>
<tr>
<td>Delta Temp (°C)</td>
<td>25%</td>
</tr>
<tr>
<td>Temp Blood (°F)</td>
<td>3%</td>
</tr>
<tr>
<td>Temp Blood (°C)</td>
<td>3%</td>
</tr>
<tr>
<td>Extreme Bradycardia</td>
<td>10% below the low heart rate</td>
</tr>
<tr>
<td>Extreme Tachycardia</td>
<td>10% above the high heart rate</td>
</tr>
</tbody>
</table>
• Passport 2 (P/N 0070-00-0649-XX)
• Passport V (P/N 0070-00-0704-XX)
• Spectrum (P/N 0070-00-0648-XX)
• Spectrum OR (P/N 0070-00-0670-XX)
• DPM 6 (P/N H-046-000184-00)
• DPM 7 (P/N H-046-000186-00)
• V 12/V 21 (P/N 0070-00-0727-XX)
• Passport 12M/Passport 17M (P/N 046-005013-00)
• Passport 8/Passport 12 (P/N 046-004417-00)
• T1 (P/N 046-002376-00)

6.2.4 Sidebar Buttons
The following sidebar buttons are used to expand the functionality of this tab.

Restore Previous Settings
Select the Restore Previous Settings sidebar button to return patient display settings to those in use when the tab was opened.

NOTE: The Restore Previous Settings sidebar button is not available at the ViewStation.

All Alarms Pause/Off (Optional)
Select the All Alarms Pause sidebar button to suspend, for a user configured period of time, all numeric physiological, lethal arrhythmia, non-lethal arrhythmia, and technical alarms at the Panorama Central Station for the selected patient tile.

• The All Alarms Pause sidebar button is shown if the All Alarms Pause option is selected, and Pause Duration is configured for a limited time in the System Alarms tab.

• The All Alarms Off sidebar button is shown if the All Alarms Pause button is selected, and Pause Duration is configured as Permanent in the System Alarms tab.

WARNING: When the All Alarms Pause/Off sidebar button is enabled, the Panorama Central Station will not monitor any lethal or non-lethal arrhythmias and will not display numeric physiological or technical alarms for the selected patient tile.

NOTE: The All Alarms Pause/Off sidebar button is not available at the ViewStation or the WorkStation.

NOTE: The amount of time an alarm is suspended is dependent on the configured System Pause Duration setting.

When the All Alarms Pause/Off sidebar button is selected:
Patient Alarm Functions

- The **All Alarms Pause/Off** sidebar button changes to white with black text.
- The **All Alarms Paused/Off** message displays in red text in the Patient Status line of the patient tile. For additional information regarding the Patient Status line, refer to the “Patient Status Line” on page 2-10.

To resume the monitoring of alarms, select the **All Alarms Pause/Off** sidebar button.

- When the **All Alarms Pause/Off** sidebar button is disabled, the button changes to blue with white text. Alarms are now re-enabled for the selected patient.

**All Non-High Alarms Pause/Off (Optional)**

Select the **All Non-High Alarms Pause/Off** sidebar button to suspend all priority 2 and 3 alarms for the selected patient tile. Priority 1 alarms continue to be monitored at the Panorama Central Station while priority 2 and 3 alarms are suspended.

- The **All Non-High Alarms Pause** sidebar button is only shown if the **All Alarms Pause** or **All Non-High Alarms Pause** option is selected, and **Pause Duration** is configured as a limited time in the **System Alarms** tab.
- The **All Non-High Alarms Off** sidebar button is only shown if the **All Alarms Pause** or **All Non-High Alarms Pause** option is selected, and **Pause Duration** is configured as **Permanent** in the **System Alarms** tab.

**WARNING:** When the **All Non-High Alarms Pause/Off** sidebar button is enabled, the Panorama Central Station will not monitor any priority 2 and priority 3 physiological alarms, and will not display numeric physiological or technical alarms for the selected patient tile.

The **All Non-High Alarms Pause/Off** sidebar button is not available at the ViewStation or the WorkStation. When the All Non-High Alarms Pause/Off button is selected:

- The **All Non-High Alarms Pause/Off** sidebar button changes to white with black text.
- The **All Non-High Alarms Paused/Off** message displays in yellow text in the Patient Status line of the patient tile. For additional information regarding the Patient Status line, refer to the “Patient Status Line” on page 2-10.

To resume the monitoring of non-high alarms, select the **All Non-High Alarms Pause/Off** sidebar button.

- When the **All Non-High Alarms Pause/Off** sidebar button is disabled, the button changes to blue with white text. Non-high alarms are now re-enabled for the selected patient.

**Relearn (Telepack Only)**

Select the **Relearn** sidebar button to relearn/refresh ST and Arrhythmia data at the Panorama Central Station.

**NOTE:** The Relearn button is only available when ST and/or arrhythmia analysis is enabled.

**NOTE:** The Relearn sidebar button is not available at the ViewStation.
Normal Screen
Select the Normal Screen sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

6.2.5 Generating a Patient Alarm Report
To review the patient’s alarm settings in a report, generate a Patient Alarm Report. This report can be generated directly from the Alarm Limits tab by selecting the Print menu bar button.

For additional information regarding the Patient Alarm Report, refer to “Patient Alarm Report” on page 8-25.

6.2.6 Testing Alarm Limits
Any alarm may be tested before use to ensure that it is functioning as intended. To test an alarm:

NOTE: For the purposes of this example, use the SpO₂ alarm limit slider. These directions apply to any parameter that uses an alarm slider control.

1. Follow the steps in the “Configuring Patient Alarm Limits” on page 6-5 to access the Alarm Limits tab.
2. Scroll to the SpO₂ alarm slider.
3. To trigger the SpO₂ alarm high limit, move the slider to the current value or below the current value to trigger the alarm, and verify that the alarm behaviours are correct.
4. To trigger the alarm low limit, move the slider to current value or above the current value to trigger the alarm, and verify that the alarm behaviours are correct.
5. Once testing is complete, reset the alarm limit to the desired value.

6.2.7 Troubleshooting
This section lists some of the potential messages and issues that may occur while using the patient Alarm Limits tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>No Patient Selected!</td>
<td>This message appears in the demographic field when either no patient tile or an empty patient tile has been selected.</td>
<td>Select an active patient tile.</td>
</tr>
<tr>
<td>Communications Lost</td>
<td>Communication between the Panorama Central Station and the Telepack for the selected patient tile has been interrupted.</td>
<td>Verify the connection between the Panorama Central Station and the monitoring device. For example, is the monitoring device out of range.</td>
</tr>
<tr>
<td>All of the sidebar buttons (except the Normal Screen button) are disabled.</td>
<td>An empty patient tile is selected.</td>
<td>Select a valid patient tile by touching the digital or waveform area of the patient tile.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>All Alarms Paused</td>
<td>The All Alarms Pause function is enabled. All numeric physiological, lethal arrhythmia, non-lethal arrhythmia and technical alarm processing for the selected tile has been suspended.</td>
<td>Resume alarm processing by selecting the All Alarms Pause sidebar button and disabling the All Alarms Pause function.</td>
</tr>
<tr>
<td>All Non-High Alarms Paused</td>
<td>The All Non-High Alarms Pause function is enabled. Alarm processing for all priority 2 and priority 3 alarms has been suspended for the selected tile.</td>
<td>Resume alarm processing by selecting the All Non-High Alarms Pause sidebar button and disabling the All Non-High Alarms Pause function.</td>
</tr>
<tr>
<td>These settings are currently being edited remotely.</td>
<td>The Alarm Limits Menu is opened at the bedside monitor while the Alarm Limits tab is active at the central station</td>
<td>No solution required. Within 20 seconds, the alarm limits from the bedside will be displayed at the central station.</td>
</tr>
<tr>
<td>The Print menu bar button is disabled.</td>
<td>The message These settings are currently being edited remotely, is displayed.</td>
<td>Select the OK button to close the message box and enable the sidebar button.</td>
</tr>
<tr>
<td>The All Alarms Pause and All Non-High Alarms Pause sidebar buttons are disabled.</td>
<td>The message These settings are currently being edited remotely, is displayed.</td>
<td>Select the OK button to close the message box and enable the sidebar button.</td>
</tr>
<tr>
<td>The patient alarm limits were cleared from the Alarm Limits tab.</td>
<td>The patient size was changed at the bedside monitor.</td>
<td>No solution required. Within 20 seconds, the alarm limits from the bedside will be displayed at the central station.</td>
</tr>
<tr>
<td>The Restore Previous Settings sidebar button not functioning.</td>
<td>The previous alarm limits were cleared because the Alarm Limits tab was open while the patient size was changed at the bedside monitor.</td>
<td>No solution required. Within 20 seconds, the alarm limits from the bedside will be displayed at the central station.</td>
</tr>
<tr>
<td>The All Alarms Pause sidebar button is not displayed.</td>
<td>The Pause option in the System Alarm Options tab option is set to OFF or All Non-High Alarms.</td>
<td>If this feature is needed, contact the System Administrator to change the System Alarm Options.</td>
</tr>
<tr>
<td>The All Non-High Alarms Pause and All Alarms Pause sidebar buttons are not displayed.</td>
<td>The Pause Alarms option in the System Alarm Options tab option is set to OFF.</td>
<td>If this feature is needed, contact the System Administrator to change the System Alarm Options.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
### Alarm Limits Tab (Patient) - Patient Alarm Functions

The **Alarm Limits** tab is disabled at the WorkStation. The central station from where the patient tile was networked is configured as View Only. No solution required.

The **Alarm Responses** tab is disabled at the WorkStation. The central station from where the patient tile was networked is configured as View Only. No solution required.

* Messages are shown in all bold text.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>The <strong>Alarm Limits</strong> tab is disabled at the WorkStation.</td>
<td>The central station from where the patient tile was networked is configured as View Only.</td>
<td>No solution required.</td>
</tr>
<tr>
<td>The <strong>Alarm Responses</strong> tab is disabled at the WorkStation.</td>
<td>The central station from where the patient tile was networked is configured as View Only.</td>
<td>No solution required.</td>
</tr>
</tbody>
</table>
6.3 Alarm Responses Tab (Patient)

The Alarm Responses tab (shown in FIGURE 6-4 and FIGURE 6-6) is used to adjust the behavior or response of an alarm during an alarm condition.

The Alarm Responses tab is available at the ViewStation (shown in FIGURE 6-7) but is for viewing purposes only. When a patient tile is assigned to the ViewStation, it will always display the alarm response settings of the associated host central station patient tile.

The patient Alarm Responses tab is available at the WorkStation and functions in the same manner as it does at the host central station but contains fewer sidebar buttons. Similar to the central station, alarm response changes made at the WorkStation, with the exception of the Print on Alarm settings, will be sent to the central station, View Only Workstation, and bedside monitor (if applicable). The Print on Alarm setting is independent of the central station settings or updates.

Automatic updating of the alarm response settings at the remote workstation will also occur when:

- The monitoring device is placed in the Standby mode at the central station
- The Telepack is not communicating with its host central station (Communications Lost)
- The patient monitor is not communicating with its host central station (Comm Loss - Check Bedside Monitor)
- Communication between the host central station and the ViewStation and/or the WorkStation is lost (Central Communications Lost)

NOTE: The host central station(s) C-LAN configuration settings and the software installed at the remote location dictates the functionality available for each patient tile. For additional information, refer to “The ViewStation and the WorkStation” on page 2-4.

6.3.1 Accessing the Alarm Responses Tab

1. From the menu bar, select the Patient Alarms menu button. If the Patient Alarm Setup tab displays, enter the password in the Password text box. The Alarm Limits and Alarm Responses tabs are displayed.
2. Select the Alarm Responses tab. The Alarm Responses tab is displayed.
FIGURE 6-4  Alarm Responses Tab (Central Station)

FIGURE 6-5  Alarm Responses Tab (WorkStation)

FIGURE 6-6  Alarm Responses Tab - Paging Option Enabled (Central Station)
6.3.2 Configuring Alarm Responses

To configure patient alarm responses at the central station, complete the following:

NOTE: Alarm response changes are effective immediately after the tab is exited.

1. Choose a patient tile by selecting its digital data or waveform area.
   - The patient’s Last Name and Bed number are shown in the Demographics field, if this information was previously entered in the Demographics tab.

2. Select the alarm to be modified.
   - The Alarm Responses list box shows the alarms that can be modified.
   - When an alarm is selected, it is shown in white with black text and is shown in the edit box.

3. Select the alarm priority for the parameter. To set an alarm priority, select the Priority button until the alarm priority setting is shown. Alarm priority is associated with the severity of an alarm condition. It determines the audio and visual indicators that occur during an alarm condition.

   Alarm priority settings may include OFF, 1, 2, and 3. Each alarm has a predetermined alarm priority settings.
   - Priority 1 are the most severe alarms (lethal alarms).
   - Priority 2 are alarms of medium severity (most numeric physiological alarms).
   - Priority 3 are the least severe alarms (technical alarms and some physiological alarms).
   - Priority OFF disables audio feedback for the selected alarm at the Panorama Central Station.

   For additional information on alarm behavior, refer to “Alarm Behavior” on page 3-1.

4. The Print on Alarm button automatically prints alarm data during an alarm condition.
   - The ON option automatically prints the alarm events when the alarm condition occurs.
   - The OFF option suspends the printing of alarm events during an alarm condition.
5. The **Save To Event** button automatically saves alarm data in the Events database when an alarm condition occurs.
   - The **ON** option automatically saves alarm data during an alarm condition.
   - The **OFF** option does not save alarm data during an alarm condition.

### 6.3.3 Sidebar Buttons

The following sidebar buttons are used to expand the functionality of this tab.

**Restore Previous Settings**

Select the **Restore Previous Settings** sidebar button to return patient display settings to those in use when the tab was opened.

**NOTE:** The **Restore Previous Settings** sidebar button is not available at the ViewStation.

**All Alarms Pause (Optional)**

Select the **All Alarms Pause** sidebar button to suspend all numeric physiological, lethal arrhythmia, non-lethal arrhythmia, and technical alarms at the Panorama Central Station for the selected patient tile. The **All Alarms Pause** sidebar button is only shown if the **Suspend Alarms** option is **ON** in the **System Alarms** tab.

**WARNING:** When the **All Alarms Pause/Off** sidebar button is enabled, the Panorama Central Station will not monitor any lethal or non-lethal arrhythmias and will not display numeric physiological or technical alarms for the selected patient tile.

**NOTE:** The **All Alarms Pause** sidebar button is not available at the ViewStation or at the WorkStation.

When the **All Alarms Pause** sidebar button is selected:

- The **All Alarms Pause** sidebar button changes to white with black text.
- The **All AlarmsPaused for xx:xx mins** message displays in red text in the Patient Status line of the patient tile. For additional information regarding the Patient Status line, refer to the “Patient Status Line” on page 2-10.

To resume the monitoring of alarms, select the **All Alarms Pause** sidebar button.

- When the **All Alarms Pause** sidebar button is disabled, the button changes to blue with white text. Alarms are now re-enabled for the selected patient.

**All Non-High Alarms Pause**

Select the **All Non-High Alarms Pause** sidebar button to suspend all Priority 2 and Priority 3 alarms for the selected patient tile. Priority 1 alarms continue to be monitored at the Panorama Central Station when Priority 2 and Priority 3 alarms are suspended.

**WARNING:** When the **All Non-High Alarms Pause/Off** sidebar button is enabled, the Panorama Central Station will not monitor any priority 2 and priority 3 physiological alarms, and will not display numeric physiological or technical alarms for the selected patient tile.
The All Non-High Alarms Pause sidebar button is not available at the ViewStation or at the WorkStation. When the All Non-High Alarms Pause sidebar button is selected:

- The **All Non-High Alarms Pause** sidebar button changes to white with black text.
- The **All Non-High Alarms Paused** message displays in yellow text in the Patient Status line of the patient tile. For additional information regarding the Patient Status line, refer to the “Patient Status Line” on page 2-10.

To resume the monitoring of non-high alarms, select the **All Non-High Alarms Pause** sidebar button.

- When the **All Non-High Alarms Pause** sidebar button is disabled, the button changes to blue with white text. Non-high alarms are now re-enabled for the selected patient.

**Alarm Setup (Telepack Only)**

Select the **Alarm Setup** sidebar button to configure **ST Alarm Delay** and **Alarm Delay**, for the selected patient tile, when using a Telepack device.

**NOTE:** The **Alarm Setup sidebar button is not available at the ViewStation.**

To configure alarm delay:

1. Select the **Alarm Setup** button. The **Alarm Setup** dialog box (shown in FIGURE 6-8) is displayed.

![Alarm Setup Dialog Box](image)

**FIGURE 6-8** Alarm Setup Dialog Box

2. Select the **ST Alarm Delay** time period for the selected patient’s alarm. ST Alarm Delay is the period of time that the Panorama Central Station delays the notification of an ST alarm. ST Alarm Delay options include 30 seconds, 45 seconds, 1 minute, 90 seconds, 2 minutes and 3 minutes.

3. Select the **Alarm Delay** time period for the selected patient’s alarm. Alarm Delay is the period of time that the Panorama Central Station delays the notification of an alarm. Alarm Delay options include None, 1 second, 2 seconds, 3 seconds, 4 seconds, 5 seconds, 6 seconds, 7 seconds and 8 seconds.

**NOTE:** The Alarm Delay setting does not affect the alarm notification for ST, Apnea, or Arrhythmia.

**NOTE:** Changing the alarm delay becomes effective immediately and affects the patient’s current alarm delay setting.

4. Select the **Done** or the **Cancel** button.
   - The **Done** button accepts the configured alarm delay settings and returns to the **Alarm Responses** tab.
The **Cancel** button discards the configured alarm delay settings and returns to the **Alarm Responses** tab.

**Normal Screen**

Select the **Normal Screen** sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.
### 6.3.4 Troubleshooting

This section lists some of the potential messages and issues that may occur while using the patient **Alarm Responses** tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>No Patient Selected!</strong></td>
<td>This message appears in the demographic field when either no patient tile or an empty patient tile has been selected.</td>
<td>Select an active patient tile.</td>
</tr>
<tr>
<td><strong>Communications Lost</strong></td>
<td>Communication between the Panorama Central Station and the Telepack for the selected patient tile has been interrupted.</td>
<td>Check the connection between the Panorama Central Station and the monitoring device. For example is the Telepack out of range?</td>
</tr>
<tr>
<td>All of the sidebar buttons (except the <strong>Normal Screen</strong> button) are disabled.</td>
<td>An empty patient tile is selected.</td>
<td>Select a valid patient tile by touching the digital or waveform area of the patient tile.</td>
</tr>
<tr>
<td><strong>All Alarms Paused for xx:xx mins</strong></td>
<td>The <strong>All Alarms Pause</strong> function is enabled. All alarm processing for the selected tile has been suspended.</td>
<td>Resume alarm processing by selecting the <strong>All Alarms Pause</strong> sidebar button and disabling the <strong>All Alarms Pause</strong> function.</td>
</tr>
<tr>
<td><strong>All Non-High Alarms Paused</strong></td>
<td>The <strong>All Non-High Alarms Pause</strong> function is enabled. Alarm processing for all priority 2 and priority 3 alarms has been suspended for the selected tile.</td>
<td>Resume alarm processing by selecting the <strong>All Non-High Alarms Pause</strong> sidebar button and disabling the <strong>All Non-High Alarms Pause</strong> function.</td>
</tr>
<tr>
<td>The <strong>All Alarms Pause</strong> sidebar button is not displayed.</td>
<td>The Pause option in the System Alarm Options tab option is set to OFF or <strong>All Non-High Alarms Pause</strong>.</td>
<td>If this feature is needed, contact the System Administrator to change the System Alarm Options.</td>
</tr>
<tr>
<td>The <strong>All Non-High Alarms Pause</strong> and <strong>All Alarms Pause</strong> sidebar buttons are not displayed.</td>
<td>The Pause option in the System Alarm Options tab option is set to OFF.</td>
<td>If this feature is needed, contact the System Administrator to change the System Alarm Options.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
This page intentionally left blank.
7.0 Views

This section describes the tabs that are available when the VIEW button is selected from the Panorama Central Station Main Screen.

- Bedside Tab
- Trends Tab
- Events Tab
- Disclosure Tab
- ST Tab
7.1 Bedside Tab

The **Bedside** tab provides an expanded view of the selected patient data. The parameters selected, and the lead set used, determines the data that is displayed in the **Bedside** tab. The data in the **Bedside** tab can be displayed in two different views, the **All Waveforms** view and the **All ECG** view. Both views are discussed in the following sections.

The **Bedside** tab (shown in FIGURE 7-2) is available at the ViewStation and the WorkStation (shown in FIGURE 7-12) and functions in the same manner as it does at the host central station. The **Bedside** tab will display the bedside data from the central station provided the patient tile is in an active state.

7.1.1 Accessing the Bedside Tab

To access the **Bedside** tab (shown in FIGURE 7-1 and FIGURE 7-2), select the **VIEW** button located on the left side of the patient tile.

**NOTE:** When using a 3-wire lead set with bedside device or a Telepack using a 3-wire or 5-wire lead set, the Bedside tab will always be in the All ECG mode (as shown in FIGURE 7-6).

**NOTE:** When in the All ECG view, the color of the All ECG sidebar button changes from dark blue with white lettering to white with black lettering. This indicates that this view will continue to display until the All ECG sidebar button is selected again.
7.1.2 Bedside Tab (All Waveforms View)
The All Waveforms view of the Bedside tab (shown in FIGURE 7-1 and FIGURE 7-2) displays an expanded view of the selected patient tile.

7.1.2.1 Components of the All Waveforms View

Patient Demographic Field
The patient demographic field displays the bed number of the selected patient, if this data was entered.

NIBP Area
The All Waveforms view includes a NIBP area that displays the NIBP systolic, diastolic, mean, and elapsed time when an active patient is selected.

If applicable, the following icons display on the right side of the NIBP area:

- Indicates that Pacer Enhancement is enabled.
- Indicates that ST analysis is enabled.
- Indicates that arrhythmia analysis is enabled.
Digital Data Area

The All Waveforms view contains a digital data area that may display the following patient parameters:

- PVC/min
- Pause/min
- QT/QTc/QT-HR
- Temperature 1
- Temperature 2
- Temperature 3
- Temp Blood
- CO/CI
- CCO/CCI
- CCO STAT/CCI STAT
- SVR
- SVRI
- SvO₂
- ScvO₂
- N₂O
- PAWP
- BIS/BIS L/BIS R

NOTE: The Delta T parameter will not display in the digital data area.

NOTE: The CO/CI, Temp 2, and Temp Blood parameters are only available if they are being monitored by the DPM 6/DPM 7, Passport 12M/Passport 17M, Passport 8/Passport 12, V 12/V 21, Spectrum/Spectrum OR External Parameter Module (EPM). These parameters are not available if they are monitored by an Edwards Vigilance® monitor except during a Bolus measurement when using an Edwards Vigilance monitor with a V 12/V 21.

NOTE: The CCO/CCI, SVR, SVRI CCO STAT/CCI STAT, SvO₂, ScvO₂, Temp 3, and BIS parameters are only available if they are being monitored by the V 12/V 21.

NOTE: The QT/QTc/QT-HR parameter is only available if it is being monitored by the Telepack and V 12/V 21.
Data Display Area
When in the All Waveforms view, the data display area of the Bedside tab displays the configured patient parameters in a waveform and digital data tile format. The waveform area of the Bedside tab displays in the left portion of the data display area and the digital representation of the waveform displays to the right. In this view, three to six waveforms/digital tiles can be displayed at one time. Parameters experiencing an alarm condition will display a different background color indicating the severity of the alarm. If ST is enabled on the ECG waveform, the digital data section of the Bedside tab includes an ST lead label that specifies the lead from which the displayed data is acquired.

The first digital tile always displays one ECG lead along with the patient heart rate (HR). The ECG waveform displayed is the same that is shown in the waveform area of the patient tile. If the selected device is a Telepack with Nonin SpO₂ module and the All Waveforms view is displayed, the SpO₂ waveform will always be positioned after the patient heart rate. When the selected device is a bedside monitor, the other waveforms/digital tiles represent the available waveforms for the selected patient tile. The waveforms/digital tiles that appear in the Bedside tab are based on the parameters configured at the patient monitoring device.

The order in which the subsequent waveforms display is determined by the order defined in the Unit Priorities tab. The color associated with the waveforms/digital tiles is determined by the configured parameter color defined in the System Setup tab. A unit of measure label and an alarm status bell icon for each parameter also displays in the digital portion of the display area. The color of the waveform/digital tiles is determined by the configured parameter color.

7.1.2.2 Navigating in the All Waveforms View
If the waveforms exceed the capacity of the viewable screen, use the navigational buttons to locate the desired waveform. The buttons are described as follows:

- The up arrow button moves the group of displayed waveforms up one position within the ordered set of waveforms
- The down arrow button moves the group of displayed waveforms down one position within the ordered set of waveforms

7.1.2.3 Performing ECG Waveform Measurements
ECG waveform measurements for time and amplitude can be performed in the All Waveforms view by using the Measure dialog box (shown in FIGURE 7-3).
The Measure dialog box can only be accessed once the **Freeze** sidebar button has been selected. When the **Freeze** sidebar button is selected, the **All ECG** sidebar button is replaced with the **Measure** sidebar button.

When the Measure dialog box is first accessed, it contains five interval buttons. Once an interval button is selected, two changes occur:

- Navigational buttons and the **Clear** and **Calculate** buttons are displayed in the Measure dialog box.
- Reference and measurement lines are displayed in the **Bedside** tab waveform area.

The navigational buttons displayed depend on the selected interval. The left and right arrow buttons are available when using the PR, QRS, QT, and R to R intervals. The up and down arrow buttons are available when using the ST interval.

**Vertical Measurements (Time)**

1. Select either the PR, QRS, QT, or R to R intervals. A dashed vertical reference line is placed to the left of the solid vertical reference line.

   **NOTE:** Adjustments can be made to both the solid vertical reference line and/or the vertical dashed reference line. Selecting the interval button a second time toggles between the two reference lines. When a reference line is active, it blinks/flashes on the screen.

2. Adjust the location of the vertical reference line as follows:

   - For coarse adjustments, select the desired location on the touch screen. The reference line will move to that point.
   - For fine adjustments, use the left/right arrow buttons to position the reference line at the desired point.

3. Once the vertical reference line appears in the desired location, select the **Calculate** button. The measurement populates the text box beside the selected interval.

   - If another vertical measurement is desired, select another interval button.
   - To delete a measurement, select the interval button that needs to be to reset, then select the **Clear** button.
Horizontal Measurements (Amplitude)

1. Select the ST interval. A dashed horizontal reference line is placed above the solid horizontal reference line.

NOTE: Adjustments can be made to both the horizontal solid reference line and/or the horizontal dashed reference line. Selecting the interval button a second time toggles between the two reference lines. When a reference line is active, it blinks/flashes on the screen.

2. Adjust the location of the dashed, horizontal reference line as follows:
   - For coarse adjustments, select the desired location on the touch screen. The reference line will move to that point.
   - For fine adjustments, use the up/down arrow buttons to position the reference line at the desired point.

3. Once the dashed, horizontal reference line appears in the desired location, select the Calculate button. The measurement populates the text box beside the ST interval.
   - If another horizontal measurement is desired, select another interval button.
   - To delete the ST measurement, select the ST interval button, then select the Clear button.

7.1.2.4 Sidebar Buttons

The following sidebar buttons are used to expand the functionality of this view.

Lead
Select the Lead sidebar button to cycle through each of the waveforms in the ECG ordered set. The Lead sidebar button is not available when using a:

- Telepack device with a 3 or 5-wire lead set
- Telepack with Nonin SpO₂ module with a 3-wire lead set

All ECG
Select the All ECG sidebar button to change to the All ECG view of the Bedside tab. The Bedside tab remains in the All ECG view until the All ECG button is selected again. The All ECG sidebar button is not available when using a:

- Telepack device with a 3 or 5-wire lead set
- Telepack with Nonin SpO₂ module with a 3-wire lead set

For additional information about the All ECG view, refer to the “Bedside Tab (All ECG View)” on page 7-8.

Freeze
Select the Freeze sidebar button to halt the updating of digital parameter data and waveforms currently displayed in the Bedside tab. The Bedside tab remains frozen until the Freeze sidebar button is selected again. Freezing the Bedside tab does not affect the waveform or any real time alarms in the upper half of the Normal Screen.
When the **Freeze** sidebar button is selected, the **All ECG** sidebar button is replaced by the **Measure** sidebar button.

**Measure**
Select the **Measure** sidebar button to activate the Measure dialog box (shown in FIGURE 7-3). The Measure dialog contains reference and measurement lines to measure the time and amplitude of an ECG waveform. For additional information regarding ECG waveform measurements, refer to "Performing ECG Waveform Measurements" on page 7-11.

**Mark Event**
Select the **Mark Event** sidebar button to save the current data to the Event database as a ‘Central User Marked Event.’ This can be useful for noting such things as the time when a medication was administered.

**NOTE:** The Mark Event sidebar button is not available at the ViewStation or the WorkStation.

**Normal Screen**
Select the **Normal Screen** sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

7.1.3 **Generating an All Strips Report**
To review a patient's real-time data, generate an All Strips Report. This report can be generated directly from the **Bedside** tab by selecting the **Print** menu bar button.

For additional information regarding the All Strips Report, refer to the “All Strips Report” on page 8-2.

7.1.3.1 **Troubleshooting**
For **All Waveforms** view troubleshooting information, refer to "Troubleshooting" on page 7-12.

7.1.4 **Bedside Tab (All ECG View)**
The **Bedside** tab **All ECG** view displays all the ECG waveforms available based on the lead set currently in use.

7.1.4.1 **Accessing the All ECG View**
To access the **All ECG** view (shown in FIGURE 7-4), select the **All ECG** sidebar button from the **Bedside** tab **All Waveforms** view.
FIGURE 7-4  Bedside Tab - All ECG View (Central Station)

FIGURE 7-5  Bedside Tab - All ECG View (ViewStation and WorkStation)
7.1.4.2 Components of the All ECG View

Patient Demographic Field
The description of this field was previously described on page 7-3.

NIBP Area
The description of this area was previously described on page 7-3.

Digital Data Area
The description of this area was previously described on page 7-4.

Data Display Area
When in the All ECG view, the waveform area of the Bedside tab displays all the ECG waveforms based on the lead set currently in use. The waveform area of the Bedside tab displays in the left portion of the data display area and the ECG lead indicator displays to the right.

The first digital tile displays one ECG lead along with the patient heart rate (HR). The ECG waveform displayed is the same waveform shown in the waveform area of the patient tile, except when a View 12™ card is in use. In this case, the first waveform displayed will always be ECG Lead I. When a 3 or a 5-wire lead is used, the ECG lead that appears in the first waveform position is selected from the Display tab. The other waveforms/lead tiles represent the available waveforms/leads for the selected patient tile. The order in which the subsequent waveforms appear is based on the order specified by the ordered set. The waveforms/lead tiles that appear are based on the lead set currently in use.
7.1.4.3 Navigating in the All ECG View
If the waveforms exceed the capacity of the viewable screen, use the navigational buttons to locate the desired waveform. The buttons are described as follows:

- The up arrow button moves the group of displayed waveforms up one position within the ordered set of waveforms
- The down arrow button moves the group of displayed waveforms down one position within the ordered set of waveforms

When a View 12™ card is in use, the double up arrow/down arrow can be used to navigate:

- The double up arrow button displays the previous set of ECG waveforms
- The double down arrow button displays the next set of ECG waveforms

7.1.4.4 Performing ECG Waveform Measurements
ECG waveform measurements for time and amplitude can be performed in the All ECG view by using the Measure dialog box (shown in FIGURE 7-3). For additional information about performing waveform measurements, refer to “Performing ECG Waveform Measurements” on page 7-5.

7.1.4.5 Sidebar Buttons
The sidebar buttons available in this view are the same as the sidebar buttons described in the All Waveforms view. Please refer to page 7-7 for a full description of each of the sidebar buttons.
## Troubleshooting

This section lists some of the potential messages and issues that may occur while using the either of the **Bedside** tab views.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE*</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>A waveform parameter is not displayed.</td>
<td>If the waveform parameter exists, it has not been scrolled into view on the display.</td>
<td>Use the navigation buttons or <strong>View More</strong> sidebar button to scroll to the desired parameter waveform.</td>
</tr>
<tr>
<td></td>
<td>The waveform parameter is not being monitored at the bedside.</td>
<td>Connect the sensor(s) for the waveform parameter to the patient.</td>
</tr>
<tr>
<td>NIBP/Digital Data area empty</td>
<td>No equipment is assigned to the selected tile.</td>
<td>Assign a monitoring device to the patient tile.</td>
</tr>
<tr>
<td></td>
<td>Patient is in the “Panorama Standby Only” mode</td>
<td>Remove patient from <strong>Panorama Standby Only</strong> by selecting the <strong>Resume Monitoring</strong> button in the <strong>Standby</strong> tab.</td>
</tr>
<tr>
<td></td>
<td>Patient is in the “Bedside and Panorama Standby Only” mode</td>
<td>Remove patient from <strong>Bedside and Panorama Standby Only</strong> by pressing the <strong>STANDBY</strong> button on front panel of the bedside monitoring device.</td>
</tr>
<tr>
<td></td>
<td>Selected tile is in the Discharge state.</td>
<td>Retrieve discharged patient.</td>
</tr>
<tr>
<td>(for NIBP area only)</td>
<td>The equipment assigned to the selected tile does not support the NIBP parameter.</td>
<td>Assign a monitoring device that supports NIBP to the selected tile.</td>
</tr>
<tr>
<td>Up arrow/down arrow button is not available or is disabled</td>
<td>View 12™ card is in use</td>
<td>Use double up/down arrow button to navigate</td>
</tr>
<tr>
<td></td>
<td>There are less than six waveforms available.</td>
<td></td>
</tr>
<tr>
<td>Double up arrow/down arrow button is not available</td>
<td>View 12™ card is not in use</td>
<td></td>
</tr>
<tr>
<td>First waveform position not affected when navigation up/down arrow button is used</td>
<td>The first waveform is not affected by the navigational buttons.</td>
<td></td>
</tr>
<tr>
<td><strong>Lead/All ECG sidebar button is unavailable</strong></td>
<td>No equipment is assigned to the selected tile.</td>
<td>Assign a monitoring device to the patient tile.</td>
</tr>
<tr>
<td></td>
<td>Patient is in the “Panorama Standby Only” mode.</td>
<td>Remove patient from <strong>Panorama Standby Only</strong> by selecting the <strong>Resume Monitoring</strong> button in the <strong>Standby</strong> tab.</td>
</tr>
<tr>
<td></td>
<td>Patient is in the “Bedside and Panorama Standby Only” mode.</td>
<td>Remove patient from <strong>Bedside and Panorama Standby Only</strong> by pressing the <strong>STANDBY</strong> button on front panel of the monitoring device.</td>
</tr>
<tr>
<td></td>
<td>Selected tile is in the Communications Lost or Comm Loss - Check Bedside Monitor state or Discharge state.</td>
<td>Verify connection to central stations and/or retrieve discharged patient.</td>
</tr>
</tbody>
</table>
### MESSAGE/ISSUE

<table>
<thead>
<tr>
<th>Reason</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Selected device is a Telepack with a 3 or 5-wire lead set.</td>
<td></td>
</tr>
<tr>
<td>Selected device is a Passport 2, Passport V, Spectrum, DPM 6/DPM 7, Passport 12M/Passport 17M, Passport 8/Passport 12, T1 monitor or V 12/V 21 with a 3 or 5-wire lead set.</td>
<td></td>
</tr>
<tr>
<td>Bedside tab is in the All ECG display or Freeze mode.</td>
<td>Select the <strong>All ECG</strong> sidebar button to change to the Normal display mode.</td>
</tr>
<tr>
<td><strong>Lead</strong> sidebar button is disabled</td>
<td></td>
</tr>
<tr>
<td>The selected device is a:</td>
<td></td>
</tr>
<tr>
<td>- Telepack device with a 3 or 5-wire lead</td>
<td></td>
</tr>
<tr>
<td>- Telepack with Nonin SpO2 module attached with a 3-wire lead set</td>
<td>No solution required.</td>
</tr>
<tr>
<td>The <strong>All ECG</strong> sidebar button is latched.</td>
<td>Select the <strong>All ECG</strong> sidebar button to enable the <strong>Lead</strong> sidebar button.</td>
</tr>
<tr>
<td>Buttons in Measure dialog box are unavailable</td>
<td></td>
</tr>
<tr>
<td>Interval is not selected.</td>
<td>Select an interval.</td>
</tr>
<tr>
<td>The <strong>Print</strong> menu bar button is disabled.</td>
<td></td>
</tr>
<tr>
<td>The patient tile is in the <strong>Bedside and Panorama Standby</strong> mode.</td>
<td>If necessary, remove the patient tile from this mode.</td>
</tr>
<tr>
<td>The patient tile is in the <strong>Panorama Standby Only</strong> mode.</td>
<td>If necessary, remove the patient tile from this mode.</td>
</tr>
<tr>
<td>The patient tile is in the <strong>Communications Lost</strong> or <strong>Comm Loss - Check Bedside Monitor</strong> state.</td>
<td>Restore communications with the monitoring device.</td>
</tr>
<tr>
<td>The patient tile is in a <strong>Waiting for Patient</strong> state.</td>
<td>Restore communications with the bedside monitor.</td>
</tr>
<tr>
<td>The patient tile is in the <strong>Bedside and Panorama Standby</strong> mode.</td>
<td>If necessary, remove the patient tile from this mode.</td>
</tr>
<tr>
<td>The patient tile is in the <strong>Panorama Standby Only</strong> mode.</td>
<td>If necessary, remove the patient tile from this mode.</td>
</tr>
<tr>
<td>The patient tile is in the <strong>Communications Lost</strong> or <strong>Comm Loss - Check Bedside Monitor</strong> state.</td>
<td>Restore communications with the monitoring device.</td>
</tr>
<tr>
<td>The patient tile is in a <strong>Waiting for Patient</strong> state.</td>
<td>Restore communications with the bedside monitor.</td>
</tr>
</tbody>
</table>
7.2 Trends Tab

The **Trends** tab provides access to a patient's trended digital data. Trend data is automatically displayed in one minute intervals unless an alternate Trend Display interval was selected. If the **Trends** tab is selected before the first interval is reached, or before the first NIBP, CO/CI, or PAWP measurement occurs, the list will be empty. All previously collected parameter data will be displayed, even if the parameter is not currently being monitored.

The monitoring device, utilized parameters, and configured settings determines the data that is displayed in the **Trends** tab. The configured trend display settings determine at what interval the trends data will be displayed in the **Trends** tab.

The **Trends** tab is available at the ViewStation and the WorkStation and functions in the same manner as it does at the host central station. The **Trends** tab will display the trend data from the central station provided the patient tile is communicating with the central station.

A patient’s trended digital data can be viewed in two formats, either as a list, or graphically. There is a configuration button in the Unit Choices tab labeled Trend Access Order that configures whether to open the Trends tab in List or Graphic format. For additional information, refer to “Setting the Unit Choices (Central Station)” on page 9-51. Both views are discussed in the following sections.

7.2.1 Accessing the Trends Tab

1. Select the **VIEW** button on the desired patient tile. The **Bedside** tab is displayed.
2. Select the **Trends** tab. The **Trends** tab is displayed in the default view. If the List view is not the default view, select the **List** sidebar button. The **Trends** tab is displayed in the **List** view (shown in FIGURE 7-7)

**FIGURE 7-7** Trends Tab - List View (Central Station, ViewStation, and WorkStation)
7.2.2 Trends Tab (List View)

The **List** view of the **Trends** tab (shown in FIGURE 7-7) provides a list of all of the trended digital data for the selected patient and is displayed first when the **Trends** tab is accessed. Each trend record is displayed as a row in the list, in descending chronological order. The most recent trend record appears at the top.

When the **List** view of the **Trends** tab is selected, the top row in the list is highlighted. Each trend record’s date and time display in the first column. All subsequent columns display parameter data in the order that was specified in the **Unit Priorities** tab. For additional information, refer to “System Setup Functions” on page 9-1.

If a priority 1 physiological alarm condition existed for a parameter during the time of trend record storage, that parameter’s data will be highlighted in red.

If a priority 2 or 3 physiological alarm condition existed for a parameter during the time of trend record storage, that parameter’s data will be highlighted in yellow.

7.2.2.1 Components of the List View

**Patient Demographic Field**

The patient demographic field displays the last name and bed number of the selected patient, if this data was entered.

7.2.2.2 Navigating in the List View

If the **List** view exceeds the capacity of the viewable screen, use the navigational buttons and scroll bars to locate the desired parameter data.

**Scrolling through the List**

There are three methods of scrolling through the list rows:

- Use the New Data button or the Old Data button, both located below the horizontal scroll bar, to display the most current or the oldest trend records filtered to the current Trend display settings for the selected patient.

- Use the Page Down (double down-arrow) and Page Up (double up-arrow) buttons, located below the horizontal scroll bar, to scroll through one page of the list at a time.

- Use the up and down arrows, located below the horizontal scroll bar, for single row scrolling.

**Scrolling through the Parameters**

Use the left and right arrows of the horizontal scroll bar for single parameter scrolling or drag the scroll box to move quickly through the parameters.

**Previous Event Button**

Select the **Previous Event** button to locate and highlight the previous event condition that occurred before the currently highlighted row.
Next Event Button
Select the Next Event button to locate and highlight an event that occurred after the currently highlighted trend in the list box.

7.2.2.3 Configuring a Trend Display Interval
Select the Trend Display sidebar button to configure the interval at which a patient’s trend data will be displayed. Regardless of which Trend Display interval is selected, NIBP, CO/CI, and PAWP trend records will be displayed.

ViewStation and WorkStation patient tiles will initially display the same trend interval settings of the associated patient tile at the host central station. For additional information regarding the transfer of trend interval settings from the host central station, refer to the “Trends Tab” on page 7-14.

The default setting for a Trend Display interval is determined by the Unit Choices tab. For additional information regarding the Unit Choices tab, refer to “Unit Choices Tab” on page 9-50.

NOTE: Trends data is automatically stored in one minute intervals. The Trend Display interval selected does not affect how trend information is stored in the database, just how it is displayed.

NOTE: Changing the trend display setting in the List Trend tab automatically adjusts the trend display setting in the Display tab. Changing the trend display setting in the Display tab automatically adjusts the trend display setting in the List Trend tab.

To select the Trend Display interval:

1. From the Display tab, select the Trend Display sidebar button. The Trend Display dialog box (shown in FIGURE 7-8) is displayed.

   [FIGURE 7-8 Trend Display Dialog Box]

2. Select the Interval button until the desired interval period is displayed. Interval periods include 1 minute, 5 minutes, 10 minutes, 15 minutes, 30 minutes, 1 hour, and 2 hours.

3. Select the Done or the Cancel button.
   - The Done button accepts the selection.
   - The Cancel button discards the selection, closes the dialog box, and returns to the Display tab.
With the exception of NIBP, CO/CI and PAWP, selecting the **Done** button will change the Trend Display interval in the Trends tab based on the standard clock time intervals. Refer to the following table:

<table>
<thead>
<tr>
<th>INTERVAL</th>
<th>TREND RECORDS DISPLAYED IN TRENDS TAB</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 minute</td>
<td>All trend records will be displayed.</td>
</tr>
<tr>
<td>5 minute</td>
<td>All trend records having a timestamp of 0, 5, 10, 20, 25, 30, 35, 40, 45, 50, and 55 minutes will be displayed.</td>
</tr>
<tr>
<td>10 minute</td>
<td>All trend records having a timestamp of 0, 10, 20, 30, 40, and 50 minutes will be displayed.</td>
</tr>
<tr>
<td>15 minute</td>
<td>All trend records having a timestamp of 0, 15, 30, and 45 minutes will be displayed.</td>
</tr>
<tr>
<td>30 minutes</td>
<td>All trend records having a timestamp of 0 and 30 minutes will be displayed.</td>
</tr>
<tr>
<td>1 hour</td>
<td>All trend records having a timestamp of 0 minutes (for example, 12:00, 1:00, 2:00) will be displayed.</td>
</tr>
<tr>
<td>2 hour</td>
<td>All trend records having a timestamp of 0 minutes for even hours only (for example, 12:00, 2:00, 4:00) will be displayed.</td>
</tr>
</tbody>
</table>

### 7.2.2.4 Printing a Trend List Report from the List View

Multiple hours of trend data can be printed to a Trends List Report directly from **List** view.

To indicate the time interval for printing this report:

1. From the **List** view, select the **Print** menu bar button. The Print Selection dialog box (shown in **FIGURE 7-9**) is displayed.

   ![Print Selection Dialog Box](image)

   **FIGURE 7-9** Print Selection Dialog Box

2. Select the **Print Current** button, a time interval button, or the **Cancel** button.
   - Select the **Print Current** to create a one page Trend List Report.
   - Select one of the time interval buttons. The choices are print 1 hour, 2 hours, 4 hours, 8 hours, 12 hours, and 24 hours.

3. Select the **Cancel** button to close the dialog box without generating the print request. For additional information regarding this report, refer to “Trend List Report” on page 8-17.

### 7.2.2.5 Sidebar Buttons

The following sidebar buttons are used to expand the functionality of this view.
Graphic
Select the Graphic sidebar button to open the Graphic view of the Trends tab. For additional information regarding the Graphic view, refer to "Accessing the Graphic View" on page 7-20.

Trend Display
This sidebar button was previously described on page 7-16.

Events
Select the Events sidebar button to navigate to the Events dialog. The date and time associated with the highlighted Trend Record or vertical marker in Graphic Trend will be used to navigate to the Events dialog. The most current Event nearest to the Trend Record time and date will be selected.

Disclosure
Select the Disclosure sidebar button to navigate to the Disclosure dialog. The date and time associated with the highlighted Trend Record or vertical marker in Graphic Trend will be used to navigate to the Disclosure dialog. The most current Disclosure data to the nearest minute will be displayed at the top of the Disclosure dialog.

Normal Screen
Select the Normal Screen sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

7.2.2.6 Troubleshooting
This section lists some of the potential messages and issues that may occur while using the List view of the Trends tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>There is no data in the List view of the Trends tab.</td>
<td>A patient has not been admitted to the selected tile.</td>
<td>Admit a patient to the selected tile or choose a tile that already has an admitted patient.</td>
</tr>
<tr>
<td></td>
<td>The Trends tab was selected before the first interval was reached or before the first NIBP measurement occurred.</td>
<td>Wait until the first interval is reached or until the first NIBP measurement occurs.</td>
</tr>
<tr>
<td>A column of parameter data is not displayed.</td>
<td>If the column of parameter data exists, it has not been scrolled into view on the display.</td>
<td>Use the scroll bar located along the bottom of the Trend list to scroll to the desired parameter column.</td>
</tr>
<tr>
<td></td>
<td>The parameter has not been monitored for this patient.</td>
<td>Connect the sensor(s) for the parameter to the patient.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
<table>
<thead>
<tr>
<th>MESSAGE/ISSUE</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>No more data found</strong> (Page Down Button)</td>
<td>There is no data in the Trend database because the <strong>Trends</strong> tab was selected before the first interval was reached or before the first NIBP measurement occurred.</td>
<td>Wait until the first interval is reached or until the first NIBP measurement occurs.</td>
</tr>
<tr>
<td></td>
<td>The last row in the list is already highlighted.</td>
<td>Select the <strong>OK</strong> button to close the message box and return to the <strong>List</strong> view of the <strong>Trends</strong> tab.</td>
</tr>
<tr>
<td><strong>No more data found</strong> (Page Up Button)</td>
<td>There is no data in the Trend database because the <strong>Trends</strong> tab was selected before the first interval was reached or before the first NIBP measurement occurred.</td>
<td>Wait until the first interval is reached or until the first NIBP measurement occurs.</td>
</tr>
<tr>
<td></td>
<td>The first row in the list is already highlighted.</td>
<td>Select the <strong>OK</strong> button to close the message box and return to the <strong>List</strong> view of the <strong>Trends</strong> tab.</td>
</tr>
<tr>
<td><strong>No more data found</strong> (Previous Event Button)</td>
<td>There is no event data in the Trend database.</td>
<td>Wait until an event is recorded to the Trend database.</td>
</tr>
<tr>
<td></td>
<td>None of the rows that are below the currently highlighted row contain a parameter with an event condition.</td>
<td>Select the <strong>OK</strong> button to close the message box and return to the <strong>List</strong> view of the <strong>Trends</strong> tab.</td>
</tr>
<tr>
<td></td>
<td>The last row in the list is already highlighted.</td>
<td>Select the <strong>OK</strong> button to close the message box and return to the <strong>List</strong> view of the <strong>Trends</strong> tab.</td>
</tr>
<tr>
<td><strong>No more events found</strong> (Next Event Button)</td>
<td>There is no event data in the Trend database.</td>
<td>Wait until an event is recorded to the Trend database.</td>
</tr>
<tr>
<td></td>
<td>None of the rows that are above the currently highlighted row contain a parameter with an event condition.</td>
<td>Select the <strong>OK</strong> button to close the message box and return to the <strong>List</strong> view of the <strong>Trends</strong> tab.</td>
</tr>
<tr>
<td></td>
<td>The first row in the list is already highlighted.</td>
<td>Select the <strong>OK</strong> button to close the message box and return to the <strong>List</strong> view of the <strong>Trends</strong> tab.</td>
</tr>
<tr>
<td>The <strong>Graphic</strong> and <strong>Trend Display</strong> sidebar buttons are disabled.</td>
<td>There is no data in the Trend database because the <strong>Trends</strong> tab was selected before the first interval was reached or before the first NIBP measurement occurred.</td>
<td>Wait until the first interval is reached or until the first NIBP measurement occurs.</td>
</tr>
<tr>
<td></td>
<td>A patient has not been admitted to the selected tile.</td>
<td>Admit a patient to the selected tile or choose a tile that already has an admitted patient.</td>
</tr>
<tr>
<td></td>
<td>The Trends List data is still loading.</td>
<td>Wait for the Trends List data to finish loading.</td>
</tr>
<tr>
<td></td>
<td>The <strong>Print Selection</strong> dialog box is open.</td>
<td>Close the <strong>Print Selection</strong> dialog box.</td>
</tr>
<tr>
<td>The <strong>Print</strong> menu bar button is disabled.</td>
<td>The trends data is loading.</td>
<td>Wait until the trends data is loaded.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
7.2.3 Trends Tab (Graphic View)

The Graphic view of the Trends tab (shown in FIGURE 7-10) provides a graphical representation of all of the trended digital data for a selected patient. All previously collected data is displayed, even if the parameter is not currently being monitored.

The Graphic view for each parameter includes a range bar, plot area and parameter area for the selected time. A maximum of three parameters can be scrolled into the viewable area of the display.

When the Graphic view is displayed, a movable reference line, that crosses the three available plot areas, is initially placed at the same date/time as the row (trend record) that was highlighted in the List view. The trend record data that is displayed in the parameter area is defined by the location of the reference line.

The Graphic view is available at the ViewStation and the WorkStation and functions in the same manner as it does at the host central station. There is a configuration button in the Unit Choices tab labeled Trend Access Order that configures whether to open the Trends Tab in List Trend or Graphic Trend format. For additional information, refer to “Setting the Unit Choices (Central Station)” on page 9-51.

7.2.3.1 Accessing the Graphic View

1. Select the VIEW button on the desired patient tile. The Bedside tab is displayed.
2. Select the Trends tab. The Trends tab is displayed in the default view. If the Graphic view is not the default view, select the Graphic sidebar button. The Trends tab is displayed in the Graphic view (shown in FIGURE 7-10).

![FIGURE 7-10 Trends Tab - Graphic View (Central Station, ViewStation, and WorkStation)](image)

7.2.3.2 Components of the Graphic View

**Patient Demographic Field**

The patient demographic field displays the last name and bed number of the selected patient, if this data was entered.
Alarm Indicator Area
Vertical lines that are displayed in the alarm indicator area denote the time at which an alarm occurred. The color of each vertical line in the alarm indicator area denotes the priority of the alarm.

Date/Time Area
The date/time area indicates the date and time associated with the location of the reference line.

Display Time Interval
The display time interval is the current setting for the viewable snapshot and is one page of the Graphic view. When the Graphic view opens, the default display time interval is two hours.

Time Line
The time line displays nine equally spaced time values based on the display time interval.

Range Bars
The range bar for each parameter, except Heart Rate, is displayed with the maximum and minimum values for that parameter.

For Heart Rate, the minimum and maximum scale is automatically adjusted based on the range of values displayed at the time, also considering the current zoom level and time range. The minimum scale value for Heart Rate is 10 bpm less than the lowest displayed value in the current view; the maximum scale value for Heart Rate is 10 bpm greater than the highest displayed value in the current view.

Each range bar is displayed in the color of the parameter as described in the Parameter Color tab.

Plot Areas
The plotted trend records are displayed in the following format:

- One plotted point for single sensor parameters
- A line connecting the inspired value and end tidal value for dual sensor parameters
- A line connecting the systolic and diastolic values and a single dot for the mean value for multi-sensor parameters

Each plotted trend record is displayed in the color of the parameter as described in the Parameter Color Tab.

Parameter Areas
The trend record data that is displayed in the parameter area is defined by the location of the reference line. For parameters that have experienced an alarm event, the background color of the parameter area is based on the alarm priority at the time of trend record storage.
If there is no data at the location of the reference line, the parameter areas will display the parameter label and/or the unit of measurement.

7.2.3.3 Navigating in the Graphic View

Scrolling through the Parameters
A vertical scroll bar is located along the right side of the parameter area. The up and down arrows on the scroll bar can be used for single parameter scrolling or the bar can be dragged to move quickly through a large number of parameters.

Scrolling through the Time Line
There are two methods of scrolling through the time line:

- Use the left arrow button to move the reference line to the previous trend record on the time line and the right arrow button to move the reference line to the next trend record on the time line.
- Use the Page Left (double left-arrow) button to scroll to the previous page and the Page Right (double right-arrow) button to scroll to the next page. Increments are based on the display time interval.

Zoom In Button
The **Zoom In** button decreases the display time interval to a minimum of 2 hours, in two hour increments.

Zoom Out Button
The **Zoom Out** button increases the display time interval to a maximum of 8 hours, in two hour increments.

7.2.3.4 Sidebar Buttons
The following sidebar buttons are used to expand the functionality of this view.

List
Select the **List** sidebar button to return to the List view of the Trends tab.

Events
Select the Events sidebar button to navigate to the Events dialog. The date and time associated with the highlighted Trend Record or vertical marker in Graphic Trend will be used to navigate to the Events dialog. The most current Event nearest to the Trend Record time and date will be selected.

Disclosure
Select the **Disclosure** sidebar button to navigate to the Disclosure dialog. The date and time associated with the highlighted Trend Record or vertical marker in Graphic Trend will be used to navigate to the Disclosure dialog. The most current Disclosure data to the nearest minute will be displayed at the top of the Disclosure dialog. Normal Screen
Select the Normal Screen sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

7.2.4 Generating a Graphic Trend Report
To review the trends for the selected patient tile in a graphical format, generate a Graphic Trend Report. This report can be generated directly from the Trends tab by selecting the Print menu bar button.

For additional information regarding the Graphic Trend Report, refer to the “Graphic Trend Report” on page 8-48.

7.2.4.1 Troubleshooting
This section lists some of the potential messages and issues that may occur while using the Graphic view of the Trends tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>No more data found (Page Left Button)</td>
<td>There is no data in the Trend database because the Trends tab was selected before the first interval was reached or before the first NIBP measurement occurred.</td>
<td>Wait until the first interval is reached or until the first NIBP measurement occurs.</td>
</tr>
<tr>
<td></td>
<td>The first recorded data has been reached. No further data exists before this time.</td>
<td>Select the OK button to close the message box and return to the Graphic view of the Trends tab.</td>
</tr>
<tr>
<td>No more data found (Page Right Button)</td>
<td>There is no data in the Trend database because the Trends tab was selected before the first interval was reached or before the first NIBP measurement occurred.</td>
<td>Wait until the first interval is reached or until the first NIBP measurement occurs.</td>
</tr>
<tr>
<td></td>
<td>The most recent recorded data has been reached. No further data exists after this time.</td>
<td>Select the OK button to close the message box and return to the Graphic view of the Trends tab.</td>
</tr>
<tr>
<td>Time out of range (Zoom In Button)</td>
<td>The minimum display time interval has been reached.</td>
<td>Select the OK button to close the message box and return to the Graphic view of the Trends tab.</td>
</tr>
<tr>
<td>Time out of range (Zoom Out Button)</td>
<td>The maximum display time interval has been reached.</td>
<td>Select the OK button to close the message box and return to the Graphic view of the Trends tab.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
### MESSAGE/ISSUE *

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Time out of range</strong></td>
<td>The minimum display time interval has been reached.</td>
<td>Select the OK button to close the message box and return to the Graphic view of the Trends tab.</td>
</tr>
<tr>
<td>(Left Arrow Button)</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Time out of range</strong></td>
<td>The maximum display time interval has been reached.</td>
<td>Select the OK button to close the message box and return to the Graphic view of the Trends tab.</td>
</tr>
<tr>
<td>(Right Arrow Button)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>The List sidebar button is</td>
<td>A message box is being displayed.</td>
<td>Close the message box.</td>
</tr>
<tr>
<td>disabled.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* *Messages are shown in all bold text.*
7.3 Events Tab

The Events tab lists all the events for a selected patient tile. The monitoring device, utilized parameters, and configured settings determine the events that are saved in the Events list. All previously collected data is displayed, even if the parameter is not being monitored.

NOTE: If a lethal arrhythmia alarm condition exists, and a transition to a different lethal arrhythmia alarm condition occurs, a new event will be stored and a printout will be generated, as configured.

The Events tab is available at the ViewStation and the WorkStation (shown in FIGURE 7-12) and functions in the same manner as it does at the host central station, with the exception of the delete event buttons. The Events tab will display the event data from the central station, provided the patient tile is communicating with the central station.

The Events tab displays events in two formats, either as a list, or as a waveform. Both views are discussed in the following sections.

7.3.1 Accessing the Events Tab

1. Select the VIEW button on the desired patient tile. The Bedside tab is displayed.

2. Select the Events tab. The Events tab is displayed in the List view (shown in FIGURE 7-11).

![FIGURE 7-11 Events Tab - List View](image-url)
7.3.2 Events Tab (List View)

The List view lists all the events for the selected tile in a column format. Each event record is displayed as a row in the list, with the most recent filtered event record at the top of the list. The remaining events display in descending chronological order. For additional information regarding events filtering, refer to "Events Filtering" on page 7-33.

7.3.2.1 Components of the List View

Patient Demographic Area

The List view patient demographic area will display the bed number of the selected patient, if this data was entered. In addition, the total number of events currently in the Events database will display beside the patient demographic information.

Events Tab List Box Area

The List view displays each event record as a row in the list box. The list box is capable of displaying a maximum of 1000 rows at one time. If the 1000 row maximum is reached, and new events occur, the oldest events will automatically be deleted from the Events database.

When the List view is first selected, the top row in the list will be highlighted. The event record date and time is displayed in the first column, the event description is displayed in the second column, and the heart rate (HR) is displayed in the third column (shown in FIGURE 7-11). All subsequent columns display parameter data in the order specified in the Unit Priorities tab.

NOTE: All previously collected parameter data is displayed in the Events tab list box, even if the parameter is not currently being monitored.
Alarm Indicators
If there was an alarm associated with an event, the specific parameter value will be highlighted in the Event list. For example, if a Priority 1 physiological or bedside alarm condition existed for a parameter at the time of event record storage, that parameter's data is highlighted in red. If a Priority 2 or 3 physiological or bedside alarm condition existed for a parameter at the time of event record storage, that parameter's data is highlighted in yellow.

7.3.2.2 Navigating in the List View
If the List view list exceeds the capacity of the viewable screen, use the navigational buttons and scroll bars to locate the desired parameter data.

Scrolling through the List
There are three methods of scrolling through the list rows:

- Use the New Data button or the Old Data button, both located below the horizontal scroll bar, to display the most current or the oldest events for the selected patient.
- Use the Page Down (double down-arrow) and Page Up (double up-arrow) buttons, located below the horizontal scroll bar, to scroll through one page of the list at a time.
- Use the up and down arrows, located below the horizontal scroll bar, for single row scrolling.

Scrolling through the Parameters
Use the left and right arrows of the horizontal scroll bar for single parameter scrolling or drag the scroll box to move quickly through the parameters.

7.3.2.3 Deleting Event Records
Remove event records from the List view by selecting the Delete Single Event button and/or the Delete Events button.

Delete Single Event Button
Select the Delete Single Event button to remove a single event record from the List view.

NOTE: The Delete Single Event button is not available at the ViewStation or the WorkStation.

To delete a single event record:

1. From the List view, select the event record to be deleted.
2. Select the Delete Single Event button. A confirmation dialog box is displayed.
3. Select Yes or No.
   - The Yes button deletes the event record. The event record is removed from the Events list.

NOTE: If the Events List is open at the ViewStation or WorkStation when the host central station confirmation dialog box “Yes” button is selected, the message “Events were deleted from the database. Press the OK button to refresh the list.” will be displayed at the ViewStation or the WorkStation.
• The **No** button does not delete the event.

**NOTE:** Not all event records can be deleted. Refer to the “Alarms and Events” chapter for more information regarding which events cannot be deleted.

### Delete Events Button
Select the **Delete Events** button to remove multiple events from the **List** view. Events are deleted from the first selected event downward, towards the older events. For example, to delete the five most recent events in the list, select the first event in the list, then select the **Delete Events** button. Verify the number of events to delete is correct, then select the **Delete** button.

**NOTE:** The **Delete Events** button is not available at the **ViewStation** or the **WorkStation**.

To delete multiple events:

1. From the **List** view, select the first event to be deleted.
2. Select the **Delete Events** button. The **Delete Events** dialog box (shown in **FIGURE 7-13**) is displayed.

   ![FIGURE 7-13 Delete Events Dialog Box](image)

3. Select the 0 in the **Number Of Events** text box. The keypad dialog box is displayed.
4. Use the numeric keypad to enter the number of events to delete (between 1-10 if filtering is turned off and 1-50 if filtering is on), then select the **OK** button. The numeric keypad closes and the **Delete Events** dialog box is displayed.
5. Select the **Delete** or **Cancel** button.
   • Select the **Delete** button to remove the events.
   • Select the **Cancel** button to close the dialog box and leave the **Events** list unmodified.
6. Selecting the **Delete** button prompts a confirmation box to display.
   • Select the **Yes** button to delete the event records. The event records are removed from the **Events** list.

**NOTE:** If the **Events List** is open at the **ViewStation** or **WorkStation** when the host central station confirmation dialog box “Yes” button is selected, the message “Events were deleted from the database. Press the OK button to refresh the list.” will be displayed at the **ViewStation** or the **WorkStation**.

   • Select the **No** button to close the dialog box and leave the **Events** list unmodified.
NOTE: Not all event records can be deleted. Refer to the “Alarms and Events” chapter for more information regarding which events cannot be deleted.

7.3.2.4 Refreshing the List View
Depending on the event type, the event may not be displayed in the List view until the list is regenerated or refreshed. To refresh the List view, select any neighboring tab, such as the Disclosure or ST tabs, then reselect the Events tab. This will refresh the List view so it will display the most current events.

7.3.3 Event Filtering
The Events tab provides an event filter as a means of controlling the events displayed in the List view. Filters can be configured based on time and on parameters. Since the Event list contains historical data, the event filter is available even when the patient tile is experiencing a Communications Lost or Comm Loss - Check Bedside Monitor condition or is in a Panorama Standby Only or Bedside and Panorama Standby mode.

The initial central station event filter settings come from the System Event Filter settings, if settings were configured. ViewStation and WorkStation patient tiles will initially display the same event filter settings of the associated patient tile at the host central station. For additional information regarding the transfer of event filter settings from the host central station, refer to the “Trends Tab” on page 7-14.

7.3.3.1 Accessing the Events Filter
To access the events filter, select the Events Filtering sidebar button from the Events tab List view.

![Event Filter Text Box](image)

**FIGURE 7-14** Event Tab (Events Filtering)

7.3.3.2 Components of the Events Filter
Event Filter Text Box
The Event Filter text box is where the selected filter settings are displayed. Upon admission, the Event Filter text box lists “All” as the filter. This means that the events (maximum of 1000) for the selected patient tile will be displayed in the Events tab. Event filters can be configured based on time and/or parameters.
7.3.3.3 Time Filters

By indicating a start time and an end time, the view will only include events within the configured time period.

Accessing the Time Filter

To access the Time Filter (shown in FIGURE 7-15), select the Time sidebar button from the Events Filter view. If any previously selected time ranges were specified, they will be displayed in the Event Filter text box beside the Time: label, and in the Start Time and End Time text boxes.

FIGURE 7-15 Time Event Filter

Clearing Time Filters

To remove previously entered time filters, select the Clear button from the Time Filter view.

Configuring a Time Filter

To enter a start time:

1. Select the Start Time text box. The Start Time dialog box (shown in FIGURE 7-16) is displayed.
2. Rotate the dial controls to select the hour and minute values.
3. Rotate the dial controls to select the month, day and year values. Refer to the following value ranges.

<table>
<thead>
<tr>
<th>TIME FRAME</th>
<th>RANGE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Month</td>
<td>Month range to be inclusive within the limits of the oldest event to the most recent event in the selected patient tile’s Events database.</td>
</tr>
<tr>
<td>Day</td>
<td>Day range to be inclusive within the limits of the oldest event to the most recent event in the selected patient tile’s Events database.</td>
</tr>
<tr>
<td>Year</td>
<td>Year range to be inclusive within the limits of the oldest event to the most recent event in the selected patient tile’s Events database.</td>
</tr>
</tbody>
</table>

4. Accept or reject the selection made in the **Start Time** dialog box.
   - The **Done** button accepts the selections. When the **Done** button is selected, the date and time selected populates the Start Time text box, End Time text box, and the Event Filter text box.
   - The **Cancel** button discards the selections, closes the dialog box, and returns to the **Time Event Filter** view.

To enter an end time:

1. Select the End Time text box. The End Time dialog box (shown in FIGURE 7-17) is displayed.
2. To enter the end time, follow steps 2, 3, and 4 from the previous section (entering a Start Time).

7.3.3.4 Parameter Filters
By selecting specific parameters, the Events tab will only display the events that contain the chosen parameter. Parameter filters can be configured for physiological and technical events. The physiological events listed are dependent on the assigned monitoring device and the technical events listed are dependent on the ECG lead wire set currently in use.

Accessing the Parameter Filter
To access the Parameter Filter (shown in FIGURE 7-18), select the Param sidebar button from the All Events view. By default, the physiological events are shown first. If any previously selected parameters were specified, they will be displayed in the Event Filter text box beside the Param: label.

![FIGURE 7-18 Parameter Event Filter View](image)

Configuring a Parameter Filter
To configure a parameter filter:

1. If a technical event filter is needed, select the Tech button.

**NOTE:** By default, the physiological events are displayed when the Param sidebar is selected.

2. Move/remove the selected events by using any of the buttons described in the following table:

<table>
<thead>
<tr>
<th>BUTTON</th>
<th>FUNCTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Remove All</td>
<td>Removes all the events listed in the Selected Events list box and from the Event Filter text box.</td>
</tr>
<tr>
<td>Add/Right Arrow</td>
<td>Adds the selected event to the Selected Events list box, and to the Event Filter text box.</td>
</tr>
<tr>
<td>Remove/Left Arrow</td>
<td>Removes the chosen event from the Selected Events list box and from the Event Filter text box.</td>
</tr>
</tbody>
</table>

**NOTE:** A maximum of five physiological and/or technical events may be added to the Selected Events list at one time.
3. Select the **Done** button to apply the selected event filter and return to the **List** view or select the **Cancel** button to restore the previously chosen filter settings.

### 7.3.3.5 Printing an Events List Report from the List View

Multiple hours of event data can be printed to an Events List Report directly from the **List** view.

To indicate the time interval for printing this report:

1. From the **List** view, select the **Print** menu bar button. The Print Selection dialog box [shown in FIGURE 7-9] is displayed.
2. Select the **Print Current** button, a time interval button, or the **Cancel** button.
   - Select the **Print Current** to create a one page Trend List Report.
   - Select one of the time interval buttons. The choices are print 1 hour, 2 hours, 4 hours, 8 hours, 12 hours, and 24 hours.
3. Select the **Cancel** button to close the dialog box without generating the print request.

For additional information about this report, refer to the “Event List Report” on page 8-21.

### 7.3.3.6 Sidebar Buttons

The following sidebar buttons can be used to expand the functionality of this view.

**View Waveforms**

For additional information regarding the functionality of this sidebar button, refer to “Accessing the Waveforms View” on page 7-35.

**Events Filtering**

For additional information regarding the functionality of this sidebar button, refer to “Event Filtering” on page 7-29.

**Trends**

Select the **Trends** sidebar button to navigate to the Trends dialog. The Trends data (Graphic view or Trends List view) displays based on the configuration of the Trend Access Order in the Unit Choices dialog.

Selecting the Trends sidebar button while in the Events tab displays the trend record with the date and time prior to the event data in time selected either in the Graphic or List trend display.

**Disclosure**

Select the **Disclosure** sidebar button to navigate to the Disclosure dialog. The date and time associated with the highlighted Trend Record or vertical marker in Graphic Trend will be used to navigate to the Disclosure dialog. The most current Disclosure data to the nearest minute will be displayed at the top of the Disclosure dialog.
Selecting the Disclosure sidebar button while in the Events tab displays disclosure data closest and most current to the selected event date and time. Normal Screen
Select the Normal Screen sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

7.3.3.7 Troubleshooting
This section lists some of the potential messages and issues that may occur while using the List view of the Events tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sidebar buttons disabled when Events tab selected.</td>
<td>Patient tile not selected.</td>
<td>Select a patient tile.</td>
</tr>
<tr>
<td></td>
<td>There is no patient admitted to the selected tile.</td>
<td>Admit a patient to the tile.</td>
</tr>
</tbody>
</table>

| No patient in the tile | Patient tile not selected. | Admit a patient to the tile. |
| | There is no patient admitted to the selected tile. | |
| | Patient tile not selected. | Admit a patient to the tile. |

| No data in Events list. | Patient tile not selected. | Select a patient tile. |
| | None of the existing event data meets the criteria of the current event filter. | Change the event filter. |
| | There is no patient admitted to the selected tile. | |

| Events list column headers not displayed. | Patient tile not selected. | Select a patient tile. |
| | There is no patient admitted to the selected tile. | |

| The View Waveform and Events Filtering sidebar buttons are disabled. | The Events List is empty or none of the existing event data meets the criteria of the current event filter. | Change the event filter. |
| | There is no patient admitted to the selected tile. | |

| Some items cannot be deleted. | Specific types of events cannot be removed from the Event list. | Refer to the Alarms and Events chapter. |

| Events were deleted from the database. Press the OK button to refresh the list. | This message will appear if the Events List is open at the ViewStation or WorkStation when an event is deleted from the host central station Events database. | Select the OK button. |

| The Print menu button is disabled. | The event data is loading. | Wait until the event data is loaded. |

| Events are filtered but an event filter was not created. | A system event filter may have been created. | Contact the System Administrator to see if a system event filter exists. |

* Messages are shown in all bold text.
7.3.4 Events Tab (Waveforms View)
The Waveforms view (shown in FIGURE 7-19) of the Events tab displays waveform data for the selected patient. In order for the event to display in the Waveforms view, there must be waveform data associated with the event. Physiological alarm events and ‘marked events’ are stored as waveform data. All previously collected data is displayed, even if the parameter is not currently being monitored.

7.3.5 Accessing the Waveforms View
1. Select the VIEW button on the desired patient tile. The Bedside tab is displayed.
2. Select the Events tab. The List view is displayed.
3. Select a physiological alarm event or marked event (for example, High Heart Rate Alarm) from the List view for the selected patient, then select the View Waveforms sidebar button. The selected event is displayed in the Waveforms view (shown in FIGURE 7-19).

7.3.5.1 Components of the Waveforms View
Patient Demographic Area
The patient demographic area displays the patient’s last name and bed number, if available.
Digital Data Area
The following parameters can display in the digital data area depending on the bedside monitor configuration.

- PVC/min
- Pause/min
- QT/QTc/QT-HR
- Temperature 1
- Temperature 2
- Temperature 3
- T Blood
- N2O
- CO/CI
- CCO/CCI
- CCO STAT/CCI STAT
- SVR
- SVRI
- SvO₂
- ScvO₂
- PAWP
- BIS/BIS L/BIS R

NOTE: The Temperature 1, Temperature 2, Temperature 3 (V 12/V 21 only), and T Blood parameters are only displayed when they are used in the calculation of the Delta Temperature parameter.

Time Interval Area
The time interval area is displayed in one second intervals. A total of nine seconds can be viewed on the screen at one time. Each second shown displays a time label over each one-second grid line in the waveform area.

Event Description Area
The event description area displays the event that was selected in the Event List view. For example, a High Heart Rate Alarm.

NIBP Area
The NIBP area displays the NIBP systolic pressure, diastolic pressure, mean pressure, and elapsed time.

Date Area
The date area indicates the date the event occurred and was stored in the Events database.
Data Display Area
If data is available, the Waveforms view displays 20 seconds of waveform data. The initial Waveforms view displays the first nine seconds of waveform data before the alarm was triggered, and can display three to four waveforms and/or digital data tiles at one time.

Colored Vertical Cursor Line
The colored vertical cursor line extends from the first waveform to the last waveform on the display. The colored vertical cursor line can be moved to indicate the specific time within the event.

7.3.5.2 Navigating in the Waveforms View of the Events Tab
Navigating in the Waveforms view is accomplished by using buttons. When the Waveforms view is first opened, the colored vertical cursor line will be positioned in the far right position.

Vertical Navigation Buttons

Double Up Arrow Button
Select the double up arrow push button to display the previous page of waveforms in the ordered set. This button will be disabled if the first waveform in the ordered set is already visible in the data display area.

Up Arrow Button
Select the up arrow button to move the group of waveforms displayed up one position within the set of available waveforms. This button will be disabled if the first waveform in the ordered set is already visible in the data display area.

Double Down Arrow Button
Select the double down arrow button to display the next page of waveforms in the ordered set. This button will be disabled if the last waveform in the ordered set is already visible in the data display area.

Down Arrow Button
Select the down arrow button to move the group of waveforms displayed down one position within the ordered set of available waveforms. This button will be disabled if the last waveform in the ordered set is already visible in the data display area.

Horizontal Navigation Buttons

Previous Event Button
Select the Previous Event to locate and display an event that occurred earlier than the current event time began. This button will be disabled when there are no more events that have a time less than the current event time.
Double Left Arrow Button
Select the **double left arrow** button to display the previous nine seconds of event data, provided this data was collected. This button will be disabled if the vertical cursor is positioned at the beginning of the stored data for the event.

Left Arrow Button
Select the **left arrow** button to move the colored vertical cursor line one second back in time. This button will be disabled if the vertical cursor is positioned at the beginning of the stored data for the event.

Right Arrow Button
Select the **right arrow** button to move the colored vertical cursor line one second forward in time. This button will be disabled if the vertical cursor is positioned at the end of the stored data for the event.

Double Right Arrow Button
Select the **double right arrow** button to display the next nine seconds of event data, provided this data was collected. This button will be disabled if the vertical cursor is positioned at the end of the stored data for the event.

Next Event Button
Select the **Next Event** button to locate and highlight an event that occurred after the currently highlighted event in the list box.

**7.3.5.3**

ST Button
Select the **ST** button to open the **Waveforms** view ST dialog box.

Waveforms View ST Dialog Box
The **Waveforms** view ST dialog box (shown in FIGURE 7-20) displays the current and reference values for all the leads where ST was performed during the selected event. The number of leads that display in this dialog box is dependent on the number of leads used during the event. If the ST option has been enabled, the ST button will be enabled even if ST is not currently being monitored. If ST was never enabled for the patient tile, the ST button will be disabled.
Date and Time Text Area
The date/time text area, located at the top of the Waveforms view ST dialog, displays the date and time the ST data displayed was acquired. When the ST dialog box navigation buttons (moving forward or backward in time) are used, this area will be updated to reflect the date and time of the ST data currently displayed in the dialog box.

Left Arrow Button
Select the left arrow button to view the ST data for the previous second of the event.

Right Arrow Button
Select the right arrow button to view the ST data for the next second of the event.

Done Button
Select the Done button to close the Waveforms view ST dialog.

7.3.5.4 Comments Dialog Box (Optional)
With the addition of system-wide paging, a paging notification can be sent to clinicians to alert them that a specific event type has occurred. If a page was sent for a selected event in the Waveforms view, an asterisk (*) will appear beside the event in the event description area (shown in FIGURE 7-19). The paging information related to the event can be viewed by selecting the Comments sidebar button. The Comments dialog box will display either a Page Sent or a Page Failed message. If the selected event does not have an asterisk beside the event description, this indicates that a page has not been sent for the selected event. In this case, the Comments sidebar button will be disabled.

7.3.5.5 Sidebar Buttons
The Waveforms view may contain up to six sidebar buttons.

View List
Select the View List sidebar button to return to the Events list.
Lead
Select the **Lead** sidebar button to cycle through each of the waveforms in the ECG lead ordered set. The **Lead** sidebar button is not available when using a:

- Telepack device with a 3 or 5-wire lead set
- Telepack with Nonin SpO2 module with a 3-wire lead set

All ECG
Select the **All ECG** sidebar button to access the All ECG mode of the **Events** tab. When the **All ECG** button is white, this indicates the **Events** tab is in the All ECG mode. The **Waveforms** view remains in the All ECG mode until the **All ECG** sidebar button is selected again. The **All ECG** sidebar button is not available when using a:

- Telepack device with a 3 or 5-wire lead set
- Telepack with Nonin SpO2 module with a 3-wire lead set

**NOTE:** If a 5-wire or 12-lead set was previously used for the selected patient, the All ECG sidebar push button will always be enabled.

Measure
Select the **Measure** sidebar button to activate the Measure dialog box (shown in FIGURE 7-3). The Measure dialog contains reference and measurement lines to calculate the time and amplitude of an ECG waveform. For instructions on the use of the Measure button, refer to “Performing ECG Waveform Measurements” on page 7-5.

**NOTE:** The Print menu bar will be enabled when the Measure dialog box is open.

Comments (Optional)
This sidebar button was previously described in the “Comments Dialog Box (Optional)” on page 7-39. If the Paging feature is unavailable at the central station, the **Comments** sidebar button will not be displayed.

Normal Screen
Select the **Normal Screen** sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

7.3.5.6 Printing an Event Zoom In Report from the Waveforms View
To print a Event Zoom In Report from the **Waveforms** view, select the **Print** button from the menu bar (shown in FIGURE 2-1).
### Troubleshooting

This section lists some of the potential messages and issues that may occur while using the **Waveforms** view of the **Events** tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sidebar buttons disabled when Events tab selected.</td>
<td>Patient tile not selected.</td>
<td>Select a patient tile.</td>
</tr>
<tr>
<td></td>
<td>There is no patient admitted to the selected tile.</td>
<td>Admit a patient to the tile.</td>
</tr>
<tr>
<td>No patient in the tile</td>
<td>There is no patient admitted to the selected tile.</td>
<td>Admit a patient to the tile.</td>
</tr>
<tr>
<td></td>
<td>Patient tile not selected.</td>
<td>Select a patient tile.</td>
</tr>
<tr>
<td>No Waveform data is collected for this event.</td>
<td>The selected event is not a physiological alarm event.</td>
<td>Refer to the Alarms and Events chapter.</td>
</tr>
<tr>
<td>The correct alarmed event is selected from the List view but no waveform is displayed in View Waveforms view.</td>
<td>A 3-lead wire set is being used and the active lead was changed from when the event originally occurred.</td>
<td>Return to the lead that was active when the event occurred, then try to view the waveform again.</td>
</tr>
<tr>
<td>Events were deleted from the database. Press the OK button to refresh the list.</td>
<td>When viewing a patient tile’s Event Zoom In tab at the ViewStation or WorkStation, pressing the Next Event and/or Previous Event button to an event that has been deleted from the database, will display the message “Events were deleted from the database. Press the OK button to refresh the list.” at the ViewStation or WorkStation.</td>
<td>Select the OK button to close the message box.</td>
</tr>
<tr>
<td>Next Event button is disabled.</td>
<td>The last event is selected in the list, therefore, there are no events to view next.</td>
<td>Select an earlier event.</td>
</tr>
<tr>
<td>Previous Event button is disabled.</td>
<td>The first event is selected in the list, therefore, there are no previous events to view.</td>
<td>Select a later event.</td>
</tr>
<tr>
<td>ST button is disabled</td>
<td>ST was never enabled for the selected patient tile.</td>
<td>If the patient tile requires ST analysis, and there is an ST license available, enable ST from the bedside monitor.</td>
</tr>
<tr>
<td>Lead sidebar button is disabled</td>
<td>The selected device is a:</td>
<td>No solution required.</td>
</tr>
<tr>
<td></td>
<td>• Telepack device using a 3 or 5-wire lead</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Telepack with Nonin SpO₂ module attached with a 3-wire lead set</td>
<td></td>
</tr>
<tr>
<td></td>
<td>The All ECG sidebar button is latched.</td>
<td>Select the All ECG sidebar button to enable the Lead sidebar button.</td>
</tr>
<tr>
<td>CO/CI data is not displayed in Waveforms view but is displayed in Events list.</td>
<td>Currently, CO/CI waveform data is not displayed in the Waveforms view.</td>
<td>No solution required.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>No more events found</strong></td>
<td>The colored vertical cursor is positioned at the first or the last event in the Events database.</td>
<td>No solution required.</td>
</tr>
<tr>
<td><strong>Comments sidebar button is not displayed in sidebar.</strong></td>
<td>The Paging option is not present on the central station licensing key.</td>
<td>Obtain a licensing key that contains the Paging option.</td>
</tr>
<tr>
<td><strong>Comments sidebar button is disabled.</strong></td>
<td>No paging information was sent for the selected event.</td>
<td>If a page should have been sent for the selected event type, verify the Page On Alarm settings in the Alarm Responses tab.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
7.4 Disclosure Tab

The Disclosure tab provides access to a patient’s disclosure database. The disclosure database is a collection of continuous, running, available waveforms that can store between 1 and 72 hours of disclosure data, depending on the type of license key that is issued for the system. License keys are issued for 1-hour, 24-hour, 48-hour, and 72-hour periods.

Each record in the database includes the time of the record, digital data, patient alarm events and all of the waveforms (with the exception of the O2 and Agent waveforms) that were available for the patient at the time of the record storage. When a patient’s disclosure database reaches the maximum number of records allowed, the next new record replaces the oldest record.

The Disclosure tab is available at the ViewStation and the WorkStation (shown in FIGURE 7-21) and functions in the same manner as it does at the host central station. The Disclosure tab will display the disclosure data from the central station, provided the patient tile is communicating with the central station.

A patient’s disclosure data can be viewed in two different formats. The default view is the Compressed view. The other is the Waveforms (magnified) view. Both views are discussed in the following sections.

7.4.1 Accessing the Disclosure Tab

1. Select the VIEW button on the desired patient tile. The Bedside tab will be displayed.
2. Select the Disclosure tab. The Disclosure tab (shown in FIGURE 7-21) Compressed view is displayed.

FIGURE 7-21 Disclosure Tab - Compressed View (Central Station, ViewStation, WorkStation)
7.4.1.1 Components of the Compressed View

Patient Demographic Field
The patient demographic field displays the last name and bed number of the selected patient, if this data was entered.

Waveform Area
The waveform area of the Disclosure tab provides a compressed view of the selected patient's historical waveform data. A maximum of 10 minutes of waveform data can be scrolled into the viewable area of the display. Each row of historical disclosure data in the waveform area is displayed in one minute intervals. Each waveform is displayed in the designated color of the parameter.

On initial entry to the Compressed view of the Disclosure tab, the ECG waveform displayed is the first waveform shown in the main screen.

The waveforms displayed in the Disclosure tab are based on the sensors currently in use. Any previously collected data is displayed, even if the parameter is not currently being monitored. Newly stored waveform data does not immediately update the disclosure waveform area. Selecting another control in the Disclosure tab causes the waveform area to refresh and updates the newly stored waveform data.

Blank sections of a disclosure waveform occur during the storage of a row of data when:

- The system time has been altered
- The device has been placed into Standby mode
- The device is in a Communications Lost or Comm Loss - Check Bedside Monitor state
- An Auto Sync event occurs
- A Break of Data event occurs

The Panorama Central Station has been restarted. The entire waveform area will be blank if the selected patient tile does not have a monitoring device attached or if there is no historical disclosure data in the patient's database.

Time, Date, and Sensor Label
The actual time the data was stored is displayed on the left side of the waveform. The format of the time display follows the System Time Format as defined in the “Date/Time Tab” on page 9-41.

The date the selected waveform data was recorded is displayed in the upper right corner of the waveform area (shown in FIGURE 7-21).

A sensor label for the selected waveform data is also displayed in the upper right corner of the waveform area, just below the date (shown in FIGURE 7-21).
Event Markers
The Disclosure tab uses event markers to indicate that an event (such as an alarm condition) has occurred. Event markers are small, color coded, horizontal lines that are drawn above the disclosure waveform at the onset of the event. The color codes for the event markers are defined as follows:

- Red event markers indicate that a Level 1 alarm occurred
- Yellow event markers indicate that either a Level 2 or a Level 3 alarm occurred
- Blue event markers indicate that a user marked event occurred at either the Panorama Central Station or the bedside monitor and/or that the Nurse Call button was pressed on the Panorama Telepack
- Orange event markers indicate that a Communications Lost or Comm Loss - Check Bedside Monitor event or a clock adjustment occurred

7.4.1.2 Navigating in the Waveform Area
If the Disclosure tab waveforms exceed the capacity of the viewable screen, use the navigational buttons to locate the desired waveform. The buttons are described as follows:

New Data Button
Select the to display the most current disclosure data for the selected patient.

Old Data Button
Select the to display the oldest disclosure data for the selected patient.

Previous Event Button
Select the Previous Event button to locate the next event in the patient’s disclosure database that occurred before the time at the upper left corner of the compressed view.

Next Event Button
Select the Next Event button to locate the next event marker in the Compressed view that occurred after the current marker time.

Zoom In Button
Select the Zoom In button to magnify a selected area from the compressed view of the patient’s disclosure waveform.

Zoom Out Button
Select the Zoom Out button to return the magnified view of the patient’s disclosure waveform to the compressed view of the waveform. The Zoom Out button is only enabled when the magnified view of the waveform is displayed.

Scrolling through the Waveforms
If the disclosure database exceeds the capacity of the viewable screen (10 minutes), there are two methods of scrolling through the waveforms. Arrow buttons located along the right side of the view (next to the sidebar) function as follows:
• Use the single up-arrow and single down-arrow buttons to scroll up or down a single row.
• Use the Page Up (double up-arrow) and Page Down (double down-arrow) buttons to scroll through one page of disclosure waveforms at a time to view earlier or later data.
### 7.4.1.3 Changing the Displayed Waveform

Select the **Display Choices** sidebar button to select the waveform that will be displayed in the **Compressed** view of the **Disclosure** tab. The **Display Choices** dialog box (shown in FIGURE 7-22) displays the choices that are available based on the monitoring device, ECG lead set, and utilized sensors. Any previously used parameters, ECG lead set or sensors are displayed, even if they are not currently in use.

**NOTE:** The sensor selected in the Display Choices dialog will remain until the Disclosure tab is exited.

**NOTE:** The O2 and Agent parameters are not available in the Display Choices list box.

To change the sensor currently displayed in the **Disclosure** tab:

1. Select the **Display Choices** sidebar button. The **Display Choices** dialog box is displayed.

![FIGURE 7-22 Display Choices Dialog Box](image)

2. Select the desired waveform to be displayed from the **Display Choices** dialog box.
   - Select the **Cancel** button to return to the **Disclosure** tab.
   - Select the **Done** button return to the **Disclosure** tab and view the selected waveform.

### 7.4.1.4 Locating Specific Disclosure Data

Select the **Skip To** sidebar button to choose a different time period from the selected patient’s disclosure database. The date and time the data was stored is displayed on the left side of the waveform.

**NOTE:** The time and date selected in the Skip To dialog will remain until the Disclosure tab is exited.

To locate a specific time in the disclosure database:

1. Select the **Skip To** sidebar button. The **Skip To** dialog box (shown in FIGURE 7-23) will be displayed.
FIGURE 7-23  Skip To Dialog Box

2. Rotate the dial controls to select the time and date to be displayed. Ensure that the selected time and date are within the timeframe of the database.
   - Select the Cancel button to return to the Disclosure tab.
   - Select the Done button to return to the Disclosure tab with the selected date displayed in the date area, and the selected time displayed in the upper left corner of the waveform area.

7.4.1.5 Printing a Full Disclosure Report from the Compressed View

Multiple hours of disclosure data can be printed to a Full Disclosure Report directly from the Compressed view.

To indicate the time interval for printing this report:

1. From the Compressed view, select the Print menu bar button. The Print Selection dialog box (shown in FIGURE 7-9) is displayed.
2. Select the Print Current button, a time interval button, or the Cancel button.
   - Select the Print Current to create a one page Trend List Report.
   - Select one of the time interval buttons. The choices are print 1 hour, 2 hours, 4 hours, 8 hours, 12 hours, and 24 hours.
3. Select the Cancel button to close the dialog box without generating the print request.

7.4.1.6 Sidebar Buttons

The following sidebar buttons are used to expand the functionality of this view.

Display Choices
Select the Display Choices sidebar button to select the waveform that will be displayed in the Compressed view of the Disclosure tab. For additional information regarding this option, refer to “Changing the Displayed Waveform” on page 7-47.

Skip To
Select the Skip To sidebar button to choose a different time period from the selected patient's disclosure database. For additional information regarding this option, refer to “Locating Specific Disclosure Data” on page 7-47.
Trends
Select the **Trends** sidebar button to navigate to the Trends dialog. The Trends data (Graphic view or Trends List view) displays based on the configuration of the Trend Access Order in the Unit Choices dialog.

When disclosure data is highlighted, the time and date associated with the start of the highlighted text disclosure data is the time and date used to access the trend data when the Trends sidebar button is selected. When no disclosure data is highlighted, the date and time associated with the top row of disclosure data is used to access the trends data when the Trends sidebar button is selected.

Events
Select the **Events** sidebar button to navigate to the Events dialog. The date and time associated with the highlighted Trend Record or vertical marker in Graphic Trend will be used to navigate to the Events dialog. The most current Event nearest to the Trend Record time and date will be selected.

When disclosure data is highlighted, the time and date associated with the start of the highlighted text disclosure data is the time (or close to the time) and date used to access the event data when the Events sidebar button is selected. When no disclosure data is highlighted, the date and time associated with the top row of disclosure data is used to access the event data when the Events sidebar button is selected.

Normal Screen
Select the **Normal Screen** sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

7.4.1.7 Troubleshooting
For **Compressed** view troubleshooting information, refer to "Troubleshooting" on page 7-55.

7.4.2 Disclosure Tab (Waveforms View)
The **Waveforms** view displays a selected area from the **Compressed** view of the patient’s disclosure waveform (shown in FIGURE 7-24).
7.4.2.1 Accessing the Waveforms View

1. Select the **VIEW** button on the desired patient tile. The **Bedside** tab is displayed.

2. Select the **Disclosure** tab. The **Waveforms** view (shown in FIGURE 7-24) is displayed.

3. Select a zoomed in area using one of the following methods:
   - Selecting the **Zoomed In** button without highlighting the waveform displays the magnified view beginning with the time at the upper left corner of the compressed waveform area.
   - A section of the waveform can be highlighted by selecting a starting point and dragging to an end point. Selecting the **Zoom In** button displays a magnified view of the highlighted section.

7.4.2.2 Components of the Waveforms View

**Patient Demographic Field**
The patient demographic area displays the patient's bed number, if available.

**Digital Data Field**
The following parameters can display in the digital data area depending on the bedside monitor configuration.

- PVC/min
- Pause/min
- QT/QTc/QT-HR
- Temperature 1
- Temperature 2
- Temperature 3
- T Blood
- N2O
- CO/CI
- CCO/CCI
- CCO STAT/CCI STAT
- SVR
- SVRI
- SvO2
- ScvO2
- PAWP
- BIS/BIS L/BIS R

**NOTE:** The Temperature 1, Temperature 2, Temperature 3 (V 12/V 21 only), and T Blood parameters are only displayed when they are used in the calculation of the Delta Temperature parameter.

**Time Interval Field**
The time interval field is displayed in one second intervals. A total of nine seconds can be viewed on the screen at one time. Each second shown displays a time label over each one-second-grid line in the waveform area.

**Event Description Field**
The event description field displays the event that was selected in the Event List view. For example, a High Heart Rate Alarm.

**NIBP Field**
The NIBP field displays the NIBP systolic pressure, diastolic pressure, mean pressure, and elapsed time.

**Date Field**
The date area indicates the date the event occurred and was stored in the Events database.

**Data Display Area**
If data is available, the Waveforms view displays 20 seconds of waveform data. The initial Waveforms view displays the first nine seconds of waveform data before the alarm was triggered, and can display three to four waveforms and/or digital data tiles at one time.

**Colored Vertical Cursor Line**
The colored vertical cursor line extends from the first waveform to the last waveform on the display. Each movement of the colored vertical cursor line indicates the specific time in the event.
7.4.2.3 Navigating in the Waveform View of the Disclosure Tab

Navigating in the Waveform view is accomplished by using buttons. When the Waveform view is first opened, the colored vertical cursor line will be positioned in the far left position.

Vertical Navigation Buttons

Double Up Arrow Button
Select the double up arrow push button to display the previous page of waveforms in the ordered set. This button will be disabled if the first waveform in the ordered set is already visible in the data display area.

Up Arrow Button
Select the up arrow button to move the group of waveforms displayed up one position within the ordered set of available waveforms. This button will be disabled if the first waveform in the ordered set is already visible in the data display area.

Double Down Arrow Button
Select the double down arrow button to display the next page of waveforms in the ordered set. This button will be disabled if the last waveform in the ordered set is already visible in the data display area.

Down Arrow Button
Select the down arrow button to move the group of waveforms displayed down one position within the ordered set of available waveforms. This button will be disabled if there are less than nine seconds of earlier data to display.

Horizontal Navigation Buttons

Previous Event Button
Select the Previous Event button to locate and display an event that occurred earlier than the current event time began. This button will be disabled when the colored vertical cursor is positioned at the start time of the first event in the database.

Double Left Arrow Button
Select the double left arrow button to display the previous nine seconds of event data, provided this data was collected. This button will be disabled if the colored vertical cursor line is positioned at the start time of the first record in the database.

Left Arrow Button
Select the left arrow button to move the colored vertical cursor line one second back in time. This button will be disabled when the colored vertical cursor is positioned at the start time of the first record in the database.
Right Arrow Button
Select the **right arrow** button to move the colored vertical cursor line one second forward in time.

Double Right Arrow Button
Select the **double right arrow** button to display the next nine seconds of event data, provided this data was collected.

Next Event Button
Select the **Next Event** button to locate and highlight an event that occurred **after** the currently highlighted event in the **Waveforms** view.

7.4.2.4 ST Button
Select the **ST** button to open the Full Disclosure Zoom In ST dialog box.

Full Disclosure Zoom In ST Dialog Box
The Full Disclosure Zoom In ST dialog box (shown in FIGURE 7-20) displays the current and reference values for all the leads where ST was performed during the selected event. The number of leads that display in this dialog box is dependent on the number of leads used during the event. If the ST option has ever been enabled for the selected patient tile, the ST button will be enabled even if ST is not currently being monitored. If ST was never enabled for the patient tile, the ST button will be disabled.

![FIGURE 7-25 Full Disclosure Zoom In ST Dialog](image)

**Date and Time Text Area**
The date/time text area, located at the top of the Full Disclosure Zoom In ST dialog, displays the date and time the ST data displayed was acquired. When the ST dialog box navigation buttons (moving forward or backward in time) are used, this area will be updated to reflect the date and time of the ST data currently displayed in the dialog box.

Left Arrow Button
Select the **left arrow** button to view the ST data for the previous event second.
Right Arrow Button
Select the right arrow button to view the ST data for the next event second.

Done Button
Select the Done button to close the Full Disclosure Zoom In ST dialog.

7.4.2.5 Comments Dialog Box (Optional)
With the addition of system-wide paging, a paging notification can be sent to clinicians to alert them that a specific event type has occurred. If a page was sent for a selected event in the Waveforms view, an asterisk (*) will appear beside the event in the event description area (shown in FIGURE 7-19). The paging information related to the event can be viewed by selecting the Comments sidebar button. The Comments dialog box will display either a Page Sent or a Page Failed message. If the selected event does not have an asterisk beside the event description, this indicates that a page has not been sent for the selected event. In this case, the Comments sidebar button will be disabled.

7.4.2.6 Sidebar Buttons
The Waveforms view may contain up to six sidebar buttons.

Zoom Out
Select the Zoom Out sidebar button to return to the Compressed view of the Disclosure tab.

Lead
Select the Lead sidebar button to cycle through each of the waveforms in the ordered set. The Lead sidebar button is not available when using:
- Telepack device with a 3 or 5-wire lead set
- Telepack with Nonin SpO2 module with a 3-wire lead set

All ECG
Select the All ECG sidebar button to access the All ECG mode of the Disclosure tab. When the All ECG button is white, this indicates the Disclosure tab is in the All ECG mode. The Waveforms view remains in the All ECG mode until the All ECG sidebar button is selected again. The All ECG sidebar button is not available when using:
- Telepack device with a 3 or 5-wire lead set
- Telepack with Nonin SpO2 module with a 3-wire lead set

NOTE: If a 5-wire or 12-lead set was previously used for the selected patient, the All ECG sidebar push button will always be enabled.
Measure
Select the **Measure** sidebar button to activate the Measure dialog box (shown in FIGURE 7-3). The Measure dialog contains reference and measurement lines to calculate the time and amplitude of an ECG waveform. For instructions on the use of the Measure button, refer to “Performing ECG Waveform Measurements” on page 7-5.

**NOTE:** The Print menu bar will be enabled when the Measure dialog box is open.

Comments (Optional)
This sidebar button was previously described in the “Comments Dialog Box (Optional)” on page 7-54. If the Paging feature is unavailable at the central station, the **Comments** sidebar button will not be displayed.

Normal Screen
Select the **Normal Screen** sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

7.4.2.7 Printing a Full Disclosure Zoom In Report from the Waveforms View
To print a Full Disclosure Zoom In Report from the **Waveforms** view, select the **Print** button from the Menu Bar (shown in FIGURE 2-1).

7.4.2.8 Troubleshooting
This section lists some of the potential messages and issues that may occur while using the **Disclosure** tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>No patient in the tile</td>
<td>The selected tile is not actively monitoring a patient.</td>
<td>Choose a tile that is actively monitoring a patient.</td>
</tr>
<tr>
<td>All displays are blank and all buttons [except <strong>Normal Screen</strong>] are disabled.</td>
<td>The selected patient tile does not have any stored historical data.</td>
<td>The selected patient tile has not been monitoring long enough to have stored any data. Wait for new data to accumulate.</td>
</tr>
<tr>
<td>All buttons [except <strong>Normal Screen</strong>] are disabled.</td>
<td>The selected patient tile does not have a device attached.</td>
<td>Choose a tile that is actively monitoring a patient.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
**TABLE:**

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>A section of the waveform is blank.</td>
<td>The system time was altered during the storage of the row of data.</td>
<td>Wait for new data to accumulate.</td>
</tr>
<tr>
<td></td>
<td>The device was placed into Standby mode during the storage of the row of data.</td>
<td>Restore the system to normal operating mode to resume data collection.</td>
</tr>
<tr>
<td></td>
<td>The device experienced a Communications Lost or Comm Loss - Check Bedside Monitor event during the storage of the row of data.</td>
<td>Determine the cause of the Communications Lost or Comm Loss - Check Bedside Monitor condition and rectify to resume data collection.</td>
</tr>
<tr>
<td></td>
<td>The Panorama was restarted during the storage of the row of data.</td>
<td>Wait for new data to accumulate.</td>
</tr>
<tr>
<td>No more events found (Previous Event Button)</td>
<td>There is no event data in the Disclosure database.</td>
<td>Wait until an event is recorded to the Disclosure database. Select the OK button to close the message box and return to the Disclosure tab.</td>
</tr>
<tr>
<td></td>
<td>There are no more events in the patient’s Disclosure database that occurred before the time at the upper left corner of the compressed view.</td>
<td>Select the OK button to close the message box and return to the Disclosure tab.</td>
</tr>
<tr>
<td></td>
<td>The oldest data is at the upper left corner of the compressed view.</td>
<td></td>
</tr>
<tr>
<td>No more events found (Next Event Button)</td>
<td>There is no event data in the Disclosure database.</td>
<td>Wait until an event is recorded to the Disclosure database. Select the OK button to close the message box and return to the Disclosure tab.</td>
</tr>
<tr>
<td></td>
<td>There are no more events in the patient’s Disclosure database that occurred after the time at the lower right corner of the compressed view.</td>
<td>Select the OK button to close the message box and return to the Disclosure tab.</td>
</tr>
<tr>
<td></td>
<td>The most recent data is at the lower right corner of the compressed view.</td>
<td></td>
</tr>
<tr>
<td>Time out of range (Page Up button)</td>
<td>There are less than 10 rows of data that are newer than the top row.</td>
<td>Select the OK button to close the message box and return to the Disclosure tab. Wait for new data to accumulate.</td>
</tr>
<tr>
<td>Time out of range (Page Down button)</td>
<td>There are less than 10 rows of data that are older than the top row.</td>
<td>Select the OK button to close the message box and return to the Disclosure tab. Data is not available.</td>
</tr>
<tr>
<td>Time out of range (single up-arrow button)</td>
<td>There is no new data to display.</td>
<td>Select the OK button to close the message box and return to the Disclosure tab. Wait for new data to accumulate.</td>
</tr>
<tr>
<td>Time out of range (single down-arrow button)</td>
<td>There is no older data to display.</td>
<td>Select the OK button to close the message box and return to the Disclosure tab. Data is not available.</td>
</tr>
</tbody>
</table>

*Messages are shown in all bold text.*
<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Time out of range</strong> (Done button in the Skip To dialog box)</td>
<td>A time and date that is older than the oldest data or newer than the most recent data was entered in the Skip To dialog box.</td>
<td>Select the OK button to close the message box. Select the Skip To sidebar button and enter a time and date that is within the time frame of the database.</td>
</tr>
<tr>
<td><strong>ST button is disabled</strong></td>
<td>ST was never enabled for the selected patient tile.</td>
<td>If the patient requires ST analysis, and there is an ST license available, enable ST for the selected patient tile.</td>
</tr>
<tr>
<td><strong>Comments sidebar button is not displayed in sidebar</strong></td>
<td>The Paging option is not present on the central station licensing key.</td>
<td>Obtain a licensing key that contains the Paging option.</td>
</tr>
<tr>
<td>There is a gap between waveform times</td>
<td>Either the central station and monitoring device were not communicating during that time (Communications Lost or Comm Loss - Check Bedside Monitor) or the system time was changed in the Date/Time tab.</td>
<td>No solution required.</td>
</tr>
<tr>
<td><strong>Comments sidebar button is disabled.</strong></td>
<td>No paging information was sent for the selected event.</td>
<td>If a page should have been sent for the selected event type, verify the Page On Alarm settings in the Alarm Responses tab.</td>
</tr>
<tr>
<td>The current event was deleted from the database. Press the OK button to refresh the data.</td>
<td>This message will be displayed when viewing a patient tile’s Full Disclosure Zoom In tab at the ViewStation or WorkStation, and the Next Event and/or Previous Event button is selected to view an event that has been deleted from the database.</td>
<td>Select the OK button to close the message box.</td>
</tr>
<tr>
<td>The Previous Event button is disabled.</td>
<td>The cursor is positioned at the first record of the Disclosure database.</td>
<td>No solution required.</td>
</tr>
</tbody>
</table>
| **Lead sidebar button is disabled** | The selected device is a:  
- Telepack device with a 3 or 5-wire lead set  
- Telepack with Nonin SpO₂ module attached with a 3-wire lead set | No solution required. |
| The All ECG sidebar button is latched. | Select the All ECG sidebar button to enable the Lead sidebar button. |

* Messages are shown in all bold text.
7.5 **ST Tab**

The ST tab is used to view and configure the current ST templates or view the ST numeric data for the selected patient tile. The number of ST templates or ST numeric data displayed depends on the type of ECG lead set and the monitoring device currently in use.

ST templates are displayed when using a hardwired or wireless 2.4 GHz bedside device, or Telepack device. When using any of these devices, the ST templates can be adjusted directly from the Panorama Central Station.

When using a WMTS 608 wireless bedside device, only numeric data is displayed (shown in FIGURE 7-27). ST configuration settings must be adjusted at the bedside device.

The ST tab is available at the ViewStation and the WorkStation but is for viewing purposes only. When a patient tile is assigned at the ViewStation and/or the WorkStation, it will always display the ST information of the associated host central station patient tile provided the patient tile is in an active state.

Automatic updating of the ST information at the remote workstation will also occur when:

- The monitoring device is placed in the Standby mode at the central station
- The Telepack is not communicating with its host central station (Communications Lost)
- The patient monitor is not communicating with its host central station (Comm Loss - Check Bedside Monitor)
- Communication between the host central station and the ViewStation and/or the WorkStation is lost (Central Communications Lost)

### 7.5.1 Accessing the ST Tab

1. Select the **VIEW** button in the selected patient tile. The **Bedside** tab is displayed.
2. Select the **ST** tab. The **ST** tab (shown in FIGURE 7-26 or FIGURE 7-27) is displayed.

![FIGURE 7-26 ST Tab - Hardwired Bedside Monitor (5-wire lead set)]
7.5.1.1 Components of the ST Tab

Patient Demographic Field
The patient demographic field displays the last name and bed number of the selected patient, if this data was entered.

ST Status Field
The ST Status field displays the status of ST analysis at the Panorama Central Station. The Status field only displays one message at any given time. The available status messages include:
No License indicates that an ST license is not available for the selected patient tile.

Disabled indicates that the ST algorithm on the bedside monitoring device is currently disabled.

Relearning indicates that the ST algorithm is enabled and is in the learning phase of ST analysis.

Real Time indicates that the ST algorithm is enabled, a successful learn phase has been completed, and the most current data is being displayed.

Patient in Standby indicates that the selected patient tile is in a standby state.

Communications Lost indicates that the Panorama Central Station is not communicating with the Telepack.

Comm Loss - Check Bedside Monitor indicates that the Panorama Central Station is not communicating with the bedside monitor.

ST Template Area (Device Dependent)
The ST template area will display ST templates and digital data for each ECG lead that is utilized when using a hardwired bedside device, a wireless 2.4 GHz bedside device, or a Telepack device. ECG lead labels are shown at the top of the ST template area. These labels represent the ST data currently displayed.

NOTE: When using a 3 or 5-wire lead set, ST is invalidated when the measured ST value exceeds range, and/or paced rhythm persists for more than 45 seconds.

When using a View 12™ card, ST is invalidated when the measured ST value exceeds range, and/or paced rhythm persists for more than 30 seconds.

For all lead wire sets, ST is also invalidated during detected episodes of Asystole, V-Fib, V-Tach, and V-Rhythm.

NOTE: ST templates are not displayed when using a WMTS 608 bedside device.

- When utilizing a 3-wire lead set, the ST tab will contain one ST template for the lead that is currently being monitored.
- When utilizing a 5-wire lead set, the ST tab will contain seven ST templates.
- When utilizing a View 12™ card, the ST tab will contain 12 ST templates.

Reference ST
- The reference ST template is displayed in white
- The digital data for the reference ST displays at the bottom of the ST template area
- The reference ST template displays dashes (---) when ST is relearning, when there is no data available, or when the data is invalid

Current ST
- The current ST template is displayed in yellow
- The digital data for the current ST will be shown at the bottom of the ST template area
- The current ST template will display dashes (---) when ST is relearning, when there is no data available, and when the data is invalid
NOTE: ST templates and digital data will not be shown in the ST template areas during an ST learn phase.

FIGURE 7-29 Point Control Arrows in ST Tab (5-wire Lead Set)

7.5.1.2 Configuring ST (Device Dependent)
Configuration settings are device dependent, except for the Telepack which is turned on at the central station, and can only be performed when using a hardwired bedside device, a wireless 2.4 GHz bedside device, or a Telepack device. When using a WMTS 608 bedside device, ST configuration settings must be performed at the bedside monitor.

NOTE: If the ECG Lead set is changed during ST configuration, the ST tab will automatically be closed and returned to the normal screen mode.

To configure ST points:

1. Select the **Config ST** sidebar button in the ST tab. The ISO, J/ST and ST Point controls shown in FIGURE 7-29 are displayed in the ST tab.
2. Select a point control arrow to configure the ISO reference line. The ISO reference line is displayed in white.
   - Select the left arrow to move the ISO reference line to the left one step setting.
   - Select the right arrow to move the ISO reference line to the right one step setting.
3. Select a point control arrow to configure the J/ST reference line. The J reference line is displayed in yellow. The ST reference line is displayed in green.
   - Select the left arrow to move the J/ST reference line to the left one step setting.
   - Select the right arrow to move the J/ST reference line to the right one step setting.

NOTE: Due to monitor resolution restrictions, adjustments to the J/ST reference lines may not always be shown.
4. Select the **ST Point** button to select the desired ST Point. Changing the ST point will move the ST reference line to the right of the J reference line for the selected number of milliseconds. The options that are available for ST Point include 40, 60, 80 and 60/80 milliseconds.

5. Select the **Done** button.

### 7.5.1.3 Simultaneous ST Configuration Editing

If using a bedside device with the Panorama Central Station, ST configuration changes cannot be made at the bedside monitor and at the central station simultaneously. When an attempt is made to do this, the bedside device will take precedence and the message **These settings are currently being edited remotely.** will be displayed.

These are the scenarios in which this message will be displayed:

- The ST Menu is active at the bedside monitor and then the ST tab is opened at the central station
- The ST tab is active at the central station and then the ST Menu is opened at the bedside monitor

To close this message dialog, select the **OK** button. Acknowledging this message will convert the ST tab at the central station to a read-only state and editing will not be permitted. Once the ST Menu is closed at the bedside device, the ST tab will become active again and editing can resume.

In both of these scenarios, unsaved changes made in the central station Config ST tab will be discarded, and the system will return to the ST tab.

For additional information regarding the bedside monitor menus, refer to the bedside monitor’s Operating Instructions:

- Passport 2 (P/N 0070-00-0649-XX)
- Passport V (P/N 0070-00-0704-XX)
- Spectrum (P/N 0070-00-0648-XX)
- Spectrum OR (P/N 0070-00-0670-XX)
- V 12/V 21 (P/N 0070-00-0727-XX)
- DPM 6 (P/N 046-000184-00)
- DPM 7 (P/N 046-000186-00)
- Passport 12M/Passport 17M (P/N 046-005013-00)
- Passport 8/Passport 12 (P/N 046-004417-00)
- T1 monitor (P/N 046-002376-00)

### 7.5.1.4 Sidebar Buttons

The following sidebar buttons are used to expand the functionality of this view.
Config ST
Select the Config ST sidebar button to configure the ST templates in the ST tab. Selecting the Config ST sidebar button will display the controls (shown in FIGURE 7-29) needed to configure the ST templates. When the Config ST sidebar button is selected, it is replaced by the Done sidebar button.

NOTE: The Config ST sidebar button is not available when using a DPM 6/DPM 7, Passport 12M/Passport 17M, Passport 8/Passport 12, T1 or WMTS 608 wireless bedside device.

NOTE: The Config ST sidebar button is not available at the ViewStation or the WorkStation.

Done
Select the Done sidebar button to confirm the selected ST configuration.

Normal Screen
Select the Normal Screen sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

7.5.2 Generating an ST Report
To review the current and reference digital values with the associated ST templates for the selected patient tile, generate an ST Report. This report can be generated directly from the ST tab by selecting the Print menu bar button.

NOTE: The Print button is not available when using a WMTS 608 wireless bedside device.

For additional information regarding the ST Report, refer to the “ST Report” on page 8-14.

7.5.3 Troubleshooting
This section lists some of the potential messages and issues that may occur while using the ST tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>No Patient Selected! is displayed in the ST tab demographics text box</td>
<td>This message appears in the demographic field when an empty patient tile has been selected.</td>
<td>Select an active patient tile.</td>
</tr>
<tr>
<td>No License is displayed in ST tab status text box</td>
<td>The monitoring device does not have an ST license or ST has been disabled.</td>
<td>Obtain an ST license and/or enable ST at the monitoring device.</td>
</tr>
<tr>
<td>Disabled is displayed in ST tab status text box</td>
<td>ST is disabled at the monitoring device.</td>
<td>Enable ST.</td>
</tr>
<tr>
<td>Relearning is displayed in ST tab status text box</td>
<td>ST was just enabled at the monitoring device.</td>
<td>Wait for a successful learn phase to complete.</td>
</tr>
<tr>
<td>Real Time is displayed in ST tab status text box</td>
<td>ST is enabled and a successful learn phase has been completed.</td>
<td>No solution required.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
### MESSAGE/ISSUE *

| **Patient in Standby** is displayed in ST tab status text box | The selected patient tile is in a standby state. | The message will be removed when monitoring is resumed. |
| **Communications Lost** or **Comm Loss - Check Bedside Monitor** is displayed in the ST tab status text box | The central station is in a Communications Lost state. | The message will be removed when communications is restored. |
| “---” displays in ST template | ST data displays as invalid because monitor may be in a learning phase or patient is in an ECG rhythm that does not allow for ST monitoring (V-Fib, Asystole, etc.) | Wait until learn phase is complete and/or re-establish normal ECG. |
| ST templates are present but contain no ST data. | Monitoring device is in relearn mode. | Wait for Real Time to display in ST tab status text box. |
| ST templates are not available in the ST tab. | A WMTS 608 wireless bedside device is in use. | View the ST templates at the bedside device. |
| The **Print** menu bar button is disabled. | The selected patient tile does not have an ST license. | If necessary, activate an ST license for the patient tile. |
| | The selected tile is in a learn phase. | Wait for the learn phase to complete. |
| | ST has been disabled. | If an ST license is available, enable a license for the patient tile. |
| | The patient tile is in the **Bedside and Panorama Standby** mode. | If necessary, remove the patient tile from this mode. |
| | The patient tile is in the **Panorama Standby Only** mode. | If necessary, remove the patient tile from this mode. |
| | The patient tile is in the **Communications Lost** or the **Comm Loss - Check Bedside Monitor** state. | Restore communications with the monitoring device. |
| | The **These settings are currently being edited remotely** message is displayed. | Select the **OK** button to close the message box and enable the sidebar button. |
| | The ST tab is displayed and the assigned device is a WMTS 608 bedside monitor. | ST numerics can be documented by generating a Trend List Report. |
| | The patient tile is in a **Waiting for Patient** state. | Restore communications with the bedside monitor. |

* Messages are shown in all bold text.
### ST Tab

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>These settings are currently being edited remotely.</strong></td>
<td>The ST Menu is opened at the bedside monitor while in the Config ST mode at the central station.</td>
<td>Wait until the changes are made at the bedside monitor. If necessary, close the ST menu at the bedside to enable the ST menu at the central station.</td>
</tr>
<tr>
<td></td>
<td>The ST tab is active at the central station and then the ST Menu is opened at the bedside monitor.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>The central station <strong>Config ST</strong> tab is active and then the ST Menu is opened at the bedside monitor.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>The Config ST tab at the central station was closed and all changes were discarded.</td>
<td>Use the bedside monitor to make the ST configuration changes. Close the ST Menu at the bedside monitor and then reenter the Config ST tab at the central station.</td>
</tr>
<tr>
<td></td>
<td>The <strong>Config ST</strong> sidebar button is disabled.</td>
<td>Select the <strong>OK</strong> button to close the message box and enable the sidebar button. Close the ST Menu at the bedside device.</td>
</tr>
</tbody>
</table>

*Messages are shown in all bold text.*
This page intentionally left blank.
8.0 Report Functions

This section outlines the types of reports that can be generated by the Panorama Central Station.

- Patient Reports Tab
- System Reports Tab
- Print Status Tab
- Additional Reports
8.1 Patient Reports Tab

The Patient Reports tab (shown in FIGURE 8-1) is used to print patient-specific reports.

The following reports can be printed from the Patient Reports tab:

- All Strips Report
- Full Disclosure Report
- ST Report
- Trend List Report
- Event List Report
- Patient Alarm Report

8.1.1 Accessing the Patient Reports Tab

1. From the menu bar, select the Report button.
2. Select the Patient Reports tab. The Patient Report tab is displayed.
3. Choose a patient tile by selecting its digital data or waveform area. If available, the patient's last name and bed number display in the patient demographic field.

**FIGURE 8-1** Patient Reports Tab

8.1.2 All Strips Report

The All Strips Report (shown in FIGURE 8-2 and FIGURE 8-3) contains real-time monitoring data, all current digital numeric values and all current configured waveforms for the selected patient tile. The report includes a header, digital data, waveform data and a footer section.

**NOTE:** This report is not available when the selected patient tile is in a Communications Lost or a Comm Loss - Check Bedside Monitor, Waiting for Patient, or Standby state.

**FIGURE 8-2** All Strips Report (Page 1)
8.1.2.1 Header
The All Strips Report’s first page header provides the title of the report and basic demographic information, if it is available. The demographic information includes the patient’s first and last name, patient ID, bed number, weight, height and doctor.

Page headers on subsequent pages of the report provide the report title and limited demographic information about the patient (first and last name, patient ID and bed number), if it is available.

If the demographic data was not previously entered in the system, only the demographic label is shown.

8.1.2.2 Digital Data
The All Strips Report’s digital data section displays measurements for the digital data parameters currently available for the selected patient tile. The displayed parameter order is determined by the system Unit Priorities tab. The digital data was collected at the time of the print request. The digital data section contains the following:

- A parameter label
- A numeric measurement
- A unit of measure label
- A heart rate (HR) value
- Dashes (--) to indicate missing or invalid digital data.
- Square brackets ([ ]) to indicate alarming parameter values

**NOTE:** Digital Data is only shown on the first page of the report.

8.1.2.3 Waveform Data
The waveforms that print in the All Strips Report’s waveform data section depend on which leads are currently in use and on the waveforms selected in the Print Setup function.

The first page of this report contains four waveforms. The middle pages of the report will contain six waveforms while the last page may contain one to six waveform strips depending on the waveforms available. Each waveform in this report represents approximately 10 seconds of data; seven (7) seconds prior to the time of the print request and three (3) seconds after the print request. A marker above each waveform strip indicates the time of the print request. A waveform label (i.e., ECG Lead II) is shown above each strip to identify the data source and the unit of measurement (i.e., %, RPM, BPM, mmHg, kPa). Pleth, ECG and Respiration do not display a unit of measurement. The waveforms displayed in the report are shown on a calibrated grid and contain a scale bar.

If the All Strips Report printed as a result of a Print on Alarm, the event that prompted the report will print below the report digital data in bold.
For pacemaker patients that have Pacer Enhancement enabled, the report displays a long vertical line crossing through the waveform or a short vertical line above the waveform (depending on the selection in the Pacer Settings dialog) to indicate firing of the pacemaker impulse.

For information on selecting the waveforms that can be printed in this report, refer to the "Print Setup Tab (Patient)" on page 5-62.

8.1.2.4 ECG Waveforms
The ECG waveforms shown in the All Strips Report are based on the ECG lead wire set currently in use and on the waveforms selected in the Print Setup tab. The ECG waveform report displays the following:

- An ECG lead label
- A wave gain scale bar and a calibration pulse for wave gains less than 30 mm/mV
- A calibrated grid
- Current ECG monitoring label
- A continuous straight line for active ECG leads that are not connected

Non-ECG Waveforms
The non-ECG waveforms shown in the All Strips Report only include the non-ECG parameters in use at the time of the print request.

All non-ECG waveforms in the report contain a parameter label and a unit of measure label. Non-ECG waveforms in the report contain a wave gain scale bar if the waveform is available at the time of the print request. If the waveform is not available, the report prints blank data for the specified time period. Also:

- If the IBP waveform was labeled at the bedside monitor, the report contains that IBP label.
- If the Anesthetic Agent was identified at the bedside monitor, the report contains that Agent label.

Footer
The All Strips Report contains a footer at the bottom of every page. The footer includes:

- Arrhythmia settings and device type.
- The date and time of the print request. The format displayed is taken from the System Date/Time tab.
- A centered page number.
- The name of the Panorama Central Station where the report was generated.

Printing an All Strips Report
The All Strips Report is printed in landscape orientation, using any of the following methods:
• Select the **Print All Strips** button in the **Patient Reports** tab. Selecting the **Print All Strips** button will post an **All Strips Report** print request for the selected patient.

• Select the **REC** button in a patient tile.

• Select the **Print** menu bar button in the **Bedside** tab.

• Enable the Print on Alarm option in the **Alarm Responses** tab when there is an alarm condition.

• Select the **Strip** button on a Passport 2, Passport V, Spectrum, Spectrum OR, or V 12/ V 21 monitor.

• Select the **Realtime Print** onscreen button on a DPM 6, DPM 7, Passport 8, Passport 12, Passport 12M or Passport 17M monitor. Select the **Print** button on a Telepack (refer to “Button Indicators” on page 10-3).

**NOTE:** To print this report for all active patients, use the **System Reports** tab **Print All Strips/All Patients** report.

**NOTE:** Remote printing is currently not available on the V 12/V 21.

Requesting this report posts a print job in the system Print Status tab. For additional information, refer to the “Print Status Tab” on page 8-40.

For report troubleshooting messages and issues, refer to “Troubleshooting” on page 8-29.

**8.1.3 Full Disclosure Report**

The Full Disclosure Report provides a compressed view of a patient’s historical ECG (shown in FIGURE 8-5 and FIGURE 8-6) and non-ECG (shown in FIGURE 8-7 and FIGURE 8-8) waveform data. This report includes a header, waveform data, and a footer.
FIGURE 8-5 Full Disclosure Report (Page 1 - ECG waveform layout)
FIGURE 8-6 Full Disclosure Report (Page 2- ECG waveform layout)
FIGURE 8-7  Full Disclosure Report (Page 1 - non-ECG waveform layout)
<table>
<thead>
<tr>
<th>Patient</th>
<th>Time</th>
<th>Location</th>
</tr>
</thead>
<tbody>
<tr>
<td>John Doe</td>
<td>12:00 PM</td>
<td>ICU 200</td>
</tr>
<tr>
<td>Jane Smith</td>
<td>2:00 PM</td>
<td>Ward 3</td>
</tr>
</tbody>
</table>

### FIGURE 8-8 Full Disclosure Report (Page 2 - non-ECG waveform layout)

#### 8.1.3.1 Header

The Full Disclosure Report's first page header provides the title of the report and basic demographic information, if it is available. The demographic information includes the patient's first and last name, patient ID, bed number, weight, height and doctor.

Page headers on subsequent pages of the report provide the report title, the date and time the data was collected, and limited demographic information about the patient (first and last name, patient ID and bed number), if it is available.

If the demographic data was not previously entered in the system, only the demographic label is shown.
8.1.3.2 Waveform Data
The ECG waveforms shown in the Full Disclosure Report are based on the ECG lead wire set currently in use and on the waveforms selected in the Print Setup tab.

ECG Waveform Format
The ECG waveform format for the Full Disclosure Report includes five columns. This format is used for reports that show ECG parameter waveforms.

- The first column contains the time (hour and minute) of the disclosure record. The time format follows the system format specified in the Date/Time tab.
- The second column contains a heart rate (HR) value. The heart rate value shown is from the last second of data that was collected during the disclosure record. If disclosure data is not available for the time period specified, dashes (--) are shown in the place of a heart rate value.
- The third column contains a PVC/min. value. The PVC value shown is the value that was obtained during the disclosure record. If disclosure data is not available for the time period specified, dashes are shown in the place of a PVC value.
- The fourth column contains a label for the waveform data being displayed in the report. The waveform label shown is the source of data for the disclosure record.
- The fifth column contains the events that occurred during the period of the disclosure record. If multiple events occur during the time period specified, the event with the highest priority is shown. If no events occur during the time period specified, this column remains empty.

For information regarding the type of events that occur in the Full Disclosure Report, refer to “Physiological Alarms and Events” on page 13-2.

Non-ECG Waveform Format
The non-ECG waveform format for the Full Disclosure Report includes four columns. This format is used for reports that show Non-ECG parameter waveforms.

- The first column contains the time (hour and minute) of the disclosure record. The time format follows the system format specified in the Date/Time tab.
- The second column contains a heart rate (HR) value. The heart rate value shown is from the last second of data that was collected during the disclosure record. If disclosure data is not available for the time period specified, dashes are shown in the place of a heart rate value.
- The third column contains a label for the waveform data being displayed in the report. The waveform label shown is the source of data for the disclosure record.
- The fourth column contains the events that have occurred during the period of the disclosure record. If multiple events occurred during the time period specified then the event with the highest priority is displayed. If no events occurred during the time period specified, this column remains empty.

For information regarding the type of events that occur in the Full Disclosure Report, refer to “Physiological Alarms and Events” on page 13-2.

8.1.3.3 Footer
The Full Disclosure Report contains a footer at the bottom of every page. The footer includes:
8.1.3.4 Printing a Full Disclosure Report (ECG and Non-ECG Layout)

The Full Disclosure Report is printed in portrait orientation, using any of the following methods:

To print one hour of full disclosure data (ECG layout):

1. Select the **Full Disclosure Report** button in the **Patient Reports** tab. The Print Selection dialog is displayed (shown in FIGURE 8-9).

   ![FIGURE 8-9 Print Selection Dialog Box](image)

2. Select the **Print 1 Hour** button. The report is sent to the printer.

To print one hour of full disclosure data (non-ECG layout):

1. From the **Disclosure** tab, select the **Display Choices** sidebar button. The Display Choices dialog box is displayed (shown in FIGURE 8-10).

   ![FIGURE 8-10 Display Choices Dialog Box](image)
2. Select a non-ECG parameter from the list box, then select the Done button. The waveforms in the tab will change to the selected non-ECG parameter.

3. From the menu bar, select the Print button. The Print Selection dialog is displayed (shown in FIGURE 8-9).

4. Select the Print 1 Hour button. The report is sent to the printer.

To print multiple hours of full disclosure data:

1. From the Patient Reports tab, select the Full Disclosure Report button.
2. From the menu bar, select the Print button. The Print Selection dialog is displayed.
3. Select one of the time interval buttons or the Cancel button.
   • The choices are print 1 hour, 2 hours, 4 hours, 8 hours, 12 hours, and 24 hours.
   • Select the Cancel button to close the dialog box without generating the print request.

To print multiple hours of full disclosure data (non-ECG layout):

1. From the Disclosure tab, select the Display Choices sidebar button. The Display Choices dialog box is displayed.
2. Select a non-ECG parameter from the list box, then select the Done button. The waveforms in the Disclosure tab will change to the selected non-ECG parameter.
3. From the menu bar, select the Print button. The Print Selection dialog is displayed (shown in FIGURE 8-9).
4. Select one of the time interval buttons or the Cancel button.
   • The choices are print 1 hour, 2 hours, 4 hours, 8 hours, 12 hours, and 24 hours.
   • Select the Cancel button to close the dialog box without generating the print request.

Requesting either report posts a print job in the system Print Status tab. For additional information, refer to the “Print Status Tab” on page 8-40.
8.1.4 ST Report

The ST Report (shown in FIGURE 8-11 and FIGURE 8-12) includes current and reference digital values with the associated ST templates, for a selected patient tile. This report includes a header, ST data and a footer section.

NOTE: This report is not available when using a WMTS 608 wireless bedside device, during a relearning phase, or when the selected patient tile is in a Communications Lost or a Comm Loss - Check Bedside Monitor, Standby, or a Waiting for Patient state.

FIGURE 8-11 ST Report (Page 1)
8.1.4.1 Header

The ST Report's first page header provides the title of the report and basic demographic information, if it is available. The demographic information includes the patient's first and last name, patient ID, bed number, weight, height and doctor.

Page headers on subsequent pages of the report provide the report title, the date and time the data was collected, and limited demographic information about the patient (first and last name, patient ID and bed number), if it is available.

If the demographic data was not previously entered in the system, only the demographic label is shown.

8.1.4.2 ST Data

The ST Report's data section includes current and reference templates for the ST data. ST data is shown in millimeter (mm) measurements. The ST templates shown in the report are limited to the ECG lead connections currently in use and selected in the Print Setup tab.
• One current and one reference template for the lead being monitored in a 3-lead cable connection.
• Three current and three reference templates for the leads being monitored in a 5-lead cable connection.
• Twelve current and twelve reference templates for all of the leads in a 12-lead cable connection.

Current Templates
The current templates shown in the ST Report are limited to the ECG lead connections currently in use and the waveforms the patient has selected in the Print Setup tab.

• Current templates in the report contain a label section above the ST data that includes CUR, the ST lead and the ST measurement.
• Current ST values display in brackets ([ ]), if the data represents an alarm condition.
• Current ST values in the report print on a calibrated grid with a scale bar, which corresponds to the ECG wave gain measurement.

Reference Templates
The reference templates shown in the ST Report are limited to the ECG lead wire set currently in use and the waveforms selected.

• Reference templates in the report contain a label section above the ST data that includes REF, the ST lead and the ST measurement.
• Reference ST values display in brackets ([ ]), if the data is in an alarm condition.
• Reference ST values in the report print on a calibrated grid with a scale bar, which corresponds to the ECG wave gain measurement.

8.1.4.3 Footer
The ST Report contains a footer at the bottom of every page. The footer includes:

• The date and time of the print request. The format displayed is taken from the System Date/Time tab.
• A centered page number.
• The name of the Panorama Central Station where the report was generated.

8.1.4.4 Printing an ST Report
The ST Report is printed in landscape orientation, using any of the following methods:

• From the Patient Reports tab, select the ST Report button.
• From the menu bar, select the Print button.

Requesting this report posts a print job in the system Print Status tab. For additional information, refer to the “Print Status Tab” on page 8-40.
8.1.5 Trend List Report

The Trend List Report (shown in FIGURE 8-13 and FIGURE 8-14) shows a patient’s historical trended digital data. The data displayed is based on the time interval selected in the Trend Display dialog box, and can be shown over a 1, 2, 4, 8, 12, or 24 hour time period. This report includes a header, trend records, and a footer.
FIGURE 8-14 Trend List Report (Page 2)
8.1.5.1 Header
The Trend List Report’s first page header provides the title of the report and basic demographic information, if it is available. The demographic information includes the patient's first and last name, patient ID, bed number, weight, height and doctor.

Page headers on subsequent pages of the report provide the report title, the date and time the data was collected, and limited demographic information about the patient (first and last name, patient ID and bed number), if it is available.

If the demographic data was not previously entered in the system, only the demographic label is shown.

8.1.5.2 Trend Records
The Trend List Report’s digital data section includes the digital data that occurred for the selected patient, based on the time interval selected in the Trend Display dialog box. The data printed in the report is determined by the device connected to the selected patient tile and the sensors in use.

The report displays digital data records for each minute of the selected time period. Digital data records are separated by a solid horizontal line. If data is not available for the selected time period, the report will not print. Each digital record in the report contains the following:

- The time format follows the system format specified in the Date/Time tab.
- Digital data values for the parameters specific to the device connected to the patient tile and the sensors used during the specified time period.
- A heart rate (HR) value. The source of the heart rate data is shown in parentheses next to the HR label. If data is not available for the specified time period, dashes are shown in the place of a heart rate value.
- Digital Data records in the report use brackets ([ ]) to distinguish data in the alarm condition.

8.1.5.3 Footer
The Trend List Report contains a footer at the bottom of every page. The footer includes:

- The date and time of the print request. The format displayed is taken from the System Date/Time tab.
- A centered page number.
- The name of the Panorama Central Station where the report was generated.

8.1.5.4 Printing a Trend List Report
The Trend List Report is printed in portrait orientation.

To print one hour of trend data:

- From the Patient Reports tab, select the Trend List Report button. The Print Selection dialog is displayed (shown in FIGURE 8-9).
- Select the Print 1 Hour button. The report is sent to the printer.
To print multiple hours of trend data from the **Patient Reports** tab:

1. Select the **Trend List Report** button.
2. Select one of the time interval buttons or the **Cancel** button.
   - The choices are print 1 hour, 2 hours, 4 hours, 8 hours, 12 hours, and 24 hours.
   - Select the **Cancel** button to close the dialog box without generating the print request.

To print multiple hours of trend data from the **Trends** tab:

1. From the menu bar, select the **Print** button. The Print Selection dialog is displayed.
2. Select the **Print Current** button, a time interval button, or the **Cancel** button.
   - Select the **Print Current** to create a one page Trend List Report.
   - Select one of the time interval buttons. The choices are print 1 hour, 2 hours, 4 hours, 8 hours, 12 hours, and 24 hours.
   - Select the **Cancel** button to close the dialog box without generating the print request.

Requesting this report posts a print job in the system Print Status tab. For additional information, refer to the “Print Status Tab” on page 8-40.
8.1.6 Event List Report

The Event List Report (shown in FIGURE 8-15 and FIGURE 8-16) shows a patient's historical physiological, system and technical alarm events for a 1, 2, 4, 8, 12 and 24 hour time period. This report includes a header, event records, and a footer.

![Event List Report](image)

**FIGURE 8-15 Event List Report (Page 1)**
8.1.6.1 Header

The Event List Report's page header provides the title of the report and basic demographic information, if it is available. The demographic information includes the patient's first and last name, patient ID, bed number, weight, height and doctor.

Page headers on subsequent pages of the report provide the report title, the date and time the data was collected, and limited demographic information about the patient (first and last name, patient ID and bed number), if it is available.
If the demographic data was not previously entered in the system, only the demographic label is shown.

### 8.1.6.2 Event Record

The Event List Report’s event record section includes the digital data that occurred during the specified time period. The data in this report is determined by the events that occurred during the specified time period.

Event data records are separated by a solid horizontal line. If data is not available for the selected time period, the report will not print.

- The first page of event records include numbers for the Total Events (the total number of events in the selected patient’s event database), the Displayed Events (the number of events shown in the report) and the Event Filter (based on the event filter setting).
- Each event record in the report contains the time (hour and minute) when the event occurred. The time format follows the system format specified in the **Date/Time** tab.
- Each Event record in the report contains the name of the event that has occurred for the selected patient.
- Each physiological event record in the report contains the digital data values for the specified time period.
- Each Event Data record in the report contains a digital heart rate (HR) value. The source of the heart rate data is shown in parentheses next to the HR label. If data is not available for the time period specified, dashes are shown in the place of a heart rate value.

For information regarding the digital data supported by each device, refer to “Supported Devices” on page 1-3. The column order in which the digital parameter data is printed in a record is based on the order selected in the **Unit Priorities** tab.

- Event records in the report use brackets ([ ]) to distinguish data in the alarm condition and dashes (- - -) to distinguish incomplete or missing data.
- System and technical event records in the report do not contain digital data values.

### 8.1.6.3 Footer

The Event List Report contains a footer at the bottom of every page. The footer includes:

- The date and time of the print request. The format displayed is taken from the **System Date/Time** tab.
- A centered page number.
- The name of the Panorama Central Station where the report was generated.

### 8.1.6.4 Printing an Event List Report

The Event List Report is printed in portrait orientation.

To print one hour of event data:

- From the **Patient Reports** tab, select the **Event List Report** button. The Print Selection dialog is displayed (shown in FIGURE 8-9).
- Select the **Print 1 Hour** button. The report is sent to the printer.
To print multiple hours of event data from the **Patient Reports** tab:

1. Select the **Event List Report** button.
2. Select one of the time interval buttons or the **Cancel** button.
   - The choices are print 1 hour, 2 hours, 4 hours, 8 hours, 12 hours, and 24 hours.
   - Select the **Cancel** button to close the dialog box without generating the print request.

To print multiple hours of trend data from the **Events** tab:

1. From the menu bar, select the **Print** button. The Print Selection dialog is displayed.
2. Select one of the **Print Current**, a time interval button, or the **Cancel** button.
   - Select the **Print Current** to create a one page Trend List Report.
   - Select one of the time interval buttons. The choices are print 1 hour, 2 hours, 4 hours, 8 hours, 12 hours, and 24 hours.
   - Select the **Cancel** button to close the dialog box without generating the print request.

**NOTE:** **If the number of hours selected exceeds the amount of data in system, the report shows only the available data.**

Requesting this report posts a print job in the system Print Status tab. For additional information, refer to the “Print Status Tab” on page 8-40.
8.1.7 Patient Alarm Report

The Patient Alarm Report (shown in FIGURE 8-17) is a single page report containing a patient's alarm settings information (unit of measure, alarm limits, priority and alarm responses). This report includes a header, patient alarm data, optional arrhythmia and paging delay settings (if enabled), signal lost delay (when using a Telepack), and ECG settings (when using a V 12/V 21 or Telepack device) and a footer.

![FIGURE 8-17 Patient Alarm Report - Paging Option Disabled](image)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
<th>High Limit</th>
<th>Low Limit</th>
<th>Priority</th>
<th>Print On Alarm</th>
<th>Save To Event</th>
</tr>
</thead>
<tbody>
<tr>
<td>HR</td>
<td>BPM</td>
<td>OFF</td>
<td>50</td>
<td>2</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Ext. Brady</td>
<td>BPM</td>
<td>---</td>
<td>---</td>
<td>1</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Ext. Tachy</td>
<td>BPM</td>
<td>---</td>
<td>---</td>
<td>1</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>NIBP Sys</td>
<td>mmHg</td>
<td>160</td>
<td>90</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>NIBP Dia</td>
<td>mmHg</td>
<td>90</td>
<td>50</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>NIBP Mean</td>
<td>mmHg</td>
<td>110</td>
<td>60</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>SpO2</td>
<td>%</td>
<td>OFF</td>
<td>90</td>
<td>2</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Resp</td>
<td>BPM</td>
<td>30</td>
<td>9</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>INSpCO2</td>
<td>%</td>
<td>0.5</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>ETcO2</td>
<td>%</td>
<td>8.6</td>
<td>3.3</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>PVC/min</td>
<td>PVC/min</td>
<td>10</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Art Sys</td>
<td>mmHg</td>
<td>160</td>
<td>90</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Art Dia</td>
<td>mmHg</td>
<td>90</td>
<td>50</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Art Mean</td>
<td>mmHg</td>
<td>110</td>
<td>70</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>PA Sys</td>
<td>mmHg</td>
<td>35</td>
<td>10</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>PA Dia</td>
<td>mmHg</td>
<td>16</td>
<td>0</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>PA Mean</td>
<td>mmHg</td>
<td>20</td>
<td>0</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>CVP Mean</td>
<td>mmHg</td>
<td>10</td>
<td>0</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>ICP Mean</td>
<td>mmHg</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>T1</td>
<td>ºF</td>
<td>100</td>
<td>95</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>T2</td>
<td>ºF</td>
<td>100</td>
<td>95</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Delta T</td>
<td>ºF</td>
<td>4</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Tis</td>
<td>ºF</td>
<td>102</td>
<td>97</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>RAL</td>
<td>%</td>
<td>---</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>ISO</td>
<td>%</td>
<td>---</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>ENF</td>
<td>%</td>
<td>---</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>SEV</td>
<td>%</td>
<td>---</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>DES</td>
<td>%</td>
<td>---</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>INSpO2</td>
<td>%</td>
<td>---</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>INSpN2O</td>
<td>%</td>
<td>---</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Apneu</td>
<td>Sec</td>
<td>20</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Failure to Sense</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Failure to Capture</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Arrhythmia</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Bigeminy</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Couples</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Irregular HR</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Run</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Torsion</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>V-Fis</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>1</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>V-Tach</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>1</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>V-Rhythm</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Multifocal PVC</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Pause</td>
<td>---</td>
<td>---</td>
<td>2</td>
<td>OFF</td>
<td></td>
<td></td>
</tr>
<tr>
<td>R-On-T PVC</td>
<td>---</td>
<td>---</td>
<td>2</td>
<td>OFF</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Print Time: 2014/12/13 05:15 PM  
Page 1  
Panorama: MR50214326PC1  

FIGURE 8-17 Patient Alarm Report - Paging Option Disabled
Patient Alarm Report

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
<th>High Limit</th>
<th>Low Limit</th>
<th>Priority</th>
<th>Print On Alarm</th>
<th>Save To Event</th>
</tr>
</thead>
<tbody>
<tr>
<td>HR</td>
<td>BPM</td>
<td>150</td>
<td>50</td>
<td>2</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Extr. Brady</td>
<td>BPM</td>
<td>---</td>
<td>45</td>
<td>1</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Extr. Tachy</td>
<td>BPM</td>
<td>165</td>
<td>---</td>
<td>1</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>SpO2</td>
<td>%</td>
<td>---</td>
<td>85</td>
<td>2</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>PV/PM</td>
<td>PVC/min</td>
<td>OFF</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Pause/PM</td>
<td>Pause/min</td>
<td>OFF</td>
<td>---</td>
<td>2</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Nurse Call</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Asystole</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>1</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Bigeminy</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Couplet</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Irregular HR</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Run</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Trigeminy</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>V-Fib</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>1</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>V-Tach</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>1</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>V-Rhythm</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Multifocal PVC</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Pause</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>2</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>R-Occ-T PVC</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>2</td>
<td>OFF</td>
<td>ON</td>
</tr>
</tbody>
</table>

Arrhythmia Settings

- Asystole Delay: 4 Sec
- V-Tach Threshold: 3 Beats
- All Arrhythmia On/Off: ON
- V-Tach Rate: 120 BPM
- ECG Noise Delay: 5 Sec
- Pause Threshold (RR): 1.8

Signal Lost Delay Settings

- Signal Lost Delay: 0 Sec

ECG Settings

- Primary Analysis Lead: II
- Secondary Analysis Lead: V
- Beat Classification Lead: I
- Pacing Rate: ---
- Minimum QRS Threshold: 0.16 mV
- QT Monitoring On/Off: OFF
- QTc Formula: Hodges

Print Time: 2014/12/13 05:20 PM
Page 1
Panorama: MR0214329PC1

**FIGURE 8-18** Patient Alarm Report - Paging Option Disabled, Telepack Device
8.1.7.1 Header

The Patient Alarm Report’s page header provides the title of the report and basic demographic information, if it is available. The demographic information includes the patient’s first and last name, patient ID, bed number, weight, height and doctor.

If the demographic data was not previously entered in the system, only the demographic label is shown.

![FIGURE 8-19 Patient Alarm Report - Paging Option Enabled]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
<th>High Limit</th>
<th>Low Limit</th>
<th>Priority</th>
<th>Print On Alarm</th>
<th>Save To Event</th>
</tr>
</thead>
<tbody>
<tr>
<td>HR</td>
<td>BPM</td>
<td>150</td>
<td>50</td>
<td>2</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Ect Brady</td>
<td>BPM</td>
<td>--</td>
<td>45</td>
<td>1</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Ect Tachy</td>
<td>BPM</td>
<td>165</td>
<td>--</td>
<td>1</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>NIBP Sys</td>
<td>mmHg</td>
<td>60</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>NIBP Dia</td>
<td>mmHg</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>NIBP Mean</td>
<td>mmHg</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Resp</td>
<td>RPM</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>INSEPCO2</td>
<td>mmHg</td>
<td>10.0</td>
<td>--</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>ETCO2</td>
<td>mmHg</td>
<td>55.0</td>
<td>15.0</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>PVC/min</td>
<td>PVC/min</td>
<td>--</td>
<td>--</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Qtc</td>
<td>ms</td>
<td>OFF</td>
<td>OFF</td>
<td>2</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Pause/min</td>
<td>Pause/min</td>
<td>--</td>
<td>--</td>
<td>2</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>IBP1 Sys</td>
<td>mmHg</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>IBP1 Dia</td>
<td>mmHg</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>IBP1 Mean</td>
<td>mmHg</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>IBP2 Dia</td>
<td>mmHg</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>IBP2 Mean</td>
<td>mmHg</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>IBP3 Dia</td>
<td>mmHg</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>IBP3 Mean</td>
<td>mmHg</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>IBP4 Dia</td>
<td>mmHg</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>IBP4 Mean</td>
<td>mmHg</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>IBP5 Dia</td>
<td>mmHg</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>IBP5 Mean</td>
<td>mmHg</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>IBP6 Dia</td>
<td>mmHg</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>IBP6 Mean</td>
<td>mmHg</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>IBP7 Dia</td>
<td>mmHg</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>IBP7 Mean</td>
<td>mmHg</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>SVR</td>
<td>mmHg</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>SVR</td>
<td>mmHg</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>CCI</td>
<td>LPM</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>CCI STAT</td>
<td>LPM/m2</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>CCI STAT</td>
<td>LPM/m2</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>T1</td>
<td>°F</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>T2</td>
<td>°F</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>T3</td>
<td>°F</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Delta T</td>
<td>°F</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>HAI</td>
<td>%</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>ISO</td>
<td>%</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>ENF</td>
<td>%</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>SEV</td>
<td>%</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>DES</td>
<td>%</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>INSEPCO2</td>
<td>%</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>INSEPCO2</td>
<td>%</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>SvO2</td>
<td>%</td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>BIS</td>
<td></td>
<td>OFF</td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>ST Single</td>
<td>mm</td>
<td>OFF</td>
<td>OFF</td>
<td>2</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>ST Dual</td>
<td>mm</td>
<td>OFF</td>
<td>OFF</td>
<td>2</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Apnea</td>
<td></td>
<td></td>
<td>OFF</td>
<td>3</td>
<td>OFF</td>
<td>ON</td>
</tr>
<tr>
<td>Available</td>
<td></td>
<td></td>
<td>OFF</td>
<td>1</td>
<td>OFF</td>
<td>ON</td>
</tr>
</tbody>
</table>

Print Time: 2014/12/11 07:52 PM  Page 1  Panorama: MR50212800PC1

FIGURE 8-19 Patient Alarm Report - Paging Option Enabled

8.1.7.1 Header

The Patient Alarm Report's page header provides the title of the report and basic demographic information, if it is available. The demographic information includes the patient's first and last name, patient ID, bed number, weight, height and doctor.

If the demographic data was not previously entered in the system, only the demographic label is shown.
8.1.7.2 Patient Alarm Data
The Patient Alarm Report’s data section includes alarm information for all of the alarms. The parameters in a Patient Alarm Report are dependent on the device connected to the selected patient tile. The columns are described as follows:

- **Parameter** displays the parameters for which the specified patient has an alarm.
- **Units** displays the unit of measure in use for the parameter specified.
- **High Limit** displays the High alarm limit for the parameter specified.
- **Low Limit** displays the Low alarm limit for the parameter specified.
- **Priority** displays the alarm priority assigned to the parameter specified.
- **Print On Alarm** displays the Print on Alarm response assigned to the parameter specified.
- **Save to Event** displays the Save To Event alarm response that has been assigned to the parameter specified.
- **Page On Alarm** displays the Page On Alarm response that has been assigned to the parameter specified.

8.1.7.3 Arrhythmia Settings
The Patient Alarm Report’s Arrhythmia Settings section includes values for the following:

- Asystole Delay
- V-Tach Threshold
- All Arrhythmia On/Off
- V-Tach Rate

8.1.7.4 Paging Delay Settings
The Patient Alarm Report’s Paging Delay section includes the system paging delay value that was configured in the System Alarms tab.

8.1.7.5 Footer
The Patient Alarm Report contains a footer at the bottom of the page. The footer includes:

- The date and time of the print request. The format displayed is taken from the System Date/Time tab.
- A centered page number.
- The name of the Panorama Central Station where the report was generated.

8.1.7.6 Printing a Patient Alarm Report
The Patient Alarm Report is printed in portrait orientation, using any of the following methods:

- From the Patient Reports tab, select the Patient Alarm Report button.
- From the Alarm Limits tab, select the Print menu bar button.

Requesting this report posts a print job in the system Print Status tab. For additional information, refer to the “Print Status Tab” on page 8-40.
### Troubleshooting

This section lists some of the potential messages and issues that may occur while using the **Patient Reports** tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>All report buttons are disabled.</td>
<td>A patient is not currently selected.</td>
<td>Select a patient before trying to print a report.</td>
</tr>
<tr>
<td>The <strong>ST Report</strong> button is disabled.</td>
<td>The selected patient does not have ST enabled.</td>
<td>Enable ST for the selected patient before trying to print an ST report.</td>
</tr>
<tr>
<td></td>
<td>A WMTS 608 wireless bedside device is in use.</td>
<td>No solution required.</td>
</tr>
<tr>
<td>The selected patient tile is in a relearning phase, or is in a Communications Lost or a Comm Loss - Check Bedside Monitor or Standby state.</td>
<td>Wait for the patient tile to resume monitoring.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>The selected patient tile is in a Waiting for Patient state.</td>
<td>Activate the destination device.</td>
</tr>
<tr>
<td>The <strong>Print All Strips</strong> report button is disabled.</td>
<td>The selected patient tile is in a Communications Lost or a Comm Loss - Check Bedside Monitor, or Standby state.</td>
<td>Wait for the patient tile to resume monitoring.</td>
</tr>
<tr>
<td></td>
<td>The selected patient tile is in a Waiting for Patient state.</td>
<td>Activate the destination device.</td>
</tr>
<tr>
<td>The report will not print</td>
<td>There are several reasons that a report would not print. There could be a communication problem between the printer and the Panorama Central Station and/or there could be a problem with the printer.</td>
<td>If there is an error that appears such as <strong>Printer out of Paper</strong> you will need to remedy that situation. If the <strong>No Printer connected or Printer not Available</strong> error is displayed, verify that the printer was properly configured with the Panorama Central Station. Refer to the Panorama Printer Configuration Manual (P/N 0070-00-0561).</td>
</tr>
<tr>
<td>The Delta T parameter does not display in the All Strips report.</td>
<td>One of the two temperature sources used to calculate the Delta T value is not valid.</td>
<td>Verify that the two temperature sources have been properly configured.</td>
</tr>
<tr>
<td>Page On Alarm column not displayed in Patient Alarm Report</td>
<td>Paging option is not enabled at the central station.</td>
<td>If available, enable Paging option. If feature unavailable, no solution required.</td>
</tr>
<tr>
<td>All Arrhythmia On/Off setting set to Off</td>
<td>Arrhythmia is disabled for a selected patient tile.</td>
<td>If available, enable arrhythmia for the selected patient tile. If feature or license is unavailable, no solution required.</td>
</tr>
</tbody>
</table>
In the Patient Alarm Report's Arrhythmia Settings section, a "---" is displayed as the value for Asystole Delay, V-Tach Threshold, and V-Tach Rate. If available, enable arrhythmia for the selected patient tile. If feature or license is unavailable, no solution required.

Paging Delay Settings section not displayed in the Patient Alarm Report. This section will only be displayed when Paging is enabled at the central station. If available, enable Paging option. If feature unavailable, no solution required.

* Messages are shown in all bold text.
8.2 System Reports Tab

The **System Reports** tab (shown in FIGURE 8-21) is used to generate reports on a system level. A patient tile does not have to be selected to generate a system report.

The following reports can be printed from the **System Reports** tab:

- Accessing the System Reports Tab
- All Strips (All Patients) Report
- Equipment Report
- Paging Report

8.2.1 Accessing the System Reports Tab

1. From the menu bar, select the **Report** button.
2. Select the **System Reports** tab. The **System Reports** tab is displayed.

![FIGURE 8-20 System Reports Tab](image)

![FIGURE 8-21 System Reports Tab - Paging Option Enabled](image)

**NOTE:** The Paging Report button will only be displayed when the Paging Option is enabled at the central station.
8.2.2 All Strips (All Patients) Report

The All Strips (All Patients) Report (shown in FIGURE 8-22 and FIGURE 8-23) contains real-time monitoring data for all the active patients monitored by the Panorama Central Station. This report contains all current numeric values and waveforms. It is similar to the All Strips Report, except that it prints for all active patients. The report includes a header, digital data, waveform data and a footer.

FIGURE 8-22 All Strips (All Patients) Report (Page 1)
8.2.2.1 Header

The All Strips (All Patients) Report’s first page header provides the title of the report and basic demographic information, if it is available. The demographic information includes the patient’s first and last name, patient ID, bed number, weight, height, and doctor.

Page headers on subsequent pages of the report provide the report title, the date and time the data was collected, and limited demographic information about the patient (first and last name, patient ID, and bed number), if it is available.

If the demographic data was not previously entered in the system, only the demographic label is shown.
8.2.2.2 Digital Data
The All Strips (All Patients) Report's digital data section displays measurements for the digital data parameters currently available for the selected patient tile. The displayed parameter order is determined by the system Unit Priorities tab. The digital data section contains the following:

- A parameter label
- A numeric measurement
- A unit of measure label
- A heart rate (HR) value
- Dashes (--) to indicate missing or invalid digital data.
- Square brackets ([ ] ) to indicate alarming parameter values

NOTE: Digital Data is only shown on the first page of the report.

8.2.2.3 Waveform Data
The waveforms that print in the All Strips (All Patients) Report's waveform data section are limited to which leads are currently in use, and on the waveforms selected in the Print Setup function.

The first page of this report contains up to two waveforms while subsequent pages contain up to three waveforms. Each waveform in this report represents approximately 10 seconds of data; seven (7) seconds prior to the time of the print request and three (3) seconds after the print request. A waveform label (i.e., ECG Lead II) is shown above each strip to identify the data source and the unit of measurement (i.e., %, RPM, BPM, mmHg, kPa). Pleth, ECG and Respiration do not display a unit of measurement. The waveforms displayed in the report are shown on a calibrated grid and contain a scale bar.

For pacemaker patients that have Pacer Enhancement enabled, the report displays a long vertical line crossing through the waveform or a short vertical line above the waveform (depending on the selection in the Pacer Settings dialog) to indicate firing of the pacemaker impulse.

ECG Waveforms
The ECG waveforms shown in the All Strips (All Patients) Report are based on the ECG lead wire set currently in use and on the waveforms selected in the Print Setup tab. The ECG waveform report displays with the following:
• An ECG lead label
• A wave gain scale bar and a calibration pulse for wave gains less than 30 mm/mV
• A calibrated grid
• Current ECG monitoring label
• A continuous straight line for active ECG leads that are not connected

Non-ECG Waveforms
The non-ECG waveforms shown in the All Strips (All Patients) Report are based on the non-ECG waveforms that are in use and the non-ECG waveforms that are selected to be printed. The report only includes non-ECG parameters if they are in use at the time of the print request.

All non-ECG waveforms in the report contain a parameter label and a unit of measure label. Non-ECG waveforms in the report may contain a wave gain scale bar. If a non-ECG waveform is unavailable at the time of the print request, the report prints blank data for the specified time period.

• If the IBP waveform was labeled at the bedside monitor, the report contains that IBP label.
• If the Anesthetic Agent was identified at the bedside monitor, the report contains that Agent label.

8.2.2.4 Footer
The report contains a footer at the bottom of every page. The footer includes:

• The date and time of the print request. The format displayed is taken from the System Date/Time tab.
• A centered page number.
• The name of the Panorama Central Station where the report was generated.

8.2.2.5 Printing an All Strips (All Patients) Report
The All Strips (All Patients) Report prints in portrait orientation, using the following method:

• From the System Reports tab, select the Print All Strips/All Patients button. Selecting this button sends a print request, for each patient, to the printer.

Requesting this report posts a print job in the system Print Status tab. For additional information, refer to the “Print Status Tab” on page 8-40.
8.2.3 Equipment Report
The Equipment Report (shown in FIGURE 8-24) contains a list of devices currently assigned to
the Panorama Central Station. This report displays a maximum of 32 devices per page and
includes a header, seven columns and a footer.

NOTE: The Equipment Report is not available at the ViewStation.

FIGURE 8-24 Equipment Report

8.2.3.1 Header
The header of the Equipment Report displays a centered report title.

8.2.3.2 Equipment Information
The Equipment Report contains seven columns. They are describes as follows:
Label displays the equipment device label. For additional information regarding Device ID's, refer to the “Equipment Setup Tab” on page 9-33.

Type displays the type of device that is attached to the central station. For example, a Telepack-WMTS or a Passport 2.

Tile displays the tile the device is attached to. Six dashes (- - - - - -) display in this column if the tile number associated with the device in the Equipment Setup dialog is None.

Device ID displays the ID of the device or unit. Six dashes display in this column if the device is a Panorama Central Station.

Patient ID displays the identification number of the patient attached to the device. Six dashes display in this column if the device attached to the patient does not have the Patient ID entered.

IP Address displays the unique IP address for the device.

8.2.3.3 Footer
The Equipment Report contains a footer at the bottom of every page. The footer includes:

- The date and time of the print request. The format displayed is taken from the System Date/Time tab.
- A centered page number.
- The name of the Panorama Central Station where the report was generated.
- The serial number of the Panorama Central Station where the report was generated.

8.2.3.4 Printing the Equipment Report
The Equipment Report is printed in portrait orientation, using any of the following methods:

- From the System Report tab, select the Equipment Report button.

NOTE: The Equipment Report button is not available at the ViewStation.

- From the Equipment Setup tab, select the Print menu bar button.

Requesting this report posts a print job in the system Print Status tab. For additional information, refer to the “Print Status Tab” on page 8-40.

8.2.4 Paging Report
The Paging Report (shown in FIGURE 8-25) prints all the currently assigned pagers, lists who each pager is assigned to, and specifies which patient tiles are assigned to which pagers. This report contains a row for every pager listed in the Paging tab and can contain a maximum of 100 pager assignments. Depending on the number of assigned pagers, this report may contain more than one page.

NOTE: The Paging Report is not available at the ViewStation.
8.2.4.1 Header
The Paging Report’s header provides the title of the report.

8.2.4.2 Pager Information
The Paging information section of the report contains all the pager information as they are configured at the central station at the time the report was generated.

The following pager information is shown in three columns:
• **Pager Number** displays the number that has been assigned to the pager. This column can display a maximum of 14 alphanumeric characters.

• **Pager Assignment** displays which person or department the pager is assigned to.

• **Tile** displays which patient tile number the pager is assigned to. This column can display a maximum of 30 alphanumeric characters. In addition, this column can display a maximum of 8-12 patient tiles when using a single display or a dual display configuration. The Tile column uses a comma (,) as a separator when more than one patient tile is assigned to a pager.

### 8.2.4.3 Footer

The Paging Report contains a footer at the bottom of the report page. The footer includes:

• The date and time of the print request. The format displayed is taken from the **System Date/Time** tab.

• A centered page number.

• The name of the Panorama Central Station where the report was generated.

### 8.2.4.4 Printing a Paging Report

The Paging Report is printed in portrait orientation, using any of the following methods:

• From the **System Reports** tab, select the **Paging Report** button.

• From the **Paging** tab, select the **Print** menu bar button.

Requesting this report posts a print job in the system Print Status tab. For additional information, refer to the “Print Status Tab” on page 8-40.
8.3 Print Status Tab

The **Print Status** tab (shown in FIGURE 8-26) contains a list of current print requests. Once the status of a print request changes or the report prints, the list updates automatically without having to refresh the list.

![Print Status Tab](image)

**FIGURE 8-26** Print Status Tab

8.3.1 Print Status Tab List Box

Once a new print request is submitted, specific information displays in the **Print Status** tab. The list box contains the following information:

- **Time** displays the time the print request was submitted. The print time format follows the system format specified in the **Date/Time** tab.
- **Job** displays the name of the requested report. For example, if the Full Disclosure Report was requested, the **Job** column displays **Full Disclosure Report**.
- **Tile ID** displays the **Tile ID** associated with the print request. For example, if the All Strips Report is requested and tile six is selected, the **Tile ID** column displays **6**.
- **Status** displays the current status of the print request. There are five possible status options available:
  - **In Job Queue** indicates that a print request was generated
  - **Printing** indicates that the print request is currently printing
  - **Deleting** indicates that the print request is currently being deleted
  - **Spooling** indicates the time between when the print request is made, and when the request is printed
  - **Error** indicates that there is an error at the printer that requires user intervention

The information available in this column is real-time. For example, once the status of a print request changes or the report prints, the list updates accordingly.

- **Printer** displays the name of the printer associated with the print request.

8.3.2 Removing Print Jobs from the Print Queue

Once print requests are submitted, they can be cancelled or deleted by using the following buttons.
Cancel Selected Print Job
Select the **Cancel Selected Print job** button to delete the selected print request from the job queue. When this button is selected, the multi-column list box Status column displays **Deleting**, and the request is removed from the multi-column list box.

Delete All Print Jobs
Select the **Delete All Print Jobs** button to cancel all print requests in the job queue. When this button is selected, the multi-column list box Status column displays **Deleting** for each print request and the requests are removed from the multi-column list box.

8.3.3 Sidebar Buttons
The following sidebar button is used to expand the functionality of this tab.

Normal Screen
Select the **Normal Screen** sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

8.3.4 Troubleshooting
This section lists some of the potential messages and issues that may occur while using the **Print Status** tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Report is not printing</td>
<td>Printer not selected at the Panorama Central Station.</td>
<td>Refer to the Panorama Printer Configuration document (P/N 0070-00-0561).</td>
</tr>
<tr>
<td>Report is not printing</td>
<td>Printer is not turned on.</td>
<td>Turn the printer on.</td>
</tr>
<tr>
<td>Report is not printing</td>
<td>Printer is offline.</td>
<td>Verify that the printer is online.</td>
</tr>
<tr>
<td>Report is not printing</td>
<td>Printer is out of paper.</td>
<td>Verify that the printer paper tray contains paper.</td>
</tr>
<tr>
<td>Report is not printing</td>
<td>Verify that the printer is connected to the Panorama Central Station.</td>
<td>Print a test page. Refer to the the Panorama Printer Configuration document (P/N 0070-00-0561).</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.*
### MESSAGE/ISSUE *

<table>
<thead>
<tr>
<th>Report is not printing from bedside device</th>
<th>Communication problem among the printer, the Panorama Central Station and bedside device.</th>
<th>Verify that the printer, Panorama Central Station and bedside device are all physically connected. Verify that the printer is properly configured with the Panorama Central Station.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Improper print setup.</td>
<td>The printer is not configured to print the report or strip. Refer to “Accessing the Print Setup Tab (Central Station)” on page 9-45.</td>
<td></td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
8.4 Additional Reports

The Panorama Central Station is capable of generating additional reports from areas other than the Reports menu button.

Reports generated by the Panorama Central Station, but cannot be generated from the Reports menu button include:

- Event Zoom In Report
- Full Disclosure Zoom In Report
- Graphic Trend Report
- 12-lead Report
- Telepack Error Log Report
- System Alarm Report

8.4.1 Accessing Additional Reports

These reports can be accessed from different areas throughout the system. For specific instructions on how to request each of these reports, refer to each report's printing section.

8.4.2 Event Zoom In Report

The Event Zoom In Report (shown in FIGURE 8-27 and FIGURE 8-28) shows all of a patient's historical digital and waveform data for a selected event. This report includes a header, digital data, waveform data and a footer.

![Event Zoom In Report](image)

**FIGURE 8-27** Event Zoom In Report (Page 1)
8.4.2.1 Header

The Event Zoom In Report’s first page header provides the title of the report and basic demographic information, if it is available. The demographic information includes the patient’s first and last name, patient ID, bed number, weight, height and doctor.

Page headers on subsequent pages of the report provide the report title, and limited demographic information about the patient (first and last name, patient ID and bed number), if it is available.

If the demographic data was not previously entered in the system, only the demographic label is shown.

8.4.2.2 Digital Data

The Event Zoom In Report’s digital data section displays measurements for the digital data parameters currently available for the selected patient tile. The displayed parameter order is determined by the system Unit Priorities tab. The digital data was collected at the time of the print request. The digital data section contains the following:
8.4.2.3 Waveform Data
The waveforms that print in the Event Zoom In Report’s waveform data section depend on which leads are currently in use and on the waveforms selected in the Print Setup function.

The first page of this report contains four waveforms. The middle pages of the report will contain six waveforms while the last page may contain two, four, or six waveform strips depending on the waveforms available. Each waveform in this report represents approximately 20 seconds of data; nine (9) seconds prior to the time of the print request and 11 seconds after the print request. A marker above each waveform strip indicates the time of the print request. A waveform label (i.e., ECG Lead II) is shown above each strip to identify the data source and the unit of measurement (i.e., %, RPM, BPM, mmHg, kPa). Pleth, ECG and Respiration do not display a unit of measurement. The waveforms displayed in the report are shown on a calibrated grid and contain a scale bar.

For pacemaker patients that have Pacer Enhancement enabled, the report displays a long vertical bar or a short vertical bar (based on configuration) to indicate the firing of the pacemaker impulse.

8.4.2.4 ECG Waveforms
The ECG waveforms shown in the Event Zoom In Report are limited to the ECG lead wire set currently in use, and on the waveforms selected in the Print Setup tab. The ECG waveform report is displayed with the following:

- An ECG lead label
- A wave gain scale bar and a calibration pulse for wave gains less than 30 mm/mV
- A calibrated grid
- Current ECG monitoring label
- A continuous straight line for active ECG leads that are not connected
8.4.2.5 Non-ECG Waveforms
The non-ECG waveforms shown in the Event Zoom In Report are limited to the non-ECG waveforms that are in use, and the non-ECG waveforms that are selected to be printed.

All non-ECG waveforms in the report contain a parameter label and a unit of measure label. Non-ECG waveforms in the report may contain a wave gain scale bar. If a non-ECG waveform is unavailable at the time of the print request, the report prints blank data for the specified time period.

- If the IBP waveform was labeled at the bedside monitor, the report contains that IBP label.
- If the Anesthetic Agent was identified at the bedside monitor, the report contains that Agent label.

8.4.2.6 Footer
The Event Zoom In Report contains a footer at the bottom of every page. The footer includes:

- Arrhythmia settings and device type.
- The date and time of the print request. The format displayed is taken from the System Date/Time tab.
- A centered page number.
- The name of the Panorama Central Station where the report was generated.

8.4.2.7 Printing an Event Zoom In Report
The Event Zoom In Report prints in landscape orientation using the following method:

- From the Waveforms View of the Events tab, select the Print menu bar button.

Requesting this report posts a print job in the system Print Status tab. For additional information, refer to the “Print Status Tab” on page 8-40.

8.4.3 Full Disclosure Zoom In Report
The Full Disclosure Zoom In Report (shown in FIGURE 8-29 and FIGURE 8-30) provides historical digital and waveform data available for the selected patient tile. The amount of disclosure data available for a patient is dependent on the amount of disclosure data allowed by the Panorama Central Station license. This report includes a header, digital data, waveform data and a footer.
FIGURE 8-29 Full Disclosure Zoom In Report (Page 1)

FIGURE 8-30 Full Disclosure Zoom In Report (Page 2)
8.4.3.1 Header
The Full Disclosure Zoom In Report’s first page header provides the title of the report and basic demographic information, if it is available. The demographic information includes the patient’s first and last name, patient ID, bed number, weight, height and doctor.

Page headers on subsequent pages of the report provide the report title, and limited demographic information about the patient (first and last name, patient ID and bed number), if it is available.

If the demographic data was not previously entered in the system, only the demographic label is shown.

8.4.3.2 Digital Data
The Full Disclosure Zoom In Report’s digital data section displays measurements for the digital data parameters currently available for the selected patient tile. The displayed parameter order is determined by the system Unit Priorities tab. The digital data was collected at the time of the print request. The digital data section contains the following:

- A parameter label
- A numeric measurement
- A unit of measure label
- A heart rate (HR) value
- Dashes (-) to indicate missing or invalid digital data.
- Square brackets ([ ]) to indicate alarming parameter values

**NOTE:** Digital Data is only shown on the first page of the report.

**NOTE:** ST digital data includes both Current (C) and Reference (R) values.

8.4.3.3 Waveform Data
The waveforms that print in this Full Disclosure Zoom In Report’s waveform data section are limited to which leads are currently in use, and on the waveforms selected in the Print Setup function.

The first page of this report contains four waveforms. The middle pages of the report contain two, four, or six waveforms depending on the waveforms available. Each waveform in this report represents approximately 10 seconds of data, seven(7) seconds prior to the time of the print request and three (3) seconds after the print request. A marker above each waveform strip indicates the time of the print request. A waveform label (i.e., ECG Lead II) is shown above each strip to identify the data source and the unit of measurement (i.e., %, RPM, BPM, mmHg, kPa). Pleth, ECG and Respiration do not display a unit of measurement. The waveforms displayed in the report are shown on a calibrated grid and contain a scale bar.
For pacemaker patients that have Pacer Enhancement enabled, the report displays a long vertical line crossing through the waveform or a short vertical line above the waveform (depending on the selection in the Pacer Settings dialog) to indicate firing of the pacemaker impulse.

**ECG Waveforms**
The ECG waveforms shown in the Full Disclosure Zoom In Report are limited to the ECG lead wire set currently in use and on the waveforms selected in the **Print Setup** tab. The ECG waveform report displays with the following:

- An ECG lead label
- A wave gain scale bar and a calibration pulse for wave gains less than 30 mm/mV
- A calibrated grid
- Current ECG monitoring label
- A continuous straight line for active ECG leads that are not connected

**Non-ECG Waveforms**
The non-ECG waveforms shown in the Full Disclosure Zoom In Report are based on the non-ECG waveforms that are in use and the non-ECG waveforms that are selected to be printed.

All non-ECG waveforms in the report contain a parameter label and a unit of measure label. Non-ECG waveforms in the report may contain a wave gain scale bar. If a non-ECG waveform is unavailable at the time of the print request, the report prints blank data for the specified time period.

- If the IBP waveform was labeled at the bedside monitor, the report contains that IBP label.
- If the Anesthetic Agent was identified at the bedside monitor, the report contains that Agent label.

### 8.4.3.4 Footer
The Full Disclosure Zoom In Report contains a footer at the bottom of every page. The footer includes:

- The date and time of the print request. The format displayed is taken from the **System Date/Time** tab.
- A centered page number.
- The name of the Panorama Central Station where the report was generated.

### 8.4.4 Printing a Full Disclosure Zoom In Report
The Full Disclosure Zoom In Report is printed in landscape orientation. Follow these steps to print this report:

1. From the **Disclosure** tab, select a portion of the compressed waveform view, then select the **Zoom In** button. The magnified view of the selected section is displayed in the **Waveforms** view.
2. Select the **Print** menu bar button. The report is sent to the printer.

Requesting this report posts a print job in the system Print Status tab. For additional information, refer to the “Print Status Tab” on page 8-40.
8.4.5 Graphic Trend Report

The Graphic Trend Report (titled Trend Graphic Report in FIGURE 8-31 and FIGURE 8-32) displays the trends for the selected patient tile in a graphical format. Graphic Trends are viewed by selecting the Graphic sidebar button in the Trends tab. This report includes a header, Digital data, Graphic data, and a footer section.

FIGURE 8-31 Graphic Trend Report (Page 1)
8.4.5.1 Header

The Graphic Trend Report’s first page header provides the title of the report and basic demographic information, if it is available. The demographic information includes the patient’s first and last name, patient ID, bed number, weight, height and doctor.

Page headers on subsequent pages of the report provide the report title, the date and time the data was collected, and limited demographic information about the patient (first and last name, patient ID and bed number), if it is available.

If the demographic data was not previously entered in the system, only the demographic label is shown.
8.4.5.2 Digital Data

The Graphic Trend Report’s digital data section displays measurements for the digital data parameters currently available for the selected patient tile. The displayed parameter order is determined by the system **Unit Priorities** tab. The digital data section contains:

- A parameter label
- A numeric measurement
- A unit of measure label
- A heart rate (HR) value
- Dashes (--) to indicate missing or invalid digital data.
- Square brackets ([ ]) to indicate alarming parameter values

**NOTE:** Digital Data is only shown on the first page of the report.

8.4.5.3 Graphic Data

The graphic data section of the Graphic Trend Report displays a cursor time, parameter list and a graphical representation of the parameter measurements that occurred during the selected time period.

- The cursor time represents the time that was selected in the Trends (Graphic) tab. The cursor time in the report follows the system Date/Time format.
- The parameters shown in this report display on the left side of the report. Each parameter has its own graphical layout which includes a parameter label, a unit of measure label, a scale bar and the scale bar range of values.
- For historical purposes, the Graphic Data section of the report shows all the selected parameters, even if the parameters are currently turned off.
- The report shows a vertical line on the graph to represent the occurrence of a physiological alarm.
- The report shows nine time periods across the top of the graph for a more representative look at the parameter data. The time periods will be shown in the system default time/date format.
- The time periods shown in the report are equal sections of the total time interval selected in the graphic report. For example, if a 4 hour interval is selected, graphical data is plotted in 30 minute increments and if an 8 hour interval is selected, data is plotted in 60 minute increments. This report is capable of plotting graphical data in 2, 4, 6 and 8 hour time intervals. The report is printed in the time interval specified in the Trends (Graphic) tab.

Time intervals are changed in the **Trends** (Graphic) tab by selecting the **Zoom In** and **Zoom Out** buttons.
- This report shows a cursor line in the graph for the Cursor Time posted on the report.
- The data plotted in the Graphic Trend Report reflects the trend data records that were collected based on the Trend Display interval configuration.
- The report shows one plotted point for each parameter sensor.
- Dual sensor parameters (such as INSP/ET CO2) are plotted individually then connected with a line (for example, inspired and end tidal values).
- Multiple sensor parameters (such as NIBP) are plotted individually then the systolic and diastolic points are connected and a single point is shown for the Mean measurement.

8.4.5.4 Footer

The Graphic Trend Report contains a footer at the bottom of every page. The footer includes:

- The date and time of the print request. The format displayed is taken from the System Date/Time tab.
- A centered page number.
- The name of the Panorama Central Station from which the report was generated.

8.4.5.5 Printing a Graphic Trend Report

The Graphic Trend Report is printed in landscape orientation, using any of the following:

- Select a valid plot point, then select the Print menu bar button from the Trends (Graphic) tab.
- From the bedside monitor, verify that Select Printer option is configured to print remotely, and that the trends graphic option is active, then press the PRINT TREND button.

Requesting this report posts a print job in the system Print Status tab. For additional information, refer to the “Print Status Tab” on page 8-40.
8.4.6 12-lead Report

The 12-lead Report (shown in FIGURE 8-33) provides analysis of 12 vectors of ECG data for the selected patient tile. This report includes a header and waveform data. All of the data contained in this report is obtained from the bedside monitor.

NOTE: This report is not available when using a WMTS 608 bedside device.

NOTE: The 12-lead Report is not available at the ViewStation.

FIGURE 8-33 12-lead Report

8.4.6.1 Header

The 12-lead Report’s page header provides the title of the report and basic demographic information, if it is available. The demographic information includes the patient’s first and last name, patient ID, bed number, weight, height and doctor.

If the demographic data was not previously entered in the system, only the demographic label is shown.

- Interpretation Area: The Interpretation area of the 12-lead report shows an analysis of the collected data.
  - If data is missing from the report, the message **Unable to Obtain Interpretation. Enter Patient Date of Birth and Gender** is displayed.
  - If the patient’s Patient Size is not set to Adult, the message **Unable to Obtain Interpretation. Patient not an adult** is displayed.
If the patient is less than 18 years old, the message **Unable to Obtain Interpretation. Patient Must Be At Least 18 Years Old** is displayed.

- **Unconfirmed Report:** The 12-lead Report is marked as an Unconfirmed Report when interpretation strings are printed on the report. This indicates that a physician must review the report for a diagnosis confirmation.

### 8.4.6.2 Waveform Data

The 12-lead Report section displays 2.5 seconds of waveform data for each of the 12 leads monitored at the bedside monitor, a 10-second ECG rhythm strip for Lead II, a calibration pulse and a wave gain setting of 1 millivolt.

### 8.4.6.3 Printing a 12-lead Report (Passport 2 and Spectrum)

The 12-lead Report prints when a print request is received from a connected bedside monitor. This report prints in landscape orientation.

Before this report is requested, the following must be configured at the bedside monitor:

- A View 12™ card must be in use
- The monitor must be in the View All ECG Mode
- Remote printing must be enabled

To print the report, select the **STRIP** button from the bedside monitor. A print request is sent from the bedside monitor to the Panorama Central Station’s selected printer.

Please refer to the Spectrum Operating Instructions (P/N 0070-00-0648-XX) for more specific printing instructions.

Requesting this report posts a print job in the system Print Status tab. For additional information, refer to the “Print Status Tab” on page 8-40.
8.4.7 Telepack Error Log Report

The Telepack Error Log Report (shown in FIGURE 8-34) is a one page report that shows all logged errors for any wireless Telepack devices connected to the Panorama Central Station in service mode. This report includes a header, error log data and a footer.

NOTE: The Telepack Error Log Report is not available at the ViewStation.

FIGURE 8-34 Telepack Error Log Report

8.4.7.1 Header

The Telepack Error Log Report header provides the title of the report and the connected Device ID for the connected wireless device.

8.4.7.2 Error Log Data

The Telepack Error Log Report’s error log data section displays all of the logged error messages for any connected Telepack device in a columnar format. These messages are used to diagnose a problem with the connected wireless device.

8.4.7.3 Footer

The Telepack Error Log Report contains a footer at the bottom the page. The footer includes:
• The date and time of the print request. The format displayed is taken from the **System Date/Time** tab. For more information regarding date/time formats, refer to the “Date/Time Tab” on page 9-41.

• A centered page number.

• The name of the Panorama Central Station where the report was generated.

### 8.4.7.4 Printing a Telepack Error Log Report

The Telepack Error Log Report prints in a landscape orientation. Follow these steps to print this report:

1. From the **Wireless** tab, select the **Error Log** sidebar button.
2. Select the **Print** menu bar button.

Requesting this report posts a print job in the system Print Status tab. For additional information, refer to the “Print Status Tab” on page 8-40.
8.4.8 System Alarm Report

The System Alarm Report (shown in FIGURE 8-35) shows all of the system default settings for the parameter alarms available in the Panorama Central Station. The report includes header, System Alarm data, optional arrhythmia, paging delay settings and ECG settings, and a footer section.

FIGURE 8-35 System Alarm Report - Paging Option Disabled
FIGURE 8-36 System Alarm Report - Paging Option Disabled (Page 2)
### FIGURE 8-37 System Alarm Report - Paging Option Enabled

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Event</th>
<th>Time</th>
<th>Date</th>
<th>Time Zone</th>
<th>Event Type</th>
<th>Event ID</th>
<th>Event ID Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alarm 1</td>
<td>001</td>
<td>OK</td>
<td>001</td>
<td>001</td>
<td>001</td>
<td>OK</td>
<td>001</td>
<td>OK</td>
</tr>
<tr>
<td>Alarm 2</td>
<td>002</td>
<td>OK</td>
<td>002</td>
<td>002</td>
<td>002</td>
<td>OK</td>
<td>002</td>
<td>OK</td>
</tr>
<tr>
<td>Alarm 3</td>
<td>003</td>
<td>OK</td>
<td>003</td>
<td>003</td>
<td>003</td>
<td>OK</td>
<td>003</td>
<td>OK</td>
</tr>
<tr>
<td>Alarm 4</td>
<td>004</td>
<td>OK</td>
<td>004</td>
<td>004</td>
<td>004</td>
<td>OK</td>
<td>004</td>
<td>OK</td>
</tr>
<tr>
<td>Alarm 5</td>
<td>005</td>
<td>OK</td>
<td>005</td>
<td>005</td>
<td>005</td>
<td>OK</td>
<td>005</td>
<td>OK</td>
</tr>
<tr>
<td>Alarm 6</td>
<td>006</td>
<td>OK</td>
<td>006</td>
<td>006</td>
<td>006</td>
<td>OK</td>
<td>006</td>
<td>OK</td>
</tr>
</tbody>
</table>

*Note: The diagram shows a table with columns for Parameter, Value, Event, Time, Date, Time Zone, Event Type, Event ID, and Event ID Description. The table includes entries for various system alarms, each with an OK status.*
8.4.8.1 Header
The System Alarm Report's header provides the title of the report.

8.4.8.2 System Alarm Information
The System Alarm section of the report will display all of the system alarm default settings, as configured at the Panorama Central Station. Alarm limits are listed for the Adult, Pediatric and Neonate patient sizes.

The following alarm information is shown in the report:

FIGURE 8-38 System Alarm Report - Paging Option Enabled (Page 2)
• **Parameter** displays the name of the parameter for which the alarm is set.
• **Units** displays the default unit of measure in which the parameter is shown. If the unit of measure is not applicable to the parameter, dashes (--) display instead of a unit of measure.
• **High Limit** displays the high limit for the alarm setting. Measurements that exceed this setting will trigger an alarm condition. If a high limit is not applicable to the parameter, dashes (--) display instead of the high limit.
• **Low Limit** displays the low limit for the alarm setting. Measurements below this setting trigger an alarm condition. If a low limit is not applicable to the parameter, dashes (--) display instead of the low limit.
• **Priority** displays the importance rating for the alarm. The alarm priorities available to each parameter may differ.
• **Print On Alarm** displays whether or not the parameter data should be printed during an alarm condition. Print On Alarm settings include ON and OFF.
• **Save To Event** displays whether or not the parameter data should be saved as an event during an alarm condition. Save To Event settings include ON and OFF.

Alarm settings can be customized to a patient using the patient Alarms tabs.

8.4.8.3 Arrhythmia Settings
The System Alarm Report's Arrhythmia Settings section includes values for the following:

- Asystole Delay
- V-Tach Threshold
- All Arrhythmia On/Off
- V-Tach Rate
- ECG Noise Delay
- Pause Threshold

**NOTE:** The arrhythmia settings in the System Alarm Report are only applicable to a Telepack device.

8.4.8.4 Paging Delay Settings
The System Alarm Report's Paging Delay section includes the system paging delay value that was configured in the System Alarms tab.

8.4.8.5 Signal Lost Delay Settings
The System Alarm Report's Signal Lost Delay section includes the signal lost delay value that was configured in the System Alarms tab.

8.4.8.6 ECG Settings
The System Alarm Report's ECG settings section includes values for the following:

- QT Monitoring On/Off
- QTc Formula

**NOTE:** The ECG settings in the System Alarm Report are only applicable to a Telepack device.
8.4.8.7 Footer
The System Alarm Report contains a footer at the bottom of the report page. The footer includes:

- The date and time of the print request. The format displayed is taken from the System Date/Time tab.
- A centered page number.
- The name of the Panorama Central Station where the report was generated.

8.4.8.8 Printing a System Alarm Report
The System Alarm Report is printed in portrait orientation. Follow these steps to print this report:

1. From the System Alarms tab, select the System Alarm Limits sidebar button.
2. Select the Print menu bar button. The report is sent to the printer.

Requesting this report posts a print job in the system Print Status tab. For additional information, refer to the “Print Status Tab” on page 8-40.

8.4.9 Troubleshooting
This section lists some of the potential messages and issues that may occur while trying to print an additional report.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>The report will not print</td>
<td>There are several reasons that would cause a report not to print. It could be a problem with the communication between the printer and the Panorama Central Station or simply a problem with the printer.</td>
<td>If there is an error that appears such as Printer out of Paper you will need to remedy that situation. If the No Printer connected or Printer not Available error is displayed check that the printer was properly configured with the Panorama Central Station. Refer to the Printer Configuration manual (P/N 0070-00-0561).</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
9.0  System Setup Functions

This section outlines the tabs and default configuration settings associated with the Panorama Central Station, the ViewStation, and the WorkStation.

- Parameter Color Tab
- Touch Screen Tab
- Installation Setup Tab
- Care Group Tab
- Volume Tab
- System Alarms Tab
- Passwords Tab
- Equipment Setup Tab
- More Tab
- Previous Tab
- Date/Time Tab
- Print Setup Tab (System)
- Unit Priorities Tab
- Unit Choices Tab
- Wave Gain Tab (System)
- Wireless Tab
9.1 Parameter Color Tab

The **Parameter Color** tab (shown in FIGURE 9-1) selects the system default colors for the parameters which will be displayed in the patient tile. This tab is available at the ViewStation and the WorkStation, and it functions in the same manner as it does at the host central station. Parameter Color tab changes made at the central station or remote workstations are independent of each other.

9.1.1 Accessing the Parameter Color Tab

- From the menu bar, select the **System Setup** button. The tabs associated with the **System Setup** button are displayed. By default, the **Parameter Color** tab is displayed first.

![FIGURE 9-1 Parameter Color Tab](image)

9.1.2 Selecting a Color for Parameter Data

This section outlines assigning a color to a parameter in the **Parameter Color** tab.

1. Select a color tile button.
   - A checkmark indicates that the colored tile is selected.
   - An empty tile indicates that a colored tile is not selected.

2. Select a parameter in the Parameter list box (use the scroll bars to view all of the parameters). When a parameter is selected, the associated label changes from a black background with white text to a white background with black text.

3. Choose the **Select** button to accept the parameter selection. The color is assigned to the selected parameter and the color icon is shown in the list box to the left of the parameter name.

9.1.3 Sidebar Button - Normal Screen

Select the **Normal Screen** sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.
9.1.4 Troubleshooting

This section lists some of the potential messages and issues that may occur while using the **Parameter Color** tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>The System Setup menu button is disabled.</td>
<td>The System Setup function is already open on another Panorama display and cannot be opened on multiple displays.</td>
<td>Close the System Setup function on the Panorama display where it is open.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
9.2 Touch Screen Tab

The **Touch Screen** tab provides a means to turn the touch screen on and off and recalibrate (or reset) the touch screen. This tab is available at the ViewStation and the WorkStation, and it functions in the same manner as it does at the host central station. Touch Screen tab changes made at the central station or remote workstations are independent of each other.

9.2.1 Accessing the Touch Screen Tab

1. From the menu bar, select the **System Setup** button. The first set of **System Setup** tabs is displayed.

2. Select the **Touch Screen** tab. A **Touch Screen On/Off** and **Recalibrate Touch Screen** button is displayed in the tab (shown in FIGURE 9-2).

3. For further instructions, refer to sections Turning the Touch Screen On or Off and Recalibrating the Touch Screen.

![FIGURE 9-2 Touch Screen tab](image)

9.2.1.1 Turning the Touch Screen On or Off

The touch screen may be turned on or off by selecting the Touch Screen On/Off button. If the touch screen is turned off, a mouse is required to navigate. When the touch screen is turned off, the message **Touch Screen Off** will be displayed in the lower left corner of the screen in the System Status line. By default, the touch screen is turned on.

9.2.1.2 Recalibrating the Touch Screen

The touch screen should be recalibrated whenever the touch interface becomes difficult to maneuver. By default the

1. From the menu bar, select the **System Setup** button. The first set of **System Setup** tabs is displayed.

2. Select the **Touch Screen** tab.

3. Select the **Recalibrate Touch Screen** button. The recalibration process is initiated (shown in FIGURE 9-3).
4. Touch the red targets on the screen as instructed. Three calibration targets will be displayed.

**NOTE:** The messages in the Recalibrate Touch Screen display are always displayed in the English language.

**NOTE:** During the recalibration process, the patient tiles, menu bar, and sidebar buttons will be replaced by the recalibration screen.

5. Once the three targets have been touched, the message **Touch the screen. Does the cursor follow your finger?** is displayed (shown in FIGURE 9-5).

- Select either the ✅ or ⬅️ button.
  - Select the ✅ button if the targets moved as they were selected. Once selected, the screen is recalibrated.
  - Select the ⬅️ button if the targets did not move as they were selected. Once selected, the recalibration process begins again.

**NOTE:** When the central station, ViewStation, or WorkStation uses a dual display configuration, recalibration will affect both touch screen displays. The first touch screen display will be calibrated and then the second.
9.2.2 Sidebar Button - Normal Screen
Select the **Normal Screen** sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

9.2.3 Troubleshooting
This section lists some of the potential messages and issues that may occur **Touch Screen** tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Recalibration settings not accepted</td>
<td>The Recalibration screen timed out because the first target was not touched within 30 seconds.</td>
<td>Try to recalibrate again and touch the first target before the 30 second time out.</td>
</tr>
<tr>
<td>Recalibration steps are repeated</td>
<td>The [✓] button was selected in error.</td>
<td>Recalibrate again and if the targets moved as they were selected, select the [✓] button.</td>
</tr>
<tr>
<td><strong>Touch Screen OFF</strong></td>
<td>The Touch Screen On button was selected.</td>
<td>Use the mouse to navigate. If touch screen functionality is needed, press the Touch Screen OFF button again to reactivate the touch screen.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.

Contact your local Service Representative if recalibration of the touch screen does not improve the performance of the touch screen display.
9.3 Installation Setup Tab

The system Installation Setup tab (shown in FIGURE 9-5) controls access to restricted central station functions. This tab is available at the ViewStation and the WorkStation, and it functions in the same manner as it does at the host central station. The Installation Setup tab is password protected.

9.3.1 Accessing the Installation Setup Tab

1. From the menu bar, select the System Setup button. The first set of System Setup tabs is displayed.
2. Select the Installation Setup tab. The Installation Setup tab is displayed.

![FIGURE 9-5 Installation Setup Tab](image)

9.3.2 Entering a Password

To gain access to the protected central station system functions, the correct password must be entered in the Installation Setup tab’s password box. The password required for the Installation Setup tab is defined in the system Password tab.

1. Select the Password text box in the Installation Setup tab. The keyboard dialog box is displayed.
2. Using the keyboard dialog box, enter the System password that was established in the Passwords tab (maximum 15 characters). For security purposes asterisks (*) are shown for each character entered.
3. Accept or reject the text entered in the keyboard dialog box. For additional information about using the keyboard dialog box, refer to “Keyboard Dialog Box” on page 2-18.

The following tabs can be accessed at the central station after the correct password has been entered in the Installation Setup tab: Care Group, Volume, System Alarms, Passwords, Equipment Setup, More/Previous, Date/Time, Unit Priorities, Unit Choices, Wave Gain, and Wireless.

The following tabs can be accessed at the ViewStation and the WorkStation after the correct password has been entered in the Installation Setup tab: Volume, System Alarms, Passwords, Date/Time, and Unit Choices.
9.3.3 Sidebar Button - Normal Screen

Select the **Normal Screen** sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

9.3.4 Troubleshooting

This section lists some of the potential messages and issues that may occur while using the **Installation Setup** tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Incorrect password. Please try again.</td>
<td>The wrong password was entered in the <strong>Password</strong> text box.</td>
<td>Select the <strong>OK</strong> button to close the dialog box. Enter the correct password in the text box.</td>
</tr>
<tr>
<td>Incorrect password. This is your last chance. Please try again.</td>
<td>The wrong password was entered two consecutive times in the <strong>Password</strong> text box.</td>
<td>Select the <strong>OK</strong> button to close the dialog box. Enter the correct password in the text box.</td>
</tr>
<tr>
<td>The <strong>Installation Setup</strong> tab closes and the Panorama Central Station is returned to a full screen monitoring display</td>
<td>The wrong password was entered three consecutive times in the <strong>Password</strong> text box.</td>
<td>Reopen the <strong>Installation Setup</strong> tab and enter the correct password in the text box.</td>
</tr>
<tr>
<td>The <strong>System Setup</strong> menu button is disabled.</td>
<td>The <strong>System Setup</strong> function is already open on another Panorama display and cannot be opened on multiple displays.</td>
<td>Close the System Setup function on the Panorama display where it is open.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
9.4 Care Group Tab

The system Care Group tab (shown in FIGURE 9-6) allows for the default assignment of selected patient tiles to specific system care groups. This enables a patient tile to automatically be assigned to a specific care group upon admission to the Panorama Central Station. The central station supports up to twelve care groups. Each care group is associated with a unique color indicator and a unique reference label. The Care Group tab and the associated colorized indicator are not available at the ViewStation or the WorkStation.

Care groups may be used to indicate a special meaning with regard to a patient tile. For example, care groups may be used to represent a specific care-giver, hospital department, or a critical monitoring condition. Contact the System Administrator before making adjustments to the system care groups.

9.4.1 Accessing the Care Group Tab

1. From the menu bar, select the System Setup button. The Password tab is displayed.
2. Follow the directions in the "Entering a Password" section on page 9-7.
3. Once the correct password has been entered, the tabs associated with the System Setup button are displayed. By default, the Care Group tab is displayed first.

![FIGURE 9-6 Care Group Tab](image)

9.4.2 Editing Care Group Labels

This section outlines editing the system Care Group reference label.

Selecting a Patient’s Care Group

1. Select a care group by choosing the colored tile below the care group label. When a care group is selected, the reference label changes from a black background with white text to a white background with black text.
2. Select the Edit Labels button. The keyboard dialog box is displayed.
3. Enter the name for the selected care group (maximum of 6 characters). The name is displayed in the white text box.
4. Accept or reject the text entered in the keyboard dialog box. For additional information about using the keyboard dialog box, refer to “Keyboard Dialog Box” on page 2-18.
9.4.3 Assigning a Patient Tile to a Care Group

The assignment of a system care group occurs when a patient is admitted to a tile at the Panorama Central Station. When a patient tile is assigned to a care group, the color associated with the care group will be shown in vertical bar beside the patient tile VIEW button. To adjust the care group assignment of an active patient tile, refer to the “Demographics Tab” on page 5-2.

This section outlines assigning system default care group settings to specific patient tiles.

Selecting/Editing a Patient’s Care Group

1. Select a care group tile by selecting the colored tile below the care group label. When a tile is selected the associated label changes from a black background with white text to a white background with black text.

2. Select the tile number to be assigned to the selected care group. When a tile is selected it changes from a black background with white text to a white background with black text.

3. Accept or reject the care group assignment.
   - Select the Add button to assign the selected patient to the care group. The care group icon is displayed next to the selected patient tile in the care group list box.
   - Select the Clear button to delete the care group currently assigned to the selected patient.

9.4.4 Sidebar Button - Normal Screen

Select the Normal Screen sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

9.4.5 Troubleshooting

This section lists some of the potential messages and issues that may occur while using the system Care Group tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>The System Setup menu button is disabled.</td>
<td>The System Setup function is already open on another Panorama display and cannot be opened on multiple displays.</td>
<td>Close the System Setup function on the Panorama display where it is open.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
9.5 Volume Tab

The system Volume tab (shown in FIGURE 9-7 and FIGURE 9-8) adjusts the system default volume levels. Volume levels can be adjusted for system alarms and physiological alarms. This tab is available at the ViewStation and the WorkStation, and it functions in the same manner as it does at the host central station. Volume tab changes made at the central station or remote workstations are independent of each other.

9.5.1 Accessing the Volume Tab

1. From the menu bar, select the System Setup button. The Parameter Color tab is displayed.
2. Select the Installation Setup tab. The Installation Setup tab is displayed.
3. Follow the directions in the "Entering a Password" section on page 9-7.
4. Once the correct password has been entered, the tabs associated with the System Setup button are displayed.
5. Select the Volume tab. The Volume tab is displayed.

FIGURE 9-7 Volume Tab (Central Station)

FIGURE 9-8 Volume Tab (ViewStation and WorkStation)

CAUTION: Do not block the speakers. Set the volume levels so that alarms can be heard at all times, as described in this Operation Manual.
Setting Reminder Tone Volume Level
The Reminder Tone volume setting determines the volume of the sound associated with the Panorama Central Station Reminder Tone settings.

Characteristics of the Reminder Tone volume setting:

- A volume level range of 1-10
- A factory default setting of 3
- It cannot be turned OFF

NOTE: Adjustments to the Reminder Tone volume setting will take effect immediately.

To adjust the Reminder Tone volume level:

- Move the volume control scroll bar box or the scroll bar arrows up (to maximize sound) or down (to minimize sound).

Setting a System Alarms Volume Level
The System Alarms volume setting determines the volume of the sound associated with the Panorama Central Station alarm settings.

Characteristics of the System Alarms volume setting:

- A volume level range of 10-100%
- A factory default setting of 30%
- It cannot be turned OFF

NOTE: Adjustments to the System Alarms volume setting will take effect immediately.

To adjust the System Alarms volume level:

- Move the volume control scroll bar box or the scroll bar arrows up (to maximize sound) or down (to minimize sound)

Setting a Physiological Alarms Volume Level
The Physiological Alarm volume level is a system setting that applies to enabled numeric and arrhythmia alarm violations. The Panorama Central Station allows up to three time specified volume settings to be configured for physiological alarms. This feature allows user-defined volume settings for physiological alarms to be specified, based on the time of day.

Characteristics of the Physiological Alarm volume setting:

- The time period format for physiological alarms will follow the system’s time format
- A volume level range of 10-100%
- A factory default setting of 100%
- It cannot be turned OFF

CAUTION: Do not block the speakers. Set the volume levels so that alarms can be heard at all times, as described in this Operation Manual.
1. Select the colored tile that corresponds to the time period for which the volume level is being adjusted.
   - A checkmark in the colored tile indicates that the tile has been selected and the From/To fields are displayed.
   - A colored tile without a checkmark indicates that the tile has not been selected.

2. Select the From/To fields to enter a time period for the selected physiological alarm. The keypad dialog box displays.
   - Delete existing hour and minute data by using the CLEAR or CE buttons on the keypad.
   - When using the 12-hour time format, the AM/PM toggle button switches between the AM and PM settings.

NOTE: Time periods are mutually exclusive. When a new time period is entered, the time periods for the other physiological alarms are automatically adjusted to prevent overlapping of time periods.

3. Set the volume level for the selected physiological alarm.
   To adjust the volume level, move the volume control scroll bar box or the scroll bar arrows up (to maximize sound) or down (to minimize sound). The colored icon to the right of the scroll bar is automatically moved when the volume level is adjusted.

4. Select the Done button to save the physiological alarm volume adjustments. The Physiological Alarm volume adjustments will take effect once the Done button is selected.

9.5.2 Sidebar Button - Normal Screen
Select the Normal Screen sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

9.5.3 Troubleshooting
This section lists some of the potential messages and issues that may occur while using the Volume tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>The <strong>System Setup</strong> menu button is disabled.</td>
<td>The <strong>System Setup</strong> function is already open on another Panorama display and cannot be opened on multiple displays.</td>
<td>Close the <strong>System Setup</strong> function on the Panorama display where it is open.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
9.6 System Alarms Tab

The System Alarms tab (shown in FIGURE 9-10, FIGURE 9-11, and FIGURE 9-11) controls the default settings for system alarm options, specific alarm limits, and alarm response functions. This tab is available at the ViewStation and the WorkStation, and it functions in the same manner as it does at host central station with fewer buttons. System Alarms tab changes made at the central station or remote workstations are independent of each other.

For information regarding individual patient alarm settings, refer to the “Patient Alarm Setup Tab (Optional)” on page 6-2.

9.6.1 System Alarms (Options) Tab

The System Alarms (System Alarm Options) tab controls the system default settings for patient alarms.

9.6.1.1 Accessing the System Alarm (Options) Tab

1. From the menu bar, select the System Setup button. The Password tab is displayed.
2. Follow the directions in the “Entering a Password” section on page 9-7.
3. Once the correct password has been entered, the tabs associated with the System Setup button are displayed.
4. Select the System Alarms tab. The System Alarms (System Alarm Options) tab is displayed.

FIGURE 9-9 System Alarms (System Alarm Options) Tab (Central Station)

FIGURE 9-10 System Alarms (System Alarm Options) Tab - Paging Option Enabled (Central Station)
9.6.1.2 Setting the System Alarms Options (Central Station)

This section outlines selecting system alarm option defaults.

1. Select the **Latch Alarms** toggle button to either enable or disable the latch alarms function. Latch Alarm options include **All**, **Lethals** and **None**. The factory default setting is **All**. Alarm latching allows physiological alarms to continue their alarm behaviors until the alarm is acknowledged by the user.

   - The **All** option indicates that all alarms are latched.
   - The **Lethals** option indicates that only lethal arrhythmia alarms are latched.
   - The **None** option indicates that no alarms are disabled.

**NOTE:** The Latch Alarms option will not affect apnea alarms.

**NOTE:** Changes to the Latch Alarms option will only affect new alarm violations.
2. Select the **Audio Off/Pause Duration** button to select the audio pause duration period for alarms in the Panorama Central Station. The audio pause duration period is the amount of time an alarm parameter will be silenced after selecting the **Alarm ACK** button. Audio pause duration periods include 2 minutes, 3 minutes, 5 minutes, and 10 minutes. When the Permanent period is selected, the button displays as **Audio Off Duration** after selecting the **Alarm ACK** button. The factory default setting is 2 minutes.

**NOTE:** Changes to the Audio Off/Pause Duration period will take effect within ten (10) seconds of the change.

**NOTE:** Using the Audio Off/Pause Duration button will not affect the visual indicators associated with an alarm.

3. Select the **Technical Events Sounds** toggle button to enable or disable sound during a technical event. **Technical Events Sounds** options include **ON** and **OFF**. The factory default setting is **ON**.
   - The **ON** option indicates that **Technical Events Sounds** will be heard during a technical event.
   - The **OFF** options indicates that **Technical Events Sounds** will not be heard during a technical event.

**NOTE:** Changes to the Technical Events Sounds option will affect all patients within ten (10) seconds of the change.

4. Select the **Apnea Latching** toggle button to either enable or disable the apnea latching function. Apnea latching is the ability to set Apnea alarms to continue the alarm behaviors until the alarm is acknowledged by the user. Apnea latching options include **ON** and **OFF**. The factory default setting is **ON**.
   - The **ON** option indicates that **Apnea Latching** will be enabled.
   - The **OFF** options indicates that **Apnea Latching** will not be enabled.

**NOTE:** Changes to the Apnea option will affect all new Apnea alarms.

5. Select the **Paging Delay** button to select a paging delay for the system. This button will only be available when the paging option has been enabled. The Paging Delay options include 5 seconds, 10 seconds, 15 seconds, 20 seconds, 25 seconds, and 30 seconds. The factory default setting is 10 seconds.

**NOTE:** Changing the Paging Delay will only be applied to newly admitted patients and will not affect patients currently being monitored or to currently alarming parameters.

**NOTE:** This option will only be displayed if the Paging option is enabled at the central station.

6. Select the **Password Protection** toggle button to either enable or disable the password protection for the **Patient Alarms** tab in the Panorama Central Station or the WorkStation. **Password Protection** options include **ON** and **OFF**. The factory default setting is **ON**.
   - The **ON** option indicates that **Password Protection** will be enabled when trying to access the **Patient Alarms** tab.
   - The **OFF** options indicates that **Password Protection** will not be enabled when trying to access the **Patient Alarms** tab.
NOTE: Changes to the Password Protection option will affect all patients upon entering the Patient Alarms tab.

7. Select the **Pause** toggle button to either enable or disable the ability to suspend all, all non-high, or disable the ability to suspend any alarms. Pause options include **All Alarms**, **OFF**, and **All Non-High Alarms**. The factory default setting is **All Alarms**.
   - The **All Alarms** option indicates that the **All Alarms Pause** and the **All Non-High Alarms Pause** sidebars will be available in the **Patient Alarms** tab.
   - The **All Non-High Alarms** option indicates that only the **All Non-High Alarms Pause** sidebar button will be available in the **Patient Alarms** tab.
   - The **OFF** option indicates that neither the **All Alarms Pause** nor the **All Non-High Alarms Pause** sidebar buttons will be accessible in the **Patient Alarms** tab.

NOTE: Changes to the **Pause** option will affect all patients upon entering the **Patient Alarms** tab.

8. Select the **ST Alarm Delay** button to set an alarm delay for ST alarms. **ST alarm delay** is the period of time that the Panorama Central Station will wait before ST alarm notification. **ST Alarm Delay** options include 30 seconds, 45 seconds, 1 minute, 90 seconds, 2 minutes, and 3 minutes. The factory default setting is 30 seconds.

NOTE: ST Alarm Delay only affects new ST alarms for Telepack devices. When using a bedside monitor, the ST alarm delay settings are transferred to the central station.

9. Select the **Alarm Delay** button to set an alarm delay for the Panorama Central Station. The alarm delay is the period of time that the Panorama Central Station will wait before alarm notification. **Alarm Delay** options include None, 1 second, 2 seconds, 3 seconds, 4 seconds, 5 seconds, 6 seconds, 7 seconds, and 8 seconds. The factory default setting is None.

NOTE: Alarm Delay only affects new alarms for Telepack devices. When using a bedside monitor, the ST alarm delay settings are transferred to the central station.

10. Select the **Pause Duration** button to temporarily suspend all alarms or all non-high alarms for a configured time period. This button is disabled when the **Pause** option is set to **Off**. When this option is enabled, and a time period is selected, either the **All Alarms Pause** or the **All Non-High Alarms Pause** sidebar button in the **Patient Alarm Limits** and **Patient Alarm Responses** tab must be selected before the alarm suspension begins. **Pause Duration** options include 2 minutes, 3 minutes, 5 minutes, 10 minutes, and Permanent. The factory default setting is 2 minutes.

11. Select the **Reminder Tone** button to enable or disable a reminder tone. The reminder tone is a sound to remind the user of audio off and all alarms off conditions. The **Reminder Tone** option includes **ON** and **OFF**. The factory default setting is **ON**.

NOTE: Changes to the **Reminder Tone** option will affect all patients on the **Central Station**.

NOTE: After the reminder tone function is enabled, the Central Station will sound the tone if a patient tile is in audio off or alarms off condition, and audio alarm are absent at the Central Station.
12. Select the **Reminder Tone Interval** button to set the reminder tone interval. The Reminder Tone Interval option includes 1 minute, 2 minutes and 3 minutes. The factory default setting is 3 minutes.

9.6.1.3 Setting the System Alarm Options (ViewStation and WorkStation)

This section outlines selecting system alarm option defaults at the ViewStation and the WorkStation.

The ViewStation and the WorkStation are independent of each other. For example, if an option is selected at the ViewStation, the WorkStation will not be affected unless the same option is selected at the WorkStation.

1. Select the **Password Protection** toggle button to either enable or disable the password protection for the **Patient Alarms** tab at the WorkStation. **Password Protection** options include **ON** and **OFF**. The factory default setting is **ON**.
   - The **ON** option indicates that **Password Protection** will be enabled when trying to access the **Patient Alarms** tab.
   - The **OFF** options indicates that **Password Protection** will not be enabled when trying to access the **Patient Alarms** tab.

**NOTE:** The Password Protection toggle button is not available at the ViewStation.

**NOTE:** Changes to the Password Protection option will affect all patients upon entering the **Patient Alarms** tab at the WorkStation.

**NOTE:** Select the **Audio Off/Pause Duration** button to select the mute duration period for alarms. The mute duration period is the amount of time an alarm parameter will be silenced. Audio pause duration periods include 10 seconds, 15 seconds, 30 seconds, 45 seconds, 60 seconds, 120 seconds, and Permanent. When the Permanent period is selected, the button displays as **Audio Off Duration**. The factory default setting is 30 seconds. The host central station will not have an Audio Off/Pause button. The Audio Off/Pause Duration button will affect all patient tiles at the ViewStation and WorkStation (if selected at both sites). The Audio Off/Pause function will not affect the patient tiles at the host central station.

**NOTE:** Selecting the Audio Off/Pause Duration button will not affect the visual indicators associated with an alarm.

2. Select the **Technical Events Sounds** toggle button to enable or disable sound during a technical event. **Technical Events Sounds** options include **ON** and **OFF**. The factory default setting is **ON**.
   - The **ON** option indicates that **Technical Events Sounds** will be heard during a technical event.
   - The **OFF** options indicates that **Technical Events Sounds** will not be heard during a technical event.

**NOTE:** Changes to the Technical Events Sounds option will affect all patients within ten (10) seconds of the change.
3. Select the **Reminder Tone** button to enable or disable a reminder tone. The reminder tone is a sound to remind the user of the audio off status. The Reminder Tone options include **ON** and **OFF**. The factory default setting is **ON**.

**NOTE:** Changes to the Reminder Tone option will affect all patients on the WorkStation or ViewStation.

4. Select the **Reminder Tone Interval** button to set the reminder tone interval. The Reminder Tone Interval option includes 1 minute, 2 minutes and 3 minutes. The factory default setting is 3 minutes.

### 9.6.2 System Event Filtering

Select the **Events Filtering** sidebar button to control the system default event filters that will initially be used when a selected patient tile’s **Events** tab is opened.

System Event filters can be configured based on time, parameters, or both. ViewStation and WorkStation patient tiles will initially display the same event filter settings of the associated patient tile at the host central station.

For information regarding patient level event filtering, refer to “Event Filtering” on page 7-29.

#### 9.6.2.1 Accessing the Events Filter

To access the system events filter view (shown in FIGURE 9-13), select the **Events Filtering** sidebar button from the **System Alarms Options** tab.

![FIGURE 9-13 System Alarms - Events Filtering (Central Station)](image)

**FIGURE 9-13** System Alarms - Events Filtering (Central Station)

#### 9.6.2.2 Components of the Events Filter

**Event Filter Text Box**

The Event Filter text box is where the selected filter settings are displayed. The factory default Event Filter text box lists “All” as the initial filter, which indicates that there are no active event filters being used.

**Done Button**

Select the **Done** button to close the **All Event Filter** view and return to the **System Alarm Options** tab.
Cancel Button
Select the **Cancel** button to close the **All Event Filter** view and return to the **System Alarm Options** tab.

9.6.2.3 Time Filters
By indicating a start time and an end time, only events included within the configured time period will be displayed.

Accessing the Time Filter
To access the Time Filter view (shown in FIGURE 9-14), select the **Time** sidebar button from the **Events Filter** view. If any previously selected time ranges were specified, they will be displayed in the Event Filter text box beside the **Time** label, and in the Start Time and End Time text boxes.

![FIGURE 9-14 Time Event Filter](image)

Clearing Time Filters
To remove previously entered time filters, select the **Clear** button from the **Time Filter** view.

Configuring a Time Filter
To enter a start time:

1. Select the Start Time text box. The Start Time dialog box (shown in FIGURE 9-15) is displayed.

![FIGURE 9-15 Start Time Dialog Box](image)
2. Rotate the dial controls to select the hour and minute values.
3. Rotate the dial controls to select the month, day, and year values.
4. Accept or reject the selection made in the Start Time dialog box.
   - The Done button accepts the selections. When the Done button is selected, the date and time selected populates the Start Time text box, End Time text box, and the Event Filter text box.
   - The Cancel button discards the selections, closes the dialog box, and returns to the Time Event Filter view.

To enter an end time:

1. Select the End Time text box. The End Time dialog box (shown in FIGURE 9-16) is displayed.

![End Time Dialog Box](image)

**FIGURE 9-16** End Time Dialog Box

2. To enter the end time, follow steps 2, 3, and 4 from the previous section (entering a start time).

**9.6.2.4 Parameter Filters**

By indicating a parameter filter, only the selected parameters will be displayed. Parameter filters can be configured for physiological and technical events.

**Accessing the Parameter Filter**

To access the Parameter Filter (shown in FIGURE 9-17), select the Param sidebar button from the All Events view. By default, the physiological events are shown first. If any previously selected parameters were specified, they will be displayed in the Event Filter text box beside the Param: label.
Configuring a Parameter Filter

To configure a parameter filter:

1. If a technical event filter is needed, select the Tech button.

**NOTE:** By default, the physiological events are displayed when the Param sidebar is selected.

2. Move/remove the selected events by using any of the buttons described in the following table:

<table>
<thead>
<tr>
<th>BUTTON</th>
<th>FUNCTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Remove All</td>
<td>Removes all the events listed in the Selected Events list box and from the Event Filter text box.</td>
</tr>
<tr>
<td>Add/Right Arrow</td>
<td>Adds the selected event to the Selected Events list box, and to the Event Filter text box.</td>
</tr>
<tr>
<td>Remove/Left Arrow</td>
<td>Removes the chosen event from the Selected Events list box and from the Event Filter text box.</td>
</tr>
</tbody>
</table>

**NOTE:** A maximum of five physiological and/or technical events may be added to the Selected Events list at one time.

3. Select the Done button to apply the selected event filter and return to the List view or select the Cancel button to restore the previously chosen filter settings.

**9.6.2.5 Sidebar Buttons**

The following sidebar buttons are used to expand the functionality of this tab.

**System Alarm Limits**

Select the System Alarm Limits sidebar button to display the System Alarm Limits tab. For additional information regarding system alarm limits, refer to “System Alarms (Alarm Limits) Tab” on page 9-24.

**NOTE:** The System Alarm Limits sidebar button is not available at the ViewStation or the WorkStation.
System Alarm Responses
Select the System Alarm Responses sidebar button to display the System Alarm Responses tab. For information regarding system alarm responses, refer to “Accessing the System Alarms (System Alarm Responses) Tab” on page 9-26.

NOTE: The System Alarm Responses sidebar button is not available at the ViewStation or the WorkStation.

Events Filtering
This sidebar button was previously described on page 9-19.

Normal Screen
Select the Normal Screen sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

9.6.2.6 Troubleshooting
This section lists some of the potential messages and issues that may occur while using the System Alarm Options tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>The System Setup menu button is disabled.</td>
<td>The System Setup function is already open on a Panorama display and cannot be opened on multiple displays.</td>
<td>Close the System Setup function on the Panorama display on which it is opened.</td>
</tr>
<tr>
<td>The Paging Delay button is not displayed.</td>
<td>The central station where the patient tile resides does not have the Paging option enabled.</td>
<td>If the Paging feature was purchased, enable it.</td>
</tr>
<tr>
<td>The System Alarm Limits and System Alarm Responses tab is not available at the ViewStation or WorkStation.</td>
<td>System alarm limits and responses must be changed at the central station.</td>
<td>Change the alarm limits and responses from the central station.</td>
</tr>
<tr>
<td>Pause Duration button is disabled.</td>
<td>The Pause option is set to OFF.</td>
<td>Change the Pause option to All or All Non-High Alarms.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
9.6.3 System Alarms (Alarm Limits) Tab

The System Alarms (System Alarm Limits) tab [shown in FIGURE 9-18] controls the system default settings for the Telepack alarm limits.

The default parameter alarm limits are taken directly from the bedside device. The System Alarms (Alarm Limits) is available at the Panorama Central Station and the WorkStation. The ViewStation does not have an Alarm Limits tab. ViewStation Alarm limit settings are transferred from the host central to the ViewStation when the patient tile is assigned.

9.6.3.1 Accessing the System Alarms (System Alarm Limits) Tab

1. From the menu bar, select the System Setup button. The Password tab is displayed.
2. Follow the directions in the "Entering a Password" section on page 9-7.
3. Once the correct password has been entered, the tabs associated with the System Setup button are displayed.
4. Select the System Alarms tab. The System Alarms (System Alarm Options) tab is displayed.
5. Select the System Alarm Limits sidebar button. The System Alarms (System Alarm Limits) tab is displayed.

![FIGURE 9-18 System Alarms (System Alarm Limits) Tab (Central Station)](image_url)

9.6.3.2 Setting the System Alarm Limits

This section outlines setting system defaults for alarm limits, according to patient size.

The following steps should be performed for each patient size.

1. Select the Patient Size button to set the system alarm limit defaults for a specific patient size. Patient Size options include Adult, Pediatric, and Neonate. The factory default setting is Adult. When a Patient Size is selected, the button will change from black text on a grey background to grey text on a grey background.
2. Use the horizontal scroll bar in the System Alarms (System Alarm Limits) tab to locate the slider controls that need to be set or modified. Each page can show a maximum of six slider controls and a minimum of one slider control on each screen.
   - Select the double arrow scroll bar buttons to display the next/previous page of available parameter slider controls (until the first/last page is reached).
   - Select the single arrow scroll bar buttons to display the next/previous slider control (until the last/first control is shown).
3. Manually set the alarm limits for the selected parameter.

   To set a high alarm limit:
   - Select the upper alarm limit button to the left of the parameters slider bar. This button is either labeled with the word **OFF** or with the high alarm limit setting.
   - To set an upper alarm limit, either select the up/down arrow buttons on the parameters vertical scroll bar or drag and drop the scroll box in the scroll bar.

   To set a low alarm limit:
   - Select the lower alarm limit button to the left of the parameters slider bar. This button is either labeled with the word **OFF** or with the low alarm limit setting.
   - To set a lower alarm limit, either select the up/down arrow button on the parameter’s vertical scroll bar or drag and drop the scroll box in the scroll bar.

   For additional information on the default alarm limit ranges, refer to the “System Alarms Tab” on page 9-14.

9.6.3.3 Sidebar Buttons

The following sidebar buttons are used to expand the functionality of this tab.

**System Alarm Options**
Select the **System Alarm Options** sidebar button to display the **System Alarm Options** tab. For additional information regarding System Alarm Options, refer to the “System Alarms Tab” on page 9-14.

**System Alarm Responses**
Select the **System Alarm Responses** sidebar button to display the **System Alarm Responses** tab. For additional information regarding System Alarm Responses, refer to the “System Alarms Tab” on page 9-14.

**Normal Screen**
Select the **Normal Screen** sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

9.6.3.4 Troubleshooting

This section lists some of the potential messages and issues that may occur while using the **System Alarm Limits** tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>The <strong>System Setup</strong> menu button is disabled.</td>
<td>The <strong>System Setup</strong> function is already open on a Panorama display and cannot be opened on multiple displays.</td>
<td>Close the <strong>System Setup</strong> function on the Panorama display on which it is opened.</td>
</tr>
<tr>
<td>The <strong>System Alarm Limits</strong> and <strong>System Alarm Responses</strong> tab is not available at the ViewStation or WorkStation.</td>
<td>System alarm limits and responses must be changed at the central station.</td>
<td>Change the alarm limits and responses from the central station.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
9.6.4 System Alarms (Alarm Responses) Tab

The System Alarms (System Alarm Responses) tab (shown in FIGURE 9-19 and FIGURE 9-20) controls the system default settings for alarm responses at the Panorama Central Station and the WorkStation. This tab is not available at the ViewStation or the WorkStation. Alarm response settings are transferred from the host central to the ViewStation and the WorkStation when the patient tile is assigned.

9.6.4.1 Accessing the System Alarms (System Alarm Responses) Tab

The System Alarms (System Alarm Responses) tab (shown in FIGURE 9-20) controls the system default settings for alarm responses.

1. From the menu bar, select the System Setup button. The Password tab is displayed.
2. Follow the directions in the "Entering a Password" section on page 9-7.
3. Once the correct password has been entered, the tabs associated with the System Setup button are displayed.
4. Select the System Alarms tab. The System Alarms (System Alarm Options) tab is displayed.
5. Select the System Alarm Responses sidebar button. The System Alarms (System Alarm Responses) tab is displayed.

FIGURE 9-19 System Alarms (System Alarm Responses) Tab (Central Station)

FIGURE 9-20 System Alarms (System Alarm Responses) Tab - Paging Option Enabled (Central Station)
9.6.4.2 Setting the System Alarms Responses

This section outlines setting system defaults for alarm responses.

1. Select a parameter in the **System Alarm Limits** (System Alarm Responses) tab to make adjustments.
   - The alarm responses list box shows a list of the available alarm conditions.
   - When an alarm is selected, it will be highlighted in white with black text and will be displayed on the left side of the Priority button.

2. Select the alarm **Priority** button to select the alarm priority for the selected alarm. The Alarm Priority setting determines the audio and visual indicators the central station will elicit when an alarm condition occurs. Alarm Priority is based on the severity of the alarm condition. Every alarm condition will have predetermined alarm priority choices. The Alarm Priorities available include Off, 1, 2, and 3, depending on the alarm.
   - Priority 1 alarms are the most severe alarms (lethal alarms). Priority 1 alarms are shown in red.
   - Priority 2 alarms include alarms of medium severity (most physiological numeric alarms). Priority 2 alarms are shown in yellow.
   - Priority 3 alarms are the least severe alarms (technical alarms and some physiological alarms). Priority 3 alarms are shown in yellow.
   - When Alarm Priorities are turned OFF there will be no audio feedback for alarms at the Panorama Central Station.

3. Select the **Print on Alarm** button to determine if the data for the alarm should be printed when the alarm occurs.
   - Select the **ON** option to automatically print the alarmed event when the alarm condition occurs.
   - Select the **OFF** option to suspend printing the alarmed event when the alarm condition occurs.

4. Select the **Save To Event** button to determine if the alarm data is saved in the Event database when the alarm condition occurs.
   - Select the **ON** option to automatically save the alarm event when the alarm condition occurs.
   - Select the **OFF** option and the alarm event will not be saved to the Events database when the alarm condition occurs.

5. Select the **Page On Alarm** button to determine if a page should be sent when the alarm condition occurs.
   - Select the **ON** option to automatically send a page when the alarm condition occurs.
   - Select the **OFF** option and a page will not be sent when the alarm condition occurs.

9.6.4.3 Sidebar Buttons

The following sidebar buttons are used to expand the functionality of this tab.

**System Alarm Options**

Select the **System Alarm Options** sidebar button to display the **System Alarm Options** tab. For additional information regarding System Alarm Options, refer to the “System Alarms Tab” on page 9-14.
System Alarm Limits
Select the System Alarm Limits sidebar button to display the System Alarm Limits tab. For additional information regarding System Alarm Limits, refer to the “System Alarms Tab” on page 9-14.

Normal Screen
Select the Normal Screen sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

9.6.4.4 Troubleshooting
This section lists some of the potential messages and issues that may occur while using the System Alarm Responses tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE*</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>The System Setup menu button is disabled.</td>
<td>The System Setup function is already open on another Panorama display and cannot be opened on multiple displays.</td>
<td>Close the System Setup function on the Panorama display where it is open.</td>
</tr>
<tr>
<td>The Page On Alarm button is not displayed.</td>
<td>The central station where the patient tile resides does not have the Paging option enabled.</td>
<td>If necessary, enable the Paging option.</td>
</tr>
<tr>
<td>The System Alarm Limits and System Alarm Responses tab is not available at the ViewStation or WorkStation.</td>
<td>System alarm limits and responses must be changed at the central station.</td>
<td>Change the alarm limits and responses from the central station.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
9.7 Passwords Tab

The system Passwords tab (shown in FIGURE 9-21 through FIGURE 9-24) establishes and modifies the passwords used to access the System Setup and Patient Alarm Setup areas of the Panorama Central Station. This tab is available at the ViewStation and the WorkStation, and it functions in the same manner as it does at the host central station but does not contain an Alarm button. Passwords tab changes made at the central station or remote workstations are independent of each other.

9.7.1 Accessing the Installation Setup Tab

1. From the menu bar, select the System Setup button. The Password tab is displayed.
2. Follow the directions in the "Entering a Password" section on page 9-7.
3. Once the correct password has been entered, the tabs associated with the System Setup button are displayed.
4. Select the Passwords tab. The Passwords tab is displayed.

**FIGURE 9-21** Passwords Tab - Paging Option Disabled (Central Station)

**FIGURE 9-22** Passwords Tab - Paging Option Enabled (Central Station)
9.7.2 Modifying a Default Password

The **Passwords** tab changes the password that provides access to the Patient Alarms and System Alarms tab.

1. Select either the **Alarm**, **System**, or **Paging** (displayed if Paging option is enabled) button. The **Old Password** text box is displayed.

   **NOTE:** The Alarm button is not available at the ViewStation.

2. Select the **Old Password** text box. The keyboard dialog box is displayed.

3. Using the keyboard dialog box, enter the alarm password that was previously established in “Recording Password Changes” on page 9-31. The Alarm and System password is case sensitive and can be a maximum of 15 characters.

4. Accept or reject the text entered in the keyboard dialog box. For additional information about using the keyboard dialog box, refer to “Keyboard Dialog Box” on page 2-18.

   **NOTE:** The factory default Alarm password is ‘alarm’. The factory default System password is ‘system’. The factory default Paging password is ‘paging’. Users are advised to change the factory default passwords once the Panorama Patient Monitoring Network is installed.

   **NOTE:** Passwords are case sensitive.
5. Select the **New Password** text box. The keyboard dialog box is displayed.

6. Using the keyboard dialog box, enter the new alarm password. The Alarm password is case sensitive and can be a maximum of 15 characters.

7. Accept the text entered in the keyboard dialog box. For additional information about using the keyboard dialog box, refer to “Keyboard Dialog Box” on page 2-18.

8. Select the **Confirm** text box. The keyboard dialog box is displayed.

9. Using the keyboard dialog box, re-enter the new alarm password. Accept or reject the text entered in the keyboard dialog box. For additional information about using the keyboard dialog box, refer to “Keyboard Dialog Box” on page 2-18.

The password is changed and the **Passwords** tab is returned to its original state.

**Recording Password Changes**

**CAUTION:** If passwords are changed from the factory defaults, users MUST ensure that they document new passwords in the space provided in this manual. For additional information refer to “Passwords Tab” on page 9-29. Failure to recall user defined passwords will render the system **UNUSABLE**.

**NOTE:** Restrict access among hospital staff to the **System password**.

Record password changes in the following table:

<table>
<thead>
<tr>
<th>PROVIDES ACCESS TO</th>
<th>DEFAULT PASSWORD</th>
<th>USER DEFINED PASSWORD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patient Alarms</td>
<td>alarm</td>
<td></td>
</tr>
<tr>
<td>System Menus</td>
<td>system</td>
<td></td>
</tr>
<tr>
<td>Paging Tab (optional)</td>
<td>paging</td>
<td></td>
</tr>
</tbody>
</table>

**9.7.3 Sidebar Button - Normal Screen**

Select the **Normal Screen** sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

**9.7.4 Troubleshooting**

This section lists some of the potential messages and issues that may occur while using the **Passwords** tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Incorrect password. Please try again.</td>
<td>The wrong password was entered in the text box.</td>
<td>Select the OK button to close the dialog box. Enter the correct password in the dialog box.</td>
</tr>
<tr>
<td></td>
<td>The password was entered in the wrong case.</td>
<td>Re-enter the password turning on/off the keyboard dialog Caps Lock key.</td>
</tr>
<tr>
<td>Illegal Password. Please choose a new password.</td>
<td>An illegal character (anything other than A-Z and/or 0-9) was entered in the keyboard dialog box.</td>
<td>Try re-entering a password that contains only the letters A-Z and/or numbers 0-9.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Illegal Password. Password should be 1 to 15 characters.</td>
<td>When entering a New Password in the keyboard dialog, the Enter key was selected in the keyboard dialog before any characters/numbers were selected.</td>
<td>Enter a password that contains at least 1 character before selecting the Enter key.</td>
</tr>
<tr>
<td>The System Setup menu button is disabled.</td>
<td>The System Setup function is already open on another central station display and cannot be opened on multiple displays.</td>
<td>Close the System Setup function on the Panorama display where it is open.</td>
</tr>
<tr>
<td>The Alarm button is not displayed in the Passwords tab.</td>
<td>System Alarm tab is being viewed from a ViewStation.</td>
<td>If password maintenance is necessary for this function, access the Alarm button from a host central station.</td>
</tr>
<tr>
<td>The Paging button is not displayed in the Passwords tab.</td>
<td>The central station where the patient tile resides does not have the Paging option enabled.</td>
<td>If necessary, enable the Paging option.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
9.8 Equipment Setup Tab

The **Equipment Setup** tab (shown in FIGURE 9-25 and FIGURE 9-26) adds, deletes, and modifies the equipment used with the Panorama Central Station. The **Equipment Setup** tab includes an equipment list that shows the types of equipment that may communicate with the central station. This tab is not available at the ViewStation or the WorkStation. Equipment maintenance can only be performed at the host central station.

9.8.1 Accessing the Equipment Setup Tab

1. From the menu bar, select the **System Setup** button. The **Password** tab is displayed.
2. Follow the directions in the "Entering a Password" section on page 9.7.
3. Once the correct password has been entered, the tabs associated with the **System Setup** button are displayed.
4. Select the **Equipment Setup** tab. The **Equipment Setup** tab is displayed.

![FIGURE 9-25 Equipment Setup Tab](image)

![FIGURE 9-26 Equipment Setup Tab (with Device Setup Area shown)](image)
The first column of the equipment list box displays the device **Label** for the equipment listed.

The second column of the equipment list box displays the **Type** of device listed.

The third column of the equipment list box displays the **Tile** number associated with the equipment.

The fourth column of the equipment list box displays the **Device ID** number for the equipment listed.

### Device ID Format

<table>
<thead>
<tr>
<th>DEVICE</th>
<th>STARTING CHARACTER</th>
<th># OF CHARACTERS</th>
<th>ENDING CHARACTER</th>
</tr>
</thead>
<tbody>
<tr>
<td>Telepack WMTS</td>
<td>M or m</td>
<td>6</td>
<td>X or x</td>
</tr>
<tr>
<td>Passport 2</td>
<td>E or e</td>
<td>6</td>
<td>X or x</td>
</tr>
<tr>
<td>Passport 2 WMTS</td>
<td>U or u</td>
<td>6</td>
<td>X or x</td>
</tr>
<tr>
<td>Passport V</td>
<td>N or n</td>
<td>6</td>
<td>X or x</td>
</tr>
<tr>
<td>Spectrum</td>
<td>S or s</td>
<td>6</td>
<td>X or x</td>
</tr>
<tr>
<td>Spectrum WMTS</td>
<td>B or b</td>
<td>6</td>
<td>X or x</td>
</tr>
<tr>
<td>Spectrum OR</td>
<td>D or d</td>
<td>6</td>
<td>X or x</td>
</tr>
<tr>
<td>DPM Series (DPM 6/DPM 7,</td>
<td>H or h</td>
<td>6</td>
<td>X or x</td>
</tr>
<tr>
<td>Passport 12M/Passport 17M,</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Passport 8/Passport 12, and T1)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>V 12/V 21</td>
<td>A or a</td>
<td>6</td>
<td>X or x</td>
</tr>
</tbody>
</table>

### 9.8.2 Entering New Equipment

This section outlines adding new equipment to the Panorama Central Station equipment list.

1. Select the **New** button below the equipment list box. The Device Setup Area (shown in FIGURE 9-26) is displayed.

2. Select the **Tile** button. In the **Tile Number** dialog box, select the tile number for the new monitoring device.

   - The tile numbers available for a single 8 tile display include **NONE** (when no tile number is assigned to the equipment) and 0 to 7.
   - The tile numbers available for a single 12 tile display include **NONE** (when no tile number is assigned to the equipment) and 0 to 11.
   - The tile numbers available for a dual 24 tile display include **NONE** (when no tile number is assigned to the equipment) and 0 to 23.

   Select the **Done** button to accept the selection.

   Select the **Cancel** button to discard the selection.

   The tile number is displayed in the **Tile** field.

**NOTE:** The **Tile** button will be disabled and display a dash (-) if the type of equipment being entered is a Panorama Server or a Panorama Central Station.
3. Select the **Type** button. In the **Device Type** dialog box, select the device type of the new monitoring device.

The equipment list supports the following types of equipment: Telepack, Telepack WMTS, Passport 2, Passport 2 WMTS, Passport V, Spectrum, Spectrum WMTS, Spectrum OR, DPM Series, and V 12/V 21.

Select the **Done** button to accept the selection or the **Cancel** button to discard the selection.

The device type is displayed in the **Type** field.

4. Select the **Label** text box to enter a label for the new equipment.

5. Use the keyboard dialog box to enter the device label for the new equipment. A device label must be entered in the equipment list and can only use alphanumeric characters and underscores (_). There is a maximum of 15 characters.

6. Accept or reject the text entered in the keyboard dialog box. For additional information about using the keyboard dialog box, refer to “Keyboard Dialog Box” on page 2-18.

NOTE: **All Device Labels in the Equipment List must be unique and cannot be duplicated.**

7. Select the **Device ID** text box.

8. Use the keyboard dialog box to enter the device identification number for the new equipment. A Device ID must be entered in the equipment list for Passport, Telepack, Spectrum, DPM Series, V 12/V 21, and Panorama Telemetry Server devices (maximum of 8 characters). For additional information about Device ID formats, refer to “Device ID Format” on page 9-34.

NOTE: **All Device IDs in the equipment list must be unique and cannot be duplicated.**

NOTE: The **Device ID** field will display a dash (-) and be disabled if the type of equipment being entered is a Panorama Central Station.

9. Accept or reject the text entered in the keyboard dialog box. For additional information about using the keyboard dialog box, refer to “Keyboard Dialog Box” on page 2-18.

10. Select the **Done** button. The equipment is added to the equipment list.

9.8.3 Modifying Equipment Included in the Equipment List

This section outlines modifying equipment added to the Panorama Central Station Equipment List.

1. From the equipment list box, select the monitoring device to be modified. A monitoring device has been selected when it changes from white text on a black background to black text on a white background.

2. Select the **Edit** button under the Equipment list box. The Device Setup Area is displayed.

3. Select the area in the Device Setup to be modified. For additional information about modifying the Device Setup area, refer to “Entering New Equipment” on page 9-34.

NOTE: **The Label text field is the only field that can be modified for a monitoring device currently monitoring a patient.**

4. Select the **Done** button. The equipment is added to the Equipment List.
9.8.4 Deleting Equipment in the Equipment List
This section outlines deleting equipment from the Panorama Central Station Equipment List.

1. From the equipment list box, select the monitoring device to be modified. A monitoring device has been selected when it changes from white text on a black background to black text on a white background.

2. Select the Delete button below the equipment list box.

NOTE: Only equipment not actively monitoring a patient can be deleted.

3. The message Are you sure you want to delete this device? message box is displayed.
   • Select the Yes button to delete the selected device and close the dialog box.
   • Select the No button to keep the selected device and close the dialog box.

NOTE: If an associated patient tile has been assigned at the ViewStation or the WorkStation, or both, the connection between the patient tiles will continue to exist, even if the equipment is deleted at the host central station.

9.8.5 Sidebar Buttons
The following sidebar buttons are used to expand the functionality of this tab.

WMTS Device
This sidebar button should not be used without assistance from a Service Representative.

WMTS All
This sidebar button should not be used without assistance from a Service Representative.

Normal Screen
Select the Normal Screen sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

9.8.6 Troubleshooting
This section lists some of the potential messages and issues that may occur while using the Equipment tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Equipment list is full</td>
<td>This message appears when the New button is pressed and the equipment list contains the maximum number of equipment list entries.</td>
<td>Select the OK button to close the dialog box. Delete a device from the equipment list.</td>
</tr>
<tr>
<td>Illegal Device Tile. Device Tile must be defined. Please try again.</td>
<td>This message appears when the Done button is pressed in the Equipment Setup tab before a tile number is selected.</td>
<td>Select the OK button to close the dialog box. Select a tile number in the Tile Number dialog box.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>The Tile button is disabled and a dash (-) is shown in the Tile field.</td>
<td>The type of equipment being entered in the equipment list is either a Panorama Server or a Panorama</td>
<td>A tile number cannot be entered for the Panorama Server or the Panorama equipment types.</td>
</tr>
<tr>
<td><strong>Invalid Device Type</strong></td>
<td>This message appears when the Done button is pressed in the Equipment Setup tab before a device type is selected.</td>
<td>Select the OK button to close the dialog box. Select a device type in the Device Type dialog box.</td>
</tr>
<tr>
<td><strong>Illegal Device Label. Device Label must be defined. Please try again.</strong></td>
<td>This message appears when the Done button is pressed in the Equipment Setup tab before a device label is selected.</td>
<td>Select the OK button to close the dialog box. Select a device label in the Device Label dialog box.</td>
</tr>
<tr>
<td><strong>Device Label already exists! Please enter a new one</strong></td>
<td>This message appears when a duplicate device label is entered in the Device Label field.</td>
<td>Select the OK button to close the dialog box. Enter a new device label in the Device Label field.</td>
</tr>
<tr>
<td><strong>Invalid characters entered. Please enter a new label.</strong></td>
<td>This message appears when invalid characters have been entered in the Device Label. The device label can only contain alphanumeric characters and underscores (_).</td>
<td>Select the OK button to close the dialog box. Enter a valid Device label.</td>
</tr>
<tr>
<td><strong>Illegal Device ID. Please enter a new one.</strong></td>
<td>This message appears when the Done button is pressed in the Equipment Setup tab before a device ID is selected.</td>
<td>Select the OK button to close the dialog box. Enter a Device ID in the Device ID dialog box.</td>
</tr>
<tr>
<td><strong>Device ID must be 8 characters in length.</strong></td>
<td>This message appears when the Device ID that was entered is less than 8 characters in length. Please refer to the Device ID setup rules in the Setting up New Equipment section.</td>
<td>Select the OK button to close the dialog box. Enter a Device ID in the Device ID dialog box.</td>
</tr>
<tr>
<td><strong>Device ID must start with the proper device prefix</strong></td>
<td>This message appears when the incorrect device prefix is entered in the Device ID box.</td>
<td>Select the OK button to close the dialog box. Enter the correct Device ID prefix.</td>
</tr>
<tr>
<td><strong>Suffix of Device ID last character must be 'x' or 'X'</strong>.</td>
<td>This message appears when the incorrect device suffix is entered in the Device ID box.</td>
<td>Select the OK button to close the dialog box. Enter the correct Device ID suffix.</td>
</tr>
<tr>
<td><strong>This Device ID already exists!</strong></td>
<td>This message appears when a duplicate Device ID is entered in the Device ID field.</td>
<td>Select the OK button to close the dialog box. Enter a new Device ID in the Device ID field.</td>
</tr>
<tr>
<td><strong>You have reached the maximum number of channel licenses available.</strong></td>
<td>This message appears when all available channels on the Panorama Central Station are occupied.</td>
<td>Select the OK button to close the dialog box. Delete a piece of equipment from an existing channel if it is no longer in use. The system administrator can also purchase additional channel licenses.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>This version of software does not support more than 16 monitored patients.</strong></td>
<td>This message appears when the maximum number of patients is being monitored on the Panorama Central Station.</td>
<td>Select the OK button to close the dialog box. Discharge a patient if the patient is no longer being monitored.</td>
</tr>
<tr>
<td><strong>System Setup menu button is disabled.</strong></td>
<td>The System Setup function is already open on another Panorama display and cannot be opened on multiple displays.</td>
<td>Close the System Setup function on the Panorama display where it is open.</td>
</tr>
<tr>
<td><strong>Cannot delete a device connected to a patient</strong></td>
<td>The device cannot be deleted because it is currently connected to a patient.</td>
<td>Select the OK button to close the dialog box. Disconnect the patient from the selected device before attempting to delete it.</td>
</tr>
<tr>
<td><strong>The Tile button, Type button and Device ID field are disabled.</strong></td>
<td>The selected equipment cannot be edited because it is currently monitoring a patient.</td>
<td>This equipment cannot be edited until a patient discharge is performed.</td>
</tr>
<tr>
<td><strong>Unable to perform operation on tile: Central to Central Transfer In in progress.</strong></td>
<td>Equipment Setup tab Delete button was selected for equipment that was not actively monitoring a patient, while a Central to Central Transfer In is in progress to that same device.</td>
<td>Wait for the Central to Central Transfer to complete, then try again.</td>
</tr>
</tbody>
</table>

*Messages are shown in all bold text.*
9.9 More Tab

The More tab (shown in FIGURE 9-27) is used for navigational purposes and when selected, will display another set of System Setup tabs at the Panorama Central Station. This tab is not available at the ViewStation or the WorkStation.

9.9.1 Accessing Additional (More) Tabs

1. From the menu bar, select the System Setup button. The Password tab is displayed.
2. Follow the directions in the "Entering a Password" section on page 9-7.
3. Once the correct password has been entered, the tabs associated with the System Setup button are displayed.
4. Select the More tab. The next set of System Setup tabs is displayed.

FIGURE 9-27 More Tab

FIGURE 9-28 Tabs available when More tab is selected
9.10 Previous Tab

The Previous tab (shown in FIGURE 9-29) is used for navigational purposes and when selected, will display another set of system setup tabs at the Panorama Central Station. This tab is not available at the ViewStation or the WorkStation.

9.10.1 Accessing Previous Tabs

1. From the menu bar, select the System Setup button. The first set of System Setup tabs is displayed.
2. Select the Installation Setup tab. The Installation Setup tab is displayed.
3. Enter the correct system password.
4. Select the More tab. The next set of System Setup tabs is displayed.
5. Select the Previous tab to display the first set of tabs (shown in FIGURE 9-30) associated with the System Setup tab.

![Previous Tab](FIGURE 9-29)

FIGURE 9-29 Previous Tab

![Previous Tabs](FIGURE 9-30)

FIGURE 9-30 Tabs available when Previous tab is selected
9.11 Date/Time Tab

The Date/Time tab (shown in FIGURE 9-31) sets the system date and time and configures the default date and time formats at the Panorama Central Station. This tab is available at the ViewStation and the WorkStation, and it functions in the same manner as it does at the host central station. Date/Time tab changes made at the central station or remote workstations are independent of each other.

NOTE: The host central station, ViewStation, and WorkStation system times are independent of each other.

9.11.1 Accessing the Date/Time Tab

1. From the menu bar, select the System Setup button. The Password tab is displayed.
2. Follow the directions in the "Entering a Password" section on page 9-7.
3. Once the correct password has been entered, the tabs associated with the System Setup button are displayed.
4. Select the More tab. The next set of System Setup tabs is displayed.
5. Select the Date/Time tab. The Date/Time tab is displayed.

FIGURE 9-31 Date/Time Tab (Central Station, not NTP configured)

FIGURE 9-32 Date/Time Tab (WorkStation, not NTP configured)
9.11.2 Set Date/Time

The settings configured in the Date/Time tab are the settings that will be used for all date/time related functions in the Panorama Central Station.

**NOTE:** When generating the Full Disclosure Report, Event List Report, etc., the Data Time indicated below the report title is based on the system time the data was collected at the host central station, not the system time displayed at the ViewStation or the WorkStation.

Setting a Date in the Date/Time Tab

**NOTE:** The Day, Month and Year fields will not be available when the central station is configured as an NTP client.

1. From the Date/Time tab, select the Day field. The keypad dialog box is displayed. The range available is shown on the bottom of the dialog box.

2. Using the keypad dialog box, enter the number that represents the current day of the month.
   - Select the Clear button to delete all of the numbers in the text box
   - Select CE button to delete a single number from the text box
   - Select Cancel button to discard changes and close the dialog box
   - Select OK button to accept changes. The Day field will remain unchanged until the OK button is pressed.

3. Select the Month field. The keypad dialog box is displayed. The range available is 1 to 12.

4. Using the keypad dialog box, enter the number that represents the current month.
   - Select the Clear button to delete all of the numbers in the text box
   - Select CE button to delete a single number from the text box
   - Select Cancel button to discard changes and close the dialog box
   - Select OK button to accept changes. The Month field will remain unchanged until the OK button is pressed.
5. Select the **Year** field. The keypad dialog box is displayed. The range available is 1970 to 2037.

6. Using the keyboard dialog box, enter the number that represents the current year.
   - Select the **Clear** button to delete all of the numbers in the text box
   - Select **CE** button to delete a single number from the text box
   - Select **Cancel** button to discard changes and close the dialog box
   - Select **OK** button to accept changes. The Month field will remain unchanged until the **OK** button is pressed.

7. Select the **Date** button to select the desired date format.
   The date formats available include DD-MM-YYYY, MM-DD-YYYY and YYYY-MM-DD (default).

8. Select the **Done** button to change the system date format.
   The Are you sure you want to change the date/time settings? confirmation box is displayed.

9. Select the **Yes** or **No** button in the dialog box.
   - Select the **Yes** button to change the system date format
   - Select the **No** button to discard the changes that were made to the system date format

**Setting a Time in the Date/Time Tab**

**NOTE:** The Hour, Min., and AM/PM fields will not be available when the central station is configured as an NTP client.

1. From the **Date/Time** tab, select the **Hour** field. The keypad dialog box is displayed. The range available is 1 to 12.

2. Using the keypad dialog box, enter the number for the current hour.
   - Select the **Clear** button to delete all of the numbers in the text box
   - Select **CE** button to delete a single number from the text box
   - Select **Cancel** button to discard changes and close the dialog box
   - Select **OK** button to accept changes. The Hour field will remain unchanged until the **OK** button is pressed.

3. Select the **Min.** field. The keypad dialog box is displayed. The range available is 0 to 59.

4. Using the keypad dialog box, enter the number that represents the current minute.
   - Select the **Clear** button to delete all of the numbers in the text box
   - Select **CE** button to delete a single number from the text box
   - Select **Cancel** button to discard changes and close the dialog box
   - Select **OK** button to accept changes. The Min. field will remain unchanged until the **OK** button is pressed.

5. Select the AM/PM toggle button to select either AM or PM.

**NOTE:** The AM/PM button is only available if the current time format is in 12 hour mode.
6. Select the **Time** button to select the desired time format.
   
   The time formats available include HH:MM (12H), HH:MM:SS (12H) (default), HH:MM (24H) and HH:MM:SS (24H).
   
   **NOTE:** Whenever the central station is set to a language other than English, it will be in the 24-hour time format.

7. Select the **Done** button to change the system time format.

   The **Are you sure you want to change the date/time settings?** confirmation box is displayed.

8. Select the **Yes** or **No** button in the dialog box.
   - Select the **Yes** button to change the system time format
   - Select the **No** button to discard the changes that were made to the system time format

9.11.3 **Sidebar Button - Normal Screen**
   
   Select the **Normal Screen** sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

9.11.4 **Troubleshooting**

   This section lists some of the potential messages and issues that may occur while using the **Date/Time** tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>The System Setup</strong> menu button is disabled.</td>
<td>The <strong>System Setup</strong> function is already open on another Panorama display and cannot be opened on multiple displays.</td>
<td>Close the <strong>System Setup</strong> function on the Panorama display where it is opened.</td>
</tr>
<tr>
<td><strong>Automatic Daylight Savings Time Disabled</strong></td>
<td>The central station is configured to not automatically update the time for daylight savings time.</td>
<td>No solution required.</td>
</tr>
<tr>
<td><strong>Automatic Daylight Savings Time Enabled!</strong></td>
<td>The central station is configured to automatically update the time for daylight savings time.</td>
<td>No solution required.</td>
</tr>
<tr>
<td><strong>Date and Time is being set automatically!</strong></td>
<td>The central station is configured to be an NTP client.</td>
<td>No solution required.</td>
</tr>
<tr>
<td><strong>Time Sync</strong> button enabled</td>
<td>The central station is configured to be an NTP client.</td>
<td>Button is used to synchronize the time at the Panorama and the NTP server.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
9.12 Print Setup Tab (System)

The **Print Setup** tab (shown in FIGURE 9-34 and FIGURE 9-35) defines the printer(s) that will be used to print standard (defined as reports that do not contain waveforms) and strip reports (such as the All Strips, Event Zoom-In, and Full Disclosure Zoom-In Reports). This tab is available at the ViewStation and WorkStation, and it functions in the same manner as the host central station. The central station and each remote workstation is capable of communicating with two network printers simultaneously.

**Print Setup** tab changes made at the central station or remote workstations are independent of each other.

**NOTE:** The network printer(s) must be properly configured to communicate with the central station.

For information regarding individual patient print setup configuration, refer to “Print Setup Tab (Patient)” on page 5-57.

For information regarding printer configuration, refer to the Panorama Printer Configuration Guide (P/N 0070-00-0561).

### 9.12.1 Accessing the Print Setup Tab (Central Station)

1. From the menu bar, select the **System Setup** button.
2. Select the **Print Setup** tab. The **Print Setup** tab is displayed.

![FIGURE 9-34 System Print Setup Tab (Central Station)](image-url)

**FIGURE 9-34** System Print Setup Tab (Central Station)
9.12.2 Accessing the Print Setup Tab (ViewStation and WorkStation)

- Follow steps from the "Accessing the Print Setup Tab (Central Station)" section on page 9-45.

![System Print Setup Tab (ViewStation and WorkStation)](image)

**FIGURE 9-35** System Print Setup Tab (ViewStation and WorkStation)

9.12.3 Setting the System Default Print Setup

This section outlines using the **Print Setup** tab to establish to which default printer print requests will be directed. To set system default print requests:

1. Select a printer’s check box button to enable/disable the printing of reports to the selected printer. The options available include Laser jet 1 and Laser jet 2. Print setup settings are changed instantly.
   - A check mark in a printer’s check box indicates that if the printer was properly configured with the Panorama Central Station, it will be capable of printing Panorama reports.
   - An empty check box button indicates the printer is not selected and will not print Panorama reports.

2. Select the printer’s check box button to enable/disable the printing of strips. The print setup settings are changed instantly.
   - A check mark in a printer’s check box button indicates that if the printer was properly configured with the Panorama Central Station, it will be capable of printing waveform strips.
   - An empty check box indicates the printer is not selected and will not print waveform strips.

9.12.4 Troubleshooting

This section lists some of the potential messages and issues that may occur while using the System **Print Setup** tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>The <strong>System Setup</strong> menu button is disabled.</td>
<td>The <strong>System Setup</strong> function is already open on another Panorama display and cannot be opened on multiple displays.</td>
<td>Close the <strong>System Setup</strong> function on the Panorama display where it is open.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
9.13  Unit Priorities Tab

The **Unit Priorities** tab (shown in FIGURE 9-36) determines the initial order in which parameters in the Main Screen, **Bedside** tab, **Trends** tab, and **Events** (List) are displayed. This tab is not available at the ViewStation or the WorkStation. Unit priority settings are transferred from the host central station to the ViewStation and the WorkStation when the patient tile is assigned.

9.13.1  Accessing the Unit Priorities Tab

1. From the menu bar, select the **System Setup** button. The first set of System Setup tabs is displayed.
2. Follow the directions in the "Entering a Password" section on page 9-7.
3. Once the correct password has been entered, the tabs associated with the **System Setup** button are displayed.
4. Select the **More** tab. The next set of System Setup tabs is displayed.
5. Select the **Unit Priorities** tab. The **Unit Priorities** tab is displayed.

![FIGURE 9-36 Unit Priorities Tab](image)

9.13.2  Setting a Default Order in the Unit Priorities Tab

This section outlines how to select the parameter order in the **Unit Priorities** tab.

**Selecting the Unit Priority Order**

1. Select a **Unit Priority** number button. The parameter with the number 1 priority will always be Heart Rate.
2. Select a parameter tile from the Parameter list. A parameter has been selected when it changes from white text with a black background to black text with a white background.

**NOTE:** Parameters show their current unit priority position in parentheses next to their name in the Parameter list.
3. Choose the **Select** button. The selected parameter is assigned its new unit priority position and will be shown in the box on the right of the selected unit priority.

4. Repeat steps 2-3 until the parameters are in the desired order.

5. Select the **Done** button. All priority changes in the **Unit Priorities** tab have now been confirmed. The changes that were made will only be applied to newly admitted patients and will not affect patients currently being monitored.

9.13.2.1 Setting the Waveform System Defaults

Select the Wave Defaults sidebar button to select the system default patient tile waveforms.

1. Select the **Wave Defaults** sidebar button in the **Unit Priorities** tab. The **Wave Defaults** dialog box (shown in FIGURE 9-37) is displayed.

![Wave Defaults Dialog Box](image)

FIGURE 9-37 Wave Defaults Dialog Box

1. Select the **Waveform 1** button until the desired ECG waveform is displayed. The settings available for the ECG waveform include ECG I through V6. The factory default setting is ECG II.

2. Select the **Waveform 2** button until the desired ECG waveform is displayed. The settings available for the ECG waveform include ECG I through V6, IBP1-IBP4, Resp, Pleth, CO2, O2, Agent, and OFF. The factory default setting is OFF.

3. Accept or reject the selections made in the dialog box.
   - Select the **Done** button to accept the selections.
   - Select the **Cancel** button to discard the selections, close the dialog box and return to the **Unit Priorities** tab.

9.13.3 Sidebar Buttons

The following sidebar buttons are used to expand the functionality of this tab.

**Wave Defaults**

This sidebar button was previously described on page 9-48.

**Normal Screen**

Select the **Normal Screen** sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.
9.13.4 Troubleshooting

This section lists some of the potential messages and issues that may occur on the Unit Priorities tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Please select a parameter button before selecting from the list</td>
<td>This message appears when the Select button is pressed before a Unit Priority button is selected.</td>
<td>Select the OK button to close the dialog box. Select a Unit Priority button.</td>
</tr>
<tr>
<td>The System Setup menu button is disabled.</td>
<td>The System Setup function is already open on another Panorama display and cannot be opened on multiple displays.</td>
<td>Close the System Setup function on the Panorama display where it is open.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
9.14 Unit Choices Tab

The **Unit Choices** tab (shown in FIGURE 9-38, FIGURE 9-39, and FIGURE 9-40) determines the initial units of measure, configures the default pacemaker settings, and selects which patient demographic information is displayed. This tab is available at the ViewStation and WorkStation, and it functions in the same manner as it does at the host central station but contains different buttons. Unit Choices tab changes made at the central station or remote workstations are independent of each other.

9.14.1 Accessing the Unit Choices Tab (Central Station)

1. From the menu bar, select the **System Setup** button. The first set of System Setup tabs is displayed.
2. Follow the directions in the "Entering a Password" section on page 9-7.
3. Once the correct password has been entered, the tabs associated with the **System Setup** button are displayed.
4. Select the **More** tab. The next set of System Setup tabs is displayed.
5. Select the **Unit Choices** tab. The **Unit Choices** tab is displayed.

![FIGURE 9-38 Unit Choices Tab - Paging Option Disabled (Central Station)](image)

![FIGURE 9-39 Unit Choices Tab - Paging Option Enabled (Central Station)](image)

9.14.2 Accessing the Unit Choices Tab (ViewStation and WorkStation)

1. Follow steps 1-4 from the "Accessing the Unit Priorities Tab" section.
2. Select the **Unit Choices** tab. The **Unit Choices** tab is displayed.
9.14.3 Setting the Unit Choices (Central Station)
This section outlines selecting system defaults for pacemaker settings, order of demographic data and units of measurement.

1. Select the **Discharge at Bedside** button to set the system default for allowing patient discharge from the central station.

   Discharge at Bedside options include **ON** and **OFF**. The factory default setting is **OFF**.
   - The **ON** option indicates that Discharge at Bedside is enabled.
   - The **OFF** option indicates that Discharge at Bedside is disabled.

**NOTE:** The Discharge at Bedside button only affects bedside monitors. This button setting has no affect when a Telepack device is in use.

**NOTE:** Changing the Discharge at Bedside option will take affect the next time the Discharge tab is entered.
2. Select the **Signal Lost Delay** button to delay the display, audible enunciation, and storage of the Signal Lost event with or without the Arrhythmia enabled by the number of configured seconds.

3. Select the **Trend Access Order** button to define which Trend view will display first. The choices are List or Graphic. The default is List view.

4. Select the **Paging Demographics** button to define which demographic information is sent to the paging system when a page is requested. Paging Demographics options include **Bed**, **Bed + Last Name**, and **Bed + ID**. The factory default setting is **Bed**. Refer to the following tables for formatting examples:

<table>
<thead>
<tr>
<th>OPTION</th>
<th>ALARM TYPE</th>
<th>FORMAT EXAMPLES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bed</td>
<td>Heart Rate</td>
<td>central name:XXXBed:ICU10 Low Heart Rate Alarm</td>
</tr>
<tr>
<td>Bed</td>
<td>Arrhythmia</td>
<td>central name:XXXBed:ICU10 Asystole</td>
</tr>
<tr>
<td>Bed</td>
<td>Not HR or arrhythmia alarm</td>
<td>central name:XXXBed:ICU10 High Temperature 1</td>
</tr>
<tr>
<td>Bed + Last Name</td>
<td>Heart Rate</td>
<td>central name:XXXBed:ICU10 Name:Patient Last Name Low Heart Rate Alarm</td>
</tr>
<tr>
<td>Bed + Last Name</td>
<td>Arrhythmia</td>
<td>central name:XXXBed:ICU10 Name:Patient Last Name Asystole</td>
</tr>
<tr>
<td>Bed + Last Name</td>
<td>Not HR or arrhythmia alarm</td>
<td>central name:XXXBed:ICU10 Name:Patient Last Name High Temperature 1</td>
</tr>
<tr>
<td>Bed + Last Name</td>
<td>ST alarm</td>
<td>central name:XXXBed:ICU10 Name:Patient Last Name ST Single Alarm</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>OPTION</th>
<th>ALARM TYPE</th>
<th>FORMAT EXAMPLES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bed + ID</td>
<td>Heart Rate</td>
<td>central name:XXXBed:ICU10 ID:Patient ID Low Heart Rate Alarm</td>
</tr>
<tr>
<td>Bed + ID</td>
<td>Arrhythmia</td>
<td>central name:XXXBed:ICU10 ID:Patient ID Asystole</td>
</tr>
<tr>
<td>Bed + ID</td>
<td>Not HR or arrhythmia alarm</td>
<td>central name:XXXBed:ICU10 ID:Patient ID High Temperature 1</td>
</tr>
<tr>
<td>Bed + ID</td>
<td>ST alarm</td>
<td>central name:XXXBed:ICU10 ID:Patient ID ST Single Alarm</td>
</tr>
</tbody>
</table>

**NOTE:** The amount of data that is displayed in the pager display window may differ slightly due to the type of pager being used.

**NOTE:** Changing the Paging Demographics option will take effect immediately.

**NOTE:** This option will only be displayed if the Paging option is enabled at the central station.

5. Select the **Paging Password Protection** button to enable and/or disable password protection for the Paging tab. Paging Protection options include **ON** and **OFF**. The factory default setting is **ON**.
   - The **ON** option indicates that Paging Password Protection is enabled.
   - The **OFF** option indicates that Paging Password Protection is disabled.
NOTE: This option will only be displayed if the Paging option is enabled at the central station.

6. Select the **Demographic Line Position 1** button to select the system default demographic data to be shown in the first demographic line position of the Patient Status line in the Main Screen of the Panorama Central Station.

   **Demographic Line Position 1** options include Last Name, First Name, Bed, Patient ID, and Demo Off. The factory default setting is Last Name.

   NOTE: Changing the Demographic Line Position 1 option will affect all patients with the next update of the Patient Status line in the patient tile.

7. Select the **Demographic Line Position 2** button to select the system default demographic data to be shown in the second demographic line position of the Patient Status line in the Main Screen of the Panorama Central Station.

   **Demographic Line Position 2** options include First Name, Bed, Patient ID, Demo Off, and Last Name. The factory default setting is First Name.

   NOTE: Changing the Demographic Line Position 2 option will affect all patients with the next update of the Patient Status line in the patient tile.

8. Select the **Demographic Line Position 3** button to select the system default demographic data to be shown in the third demographic line position of the Patient Status line in the Main Screen of the Panorama Central Station.

   **Demographic Line Position 3** options include Bed, Patient ID, Demo Off, Last Name, and First Name. The factory default setting is Bed.

   NOTE: Changing the Demographic Line Position 3 option will affect all patients with the next update of the Patient Status line in the patient tile.

9. Select the **Demographic Line Position 4** button to select the system default demographic data to be shown in the fourth demographic line position of the Patient Status line in the Main Screen of the Panorama Central Station.

   **Demographic Line Position 4** options include Patient ID, Demo Off, Device ID, Last Name, First Name, and Bed. The factory default setting is Patient ID.

   NOTE: Changing the Demographic Line Position 4 option will affect all patients with the next update of the Patient Status line in the patient tile.

10. Select the **Disable ECG** button (available on version 8.7.1 and above) to select the system default for displaying the **Disable ECG** sidebar button on the unit Display Tab. If **ON** is selected, the **Disable ECG** sidebar button is displayed on the unit Display tab if a Telepack with SpO₂ module is connected. If **OFF** is selected, the **Disable ECG** sidebar button is not displayed, and the ECG waveforms will be re-enabled. The factory default setting is **OFF**.

11. Select the **Height Units** button to select the system default unit of measurement to be used for the height variable in the Panorama Central Station. **Height Unit** options include cm (centimeters) and in (inches). The factory default setting is cm.

   NOTE: Changing the **Height Units** option will affect new Telepack patients upon entering the Demographics tab.
NOTE: The Height Units option is not available at the ViewStation or the WorkStation.

12. Select the **Weight Units** button to select the system default unit of measurement to be used for measuring weight at the Panorama Central Station. Weight Unit options include lbs (pounds) and kg (kilograms). The factory default setting is lbs.

NOTE: Changing the Weight Units option will affect new Telepack patients upon entering the Demographics tab.

NOTE: The Weight Units option is not available at the ViewStation or the WorkStation.

13. Select the **Temperature Units** button to select the system default temperature unit to be used for measuring temperature at the Panorama Central Station. Temperature unit options include F (Fahrenheit) and C (Celsius). The factory default setting is F.

NOTE: Changing the Temperature Units option will affect all new patients upon entering the System Alarm Limits tab.

NOTE: The Temperature Units option is not available at the ViewStation or the WorkStation.

14. Select the **CO2** button to select the system default CO2 unit to be used for measuring CO2 at the Panorama Central Station. CO2 unit options include %, kPa, and mmHg. The factory default setting is %.

NOTE: Changing the CO2 Units option will affect all new patients upon entering either the System Alarm Limits or System Wave Gain tabs.

NOTE: The CO2 Units option is not available at the ViewStation or the WorkStation.

15. Select the **Trend Display** button to configure the interval at which a patient's trend data will be displayed in the Trends tab.

Interval periods include 1 minute, 5 minutes, 10 minutes, 15 minutes, 30 minutes, 1 hour, and 2 hours. The factory default setting is 1 minute.

NOTE: Trend data is automatically stored in one minute intervals. The Trend Display interval selected does not affect how trend information is stored in the database, just how it is displayed.

NOTE: The Trend Display option is not available at the ViewStation or the WorkStation.

9.14.4 Setting the Unit Choices (ViewStation and WorkStation)

- Follow steps 6 - 9 from the "Setting the Unit Choices (Central Station)" section on page 9-51.

9.14.5 Arrhythmia Setup (Telepack)

Select the **Arrhythmia** sidebar button to:

- Enable or disable arrhythmia analysis for a Telepack device
- Enable or disable QT monitoring for a Telepack device
NOTE: If using a Passport 2, Passport V, Spectrum, Spectrum OR, DPM 6/DPM 7, Passport 12M/Passport 17M, Passport 8/Passport 12, T1 or V 12/V 21, arrhythmia analysis must be enabled or disabled at the bedside monitor.

NOTE: If using a V 12/V 21, QT monitoring must be enabled or disabled at the bedside monitor.

- Adjust the default V-Tach Rate, Asystole Delay, V-Tach Thresholds, ECG Noise Delay, Pause Threshold, and QTc Formula for a Telepack device

NOTE: The Arrhythmia sidebar button is not available at the ViewStation or the WorkStation.

NOTE: When using a bedside device, the arrhythmia settings are taken from the bedside monitor upon admission.

1. Select the All Arrhythmia On/Off check box button in the Arrhythmia Menu dialog box (shown in FIGURE 9-41) to enable or disable arrhythmia analysis for a Telepack device.

   FIGURE 9-41 Arrhythmia Menu Dialog Box

   - A checkmark in the All Arrhythmia On/Off box indicates that arrhythmia analysis is enabled.
   - An empty box indicates that arrhythmia analysis is disabled.

NOTE: Arrhythmia analysis can be enabled for Telepack devices if an Arrhythmia license is available. When arrhythmia analysis is enabled, a license is automatically assigned to a patient.

2. Select the QT Monitoring On/Off check box button to enable or disable QT monitoring for a Telepack device.

   - A checkmark in the QT Monitoring On/Off box indicates that QT monitoring is enabled.
   - An empty box indicates that QT monitoring is disabled.
3. Select the **QTc Formula** button to change the formula used to calculate the QTc for the Telepack device. The available options are Hodges, Bazett, Fridericia and Framingham.

4. Use the dial control to change the system default V-Tach Rate. The range available is 100-180 bpm, in 5 bpm increments. The factory default setting is 120 bpm.

5. Select the **Asystole Delay** button to change the system default asystole delay time for Arrhythmia processing. The range available is 3-10 seconds. The factory default setting is 4 seconds.

6. Select the **V-Tach Threshold** button to change the system default V-Tach threshold settings. The range available is 3-15 beats. The factory default setting is 3 beats.

7. Select the **Pause Threshold** button to change the Pause threshold settings. The range available is 1.2-2.0 RR%.

8. Select the **ECG Noise Delay** button to change the system default noise delay setting. The range available is 3-30 seconds. The factory default setting is 6 seconds.

**NOTE:** The ECG Noise Delay and Pause Threshold settings only affect Telepack monitoring devices.

**NOTE:** When using a bedside device, these settings are taken from the bedside upon admission.

9. Accept or reject the selections made in the dialog box.
   - Select the **Done** button to accept the selections.
   - Select the **Cancel** button to discard the selections, close the dialog box and return to the **Unit Choices** tab.

### 9.14.6 ST Settings (Telepack)

Select the **ST Settings** sidebar button to establish the default ECG monitoring filter that will be used by the Telepack, and to select the ECG leads for ST monitoring.

**NOTE:** The ST Settings sidebar button is not available at the ViewStation or the WorkStation.

- The Monitor mode for ECG filtering establishes the frequency response range for non-ST ECG monitoring.
- The ST mode for ECG filtering establishes the frequency response range for ST diagnostic monitoring.

**NOTE:** The Telepack can only have one type of ECG filtering enabled at any given time (Monitor mode or ST mode).

To configure the default ECG monitoring filter:

1. Select the **ST Settings** sidebar button in the **Unit Choices** tab. The **ST Settings** dialog box (shown in FIGURE 9-42) is displayed.

**FIGURE 9-42** ST Settings Dialog Box

2. Select the **ST** check box to enable or disable ST analysis for the selected patient.
   - A checkmark in the ST box indicates that ST analysis is enabled.
   - An empty box indicates that ST analysis is disabled.
   - When the ST check box is empty, the Monitor filter is automatically enabled.
NOTE: If ST licenses are available, an ST license is automatically assigned to the patient when ST is enabled.

NOTE: When using a 5-wire lead set or a View 12™ card, ST is performed on the leads available. When using a 3-wire lead set, ST is performed on the lead being viewed.

3. Select the Monitor or ST filter mode.

NOTE: One filter box must always be populated. When a filter mode is selected, the other mode is automatically disabled.

- A checkmark in the Monitor filter box indicates that the Monitor filter is enabled and ST analysis is disabled.
- A checkmark in the ST filter box indicates that the ST analysis is enabled and the Monitor filter is disabled.

4. Accept or reject the selections made in the dialog box.

- Select the Done button to accept the selections.
- Select the Cancel button to discard the selections, close the dialog box and return to the Unit Choices tab.

9.14.7 Pacer

Select the Pacer sidebar button to adjust default Pacer Enhancement, and Pacer Filter settings.

- **Pacer Enhancement** visually marks pacemaker-induced ECG complexes with a colored vertical spike. It can be enabled or disabled in the Pacer Setting dialog box (shown in FIGURE ).

NOTE: A yellow vertical spike will be displayed when a paced beat has been detected and the ECG waveform color is not set to yellow.

A green vertical spike will be displayed when a paced beat has been detected and the ECG waveform color is set to yellow.
- **Pacer Indicator** controls whether the pacer indicator displays as a long vertical line crossing through the waveform or a short vertical line above the waveform (depending on the selection in the Pacer Settings dialog) to indicate firing of the pacemaker impulse.

- **Pacer Filtering** adjusts the time period during which pacemaker-induced ECG complexes are filtered for ECG sampling. The preset options for the Pacer Filter are 0 to 80 ms (in increments of 20). The Pacer Filter is selected in the **Pacer Settings** dialog box (shown in FIGURE ).

![Pacer Setting Dialog Box](image)

**FIGURE 9-43** Pacer Setting Dialog Box

To enter Pacer Settings:

1. Select the **Pacer Enhancement** button to enable or disable this setting for the patient.
   - A checkmark indicates that Pacer Enhancement is enabled.
   - An empty box indicates that Pacer Enhancement is disabled.

2. Select the **Pacer Filter** button until the desired setting is displayed. The selections for **Pacer Filter** range from 0 to 80 ms (in increments of 20).

3. Accept or reject the selections made in the **Pacer Setting** dialog box.
   - The **Done** button accepts the selections.
   - The **Cancel** button discards the selections, closes the dialog box and returns to the **Unit Choices** tab.

### 9.14.8 Configuring Default Report Waveform Parameters

Select the **Strip Setup** sidebar button (shown in FIGURE 9-38) to select the default waveform parameters that will print in the All Strips, Event Zoom-In, and Full Disclosure Zoom-In reports.

**NOTE:** The Strip Setup sidebar button is not available at the ViewStation or the WorkStation.

To select the default waveform parameters:

1. From the **Unit Choices** tab, select the **Strip Setup** sidebar button.
2. Select the waveform check box button to enable/disable the printing of waveforms in strip reports. The print setup settings are changed instantaneously.
   - A check mark in a waveform’s check box button indicates the waveform is selected to print in reports when that parameter is in use.
   - An empty check box button indicates the waveform is not selected and will not print in reports.

3. To return to the **Unit Choices** tab, select the **Previous** sidebar button.

For additional information about customizing the waveform strips printed for a particular patient, refer to the “Print Setup Tab (Patient)” on page 5-57.

9.14.9 Sidebar Buttons

The following sidebar buttons are used to expand the functionality of this tab.

**Strip Setup**
This sidebar button was previously described on page 9-58.

**Previous**
This sidebar button was previously described on page 9-58.

**Arrhythmia**
This sidebar button was previously described on page 9-54.

**Settings (Telepack)**
This sidebar button was previously described on page 9-56.

**Normal Screen**
Select the **Normal Screen** sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.
9.14.10 Troubleshooting

This section lists some of the potential messages and issues that may occur while using the **Unit Choices** tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>The <strong>System Setup</strong> menu button is disabled.</td>
<td>The <strong>System Setup</strong> function is already open on a Panorama display and cannot be opened on multiple displays.</td>
<td>Close the <strong>System Setup</strong> function on the Panorama display on which it is opened.</td>
</tr>
<tr>
<td>The <strong>Strip Setup</strong> sidebar button is not displayed.</td>
<td><strong>Unit Choices</strong> tab is being viewed from a ViewStation.</td>
<td>If strip setup is necessary, access the <strong>Strip Setup</strong> button from a central station.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
9.15 Wave Gain Tab (System)

The Wave Gain tab (shown in FIGURE 9-45) is used to change the initial wave gain settings. This tab is not available at the ViewStation or the WorkStation. Wave gain settings are transferred from the host central station to the ViewStation and the WorkStation when the patient tile is assigned.

For information regarding individual patient tile wave gain settings, refer to the “Wave Gain Tab (Patient)” on page 5-54.

NOTE: The changes made in the Wave Gain tab (System) will apply to newly admitted patients.

9.15.1 Accessing the Wave Gain Tab

1. From the menu bar, select the System Setup button. The first set of System Setup tabs is displayed.
2. Follow the directions in the "Entering a Password" section on page 9-7.
3. Once the correct password has been entered, the tabs associated with the System Setup button are displayed.
4. Select the More tab. The next set of System Setup tabs is displayed.
5. Select the Wave Gain tab. The Wave Gain tab is displayed.

FIGURE 9-45 Wave Gain Tab (System)

9.15.2 Changing the System Default Wave Gain Settings

This section outlines entering and modifying the system default wave gain settings.

1. Select the ECG mm/mV button until the desired ECG wave gain is displayed.
   The settings available for the ECG wave gain include 2.5, 5.0, 10.0, 20.0, and 30.0 mm/mV. The factory default setting is 10.0 mm/mV.
2. Select the IBPx mmHg buttons until the desired IBPx wave gains is displayed. There are four IBP buttons.
   The settings available for the IBP wave gains include 0-20, 0-40, 0-80, 60-140, 0-160, 0-225, 0-320, and -10 mmHg. The factory default setting is 0-320 mmHg.

NOTE: If the IBPx button(s) have been labeled at the bedside monitor, the same label(s) will be displayed on the individual wave gain button(s).
NOTE: Changing the ECG wave gain does not affect the arrhythmia analysis.

3. Select the **Resp Gain** button until the desired Resp Gain wave gain is displayed.
   The settings available for the Resp Gain wave gain include 1, 2, 3, 4, and 5. The factory default setting is 1.

4. Select the **CO2 (%)**, **kPa**, **mmHg** button until the desired CO2 wave gain is displayed.
   • The settings available for the CO2 % wave gain include 5.0, 7.5, and 10.0 %.
     The factory default setting is 10.0 %.
   • The settings available for the CO2 kPa wave gain include 5.0, 7.5, and 10.0 kPa.
     The factory default setting is 10.0 kPa.
   • The settings available for the CO2 mmHg wave gain include 40, 60, and 100 mmHg.
     The factory default setting is 100 mmHg.

5. Select the **O2 %** button until the desired O2 wave gain is displayed.
   The settings available for the O2 wave gain include 18 - 30, 18 - 60, and 18 - 100 %.
   The factory default setting is 18 - 100 %.

6. Select the **AGENT %** button until the desired Agent wave gain is displayed.
   The settings available for the agent wave gains include 0 - 0.1, 0 - 0.25, 0 - 0.5, 0 - 1.0, 0 - 1.5, and 0 - 2.0 %.
   The factory default setting is 0 - 2.0 %.

**Sidebar Button - Normal Screen**
Select the **Normal Screen** sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.
### 9.15.4 Troubleshooting

This section lists some of the potential messages and issues that may occur while using the System **Wave Gain** tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>The System Setup menu button is disabled.</td>
<td>The System Setup function is already open on a Panorama display and cannot be opened on multiple displays.</td>
<td>Close the System Setup function on the Panorama display on which it is opened.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
9.16 Wireless Tab

The Wireless tab manages the devices on the Panorama wireless network. It can be used to retrieve information from a wireless device as well as to program it. The Wireless tab should only be used when a wireless monitoring device is assigned to the selected patient tile. The Wireless tab is not available at the ViewStation or the WorkStation. Wireless device maintenance can only be performed from the host central station.

NOTE: The layout and functionality of the Wireless tab changes, based on the type of monitoring equipment attached to the programming cable.

9.16.1 Accessing the Wireless Tab

1. From the menu bar, select the System Setup button. The first set of System Setup tabs is displayed.
2. Follow the directions in the "Entering a Password" section on page 9-7.
3. Once the correct password has been entered, the tabs associated with the System Setup button are displayed.
4. Select the More tab. The next set of System Setup tabs is displayed.
5. Select the Wireless tab. The Wireless tab is displayed.

![Wireless Tab (device unread)](image)

FIGURE 9-46 Wireless Tab (device unread)

9.16.2 Entering Wireless Settings (Telepack)

This section outlines the settings available in the Wireless tab. The settings in the Wireless tab are shown by selecting the Read button and configured by selecting the Program button.

NOTE: Programming and assigning of the Telepack must be performed by authorized service personnel only.

The fields shown in these images are described on the following pages.
Device ID
The Device ID field displays the identification number that was assigned to the connected wireless device. A Device ID has a maximum of 8 alphanumeric characters. Data is shown in the Device ID field when the Wireless tab’s Read sidebar button is selected. If the read was not successful the Device ID field will remain blank. To view the connection status of the wireless device check the message in the Connect field. The Device ID field cannot be edited and has a factory default setting of blank.

The Device ID for a monitoring device is set up in the Equipment List tab. For additional information on setting up a Device ID, refer to the “Equipment Setup Tab” on page 9-33.

Connect Field
The Connect field displays the status of the connection between the Panorama Central Station and the wireless device. A connection status is shown in the Connect field when the Wireless tab’s Read sidebar button is selected. The status in the Connect field cannot be edited and has a factory default setting of blank.

The messages shown in the Connect field include:
Software Version

The **Software Version** field displays the version of software currently in use by the wireless device. Data is shown in the Software Version field when the Wireless tab’s **Read** sidebar button is selected. The factory default setting is blank. If the read was not successful the Software Version field will remain blank. To view the connection status of the wireless device check the message in the Connect field.

Hardware Version

The **Hardware Version** field displays the version of hardware in use by the connected device. Data is shown in the Hardware Version field when the Wireless tab’s **Read** sidebar button is selected and has a factory default setting of blank. If the read was not successful the Hardware Version field will remain blank. To view the connection status of the wireless device check the message in the Connect field.

Lead Selection

The **Lead Selection** button displays the ECG Lead to be monitored by the assigned device. It sets the lead to be monitored by the attached device. ECG lead selection options include Lead I, Lead II and Lead III.

- Data is shown in the Lead Selection field when the Wireless tab’s **Read** sidebar button is selected. Lead Selection has a factory default setting of blank. If the read was not successful the Lead Selection field will remain blank. To view the connection status check the message in the Connect field.
- The Lead to be monitored by a connected device can be assigned by selecting the **Program** sidebar button.

**NOTE:** The Lead Selection button will not be shown when the device cannot be recognized or there is a Read error.

Protocol Selection

The **Protocol Selection** button displays the protocol selection for the assigned device. It sets the protocol of the attached device.

- Data is shown in the Protocol Selection field when the Wireless tab’s **Read** sidebar button is pressed and has a factory default setting of blank. If the read was not successful the Protocol Selection field will remain blank. To view the connection status check the message in the Connect field.
- A Protocol Selection data can be programmed in a connected device by selecting the **Program** sidebar button.

### MESSAGE | REASON
--- | ---
Connected to 608 Telemetry | The connection with the Telepack was successful.
Connected to 608 Instrument Telemetry | The connection with the 608 Instrument Radio was successful.
Not Connected | The connection with the wireless device was not successful.
Error: Could not receive data | The connection with the wireless device was not successful. Either there was no response from the device or the device is not connected.
NOTE: The Protocol Selection button will not be shown when the device cannot be recognized or there is a Read error.

Channel Type
The Channel Type button displays the selected channel for the assigned device. It selects the channel type of the attached device. Channel type options include Medium and High.

- Data is shown in the Channel Type field when the Wireless tab’s Read sidebar button is pressed and has a factory default setting of blank. If the read was not successful the Channel Type field will remain blank. To view the connection status check the message in the Connect field.
- A Channel type can be programmed in a connected device by selecting the Program sidebar button.

NOTE: The Channel Type button will not be shown when the device cannot be recognized or there is a Read error.

DL Pair
The DL Pair button displays the channel that will be used for transferring data (i.e., demographics, alarms, arrhythmia) from the central station to the wireless bedside device. DL Pair options include 1, 2, 3, and Auto. Options 1, 2, and 3 should be used for networks that are configured for roaming. The Auto setting should be used for networks that are not configured for roaming.

NOTE: The DL Pair option affects wireless bedside devices using instrument radios.

- Data is shown in the DL Pair field when the instrument radio has previously been programmed.
- A DL Pair can be programmed in a connected device by selecting the Program sidebar button.

9.16.3 Sidebar Buttons
The following sidebar buttons are used to expand the functionality of this tab.

9.16.3.1 Read
Select the Read sidebar button to retrieve status parameters from the connected wireless device when it is connected to the Panorama Central Station in Service mode.

Selecting the Read button will attempt to obtain a reading from the connected wireless device.

- A successful read from a known device will display the fields and sidebar buttons appropriate to the attached device.
- An unsuccessful read will only display a connect status in the Connect field. The Device ID, Software Version, and Hardware Version fields will be blank.

9.16.3.2 Program
Select the Program sidebar button to program information in a wireless device when it is connected to the Panorama Central Station in Service mode.
NOTE: The Program sidebar button will be disabled until a valid device has been detected by the Panorama Central Station.

Selecting the Program sidebar button will display the message **Are you sure you want to program the wireless device?**.

- Select the Yes button to program the wireless device. The message **Wireless device programmed successfully.** is displayed.
- Select the No button to cancel the programming of the connected device.

### 9.16.4 Error Log Sidebar Button

Select the Error Log sidebar button to display the log of error messages for the connected wireless device. Selecting the Error Log sidebar button will bring up the Error log for the connected wireless device. When the Error Log sidebar button is selected the button is changed to the Telepack sidebar button. FIGURE 9-49 shows the Error Log for the Wireless tab.

![FIGURE 9-49 Error Log for the Wireless Tab](image)

NOTE: The Error Log sidebar button will be disabled until a valid device has been detected by the Panorama Central Station.

### Printing a Telepack Error Log Report

A Telepack Error Log Report can be printed from this tab by selecting the Print menu bar button.

For additional information regarding this report, refer to the “Telepack Error Log Report” on page 8-54.

### 9.16.5 Device ID

The Device ID field displays the identification number that was assigned to the connected wireless device. A Device ID has a maximum of 8 alphanumeric characters. Data is shown in the Device ID field when the Wireless tab’s Read sidebar button is selected. If the read was not successful the Device ID field will remain blank. To view the connection status of the wireless device check the message in the Connect field. The Device ID field cannot be edited and has a factory default setting of blank.
Device ID is set up in the **Equipment List** tab. For additional information on setting up a Device ID and the requirements of a Device ID, refer to the “Equipment Setup Tab” on page 9-33.

### 9.16.6 Connect Field

The **Connect** field displays the status of the connection between the Panorama Central Station and the wireless device. A connection status is shown in the Connect field when the Wireless tab’s **Read** sidebar button is selected. The status in the Connect field cannot be edited and has a factory default setting of blank.

The messages shown in the **Connect** field include:

<table>
<thead>
<tr>
<th>MESSAGE</th>
<th>REASON</th>
</tr>
</thead>
<tbody>
<tr>
<td>Connected to 608 Telemetry</td>
<td>The connection with the Telepack was successful.</td>
</tr>
<tr>
<td>Connected to 608 Instrument Telemetry</td>
<td>The connection with the 608 Instrument Radio was successful.</td>
</tr>
<tr>
<td>Not Connected</td>
<td>The connection with the wireless device was not successful.</td>
</tr>
<tr>
<td>Error: Could not receive data</td>
<td>The connection with the wireless device was not successful. Either there was no response from the device or the device is not connected.</td>
</tr>
</tbody>
</table>

### List Box

The Error Log list box contains the error messages that occurred at the connected wireless device. The following information is collected in an Error log:

- **Filename column**
  
  The Filename column shows the names of the files in the error list.

- **Line Number column**
  
  The Line Number column shows the line numbers for the items in the error list.

- **Message column**
  
  The Message column shows the messages that were sent to the Panorama system by the Telepack device.

- **Arg1 column**
  
  The Arg 1 column is populated by the text fields sent to the Panorama system by the Telepack device.

- **Arg2 column**
  
  The Arg 1 column is populated by the text fields sent to the Panorama system by the Telepack device.

### Error Log Read Sidebar Button

Select the **Read** sidebar button to retrieve error log messages from the connected wireless device when it is connected to the Panorama Central Station in Service mode. A device is in Service mode when it is connected to the Panorama Central Station’s serial port 2 via the device’s programming cable.
Selecting the **Read** button will attempt to read the Error log from the connected wireless device.

- A successful read from a known device will display the fields and sidebar buttons appropriate to the attached device.
- An unsuccessful read will only display a connect status in the **Connect** field.

**Clear Sidebar Button**
The Error Log contains a **Clear** sidebar button. Select the **Clear** sidebar button to delete the contents of the Telepack Error log. Selecting the **Clear** sidebar button produces a pop-up confirmation dialog on the screen that states **Are you sure you want to clear the error log? Selecting Yes will Delete all Data.**

- The choices for the Error Log confirmation dialog are No and Yes.
- Selecting the **No** button in the Error Log confirmation dialog does not clear the error log.
- Selecting the **Yes** button in the Error Log confirmation dialog deletes the contents of the Error log.

**Telepack Sidebar Button**
The Error Log contains a **Telepack** sidebar button. Select the **Telepack** button to return the tab to the **Wireless** tab display.

For additional information regarding printer setup, refer to the Panorama Printer Configuration Guide (P/N 0070-00-0561).

For additional information regarding the Telepack Error Log Report, refer to “Additional Reports” on page 8-41.

**Sidebar Button - Normal Screen**
Select the **Normal Screen** sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

**9.16.6.1 Free Channel Sidebar Button**
Select the **Free Channel** sidebar button to free an allocated channel for a wireless device. Selecting the **Free Channel** sidebar button will request a list of the wireless devices on the wireless network.

**NOTE:** The Free Channel sidebar button will be disabled until a valid device has been detected by the Panorama Central Station.

**9.16.6.2 Free Connected Channel Sidebar Button**
Select the **Free Connected Channel** sidebar button to free a channel from a connected device that supports channels.

**NOTE:** The Free Connected Channel sidebar button will be disabled until a valid device has been detected by the Panorama Central Station.
9.16.6.3 Sidebar Button - Normal Screen
Select the **Normal Screen** sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

9.16.7 Troubleshooting
This section lists some of the potential messages and issues that may occur while using the **Wireless** tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>The System Setup menu button is disabled.</td>
<td>The System Setup function is already open on a Panorama display and cannot be opened on multiple displays.</td>
<td>Close the System Setup function on the Panorama display on which it is opened.</td>
</tr>
<tr>
<td>You must enter all fields to program the wireless device</td>
<td>All of the fields in the Wireless tab were not completed before the Program button was pressed.</td>
<td>Select the OK button to close the error message dialog box. Complete all of the fields in the Wireless tab before selecting the Program button.</td>
</tr>
<tr>
<td>Program Failure: Invalid Data</td>
<td>The wireless device was not successfully programmed.</td>
<td>Select the OK button to close the error message dialog box. Recheck the accuracy of the data to be entered in the wireless device.</td>
</tr>
</tbody>
</table>
| Warning: Attached device is already programmed with a channel. Allocated channel will be lost if not freed first. Do you want to continue | The connected device has already been programmed and allocated with a channel. | Select the OK button to close the error message dialog box.  
  • Select the YES button to discard the previously allocated channel and try to obtain a new channel.  
  • Select the NO button to keep the previously obtained channel. |
| Program Failure: Unable to obtain a channel. | The system tried to obtain a channel from the Panorama Central Station but was unsuccessful. | Select the OK button to close the error message dialog box. Free up any unused channels. Try again to obtain a channel from the system. The system administrator can also purchase additional channels. |
| Network Error: Unable to obtain device list. | The system requested a device list from the wireless network but did not receive one within the allocated request time. | Select the OK button to close the error message dialog box. Request a device list from the wireless network again. |
| There are no devices available | An empty device list was received from the wireless network. | Select the OK button to close the error message dialog box. Enter the device in the Panorama Equipment list. |

* Messages are shown in all bold text.
### MESSAGE/ISSUE *

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Free Error: Device not found</td>
<td>The device channel was not successfully freed.</td>
<td>Select the <strong>OK</strong> button to close the error message dialog box. Try to free a different channel from the system. The system administrator can also purchase additional channels.</td>
</tr>
<tr>
<td>Program Failure: The channel was successfully freed but an error was encountered when programming the device. The device must now be programmed over its existing channel.</td>
<td>The channel was successfully freed but there was a problem with the programming of the device.</td>
<td>Select the <strong>OK</strong> button to close the error message dialog box. Program the device over its existing channel.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
This section outlines the Panorama Telepack:

- Introduction
- Button Indicators
- Indicator Lights
- Operating Instructions
- Telepack with Nonin SpO2 Module (Optional)
- Approved Telepack Accessories
- Troubleshooting
10.1 Introduction

The Panorama Telepack is a battery powered ambulatory device that acquires ECG information from an adult patient and transmits it to the Panorama Central Station for analysis. For the Telepack, the transmission to the Panorama Central Station is via the Panorama Telemetry Server. The Panorama Central Station is capable of performing data analysis functions such as Heart Rate determination, R-wave detection, and ECG filtering.

The Telepack is controlled through the use of buttons which are located on the front and side of the device. The status of the unit is shown by the LED indicator lights. The Telepack will not turn off unless the batteries are removed from the unit or when the batteries are depleted.

The Telepack has the following key features:

- Uni-directional data transmission (between ambulatory patient and Panorama Central Station) using wireless communication technology in the WMTS band or ISM band
- Functions such as Nurse Call, Attendant Present, Lead Select, Remote Print, and Lead/Battery/Link tests are easily performed using buttons
- LED lights are used to indicate link status, battery status, and ECG lead connection status
- 3-wire and 5-wire Lead ECG support

**CAUTION:** The Panorama Telepack and Instrument Radio are wireless devices and may be subject to intermittent signal dropout. Panorama wireless devices are designed to perform with a maximum cumulative signal dropout of one minute in a 24-hour period under ideal conditions. A crowded wireless environment or interference from other medical or non-medical devices, either intentional (e.g., VOIP phones, pagers, asset tracking systems, etc.) or unintentional (e.g., AC motors, microwave ovens, paper shredders, etc.), may result in a significantly increased amount of signal dropout experienced by any one or multiple wireless device(s).

See “Technical Alarms and Events” on page 13-19 for troubleshooting intermittent signal dropout of the wireless device.

**CAUTION:** Changes or modifications not expressly approved by Mindray DS USA, Inc. could void the user's authority to operate the equipment.

**NOTE:** After the power of the Panorama Central Station is interrupted, the Panorama Central Station saves and uses the latest configuration.

10.2 Applied Parts

The Telepack has the following applied parts:

- ECG electrodes and lead wires
- SpO₂ sensor and cable
10.3 Button Indicators

The following buttons are located on the Telepack device:

- One Nurse Call button: 

- Two Attendant Present button: 

- One LEAD SELECT button: 

- One CHECK button: 

- One RECORD button: 

10.3.1 Nurse Call Button

Press the Nurse Call button to request the presence of a clinician. When this button is pressed:

- A message displays on the Panorama Central Station indicating that the Nurse Call button was pressed.
- If configured, an event is recorded and posted to the Events tab Event list. For additional information about configuring this option, refer to “Alarm Responses Tab (Patient)” on page 6-13.

10.3.2 Attendant Present Buttons

Press the Attendant Present buttons simultaneously to indicate the presence of a clinician at the patient’s bedside. When these buttons are pressed:

- A message posts to the Events list indicating that the Attendant Present buttons were pressed.
- The Nurse Call alarm is acknowledged, if the Nurse Call button was pressed.

10.3.3 LEAD SELECT Button

Press the ECG LEAD SELECT button to:

- Change the active lead when using a 3-wire lead set
- The ECG LED that is active will turn on for 2 seconds
10.3.4 **CHECK Button**  
Press the **CHECK** button to verify LED indicator lights are functioning, ECG Lead connection, link status, and battery status.

For information on interpreting the status of the ECG Lead, Link Status and Battery Status LED indicator lights, refer to the following sections: “ECG Lead Diagram LED Indicator Lights” on page 10-4, “Link Status LED Indicator Light” on page 10-6, and “Battery Status LED Indicator Light” on page 10-6.

10.3.5 **RECORD Button**  
Press the **RECORD** button to remotely print the “All Strips Report” to the Panorama Central Station default printer.

10.4 **Indicator Lights**  
The Telepack uses LED indicator lights to verify the functionality of the device. The indicator lights on the device include:

- 3-wire and 5-wire ECG lead indicator lights
- 1 Link Status indicator light
- 1 Battery Status indicator light

10.4.1 **ECG Lead Diagram LED Indicator Lights**  
The Telepack ECG Lead diagram’s LED indicator lights show the status of the lead connections with the patient. The Telepack’s ECG Lead LED indicator lights (shown in FIGURE 10-1 and FIGURE 10-2), include five green LED indicators that correspond to the following lead connections:
Telepack (Optional)  

Indicator Lights

The ECG Lead LED indicator lights remain off except during the initial power up phase and during a lead test sequence. For additional information, refer to “Lead Placement Procedures” on page 4-2.

Press the CHECK button to verify the Telepack’s ECG lead connections.

**NOTE:** When the CHECK button is pressed, all of the device LED indicator lights are tested simultaneously.

This test determines if the patches are connected based on thoracic impedance. If a patch adhesive is not sticking, a patch is dry, or the skin gel is dry, the test fails.

The Telepack’s ECG Lead LED indicator lights remain on for approximately 10 seconds during the test sequence, indicating that the Telepack has good lead connectivity. An unlit ECG Lead LED light indicates that the Telepack has no lead connectivity.

---

**FIGURE 10-1** Front View of Telepack (AHA) with Domestic Overlay  
**FIGURE 10-2** Front View of Telepack (AHA) with International Overlay

**Domestic Overlay**
- White RA (right arm)
- Black LA (left arm)
- Brown V (chest)
- Green RL (right leg)
- Red LL (left leg)

**International Overlay**
- Red R (right arm)
- Yellow L (left arm)
- White C (chest)
- Black N (right leg)
- Green F (left leg)
NOTE: If a 3-wire Lead set is used, a maximum of three lights will illuminate. If a 5-wire Lead set is used, a maximum of five lights will illuminate.

10.4.2 Link Status LED Indicator Light
The Telepack Link Status LED indicator light verifies the Telepack’s connection to the Panorama Patient Monitoring Network.

The Link Status LED indicator light remains on for approximately 10 seconds during the initial power up and during the test sequence.

Press the CHECK button to verify the Telepack’s link status with the wireless network.

NOTE: When the CHECK button is pressed, all of the device LED indicator lights are tested simultaneously.

The Telepack’s Link Status LED indicator light shows the results of the link status test as follows:

<table>
<thead>
<tr>
<th>TELEPACK</th>
<th>LED STATUS</th>
<th>DESCRIPTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Telepack</td>
<td>On</td>
<td>The Telepack has been configured for operation. This is not an indication of being in range of the wireless network.</td>
</tr>
<tr>
<td></td>
<td>Flashing</td>
<td>The Telepack has not been configured for operation.</td>
</tr>
<tr>
<td></td>
<td>Off</td>
<td>The Telepack batteries need replacement.</td>
</tr>
</tbody>
</table>

10.4.3 Battery Status LED Indicator Light
The Telepack Battery Status LED indicator light verifies the status of the Telepack’s batteries.

Press the CHECK button to verify the Telepack’s battery status.

NOTE: When the CHECK button is pressed, all of the device LED indicator lights are tested simultaneously.

During this test sequence, the Battery Status LED indicator light will remain on for approximately 10 seconds if the Telepack batteries have more than 2 hours of battery life remaining. A flashing Battery Status LED light indicates the Telepack batteries have less than 2 hours of battery life remaining. An unlit Battery Status LED light indicates the Telepack batteries should be replaced immediately, or that they have been incorrectly inserted.

NOTE: The Telepack device should not be used for battery storage purposes.

NOTE: When a Telepack is assigned to a patient tile, the battery charge percentage integer, and percent sign (%) are displayed in the digital data tile at the central station.

NOTE: The Telepack batteries used may be subject to local regulations regarding disposal. At the end of the battery life, never dispose of the batteries in a fire. Batteries should be disposed of in accordance with any local regulations.
10.5 Operating Instructions
This section of the manual provides instructions for programming the Telepack device for use with the Panorama Central Station.

10.5.1 Initial Setup of Telepack
The Panorama Central Station should be installed and running before you begin to set up the Telepack device. For additional information, refer to the Panorama Service Manual (P/N 0070-00-0634).

The following items are required to initially install and program the Telepack:

- Telepack device
- 2 AA size alkaline batteries (P/N 0146-00-0077-01)
- Telepack service mode cable

10.5.1.1 Inserting Batteries
To activate the Telepack device, insert the 2 AA size alkaline batteries in the Telepack battery compartment. Use the label on the back of the Telepack as a guide. Secure the end cap of the battery compartment by twisting firmly in the clockwise direction.

The Telepack automatically performs an initial system check if the batteries are installed properly.

FIGURE 10-3 Telepack, rear panel (right side of image intentionally blurred)

WARNING: Do not incinerate batteries, possible explosion may occur.

CAUTION: Visually inspect the battery compartment for any foreign object prior to use.
CAUTION: The Telepack may not function if the batteries installed in the device are corroded. Use only approved batteries (P/N 0146-00-0077-01) in the Telepack and observe proper polarity. Installing batteries with the incorrect polarity may cause the batteries to become excessively warm.

The Telepack will not transmit data to the Panorama Central Station once the batteries have become depleted. To test the batteries, press the CHECK button and confirm that both the Link LED and the Battery LED is lit.

When not in use, the Telepack device should be stored without batteries.

10.5.1.2 Programming the Telepack
This section describes how to program the Telepack. For additional information, refer to the “Wireless Tab” on page 9-64.

NOTE: Programming and assigning of the Telepack must be performed by authorized service personnel only.

CAUTION: The Telepack should not be attached to the patient when the device is being programmed into the system.

To program the Telepack:

1. Connect the Telepack service mode cable from the Panorama Central Station to the serial port provided on the base of the Telepack device.
2. Insert the batteries into the device. For additional information, refer to “Inserting Batteries” on page 10-7.
3. From the central station main screen, select the System Setup menu button.
4. Select the Installation Setup tab, and enter the System password. For additional information, refer to “Entering a Password” on page 6-2.
5. Select the More tab, then select the Wireless tab. The Wireless tab is displayed (shown in FIGURE 10-4).

![FIGURE 10-4 Wireless Tab]
6. Select the Read sidebar button. The information from the Telepack device is displayed in the appropriate fields.

7. Select the Program sidebar button. A dialog box appears.

8. Change any editable fields as necessary.

9. Select the Yes button to confirm. The message Wireless device programmed successfully appears is displayed confirming that the Telepack was programmed correctly.

NOTE: If the Telepack device is not connected properly or if the Panorama Central Station is unable to detect the device, the message, “Error: Could not receive data” is displayed at the Panorama Central Station.

10.5.1.3 Entering the Telepack to the Equipment List
The Telepack device should be entered to the Panorama Central Station Equipment List using the Equipment Setup tab. For additional information about entering a Telepack device to the Panorama Central Station, refer to “Entering New Equipment” on page 9-34.

The Device ID is found on the back of the Telepack device. After assigning the device to the Panorama Central Station, the Telepack is ready for patient use.

10.5.1.4 Telepack Power Up Sequence
The Telepack goes through a power up sequence each time the batteries are inserted in the device. The power up sequence includes a hardware and software check, an ECG Lead connection status test, a link status test, and a battery status test. A successful power up sequence is indicated when the LED indicator lights illuminate for approximately 10 seconds.

For additional information on the status tests performed by the Telepack during the start up sequence, refer to the “LEAD SELECT Button” on page 10-3.

After the Telepack device powers up, it is ready to be programmed and assigned to a patient.

10.5.1.5 Telepack Communication Range
The range for the Telepack device to communicate and transfer data is generally within 30 feet of a Panorama Antenna. This distance is subject to site survey, and is affected by building materials. The device is for indoor use only.

10.5.2 Securing the Telepack Device
Secure the Telepack to the patient after the ECG leads are connected to the device. The device must be secured in a telemetry pouch or in the pocket of a patient gown.
10.6 Telepack with Nonin SpO2 Module (Optional)

This section of the manual provides instructions for installing the Nonin SpO2 module in the Telepack.

10.6.1 Introduction

The Telepack with Nonin SpO2 module is an optional accessory used to monitor and display SpO2 data for a patient that is being monitored at the Panorama Central Station. The SpO2 data includes SpO2 waveform, SpO2 numeric data, and numeric pulse rate.

**WARNING:** The connector cover on the bottom of the Telepack must remain in place while in use without an SpO2 module connected. It should only be removed to program the device and connect an SpO2 module. When connected to an SpO2 module, the cover should be stored on the bottom of the SpO2 module and placed back on the Telepack when the SpO2 module is removed.

10.6.2 Installing the SpO2 Module on the Telepack

1. Using care, remove the flexible cover from the connector on the bottom of the Telepack by using a small flat-blade screw driver or other similar tool (shown in FIGURE 10-5).

2. Install the flexible cover (removed in step 1) on the connector on the bottom of the SpO2 Module.

**FIGURE 10-5** Removing Telepack Flexible Cover

**FIGURE 10-6** Installing Flexible Cover in SpO2 Module
3. Turn the Telepack off by rotating the battery cap in a counter-clockwise direction. The Telepack is turned off.

![FIGURE 10-7 Removing Telepack Battery Cap](image)

4. Align the two devices (shown in FIGURE 10-8) and then slide the SpO₂ Module into the Telepack. First, the connectors are mated; Continue pushing the devices together firmly until a loud audible “click” is heard as the mechanical latch is engaged.

![FIGURE 10-8 Inserting Telepack into SpO₂ Module](image)

5. Turn the Telepack on by securing the end cap of the battery compartment by twisting firmly in the clockwise direction.

10.6.3 Removing the SpO₂ Module from the Telepack

1. Turn the Telepack off by rotating the battery cap in a counter-clockwise direction. The Telepack is turned off.
2. Press the large button on the right side of the SpO₂ module (shown in FIGURE 10-9) until a loud audible click is heard. The click sound indicates that the latch has been properly disengaged from the Telepack.

![FIGURE 10-9 SpO₂ Module Release Button](image)

NOTE: As a safety feature, the connectors should remain connected even after the mechanical latch is disengaged.

3. Separate the SpO₂ Module from the Telepack.
4. Remove the flexible cover from the bottom of the SpO₂ Module.
5. To avoid electrostatic sensitivity, install the flexible cover on the connector on the bottom of the Telepack.

NOTE: The Telepack may automatically reset due to an electrostatic event if the flexible cover is not replaced.

10.6.4 Installing the Patient Cable
A patient cable is needed to collect the SpO₂ data.

1. Mate the 8-pin connector on the cable with the matching connector on the bottom of the SpO₂ Module. The connector should be firmly pressed in place to ensure that it is properly placed.

![FIGURE 10-10 Inserting Patient Cable in SpO₂ Module](image)
2. Route the cable in the channel on the back of the unit. The channel regions (shown in FIGURE 10-11) should also be pressed firmly to ensure that they are properly placed.

![FIGURE 10-11 Securing the Patient Cable](image)

10.6.5 Attaching the Finger Clip Sensor

1. Insert the finger (preferably the index, middle, or ring finger) into the Nonin finger clip until the end of the finger reaches the finger stop (shown in FIGURE 10-12). Keep the fingernail facing the sensor top. Ensure that long fingernails are not interfering with the proper finger position.

**NOTE:** Disposable finger sensors (shown in FIGURE 10-13) may also be used. Follow the packaging directions carefully to ensure proper application.

![FIGURE 10-12 Reusable finger clip sensor](image) ![FIGURE 10-13 Disposable sensor](image)

**NOTE:** For best results, when using the sensor for data collection, secure the sensor cable independently from the sensor with medical device tape, preferably around the base of the finger. Be sure that the tape securing the cable does not restrict the blood flow.

**CAUTION:** Tissue damage or inaccurate SpO2 measurements may be caused by incorrect sensor application or use, such as wrapping too tightly, applying supplemental tape, failing to inspect the sensor site periodically, or failing to position appropriately. Carefully read the sensor directions for use, the Panorama Operating Instructions, and all precautionary information before use.
CAUTION: Inaccurate SpO2 measurements may be caused by:

- Incorrect sensor application or use
- Significant levels of dysfunctional hemoglobins, (e.g., carboxyhemoglobin or methemoglobin)
- Intra-vascular dyes such as indocyanine green or methylene blue
- Exposure to excessive illumination such as surgical lamps (especially ones with a xenon light source), bilirubin lamps, fluorescent lights, infrared heating lamps, or excessive ambient light. In such cases, cover the sensor site with opaque material.
- Excessive patient movement
- Venous pulsations
- Electro-surgical interference
- Placement of a sensor on an extremity that has a blood pressure cuff, arterial catheter, or intra-vascular line.
- Cold extremities due to reduced circulation.
- Nail polish or fungus

CAUTION: In certain situations in which perfusion and signal strength are low, such as in patients with thick or pigmented skin, inaccurately low SpO2 readings will result. Verification of oxygenation should be made, especially in patients with chronic lung disease, before instituting any therapy or intervention.

CAUTION: Many patients suffer from poor peripheral perfusion due to hypothermia, hypovolemia, severe vasoconstriction, reduced cardiac output, etc. These symptoms may cause a loss in vital sign readings.

CAUTION: Prolonged and continuous monitoring may increase the risk of skin erosion and pressure necrosis at the site of the sensor. Check the SpO2 sensor site frequently to ensure proper positioning, alignment and skin integrity at least every eight (8) hours; with the Adult and Pediatric re-usable finger sensor, check every four (4) hours; for patients of poor perfusion or with skin sensitive to light, check every 2 - 3 hours; more frequent examinations may be required for different patients. Change the sensor site if signs of circulatory compromise occur.

10.6.6 Removing the Patient Cable

- The patient cable is removed by gently pulling the cable from its channel, and then gripping and pulling the pull tab, to disconnect the connectors.
### 10.7 Approved Telepack Accessories

The accessory material that contacts the patients has undertaken the bio-compatibility test and are verified to be in compliance with ISO10993-1.

<table>
<thead>
<tr>
<th>DESCRIPTION</th>
<th>PART NUMBER</th>
</tr>
</thead>
<tbody>
<tr>
<td>Telepack</td>
<td>0998-00-0191-06 (Domestic) 0998-00-0191-07 (International)</td>
</tr>
<tr>
<td>Electrodes</td>
<td>0681-00-0100-01 (Box of 600) 0681-00-0100-02 (Box of 60)</td>
</tr>
<tr>
<td>ECG Lead Set for Telepack, 18”</td>
<td>0012-00-1503-04 (3-wire lead set, domestic) 0012-00-1503-01 (5-wire lead set, domestic) 0012-00-1503-13 (3-wire lead set, international) 0012-00-1503-10 (5-wire lead set, international)</td>
</tr>
<tr>
<td>ECG Lead Set for Telepack, 24”</td>
<td>0012-00-1503-05 (3-wire lead set, domestic) 0012-00-1503-02 (5-wire lead set, domestic) 0012-00-1503-14 (3-wire lead set, international) 0012-00-1503-11 (5-wire lead set, international)</td>
</tr>
<tr>
<td>ECG Lead Set for Telepack, 36”</td>
<td>0012-00-1503-06 (3-wire lead set, domestic) 0012-00-1503-03 (5-wire lead set, domestic) 0012-00-1503-15 (3-wire lead set, international) 0012-00-1503-12 (5-wire lead set, international)</td>
</tr>
<tr>
<td>ECG Plug</td>
<td>0012-00-1527-32 (RL-Green plug, domestic) 0012-00-1527-34 (V-Brown plug, domestic) 0012-00-1527-33 (N-Black plug, international) 0012-00-1527-35 (C-White plug, international)</td>
</tr>
<tr>
<td>Kit, Telepack Programming</td>
<td>0020-00-0485-01</td>
</tr>
<tr>
<td>Telepack SpO2 Module</td>
<td>0998-00-0192-01</td>
</tr>
<tr>
<td>Adult Single Patient Use Nonin Sensor, 10 pack</td>
<td>0600-00-0139-10</td>
</tr>
<tr>
<td>Adult Reusable Nonin SpO2 Sensor, 1 each</td>
<td>0600-00-0140-01</td>
</tr>
<tr>
<td>Interface Cable for Single Patient Use Adult SpO2 Sensor</td>
<td>0012-00-1542-01</td>
</tr>
<tr>
<td>Kit, Adult Nonin SpO2 Sensors</td>
<td>0020-00-0167</td>
</tr>
<tr>
<td>SpO2 Module Cleaning Plug</td>
<td>0003-00-0055</td>
</tr>
<tr>
<td>Batteries (Alkaline, size AA)</td>
<td>0146-00-0077-01</td>
</tr>
<tr>
<td>Panorama Telepack Pouch</td>
<td>0202-00-0179-02 (Bag of 25)</td>
</tr>
</tbody>
</table>

### ECG AAMI (DOMESTIC) LEAD WIRES

<p>| Lead Wire, LA-Black, 18” | 0012-00-1527-04 |
| Lead Wire, LL-Red, 18” | 0012-00-1527-03 |
| Lead Wire, RA-White, 18” | 0012-00-1527-02 |
| Lead Wire, RL-Green, 18” | 0012-00-1527-05 |
| Lead Wire, V-Brown, 18” | 0012-00-1527-01 |
| Lead Wire, LA-Black, 24” | 0012-00-1527-14 |
| Lead Wire, LL-Red, 24” | 0012-00-1527-13 |
| Lead Wire, RA-White, 24” | 0012-00-1527-12 |
| Lead Wire, RL-Green, 24” | 0012-00-1527-15 |
| Lead Wire, V-Brown, 24” | 0012-00-1527-11 |
| Lead Wire, LA-Black, 36” | 0012-00-1527-24 |</p>
<table>
<thead>
<tr>
<th>DESCRIPTION</th>
<th>PART NUMBER</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lead Wire, LL-Red, 36&quot;</td>
<td>0012-00-1527-23</td>
</tr>
<tr>
<td>Lead Wire, RA-White, 36&quot;</td>
<td>0012-00-1527-22</td>
</tr>
<tr>
<td>Lead Wire, RL-Green, 36&quot;</td>
<td>0012-00-1527-25</td>
</tr>
<tr>
<td>Lead Wire, V-Brown, 36&quot;</td>
<td>0012-00-1527-21</td>
</tr>
</tbody>
</table>

**ECG IEC (INTERNATIONAL) LEAD WIRES**

<table>
<thead>
<tr>
<th>DESCRIPTION</th>
<th>PART NUMBER</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lead Wire, N-Black, 18&quot;</td>
<td>0012-00-1527-10</td>
</tr>
<tr>
<td>Lead Wire, R-Red, 18&quot;</td>
<td>0012-00-1527-07</td>
</tr>
<tr>
<td>Lead Wire, C-White, 18&quot;</td>
<td>0012-00-1527-06</td>
</tr>
<tr>
<td>Lead Wire, F-Green, 18&quot;</td>
<td>0012-00-1527-08</td>
</tr>
<tr>
<td>Lead Wire, L-Yellow, 18&quot;</td>
<td>0012-00-1527-09</td>
</tr>
<tr>
<td>Lead Wire, N-Black, 24&quot;</td>
<td>0012-00-1527-20</td>
</tr>
<tr>
<td>Lead Wire, R-Red, 24&quot;</td>
<td>0012-00-1527-17</td>
</tr>
<tr>
<td>Lead Wire, C-White, 24&quot;</td>
<td>0012-00-1527-16</td>
</tr>
<tr>
<td>Lead Wire, F-Green, 24&quot;</td>
<td>0012-00-1527-18</td>
</tr>
<tr>
<td>Lead Wire, L-Yellow, 24&quot;</td>
<td>0012-00-1527-19</td>
</tr>
<tr>
<td>Lead Wire, N-Black, 36&quot;</td>
<td>0012-00-1527-30</td>
</tr>
<tr>
<td>Lead Wire, R-Red, 36&quot;</td>
<td>0012-00-1527-27</td>
</tr>
<tr>
<td>Lead Wire, C-White, 36&quot;</td>
<td>0012-00-1527-26</td>
</tr>
<tr>
<td>Lead Wire, F-Green, 36&quot;</td>
<td>0012-00-1527-28</td>
</tr>
<tr>
<td>Lead Wire, L-Yellow, 36&quot;</td>
<td>0012-00-1527-29</td>
</tr>
</tbody>
</table>

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

This section lists some of the potential messages and issues that may occur while using a Telepack device.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Telepack/LED’s not turning on</td>
<td>Batteries are not installed correctly, batteries are depleted, or batteries corroded.</td>
<td>Visually inspect the battery compartment for any foreign object. Replace the batteries.</td>
</tr>
<tr>
<td>Telepack not transmitting data to the Panorama Central Station.</td>
<td>Batteries are discharged.</td>
<td>Press the CHECK button to verify that the batteries are depleted. Replace as necessary.</td>
</tr>
<tr>
<td>Flashing Battery Status LED indicator light</td>
<td>Batteries in the Telepack are weakening.</td>
<td>Replace batteries</td>
</tr>
<tr>
<td>The message <strong>Error: Could not receive data</strong> is displayed, while attempting to program the Telepack</td>
<td>Telepack is not connected properly, or the Panorama Central Station is unable to detect the Telepack.</td>
<td>Verify that the service mode cable is properly connected to the device. Retry programming. Check batteries.</td>
</tr>
<tr>
<td>Unlit ECG Lead LED indicator light</td>
<td>Telepack has no lead connectivity.</td>
<td>Connect the electrode. Check batteries.</td>
</tr>
<tr>
<td><strong>Communications Lost</strong></td>
<td>Telepack’s radio signal is out of range with the wireless network.</td>
<td>For Telepack, verify the Telepack is within 30 feet of a network access point. Check batteries.</td>
</tr>
<tr>
<td>Flashing Link Status LED indicator light</td>
<td>Telepack is not communicating with the Panorama Central Station.</td>
<td>Verify that Device ID is registered on the Panorama Central Station. Verify that Panorama Central Station is functioning.</td>
</tr>
<tr>
<td>SpO2 waveform not displayed at the Panorama Central Station patient tile</td>
<td>SpO2 waveform has not been selected in the Display tab.</td>
<td>Go to the Display tab and select the SpO2 waveform from the Parameter list to display in the Waveform 2 position. For additional information regarding the Display tab, refer to “Configuring the Display Tab” on page 5-52.</td>
</tr>
<tr>
<td>SpO2 module is not properly connected to the Telepack.</td>
<td>Power down the Telepack and reconnect the SpO2 module to the Telepack.</td>
<td></td>
</tr>
<tr>
<td>Battery cap on the Telepack was not secured.</td>
<td>Verify that the Telepack battery cap is secured.</td>
<td></td>
</tr>
<tr>
<td><strong>SpO2 Check Sensor</strong></td>
<td>An SpO2 sensor issue was detected.</td>
<td>Select the digital tile or waveform tile area, or select the View button to acknowledge the technical event.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
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11.0  User Maintenance

This section of the manual outlines proper maintenance of the Panorama Central Station touch screen and Telepack device.

- Touch Screen User Maintenance
- Telepack User Maintenance
11.1 Touch Screen User Maintenance

The Panorama display uses a touch screen overlay for more simple navigation of the system. (A mouse can also be used for point-and-click navigation of the system.)

For instructions on turning the touch screen on and off, refer to “Touch Screen Tab” on page 9.4.

11.1.1 Care and Cleaning of the Touch Screen

For the best performance, it is recommended that the Panorama Central Station touch screen be kept clean. Observe the following precautions when cleaning the touch screen:

- **CAUTION:** To avoid potential system issues, deactivate the touch screen before cleaning.
- **CAUTION:** DO NOT spray glass cleaner directly on a display as it could possibly leak inside a non-sealed unit and cause damage.
- **CAUTION:** Follow your hospital protocol for handling of blood and body fluids.

- Always remember to use a cloth or towel to apply glass cleaner to the touch screen.
- Any standard glass cleaner can be used to clean the touch screen. DO NOT use abrasive cleaning materials to clean a touch screen. DO NOT use alcohol or solvents containing chlorinated hydrocarbons.
- Remove fingerprints and stains by using a liquid lens cleaner and a soft cloth.
- Use a fine soft-hair brush to carefully brush away dust and dirt particles.
11.2 Telepack User Maintenance
The following sections provide information about storing and cleaning the Telepack device and the Nonin SpO2 module.

11.2.1 Storing the Telepack
When not in use, carefully store the Telepack by wrapping the lead set loosely around the casing of the Telepack device, or with the lead set attached and hanging freely. Do not wrap the lead set tightly around the casing of the Telepack as it may damage the wires.

CAUTION: Ensure that the batteries are removed when the Telepack is being stored.

To remove the batteries, twist the endcap of the battery compartment in a counter-clockwise direction. Removing the batteries helps preserve battery life and could prevent potential damage to the Telepack due to battery leakage.

NOTE: The Telepack remains on until the batteries are removed or depleted.

The Telepack has been tested and found to comply with the limits for a Class B digital device, pursuant to part 15 of the FCC Rules. These limits are designed to provide reasonable protection against harmful interference in a residential installation.

The Telepack generates, uses and can radiate radio frequency energy and if not installed and used in accordance with the instructions, may cause harmful interference to radio communications. However, there is no guarantee that interference will not occur in a particular installation.

If the Telepack does cause harmful interference to radio or television reception, which can be determined by turning the Telepack off and on, the user is encouraged to try and correct the interference by one or more of the following measures:

- Reorient or relocate the receiving antenna.
- Increase the separation between the equipment and receiver.
- Consult the dealer or an experienced radio/TV technician for help.

11.2.2 Cleaning the Telepack Device
The information in this section pertains to cleaning the Telepack and its accessories. The components that need to be cleaned are:

- Telepack device
- Battery compartment
- ECG Lead sets
- Nonin SpO2 module
- SpO2 Finger Clip Sensor

NOTE: The Telepack should be cleaned before and after each patient’s use.
The solutions used to clean the Telepack should be compatible with the cleaning solutions used to clean the ECG lead sets, if they are going to remain attached while the Telepack is cleaned. If the cleaning solution is not compatible, ensure that the ECG leads do not come in contact with the cleaning solution used to clean the Telepack.

The cleaning solutions that can be used are:

- Soap and water

11.2.2.1 Guidelines for Cleaning the Telepack

- Remove the batteries, and reattach the end cap of the battery compartment firmly. Clean the empty Telepack using a cloth moistened with cleaning solution, by gently wiping the outside of the Telepack.

- Remove the ECG lead wire from the device to clean around the ECG connector.

**NOTE:** Do not immerse the Telepack in the cleaning solution.

- Any excess cleaning solution should be wiped away with a cloth moistened with distilled water.

- A lint free cloth should be used to dry the Telepack unit thoroughly.

11.2.2.2 Cleaning the Battery Compartment

To clean the battery compartment, follow these guidelines:

- Remove the batteries from the battery compartment by twisting the endcap in a counter-clockwise direction. Visually inspect for fluids or foreign matter.

- Use a gauze pad or long handle swab moistened with soap and water to clean the battery compartment.

- Excess cleaning solution should be wiped away with a cloth moistened with distilled water.

- A lint free cloth should be used to wipe the inside of the battery compartment. Air dry the battery compartment prior to re-inserting the batteries.

**NOTE:** Ensure that the battery contact leads and the battery compartment are dry prior to re-inserting the batteries for use.

11.2.2.3 Cleaning the ECG Lead Sets

- Refer to the “Recommendation regarding cleaning and disinfection of cables” included in the lead set packaging.

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

11.2.2.4 Cleaning the SpO2 Module

In the event that the SpO2 Module becomes soiled, a cleaning plug (P/N 0003-00-0055) should be used to cover the serial connector on the SpO2 Module before it is cleaned.
11.2.2.4.1 Approved Cleaning Solutions

Please refer to the approved cleaning solutions described in “Cleaning the Telepack Device” on page 11-3.

11.2.2.4.2 Inserting the Cleaning Plug

1. Carefully cover the connector by aligning the cleaning plug with the lower portion of the SpO₂ Module.

2. Press downward while angling the top arm of the plug into the shallow circular recess.

![Inserting the Cleaning Plug]

**FIGURE 11-1** Inserting the Cleaning Plug

![Securing the Cleaning Plug]

**FIGURE 11-2** Securing the Cleaning Plug

**NOTE:** The cleaning plug should remain in place during cleaning.

11.2.2.4.3 Removing the Cleaning Plug

Remove the cleaning plug by pulling the tip of the top arm from the recess in the SpO₂ module.
11.2.2.5 Cleaning the SpO₂ Finger Clip Sensor and Cables
Refer to the cleaning instructions provided in the SpO₂ Finger Sensor and cables packaging.

11.2.3 Disinfecting the Telepack Device
The information in this section pertains to disinfecting the Telepack and its accessories.

The components that need to be disinfected are:

- Telepack device
- ECG Lead sets
- Nonin SpO₂ module
- SpO₂ Finger Clip Sensor

**NOTE:** The Telepack should be disinfected before and after each patient’s use.

The solutions used to disinfect the Telepack should be compatible with the solutions used to disinfect the ECG lead sets, if they are going to remain attached while the Telepack is disinfected. If the disinfecting solution is not compatible, ensure that the ECG leads do not come in contact with the disinfecting solution used to disinfect the Telepack. Only hospital-approved disinfecting procedures like those recommended by AAMI or AORN should be used to disinfect the Telepack unit.

The disinfecting solutions that can be used are:

- Dilute chlorine bleach (sodium hypochlorite), 10% solution
- Glutaraldehyde 2% (for example, Cidex)
- Isopropyl alcohol 70%
- Ethyl alcohol

**Guidelines for Disinfecting the Telepack**

- Remove the batteries, and reattach the end cap of the battery compartment firmly. Disinfect the empty Telepack using a cloth moistened with disinfecting solution, by gently wiping the outside of the Telepack.
- Remove the ECG lead wire from the device to disinfect around the ECG connector.

**NOTE:** Do not immerse the Telepack in the disinfecting solution.

- Any excess disinfecting solution should be wiped away with a cloth moistened with distilled water.
- A lint free cloth should be used to dry the Telepack unit thoroughly.

11.2.3.1 Disinfecting the Battery Compartment
To clean the battery compartment, follow these guidelines:
• Remove the batteries from the battery compartment by twisting the endcap in a counterclockwise direction.
• Visually inspect for fluids or foreign matter.
• Use a gauze pad or long handle swab moistened with Ethyl alcohol to disinfect the battery compartment.
• Excess solution should be wiped away with a cloth moistened with distilled water.
• A lint free cloth should be used to wipe the inside of the battery compartment. Air dry the battery compartment prior to re-inserting the batteries.

NOTE: Ensure that the battery contact leads and the battery compartment are dry prior to re-inserting the batteries for use.

11.2.3.2 Disinfecting the ECG Lead Sets
Refer to the “Recommendation regarding cleaning and disinfection of cables” included in the lead set packaging.

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

11.2.3.3 Disinfecting the SpO₂ Module
In the event that the SpO₂ Module becomes soiled, a cleaning plug (P/N 0003-00-0055) should be used to cover the serial connector on the SpO₂ Module before it is disinfected.

CAUTION: Use the designated cleaning plug (P/N 0003-00-0055) when cleaning and disinfecting the Nonin SpO₂ module. If fluids enter the device, it may affect the functionality of the module or render it unusable.

Approved Disinfecting Solutions
Please refer to the approved cleaning solutions described in “Disinfecting the Telepack Device” on page 11-6.

Refer to “Inserting the Cleaning Plug” on page 11-5 and “Removing the Cleaning Plug” on page 11-5 for inserting and removing the Cleaning Plug.

11.2.3.4 Disinfecting the SpO₂ Finger Clip Sensor and Cables
Refer to the cleaning instructions provided in the SpO₂ Finger Sensor and cables packaging.

11.2.4 Disposal of the Telepack Device
Adhere to the following guidelines when disposing of any Telepack device and its accessories.
• The Telepack should be disposed of in a manner adhering to local or country regulations regarding disposal of electronic and computer accessories.

• If your state/country laws recommend recycling, you can dispose of the device by following the recycling program offered by state/federal run agencies.

11.2.5 Disposing of Batteries
Adhere to the following note when disposing the batteries.

NOTE: The Telepack batteries used may be subject to local regulations regarding disposal. At the end of the battery life, never dispose of the batteries in a fire. Batteries should be disposed of in accordance with any local regulations.
This section is for reference only. The Parameter Specifications section lists the allowable range specifications for all of the parameters measured by the Panorama Central Station and the invalidation of parameter data. It includes the following items:

- Parameter Specifications
- Invalid Data Display
12.1 Parameter Specifications

The parameters displayed on the Panorama Central Station are dependent on the type of equipment connected to the Panorama system.

12.1.1 Heart Rate: HR

The Heart Rate parameter obtains data from ECG and IBP sources.

Unit of Measure: Beats Per Minute (bpm)

Resolution: 1 bpm

Heart Rate Source: ECG

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (bpm)</th>
<th>MAXIMUM VALUE (bpm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>30</td>
<td>350</td>
</tr>
<tr>
<td></td>
<td>20 (V 12/V 21 and Telepack only)</td>
<td></td>
</tr>
<tr>
<td>Pediatric</td>
<td>30</td>
<td>300</td>
</tr>
<tr>
<td></td>
<td>20 (V 12/V 21 and Telepack only)</td>
<td></td>
</tr>
<tr>
<td>Adult</td>
<td>30</td>
<td>300</td>
</tr>
<tr>
<td></td>
<td>20 (V 12/V 21 and Telepack only)</td>
<td></td>
</tr>
</tbody>
</table>

The Panorama Central Station displays ECG waveforms when using 3-wire, 5-wire, and 12 lead cable. The ECG leads available at the Panorama Central Station are dependent on the lead cable connected to the patient.

<table>
<thead>
<tr>
<th>LEAD</th>
<th>WAVEFORM LEADS AVAILABLE</th>
</tr>
</thead>
<tbody>
<tr>
<td>3-wire lead set</td>
<td>I, II, III</td>
</tr>
<tr>
<td>5-wire lead set</td>
<td>I, II, III, aVR, aVL, aVF, V</td>
</tr>
<tr>
<td>6-wire lead set</td>
<td>I, II, III, aVR, aVL, aVF, Vx, Vy</td>
</tr>
<tr>
<td>View 12™ card</td>
<td>I, II, III, aVR, aVL, aVF, V1, V2, V3, V4, V5, V6</td>
</tr>
</tbody>
</table>

Unit of Measure: mV
Heart Rate Source: IBP

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (bpm)</th>
<th>MAXIMUM VALUE (bpm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>30</td>
<td>350</td>
</tr>
<tr>
<td></td>
<td>20 (V 12/V 21 and Telepack only)</td>
<td></td>
</tr>
<tr>
<td>Pediatric</td>
<td>30</td>
<td>300</td>
</tr>
<tr>
<td></td>
<td>20 (V 12/V 21 and Telepack only)</td>
<td></td>
</tr>
<tr>
<td>Adult</td>
<td>30</td>
<td>300</td>
</tr>
<tr>
<td></td>
<td>20 (V 12/V 21 and Telepack only)</td>
<td></td>
</tr>
</tbody>
</table>

12.1.2 Respiration: Resp

The Respiration parameter obtains data from ECG, CO2 or Gas Module sources.

Unit of Measure: Respirations Per Minute (rpm)
Resolution: 1 rpm

Respiration Source: ECG Thoracic Impedance

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (rpm)</th>
<th>MAXIMUM VALUE (rpm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>4</td>
<td>199</td>
</tr>
<tr>
<td>Pediatric</td>
<td>4</td>
<td>199</td>
</tr>
<tr>
<td>Adult</td>
<td>4</td>
<td>199</td>
</tr>
</tbody>
</table>

Respiration Source: CO2

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (rpm)</th>
<th>MAXIMUM VALUE (rpm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>0</td>
<td>150</td>
</tr>
<tr>
<td>Pediatric</td>
<td>0</td>
<td>150</td>
</tr>
<tr>
<td>Adult</td>
<td>0</td>
<td>150</td>
</tr>
</tbody>
</table>

Respiration Source: Gas Module

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (rpm)</th>
<th>MAXIMUM VALUE (rpm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>4</td>
<td>60</td>
</tr>
<tr>
<td>Pediatric</td>
<td>4</td>
<td>60</td>
</tr>
<tr>
<td>Adult</td>
<td>4</td>
<td>60</td>
</tr>
</tbody>
</table>

The Panorama Central Station supports respiration waveforms.

12.1.3 NIBP

The NIBP parameter includes measurements for the Systolic (Sys), Diastolic (Dia), Mean and Elapsed Time (ET).

Unit of Measure: mmHg
Resolution: 1 mmHg
ET

Unit of Measure: minute
Resolution: 1 minute

NIBP (Sys)

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (mmHg)</th>
<th>MAXIMUM VALUE (mmHg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>45</td>
<td>120</td>
</tr>
<tr>
<td>Pediatric</td>
<td>55</td>
<td>160</td>
</tr>
<tr>
<td>Adult</td>
<td>55</td>
<td>235</td>
</tr>
</tbody>
</table>

NIBP (Dia)

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (mmHg)</th>
<th>MAXIMUM VALUE (mmHg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>20</td>
<td>100</td>
</tr>
<tr>
<td>Pediatric</td>
<td>30</td>
<td>150</td>
</tr>
<tr>
<td>Adult</td>
<td>30</td>
<td>200</td>
</tr>
</tbody>
</table>

NIBP Mean

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE</th>
<th>MAXIMUM VALUE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>mmHg greater than the NIBP (Dia) value</td>
<td>mmHg less than the NIBP (Sys) value</td>
</tr>
<tr>
<td>Pediatric</td>
<td>mmHg greater than the NIBP (Dia) value</td>
<td>mmHg less than the NIBP (Sys) value</td>
</tr>
<tr>
<td>Adult</td>
<td>mmHg greater than the NIBP (Dia) value</td>
<td>mmHg less than the NIBP (Sys) value</td>
</tr>
</tbody>
</table>

NIBP (ET)

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (MIN.)</th>
<th>MAXIMUM VALUE (MINUTES)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>0</td>
<td>999</td>
</tr>
<tr>
<td>Pediatric</td>
<td>0</td>
<td>999</td>
</tr>
<tr>
<td>Adult</td>
<td>0</td>
<td>999</td>
</tr>
</tbody>
</table>

12.1.4 SpO₂

The SpO₂ parameter includes measurements from Masimo SET® and Nellcor® SpO₂ sensors.

Unit of Measure: %
Resolution: 1%

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (%)</th>
<th>MAXIMUM VALUE (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>–</td>
<td>100</td>
</tr>
<tr>
<td>Pediatric</td>
<td>–</td>
<td>100</td>
</tr>
<tr>
<td>Adult</td>
<td>–</td>
<td>100</td>
</tr>
</tbody>
</table>

The Panorama Central Station supports SpO₂ waveforms.
12.1.5 Nonin SpO2
The Nonin SpO2 parameter includes measurements from Nonin sensors.

Unit of Measure: %
Resolution: 1%

The Panorama Central Station supports SpO2 waveforms.

12.1.6 IBP: mmHg
The Panorama Central Station supports up to four invasive blood pressure parameters. The measurements shown for each pressure include Systolic (Sys), Diastolic (Dia) and Mean.

**NOTE:** IBP parameter labels include IBP, Art, UA, LV, PA, CVP, ICP, LA, and RA. The label used is based on the label selected at the bedside monitor.

Resolution: 1 mmHg

**IBP (Sys)**

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (mmHg)</th>
<th>MAXIMUM VALUE (mmHg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>-30</td>
<td>300</td>
</tr>
<tr>
<td>Pediatric</td>
<td>-30</td>
<td>300</td>
</tr>
<tr>
<td>Adult</td>
<td>-30</td>
<td>300</td>
</tr>
</tbody>
</table>

**IBP (Dia)**

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (mmHg)</th>
<th>MAXIMUM VALUE (mmHg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>-30</td>
<td>300</td>
</tr>
<tr>
<td>Pediatric</td>
<td>-30</td>
<td>300</td>
</tr>
<tr>
<td>Adult</td>
<td>-30</td>
<td>300</td>
</tr>
</tbody>
</table>

**IBP Mean**

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (mmHg)</th>
<th>MAXIMUM VALUE (mmHg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>-30</td>
<td>300</td>
</tr>
<tr>
<td>Pediatric</td>
<td>-30</td>
<td>300</td>
</tr>
<tr>
<td>Adult</td>
<td>-30</td>
<td>300</td>
</tr>
</tbody>
</table>

The Panorama Central Station supports IBP waveforms. The Panorama Central Station will not display an IBP waveform if the IBP cable is not connected to the system.
12.1.7 CO₂

The Panorama Central Station supports the mmHg, kPa and % measurements for CO₂.

**NOTE:** The unit of measure shown for CO₂ is based on the selection made at the bedside monitor.

Units of Measure:       mmHg, %, kPa
Resolution:            1 mmHg, 1 %, 1 kPa

**INSP/ET CO₂**

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (mmHg)</th>
<th>MAXIMUM VALUE (mmHg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>0</td>
<td>99</td>
</tr>
<tr>
<td>Pediatric</td>
<td>0</td>
<td>99</td>
</tr>
<tr>
<td>Adult</td>
<td>0</td>
<td>99</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (%)</th>
<th>MAXIMUM VALUE (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>0</td>
<td>13</td>
</tr>
<tr>
<td>Pediatric</td>
<td>0</td>
<td>13</td>
</tr>
<tr>
<td>Adult</td>
<td>0</td>
<td>13</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (KPA)</th>
<th>MAXIMUM VALUE (KPA)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>0</td>
<td>13</td>
</tr>
<tr>
<td>Pediatric</td>
<td>0</td>
<td>13</td>
</tr>
<tr>
<td>Adult</td>
<td>0</td>
<td>13</td>
</tr>
</tbody>
</table>

The Panorama Central Station supports a CO₂ waveform.

12.1.8 O₂

The Panorama Central Station supports INSP/ET O₂.

Unit of Measure:       %
Resolution:            1 %

**INSP/ET O₂**

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (%)</th>
<th>MAXIMUM VALUE (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>0</td>
<td>99</td>
</tr>
<tr>
<td>Pediatric</td>
<td>0</td>
<td>99</td>
</tr>
<tr>
<td>Adult</td>
<td>0</td>
<td>99</td>
</tr>
</tbody>
</table>

The Panorama Central Station supports an O₂ waveform.
12.1.9  **N₂O**  
The Panorama Central Station supports INSP/ET N₂O.

Unit of Measure: %  
Resolution: 1 %

**INSP/ET N₂O**

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (%)</th>
<th>MAXIMUM VALUE (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>0</td>
<td>99</td>
</tr>
<tr>
<td>Pediatric</td>
<td>0</td>
<td>99</td>
</tr>
<tr>
<td>Adult</td>
<td>0</td>
<td>100</td>
</tr>
</tbody>
</table>

12.1.10 **Anesthetic Agents: Iso, Enf, Des, Sev, Hal, Agent**  
The Panorama Central Station supports the INSP and ET measurements of the Isoflurane (Iso), Enflurane (Enf), Desflurane (Des), Sevoflurane (Sev), and Halothane (Hal) anesthetic agents.

**NOTE:** The Panorama Central Station will display general Agent values when the Gas Module cannot identify the agent.

The Anesthetic Agent label used is based on the labeling provided by the bedside monitor.

Unit of Measure: %  
Resolution: 1 %

**Iso, Enf and Hal INSP/ET Ranges**

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (%)</th>
<th>MAXIMUM VALUE (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>0</td>
<td>Not Applicable</td>
</tr>
<tr>
<td>Pediatric</td>
<td>0</td>
<td>Not Applicable</td>
</tr>
<tr>
<td>Adult</td>
<td>0</td>
<td>Not Applicable</td>
</tr>
</tbody>
</table>

**Des INSP/ET Ranges**

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (%)</th>
<th>MAXIMUM VALUE (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>0</td>
<td>Not Applicable</td>
</tr>
<tr>
<td>Pediatric</td>
<td>0</td>
<td>Not Applicable</td>
</tr>
<tr>
<td>Adult</td>
<td>0</td>
<td>Not Applicable</td>
</tr>
</tbody>
</table>

**Sev INSP/ET Ranges**

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (%)</th>
<th>MAXIMUM VALUE (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>0</td>
<td>Not Applicable</td>
</tr>
<tr>
<td>Pediatric</td>
<td>0</td>
<td>Not Applicable</td>
</tr>
<tr>
<td>Adult</td>
<td>0</td>
<td>Not Applicable</td>
</tr>
</tbody>
</table>
• The Panorama Central Station supports Iso, Enf, Sev, Hal and Des waveforms.
• The Panorama Central Station displays a Generic Agent waveform when the Gas Module cannot identify a specific agent.

12.1.11 Premature Ventricular Contraction (PVC)
The Panorama Central Station supports PVC data.

Unit of Measure: PVC/min
Resolution: 1 PVC/min

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>LEADS</th>
<th>MINIMUM VALUE</th>
<th>MAXIMUM VALUE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>3 lead, 5 lead, or 12 lead</td>
<td>Not Applicable</td>
<td>Not Applicable</td>
</tr>
<tr>
<td>Pediatric</td>
<td>3 lead, 5 lead, or 12 lead</td>
<td>0 PVC/min</td>
<td>Less than the Current HR value (PVCs/min)</td>
</tr>
<tr>
<td>Adult</td>
<td>3 lead, 5 lead, or 12 lead</td>
<td>0 PVC/min</td>
<td>Less than the Current HR value (PVCs/min)</td>
</tr>
</tbody>
</table>

12.1.12 QT/QTc
The Panorama Central Station supports QT/QTc data on the V 12/V 21 and Telepack.

Unit of Measure: ms
Resolution: 2 ms

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (MS)</th>
<th>MAXIMUM VALUE (MS)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>Not Applicable</td>
<td>Not Applicable</td>
</tr>
<tr>
<td>Pediatric</td>
<td>Not Applicable (Telepack) 300 (V 12/V 21)</td>
<td>Not Applicable (Telepack) 600 (V 12/V 21)</td>
</tr>
<tr>
<td>Adult</td>
<td>300</td>
<td>600</td>
</tr>
</tbody>
</table>

NOTE: The QT/QTc value consistently displays even if the measurement is out of the range.

NOTE: No alarm is provided when the measurement is lower than the minimum value.

12.1.13 Pause per Minute (Pause/min)
The Panorama Central Station supports Pause/min data.

Unit of Measure: Pause/min
Resolution: 1 PVC/min

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>LEADS</th>
<th>MINIMUM VALUE</th>
<th>MAXIMUM VALUE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pediatric</td>
<td>3 lead, 5 lead, or 12 lead</td>
<td>0 Pause/min</td>
<td>200 Pause/min</td>
</tr>
<tr>
<td>Adult</td>
<td>3 lead, 5 lead, or 12 lead</td>
<td>0 PVC/min</td>
<td>200 Pause/min</td>
</tr>
</tbody>
</table>
12.1.14 Temp: T1, T2, T3, Delta T, and T Blood
The Panorama Central Station supports five temperature measurements.

Units of Measure: F° and C°
Resolution: 0.1 F° and 0.1 C°

T1, T2, and T3

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (F°)</th>
<th>MAXIMUM VALUE (F°)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>59</td>
<td>113</td>
</tr>
<tr>
<td>Pediatric</td>
<td>59</td>
<td>113</td>
</tr>
<tr>
<td>Adult</td>
<td>59</td>
<td>113</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (C°)</th>
<th>MAXIMUM VALUE (C°)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>15</td>
<td>45</td>
</tr>
<tr>
<td>Pediatric</td>
<td>15</td>
<td>45</td>
</tr>
<tr>
<td>Adult</td>
<td>15</td>
<td>45</td>
</tr>
</tbody>
</table>

NOTE: T3 is currently supported by the V 12/V 21 only.

Delta T Values

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (F°)</th>
<th>MAXIMUM VALUE (F°)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>0</td>
<td>9.9</td>
</tr>
<tr>
<td>Pediatric</td>
<td>0</td>
<td>9.9</td>
</tr>
<tr>
<td>Adult</td>
<td>0</td>
<td>9.9</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (C°)</th>
<th>MAXIMUM VALUE (C°)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>0</td>
<td>5.5</td>
</tr>
<tr>
<td>Pediatric</td>
<td>0</td>
<td>5.5</td>
</tr>
<tr>
<td>Adult</td>
<td>0</td>
<td>5.5</td>
</tr>
</tbody>
</table>

T Blood Values

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (F°)</th>
<th>MAXIMUM VALUE (F°)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>Not Applicable</td>
<td>Not Applicable</td>
</tr>
<tr>
<td>Pediatric</td>
<td>63.7</td>
<td>109.1</td>
</tr>
<tr>
<td>Adult</td>
<td>63.7</td>
<td>109.1</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (C°)</th>
<th>MAXIMUM VALUE (C°)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>Not Applicable</td>
<td>Not Applicable</td>
</tr>
<tr>
<td>Pediatric</td>
<td>17.5</td>
<td>43</td>
</tr>
<tr>
<td>Adult</td>
<td>17.5</td>
<td>43</td>
</tr>
</tbody>
</table>
12.1.15 **ST**

The ST parameter provides an ST measurement using ECG leads to acquire data.

**PARAMETER LABELS**

<table>
<thead>
<tr>
<th>Lead Set</th>
<th>Label</th>
</tr>
</thead>
<tbody>
<tr>
<td>3-wire lead set</td>
<td>ST - I, ST - II, ST - III</td>
</tr>
<tr>
<td>5-wire lead set</td>
<td>ST - I, ST - II, ST - III, ST - aVR, ST - aVL, ST - aVF, ST - V</td>
</tr>
<tr>
<td>6-wire lead set</td>
<td>ST - I, ST - II, ST - III, ST - aVR, ST - aVL, ST - aVF, ST - Vx, ST - Vy</td>
</tr>
</tbody>
</table>

| Unit of Measure: | mm |
| Resolution:     | 0.1 mm |

<table>
<thead>
<tr>
<th>Patient Size</th>
<th>MINIMUM VALUE (MM)</th>
<th>MAXIMUM VALUE (MM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>Not Applicable</td>
<td>Not Applicable</td>
</tr>
<tr>
<td>Pediatric</td>
<td>-10.0</td>
<td>10.0</td>
</tr>
<tr>
<td>Adult</td>
<td>-10.0</td>
<td>10.0</td>
</tr>
</tbody>
</table>

**NOTE:** A (C) next to an ST measurement indicates that the value represents a Current measurement.

**NOTE:** An (R) next to an ST measurement indicates that the value represents a Reference measurement.

12.1.16 **Cardiac Output**

Unit of Measure: lpm

Resolution: 0.1 lpm

<table>
<thead>
<tr>
<th>Patient Size</th>
<th>MINIMUM VALUE (LPM)</th>
<th>MAXIMUM VALUE (LPM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adult</td>
<td>0.2</td>
<td>20.0</td>
</tr>
</tbody>
</table>

12.1.17 **Cardiac Index**

Unit of Measure: lpm/m2

Resolution: 0.1 lpm/m2

<table>
<thead>
<tr>
<th>Patient Size</th>
<th>MINIMUM VALUE (LPM/M2)</th>
<th>MAXIMUM VALUE (LPM/M2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adult</td>
<td>0.1</td>
<td>20.0</td>
</tr>
</tbody>
</table>
12.1.18 Continuous Cardiac Output (CCO)

Unit of Measure: lpm
Resolution: 0.1 lpm

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (LPM)</th>
<th>MAXIMUM VALUE (LPM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pediatric</td>
<td>0.2 (non-DPM)</td>
<td>25.0 (non-DPM)</td>
</tr>
<tr>
<td></td>
<td>1.0 (DPM)</td>
<td>20.0 (DPM)</td>
</tr>
<tr>
<td>Adult</td>
<td>0.2 (non-DPM)</td>
<td>25.0 (non-DPM)</td>
</tr>
<tr>
<td></td>
<td>1.0 (DPM)</td>
<td>20.0 (DPM)</td>
</tr>
</tbody>
</table>

12.1.19 Continuous Cardiac Output STAT (CCO STAT)

Unit of Measure: lpm
Resolution: 0.1 lpm

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (LPM)</th>
<th>MAXIMUM VALUE (LPM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pediatric</td>
<td>0.2 (non-DPM)</td>
<td>25.0 (non-DPM)</td>
</tr>
<tr>
<td></td>
<td>1.0 (DPM)</td>
<td>20.0 (DPM)</td>
</tr>
<tr>
<td>Adult</td>
<td>0.2 (non-DPM)</td>
<td>25.0 (non-DPM)</td>
</tr>
<tr>
<td></td>
<td>1.0 (DPM)</td>
<td>20.0 (DPM)</td>
</tr>
</tbody>
</table>

12.1.20 Continuous Cardiac Index (CCI)

Unit of Measure: lpm
Resolution: 0.1 lpm

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (LPM)</th>
<th>MAXIMUM VALUE (LPM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pediatric</td>
<td>0.2</td>
<td>25.0</td>
</tr>
<tr>
<td>Adult</td>
<td>0.2</td>
<td>25.0</td>
</tr>
</tbody>
</table>

12.1.21 Continuous Cardiac Index STAT (CCI STAT)

Unit of Measure: lpm
Resolution: 0.1 lpm

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (LPM)</th>
<th>MAXIMUM VALUE (LPM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pediatric</td>
<td>0.2</td>
<td>25.0</td>
</tr>
<tr>
<td>Adult</td>
<td>0.2</td>
<td>25.0</td>
</tr>
</tbody>
</table>
### 12.1.22 System Vascular Resistance (SVR)

- **Unit of Measure:** dynes*sec/cm^5
- **Resolution:** 1 dynes*sec/cm^5

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (DYNES*SEC/CM^5)</th>
<th>MAXIMUM VALUE (DYNES*SEC/CM^5)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pediatric</td>
<td>0</td>
<td>3000</td>
</tr>
<tr>
<td>Adult</td>
<td>0</td>
<td>3000</td>
</tr>
</tbody>
</table>

### 12.1.23 System Vascular Resistance Index (SVRI)

- **Unit of Measure:** dynes*sec/cm^5/m^2
- **Resolution:** 1 dynes*sec/cm^5/m^2

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (DYNES*SEC/CM^5/M^2)</th>
<th>MAXIMUM VALUE (DYNES*SEC/CM^5/M^2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pediatric</td>
<td>0</td>
<td>6000</td>
</tr>
<tr>
<td>Adult</td>
<td>0</td>
<td>6000</td>
</tr>
</tbody>
</table>

### 12.1.24 Mixed Venous Saturation (SvO$_2$)

- **Unit of Measure:** %
- **Resolution:** 1%

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (%)</th>
<th>MAXIMUM VALUE (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pediatric</td>
<td>0</td>
<td>99</td>
</tr>
<tr>
<td>Adult</td>
<td>0</td>
<td>99</td>
</tr>
</tbody>
</table>

### 12.1.25 Central Venous Oxygen Saturation (ScvO$_2$)

- **Unit of Measure:** %
- **Resolution:** 1%

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (%)</th>
<th>MAXIMUM VALUE (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pediatric</td>
<td>10</td>
<td>99</td>
</tr>
<tr>
<td>Adult</td>
<td>10</td>
<td>99</td>
</tr>
</tbody>
</table>
12.1.26 Pulmonary Artery Wedge Pressure (PAWP)

Unit of Measure: mmHg
Resolution: 1 mmHg

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE (mmHg)</th>
<th>MAXIMUM VALUE (mmHg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pediatric</td>
<td>-30</td>
<td>300</td>
</tr>
<tr>
<td>Adult</td>
<td>-30</td>
<td>300</td>
</tr>
</tbody>
</table>

12.1.27 Bispectral Index (BIS/BIS L/BIS R)

The Panorama Central Station supports BIS data on the V 12/V 21.

Unit of Measure: Not Applicable
Resolution: 1

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE</th>
<th>MAXIMUM VALUE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>Not Applicable</td>
<td>Not Applicable</td>
</tr>
<tr>
<td>Pediatric</td>
<td>0</td>
<td>100</td>
</tr>
<tr>
<td>Adult</td>
<td>0</td>
<td>100</td>
</tr>
</tbody>
</table>

12.1.28 Battery Percentage

Unit of Measure: %
Resolution: 1 %

<table>
<thead>
<tr>
<th>PATIENT SIZE</th>
<th>MINIMUM VALUE</th>
<th>MAXIMUM VALUE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adult</td>
<td>5 %</td>
<td>100 %</td>
</tr>
</tbody>
</table>
12.2 Invalid Data Display

The Panorama Central Station uses dashes (---), Xs (XX) or a straight line (waveform) to show invalid data.

Dashes (---) Digital Data
Digital data is replaced with dashes (---) for the following reasons:

- Data is above the maximum value
- Data is below the minimum value
- Parameter is not available
- Lead or a sensor is off
- Sensor is not zeroed
- Parameter is zeroing
- Measurement has not been taken
- Measurement is being taken
- Measurement has timed out
- Measurement is being retried
- Parameter is turned off
- View 12™ card is relearning
- PVC/min counter reset
- Gas module or the source agent is occluded
- Gas module or the source agent is purging
- Gas module is turned off
- ST license is not available
- Learning ST Values
- SQI is below 15%

Xs (XX) Digital Data
Digital data is replaced with Xs (XX) for the following reasons:

- IBP parameter is not zeroed
- NIBP parameter failed to obtain a reading
Straight Line (Waveforms)
Waveform data is replaced with a continuous straight line for the following reasons:

- ECG leads are disconnected
- ECG cable is not connected to the system
- Respiration parameter is disabled at the bedside monitor and the source is ECG
- Respiration parameter is using a View 12™ card at the bedside monitor and the source is ECG
- SpO₂ cable is connected but not in use
- IBP cable is connected but not zeroed or is zeroing
- CO₂ filter line is not in use or is zeroing
- O₂ parameter is not in use
- Gas module is connected but the agent is not in use
- Gas module is connected but the agent is not detected
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13.0 **Alarms and Events**

The alarms available at the Panorama Central Station are dependent on the patient monitoring devices in use and on how the alarm options are configured at the Panorama Central Station.

This section outlines the alarm types and briefly describes the events that cause alarm conditions.

- Physiological Alarms and Events
- Physiological Alarms and Events: Arrhythmias
- Technical Alarms and Events
- System Event Messages and Error Codes
13.1 Physiological Alarms and Events
The Panorama Central Station uses physiological alarms to indicate that a patient’s physiological alarm thresholds have been violated.

13.1.1 Alarm Limit Settings
This section identifies high and low alarm limit settings, and available alarm priorities for each physiological alarm, by patient size. It also differentiates which alarm limit settings are available at each monitoring device.

**WARNING:** Setting alarm limits to extreme values can render the alarm system useless.
## Alarms and Events

### Physiological Alarms and Events

#### Heart Rate Units: bpm

<table>
<thead>
<tr>
<th>ALARM</th>
<th>ALARM PRIORITY</th>
<th>ALARM LIMIT</th>
<th>NEONATE</th>
<th>PEDIATRIC</th>
<th>ADULT</th>
<th>DEVICES</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>High Limit</strong></td>
<td>1 or 2</td>
<td>Off, 100-350</td>
<td>Off, 100-300</td>
<td>Off, 100-180</td>
<td>Off, 60-250</td>
<td>Passport 2, Passport V Spectrum/Spectrum OR, Telepack</td>
</tr>
<tr>
<td><strong>Low Limit</strong></td>
<td>1 or 2</td>
<td>Off, 30-200</td>
<td>Off, 30-150</td>
<td>Off, 30-80</td>
<td>Off, 30-120</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Factory Default:</td>
<td></td>
<td>Factory Default:</td>
<td>Factory Default:</td>
<td></td>
</tr>
</tbody>
</table>

**Factory Default:**
- High Limit: 2
- Low Limit: 2

**NOTE:** When the central station language is configured for French, the Heart Rate (HR) high/low alarm limit cannot be turned Off.

**NOTE:** When the selected device is a Telepack, the Low Heart Rate alarm will be called when the heart rate becomes invalid because the rate is underrange.

#### Extreme Bradycardia

**Units: bpm**

<table>
<thead>
<tr>
<th>ALARM</th>
<th>ALARM PRIORITY</th>
<th>ALARM LIMIT</th>
<th>NEONATE</th>
<th>PEDIATRIC</th>
<th>ADULT</th>
<th>DEVICES</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>High Limit</strong></td>
<td>1 or 2</td>
<td>Off, 25-200</td>
<td>N/A</td>
<td>N/A</td>
<td>Off, 60-250</td>
<td>Telepack with Nonin SpO2 module</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Factory Default:</td>
<td></td>
<td></td>
<td>Factory Default:150</td>
<td></td>
</tr>
<tr>
<td><strong>Low Limit</strong></td>
<td>1 or 2</td>
<td>Off, 25-200</td>
<td>N/A</td>
<td>N/A</td>
<td>Off, 25-120</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Factory Default:</td>
<td></td>
<td></td>
<td>Factory Default:50</td>
<td></td>
</tr>
</tbody>
</table>

**Factory Default:**
- High Limit: 1
- Low Limit: 1

**NOTE:** The Resp tile will flash when apnea is occurring and the respiration source is ECG.

#### Extreme Tachycardia

**Units: bpm**

<table>
<thead>
<tr>
<th>ALARM</th>
<th>ALARM PRIORITY</th>
<th>ALARM LIMIT</th>
<th>NEONATE</th>
<th>PEDIATRIC</th>
<th>ADULT</th>
<th>DEVICES</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>High Limit</strong></td>
<td>1 or 2</td>
<td>Off, 50-350</td>
<td>N/A</td>
<td>N/A</td>
<td>Off, 60-300</td>
<td>Telepack</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Factory Default:</td>
<td></td>
<td></td>
<td>Factory Default:165</td>
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<tr>
<td><strong>Low Limit</strong></td>
<td>1 or 2</td>
<td>Off, 50-200</td>
<td>N/A</td>
<td>N/A</td>
<td>Off, 25-120</td>
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<tr>
<td></td>
<td></td>
<td>Factory Default:</td>
<td></td>
<td></td>
<td>Factory Default:45</td>
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</tr>
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</table>

**Factory Default:**
- High Limit: 1
- Low Limit: 1

**NOTE:** The Resp tile will flash when apnea is occurring and the respiration source is ECG.

N/A - Not Applicable
<table>
<thead>
<tr>
<th>ALARM</th>
<th>ALARM PRIORITY</th>
<th>ALARM LIMIT</th>
<th>NEONATE</th>
<th>PEDIATRIC</th>
<th>ADULT</th>
<th>DEVICES</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Respiration</strong></td>
<td>1, 2, and 3</td>
<td>High Limit</td>
<td>Off, 30-200</td>
<td>Off, 15-150</td>
<td>Off, 10-100</td>
<td>Passport 2, Passport V, Spectrum/Spectrum OR</td>
</tr>
<tr>
<td>Units: rpm</td>
<td>Factory Default: 3</td>
<td>Off, 5-50</td>
<td>Factory Default: 5</td>
<td>Off, 5-30</td>
<td>Off, 5-50</td>
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<tr>
<td></td>
<td>Low Limit</td>
<td>Off, 2-150</td>
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<td>Off, 2-100</td>
<td>Factory Default: 100</td>
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<tr>
<td></td>
<td>Factory Default: 3</td>
<td>Off, 0-148</td>
<td>Factory Default: 30</td>
<td>Off, 0-98</td>
<td>Factory Default: 8</td>
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<td></td>
<td>Low Limit</td>
<td>Off, 10-190</td>
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<td>Off, 10-190</td>
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<td>Factory Default: 3</td>
<td>Off, 5-50</td>
<td>Factory Default: Off</td>
<td>Off, 5-50</td>
<td>Factory Default: Off</td>
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<tr>
<td><strong>SpO2</strong></td>
<td>1 and 2</td>
<td>High Limit</td>
<td>Off, 80-100%</td>
<td>Off, 80-100%</td>
<td>Off, 80-100%</td>
<td>Passport 2, Passport V, Spectrum/Spectrum OR</td>
</tr>
<tr>
<td>Units: %</td>
<td>Factory Default: 2</td>
<td>Off, 50-99%</td>
<td>Factory Default: 92%</td>
<td>50-99%</td>
<td>Factory Default: 85%</td>
<td></td>
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<tr>
<td></td>
<td>Low Limit</td>
<td>Off, (Desat+2)-100%</td>
<td>Factory Default: 95%</td>
<td>Off, (Desat+2)-100%</td>
<td>Factory Default: Off</td>
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<tr>
<td></td>
<td>Factory Default: 2</td>
<td>Off, Desat-98%</td>
<td>Factory Default: 90%</td>
<td>Desat-98%</td>
<td>Factory Default: 90%</td>
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</tr>
<tr>
<td></td>
<td>Low Limit</td>
<td>Off, 75-101%</td>
<td>Factory Default: 101%</td>
<td>Off, 75-101%</td>
<td>Factory Default: 101%</td>
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<tr>
<td></td>
<td>Factory Default: 2</td>
<td>Off, 70-99%</td>
<td>Factory Default: 92%</td>
<td>70-99%</td>
<td>Factory Default: 88%</td>
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<tr>
<td><strong>SpO2 (Nonin)</strong></td>
<td>1 and 2</td>
<td>High Limit</td>
<td>N/A</td>
<td>N/A</td>
<td>Off, 80-100%</td>
<td>Telepack with Nonin SpO2 module</td>
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<td>Units: %</td>
<td>Factory Default: 2</td>
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<td>N/A</td>
<td>50-99%</td>
<td>Factory Default: 85%</td>
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<tr>
<td></td>
<td>Low Limit</td>
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<td></td>
<td>N/A</td>
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</tr>
<tr>
<td><strong>IBPx: Systolic</strong></td>
<td>2 or 3</td>
<td>High Limit</td>
<td>Off, 5-180</td>
<td>Off, 5-240</td>
<td>Off, 5-300</td>
<td>Passport 2, Passport V, Spectrum/Spectrum OR</td>
</tr>
<tr>
<td>Units: mmHg*</td>
<td>Factory Default: 3</td>
<td>Off, 0-130</td>
<td>Factory Default: Off</td>
<td>Off, 0-130</td>
<td>Off, 0-150</td>
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</tr>
</tbody>
</table>

**NOTE:** The SpO2 high/low alarm limit will always be displayed in the digital tile portion of the Bedside tab All Waveforms view.
### Physiological Alarms and Events

#### IBPx Diastolic
*Units: mmHg*

<table>
<thead>
<tr>
<th>Alarm</th>
<th>Alarm Priority</th>
<th>Alarm Limit</th>
<th>Neonate</th>
<th>Pediatric</th>
<th>Adult</th>
<th>Devices</th>
</tr>
</thead>
<tbody>
<tr>
<td>IBPx Diastolic</td>
<td>2 or 3</td>
<td>High Limit</td>
<td>Off, 0-70</td>
<td>Off, 0-100</td>
<td>Off, 0-140</td>
<td>Passport 2, Passport V, Spectrum/Spectrum OR</td>
</tr>
<tr>
<td></td>
<td>Low Limit</td>
<td>Off, 0-50</td>
<td>Factory Default: Off</td>
<td>Off, 0-100</td>
<td>Off, 0-120</td>
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<tr>
<td></td>
<td>Factory Default: Off</td>
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<td></td>
<td>Factory Default: Off</td>
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<td></td>
</tr>
</tbody>
</table>

#### IBPx Mean
*Units: mmHg*

<table>
<thead>
<tr>
<th>Alarm</th>
<th>Alarm Priority</th>
<th>Alarm Limit</th>
<th>Neonate</th>
<th>Pediatric</th>
<th>Adult</th>
<th>Devices</th>
</tr>
</thead>
<tbody>
<tr>
<td>IBPx Mean</td>
<td>2 or 3</td>
<td>High Limit</td>
<td>Off, 5-100</td>
<td>Off, 5-100</td>
<td>Off, 5-150</td>
<td>Passport 2, Passport V, Spectrum/Spectrum OR</td>
</tr>
<tr>
<td></td>
<td>Low Limit</td>
<td>Off, 2-50</td>
<td>Factory Default: Off</td>
<td>Off, 2-50</td>
<td>Off, 2-100</td>
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<tr>
<td></td>
<td>Factory Default: Off</td>
<td></td>
<td></td>
<td>Factory Default: Off</td>
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</tr>
</tbody>
</table>

*All IBPx alarm limits apply to IBP1, IBP2, IBP3, IBP4, ART, UA, LV, PA, CVP, ICP, LA, and RA. The Panorama Central Station is capable of monitoring up to four IBP measurements.*

**NOTE:** x = 1, 2, 3 or 4 (IBP3 and 4 only available if Spectrum/Spectrum OR External Parameter Module (EPM) is installed)

#### IBP1-8 Systolic
*Units: mmHg*

<table>
<thead>
<tr>
<th>Alarm</th>
<th>Alarm Priority</th>
<th>Alarm Limit</th>
<th>Neonate</th>
<th>Pediatric</th>
<th>Adult</th>
<th>Devices</th>
</tr>
</thead>
<tbody>
<tr>
<td>IBP1-8 Systolic</td>
<td>2 or 3</td>
<td>High Limit</td>
<td>Off, 5-300</td>
<td>Off, 5-300</td>
<td>Off, 5-300</td>
<td>V 12/V 21</td>
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<tr>
<td></td>
<td>Low Limit</td>
<td>Off, 0-150</td>
<td>Factory Default: Off</td>
<td>Off, 0-150</td>
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<td>Factory Default: Off</td>
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</table>

#### IBP1-8 Diastolic
*Units: mmHg*

<table>
<thead>
<tr>
<th>Alarm</th>
<th>Alarm Priority</th>
<th>Alarm Limit</th>
<th>Neonate</th>
<th>Pediatric</th>
<th>Adult</th>
<th>Devices</th>
</tr>
</thead>
<tbody>
<tr>
<td>IBP1-8 Diastolic</td>
<td>2 or 3</td>
<td>High Limit</td>
<td>Off, 5-200</td>
<td>Off, 5-200</td>
<td>Off, 5-200</td>
<td>V 12/V 21</td>
</tr>
<tr>
<td></td>
<td>Low Limit</td>
<td>Off, 0-150</td>
<td>Factory Default: Off</td>
<td>Off, 0-150</td>
<td>Off, 0-150</td>
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</tr>
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<td>Factory Default: Off</td>
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<td>Factory Default: Off</td>
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</tbody>
</table>

#### IBP1-8 Mean
*Units: mmHg*

<table>
<thead>
<tr>
<th>Alarm</th>
<th>Alarm Priority</th>
<th>Alarm Limit</th>
<th>Neonate</th>
<th>Pediatric</th>
<th>Adult</th>
<th>Devices</th>
</tr>
</thead>
<tbody>
<tr>
<td>IBP1-8 Mean</td>
<td>2 or 3</td>
<td>High Limit</td>
<td>Off, 5-200</td>
<td>Off, 5-200</td>
<td>Off, 5-200</td>
<td>V 12/V 21</td>
</tr>
<tr>
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<td>Low Limit</td>
<td>Off, 0-150</td>
<td>Factory Default: Off</td>
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<td>Off, 0-150</td>
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<tr>
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<td>Factory Default: Off</td>
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<td>Factory Default: Off</td>
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<td></td>
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</tbody>
</table>

#### Art, UAP, LV, PA, CVP, ICP, LAP, RAP, Ao, BAP, FAP, UVP Systolic
*Units: mmHg*

<table>
<thead>
<tr>
<th>Alarm</th>
<th>Alarm Priority</th>
<th>Alarm Limit</th>
<th>Neonate</th>
<th>Pediatric</th>
<th>Adult</th>
<th>Devices</th>
</tr>
</thead>
<tbody>
<tr>
<td>Art, UAP, LV, PA, CVP, ICP, LAP, RAP, Ao, BAP, FAP, UVP Systolic</td>
<td>2 or 3</td>
<td>High Limit</td>
<td>Off, -48-300</td>
<td>Off, -48-300</td>
<td>Off, -48-300</td>
<td>DPM 6/DPM 7, Passport 12M/Passport 17M, Passport 8/Passport 12, T1 monitor</td>
</tr>
<tr>
<td></td>
<td>Factory Default: Off</td>
<td></td>
<td></td>
<td>Factory Default: Off</td>
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<td></td>
</tr>
</tbody>
</table>

#### Art, UAP, LV, PA, CVP, ICP, LAP, RAP, Ao, BAP, FAP, UVP Diastolic
*Units: mmHg*

<table>
<thead>
<tr>
<th>Alarm</th>
<th>Alarm Priority</th>
<th>Alarm Limit</th>
<th>Neonate</th>
<th>Pediatric</th>
<th>Adult</th>
<th>Devices</th>
</tr>
</thead>
<tbody>
<tr>
<td>Art, UAP, LV, PA, CVP, ICP, LAP, RAP, Ao, BAP, FAP, UVP Diastolic</td>
<td>2 or 3</td>
<td>High Limit</td>
<td>Off, -48-300</td>
<td>Off, -48-300</td>
<td>Off, -48-300</td>
<td>DPM 6/DPM 7, Passport 12M/Passport 17M, Passport 8/Passport 12, T1 monitor</td>
</tr>
<tr>
<td></td>
<td>Factory Default: 3</td>
<td>Factory Default: 60</td>
<td>Factory Default: 70</td>
<td>Factory Default: 90</td>
<td>Factory Default: 90</td>
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</tr>
<tr>
<td></td>
<td>Low Limit</td>
<td>Off, -50-298</td>
<td>Factory Default: 40</td>
<td>Off, -50-298</td>
<td>Off, -50-298</td>
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<tr>
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<td>Factory Default: Off</td>
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<td></td>
<td>Factory Default: Off</td>
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</tbody>
</table>

*N/A - Not Applicable*
<table>
<thead>
<tr>
<th>ALARM</th>
<th>ALARM PRIORITY</th>
<th>ALARM LIMIT</th>
<th>NEONATE</th>
<th>PEDIATRIC</th>
<th>ADULT</th>
<th>DEVICES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Art, UAP, LV, PA, CVP, ICP, LAP, RAP, Ao, BAP, FAP, UVP Mean Units: mmHg</td>
<td>2 or 3</td>
<td>High Limit</td>
<td>Off, -48-300</td>
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<td>Off, -48-300</td>
<td>Factory Default: 110</td>
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<td>Factory Default: 70</td>
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<tr>
<td></td>
<td></td>
<td>Low Limit</td>
<td>Off, -50-298</td>
<td>Factory Default: 35</td>
<td>Off, -50-298</td>
<td>Factory Default: 70</td>
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<td>Factory Default: 35</td>
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<tr>
<td>NIBP Systolic Units: mmHg</td>
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<td>High Limit</td>
<td>Off, 40-180</td>
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<td>Off, 40-180</td>
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<tr>
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<td>Low Limit</td>
<td>Off, 15-130</td>
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<td>Factory Default: Off</td>
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<tr>
<td>NIBP Diastolic Units: mmHg</td>
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<td>Off, 10-50</td>
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<td>Factory Default: Off</td>
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<tr>
<td>NIBP Mean Units: mmHg</td>
<td>2 or 3</td>
<td>High Limit</td>
<td>Off, 40-160</td>
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<td>Off, 50-180</td>
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<td>Low Limit</td>
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<tr>
<td>NIBP Systolic Units: mmHg</td>
<td>2 or 3</td>
<td>High Limit</td>
<td>Off, 45-135</td>
<td>Factory Default: 90</td>
<td>Off, 45-200</td>
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<td>Factory Default: 90</td>
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<td>Off, 40-130</td>
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<td>Off, 40-195</td>
<td>Factory Default: 70</td>
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<td>NIBP Diastolic Units: mmHg</td>
<td>2 or 3</td>
<td>High Limit</td>
<td>Off, 15-100</td>
<td>Factory Default: 60</td>
<td>Off, 15-150</td>
<td>Factory Default: 90</td>
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<td>Off, 10-95</td>
<td>Factory Default: 20</td>
<td>Off, 10-145</td>
<td>Factory Default: 50</td>
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<tr>
<td>NIBP Mean Units: mmHg</td>
<td>2 or 3</td>
<td>High Limit</td>
<td>Off, 25-110</td>
<td>Factory Default: 70</td>
<td>Off, 25-165</td>
<td>Factory Default: 90</td>
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<tr>
<td>NIBP Systolic Units: mmHg</td>
<td>2 or 3</td>
<td>High Limit</td>
<td>Off, 30-100</td>
<td>Factory Default: Off</td>
<td>Off, 30-140</td>
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<td>Factory Default: Off</td>
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<td>Factory Default: Off</td>
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N/A - Not Applicable
<table>
<thead>
<tr>
<th>ALARM</th>
<th>ALARM PRIORITY</th>
<th>ALARM LIMIT</th>
<th>NEONATE</th>
<th>PEDIATRIC</th>
<th>ADULT</th>
<th>DEVICES</th>
</tr>
</thead>
<tbody>
<tr>
<td>NIBP Diastolic Units: mmHg</td>
<td>2 or 3</td>
<td>High Limit Off, 30-90 Factory Default: Off</td>
<td>Off, 30-120 Factory Default: Off</td>
<td>Off, 40-150 Factory Default: Off</td>
<td>V 12/V 21</td>
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</tr>
<tr>
<td>NIBP Mean Units: mmHg</td>
<td>2 or 3</td>
<td>High Limit Off, 30-90 Factory Default: Off</td>
<td>Off, 30-140 Factory Default: Off</td>
<td>Off, 60-200 Factory Default: Off</td>
<td>V 12/V 21</td>
<td></td>
</tr>
</tbody>
</table>

**NOTE:** The CO2 tile will flash when apnea is occurring and CO2 is the respiration source.

| INSP CO2 (mmHg)        | 2 or 3         | High Limit Off, 5-30 Factory Default: Off | Off, 5-30 Factory Default: Off | Off, 5-30 Factory Default: Off | Passport 2, Passport V, Spectrum/Spectrum OR |
|                       |                | Low Limit N/A | N/A | N/A | N/A |
|                       |                | Low Limit N/A | N/A | N/A | N/A |
| INSP CO2 (kPa)         | 2 or 3         | High Limit Off, 1-4 Factory Default: Off | Off, 1-4 Factory Default: Off | Off, 1-4 Factory Default: Off | Passport 2, Passport V, Spectrum/Spectrum OR |
|                       |                | Low Limit N/A | N/A | N/A | N/A |
| INSP CO2 (mmHg)        | 2 or 3         | High Limit Off, 5-30 Factory Default: 10 | Off, 5-30 Factory Default: 10 | Off, 5-30 Factory Default: 10 | V 12/V 21 |
|                       |                | Low Limit N/A | N/A | N/A | N/A |
| INSP CO2 (%)           | 2 or 3         | High Limit Off, 1-4 Factory Default: 1 | Off, 1-4 Factory Default: 1 | Off, 1-4 Factory Default: 1 | V 12/V 21 |
|                       |                | Low Limit N/A | N/A | N/A | N/A |
| INSP CO2 (kPa)         | 2 or 3         | High Limit Off, 1-4 Factory Default: 1 | Off, 1-4 Factory Default: 1 | Off, 1-4 Factory Default: 1 | V 12/V 21 |
|                       |                | Low Limit N/A | N/A | N/A | N/A |
| FiCO2 (mmHg)           | 2 or 3         | High Limit Off, 1-99 Factory Default: 4 | Off, 1-99 Factory Default: 4 | Off, 1-99 Factory Default: 4 | DPM 6/DPM 7, Passport 12M/Passport 17M, Passport 8/Passport 12, T1 monitor |
|                       |                | Low Limit N/A | N/A | N/A | N/A |

*N/A - Not Applicable*
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NOTE: Only available if Spectrum/Spectrum OR External Parameter Module (EPM) is installed and sensor is attached.

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<tr>
<th>ALARM</th>
<th>ALARM PRIORITY</th>
<th>NEONATE</th>
<th>PEDIATRIC</th>
<th>ADULT</th>
<th>DEVICES</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Low Limit</td>
<td>N/A</td>
<td>Off, 80-100°</td>
<td>Off, 80-100°</td>
</tr>
<tr>
<td></td>
<td>2 or 3 Factory Default: 3</td>
<td>High Limit</td>
<td>N/A</td>
<td>N/A</td>
<td>Off, 75-109°</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low Limit</td>
<td>N/A</td>
<td>N/A</td>
<td>Off, 74-108°</td>
</tr>
<tr>
<td>T Blood</td>
<td>2 or 3 Factory Default: 3</td>
<td>High Limit</td>
<td>N/A</td>
<td>Off, 35-43°</td>
<td>Off, 35-43°</td>
</tr>
<tr>
<td>Units: °C</td>
<td></td>
<td>Low Limit</td>
<td>N/A</td>
<td>Off, 26-38°</td>
<td>Off, 26-38°</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low Limit</td>
<td>N/A</td>
<td>Off, 24.0-43.0°</td>
<td>Off, 24.0-43.0°</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low Limit</td>
<td>N/A</td>
<td>Off, 23.0-42.0°</td>
<td>Off, 23.0-42.0°</td>
</tr>
<tr>
<td>ST Single</td>
<td>1, 2 or 3 Factory Default: 2</td>
<td>High Limit</td>
<td>N/A</td>
<td>Off, 0.50 to 10.0</td>
<td>Off, 0.50 to 10.0</td>
</tr>
<tr>
<td>Units: mm</td>
<td></td>
<td>Low Limit</td>
<td>N/A</td>
<td>Off, (0.5) to (-10.0)</td>
<td>Off, (0.5) to (-10.0)</td>
</tr>
<tr>
<td>ST Dual</td>
<td>1, 2 or 3 Factory Default: 2</td>
<td>High Limit</td>
<td>N/A</td>
<td>Off, 0.50-10.0</td>
<td>Off, 0.50-10.0</td>
</tr>
<tr>
<td>Units: mm</td>
<td></td>
<td>Low Limit</td>
<td>N/A</td>
<td>Off, (0.5) - (-10.0)</td>
<td>Off, (0.5) - (-10.0)</td>
</tr>
</tbody>
</table>

**NOTE:** Only available if Spectrum/Spectrum OR External Parameter Module (EPM) is installed and sensor is attached.
<table>
<thead>
<tr>
<th>ALARM</th>
<th>ALARM PRIORITY</th>
<th>ALARM LIMIT</th>
<th>NEONATE</th>
<th>PEDIATRIC</th>
<th>ADULT</th>
<th>DEVICES</th>
</tr>
</thead>
<tbody>
<tr>
<td>PVC</td>
<td>2 or 3 Factory Default: 3</td>
<td>High Limit</td>
<td>N/A</td>
<td>Off, 1-30 Factory Default: Off</td>
<td>Off, 1-30 Factory Default: Off</td>
<td>Passport 2, Passport V, Spectrum/Spectrum OR, Telepack</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low Limit</td>
<td>N/A</td>
<td>N/A Factory Default: 3</td>
<td>N/A Factory Default: 3</td>
<td></td>
</tr>
<tr>
<td>PVC</td>
<td>2 or 3 Factory Default: 3</td>
<td>High Limit</td>
<td>N/A</td>
<td>Off, 1-100 Factory Default: Off</td>
<td>Off, 1-100 Factory Default: Off</td>
<td>DPM 6/DPM 7, Passport 12M, Passport 17M, Passport 8, Passport 12, T1 monitor</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low Limit</td>
<td>N/A</td>
<td>N/A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Apnea Delay</td>
<td>2 or 3 Factory Default: 3</td>
<td>High Limit</td>
<td>10, 20 Factory Default: 15</td>
<td>10-20 Factory Default: 15</td>
<td>10-60 Factory Default: 60</td>
<td>Passport 2, Passport V, Spectrum/Spectrum OR</td>
</tr>
<tr>
<td>Units: Seconds</td>
<td></td>
<td>Low Limit</td>
<td>N/A Factory Default: 3</td>
<td>N/A Factory Default: 3</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low Limit</td>
<td>N/A N/A</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Apnea Delay</td>
<td>2 or 3 Factory Default: 3</td>
<td>High Limit</td>
<td>10-20 Factory Default: 15</td>
<td>10-20 Factory Default: 15</td>
<td>10-60 Factory Default: 30</td>
<td>V 12/V 21</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low Limit</td>
<td>N/A N/A</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>BIS/BIS R/BIS L</td>
<td>2 or 3 Factory Default: 3</td>
<td>High Limit</td>
<td>N/A</td>
<td>Off, 10-99 Factory Default: Off</td>
<td>Off, 10-99 Factory Default: Off</td>
<td>V 12/V 21</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low Limit</td>
<td>N/A</td>
<td>Off, 1-90 Factory Default: Off</td>
<td></td>
<td></td>
</tr>
<tr>
<td>QTc</td>
<td>2 or 3 Factory Default: 2</td>
<td>High Limit</td>
<td>N/A</td>
<td>Off, 300-600 Factory Default: Off</td>
<td>Off, 300-600 Factory Default: Off</td>
<td>V 12/V 21</td>
</tr>
<tr>
<td>Units: ms</td>
<td></td>
<td>Low Limit</td>
<td>N/A</td>
<td>N/A</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

N/A - Not Applicable
### 13.1.2 Physiological Alarm Event Messages

This section lists the alarm messages that may appear at the Panorama Central Station when an Physiological alarm event occurs.

<table>
<thead>
<tr>
<th>PARAMETER</th>
<th>PANORAMA CENTRAL STATION MESSAGES</th>
<th>REASON</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heart Rate (HR)</td>
<td>High Heart Rate Alarm Low Heart Rate Alarm Extreme Bradycardia Alarm Extreme Tachycardia Alarm</td>
<td>Current value is either equal to or higher/lower than the set alarm limits.</td>
</tr>
<tr>
<td>ECG</td>
<td>Pacer Failure to Capture</td>
<td>No QRS complex detected within 300ms after a PM event.</td>
</tr>
<tr>
<td></td>
<td>Pacer Failure to Sense</td>
<td>No QRS complex detected or within (60*1000/pacemaker pacing rate +90)ms, after a PM event or QRS detected event.</td>
</tr>
<tr>
<td>Respiration (ECG Thoracic Impedance, CO₂, and Gas Module)</td>
<td>High Respiration Rate Alarm Low Respiration Rate Alarm</td>
<td>Current value is either equal to or higher/lower than the set alarm limits.</td>
</tr>
<tr>
<td>SpO₂ (Masimo SET®, Nellcor®, Nonin)</td>
<td>High SpO₂ Alarm Low SpO₂ Alarm</td>
<td>Current value is either equal to or higher/lower than the set alarm limits.</td>
</tr>
<tr>
<td>IBPx (Sys) Labels include: IBP1, IBP2, IBP3, IBP4, IBP5, IBP6, IBP7, IBP8, ART, UA, LV, PA, CVP, ICP, LA, RA</td>
<td>High IBPx Systolic Alarm Low IBPx Systolic Alarm</td>
<td>Current value is either equal to or higher/lower than the set alarm limits.</td>
</tr>
<tr>
<td>IBPx (Dia) Labels include: IBP1, IBP2, IBP3, IBP4, IBP5, IBP6, IBP7, IBP8, ART, UA, LV, PA, CVP, ICP, LA, RA</td>
<td>High IBPx Diastolic Alarm Low IBPx Diastolic Alarm</td>
<td>Current value is either equal to or higher/lower than the set alarm limits.</td>
</tr>
<tr>
<td>IBPx (Mean) Labels include: IBP1, IBP2, IBP3, IBP4, IBP5, IBP6, IBP7, IBP8, ART, UA, LV, PA, CVP, ICP, LA, RA</td>
<td>High IBPx Mean Alarm Low IBPx Mean Alarm</td>
<td>Current value is either equal to or higher/lower than the set alarm limits.</td>
</tr>
<tr>
<td>INSP CO₂ [mmHg, %, kPa]</td>
<td>High CO₂ Inspired Alarm</td>
<td>Current value is either equal to or higher than the set alarm limits.</td>
</tr>
<tr>
<td>ET CO₂ [mmHg, %, kPa]</td>
<td>High CO₂ ET Alarm Low CO₂ ET Alarm</td>
<td>Current value is either equal to or higher/lower than the set alarm limits.</td>
</tr>
<tr>
<td>INSP O₂</td>
<td>O₂ Inspired Alarm</td>
<td>There has been a violation of the O₂ Inspired alarm.</td>
</tr>
<tr>
<td>ET O₂</td>
<td>O₂ ET Alarm</td>
<td>There has been a violation of the O₂ ET alarm.</td>
</tr>
<tr>
<td>INSP N₂O</td>
<td>N₂O Inspired Alarm</td>
<td>There has been a violation of the N₂O Inspired alarm.</td>
</tr>
<tr>
<td>ET N₂O</td>
<td>N₂O ET Alarm</td>
<td>There has been a violation of the N₂O ET alarm.</td>
</tr>
<tr>
<td>PARAMETER</td>
<td>PANORAMA CENTRAL STATION MESSAGES</td>
<td>REASON</td>
</tr>
<tr>
<td>-----------</td>
<td>----------------------------------</td>
<td>--------</td>
</tr>
<tr>
<td>Agent</td>
<td>Agent Inspired Alarm</td>
<td>There has been a violation of the Agent Inspired alarm.</td>
</tr>
<tr>
<td></td>
<td>Agent ET Alarm</td>
<td>There has been a violation of the Agent ET alarm.</td>
</tr>
<tr>
<td>INSP Iso</td>
<td>Agent Iso Inspired Alarm</td>
<td>There has been a violation of the Iso Inspired alarm.</td>
</tr>
<tr>
<td>ET Iso</td>
<td>Agent Iso ET Alarm</td>
<td>There has been a violation of the Iso ET alarm.</td>
</tr>
<tr>
<td>INSP Enf</td>
<td>Agent Enf Inspired Alarm</td>
<td>There has been a violation of the Enf Inspired alarm.</td>
</tr>
<tr>
<td>ET Enf</td>
<td>Agent Enf ET Alarm</td>
<td>There has been a violation of the Enf ET alarm.</td>
</tr>
<tr>
<td>INSP Des</td>
<td>Agent Des Inspired Alarm</td>
<td>There has been a violation of the Des Inspired alarm.</td>
</tr>
<tr>
<td>ET Des</td>
<td>Agent Des ET Alarm</td>
<td>There has been a violation of the Des ET alarm.</td>
</tr>
<tr>
<td>INSP Sev</td>
<td>Agent Sev Inspired Alarm</td>
<td>There has been a violation of the Sev Inspired alarm.</td>
</tr>
<tr>
<td>ET Sev</td>
<td>Agent Sev ET Alarm</td>
<td>There has been a violation of the Sev ET alarm.</td>
</tr>
<tr>
<td>INSP Hal</td>
<td>Agent Hal Inspired Alarm</td>
<td>There has been a violation of the Hal Inspired alarm.</td>
</tr>
<tr>
<td>ET Hal</td>
<td>Agent Hal ET Alarm</td>
<td>There has been a violation of the Hal ET alarm.</td>
</tr>
<tr>
<td>NIBP (Sys)</td>
<td>High NIBP Sys Alarm</td>
<td>Current value is either equal to or higher/lower than the set alarm limits.</td>
</tr>
<tr>
<td></td>
<td>Low NIBP Sys Alarm</td>
<td></td>
</tr>
<tr>
<td>NIBP (Dia)</td>
<td>High NIBP Dia Alarm</td>
<td>Current value is either equal to or higher/lower than the set alarm limits.</td>
</tr>
<tr>
<td></td>
<td>Low NIBP Dia Alarm</td>
<td></td>
</tr>
<tr>
<td>NIBP (Mean)</td>
<td>High NIBP Mean Alarm</td>
<td>Current value is either equal to or higher/lower than the set alarm limits.</td>
</tr>
<tr>
<td></td>
<td>Low NIBP Mean Alarm</td>
<td></td>
</tr>
<tr>
<td>CCI</td>
<td>High CCI Alarm</td>
<td>Current value is either equal to or higher/lower than the set alarm limits.</td>
</tr>
<tr>
<td></td>
<td>Low CCI Alarm</td>
<td></td>
</tr>
<tr>
<td>CCI STAT</td>
<td>High CCI STAT Alarm</td>
<td>Current value is either equal to or higher/lower than the set alarm limits.</td>
</tr>
<tr>
<td></td>
<td>Low CCI STAT Alarm</td>
<td></td>
</tr>
<tr>
<td>CCO</td>
<td>High CCO Alert</td>
<td>Current value is either equal to or higher/lower than the set alarm limits.</td>
</tr>
<tr>
<td></td>
<td>Low CCO Alert</td>
<td></td>
</tr>
<tr>
<td>CCO STAT</td>
<td>High CCO STAT Alarm</td>
<td>Current value is either equal to or higher/lower than the set alarm limits.</td>
</tr>
<tr>
<td></td>
<td>Low CCO STAT Alarm</td>
<td></td>
</tr>
<tr>
<td>SvO₂</td>
<td>High SvO₂ Alarm</td>
<td>Current value is either equal to or higher/lower than the set alarm limits.</td>
</tr>
<tr>
<td></td>
<td>Low SvO₂ Alarm</td>
<td></td>
</tr>
<tr>
<td>SVR</td>
<td>High SVR Alarm</td>
<td>Current value is either equal to or higher/lower than the set alarm limits.</td>
</tr>
<tr>
<td></td>
<td>Low SVR Alarm</td>
<td></td>
</tr>
<tr>
<td>SVRI</td>
<td>High SVRI Alarm</td>
<td>Current value is either equal to or higher/lower than the set alarm limits.</td>
</tr>
<tr>
<td></td>
<td>Low SVRI Alarm</td>
<td></td>
</tr>
<tr>
<td>T1 (F/C)</td>
<td>High Temp 1 Alarm</td>
<td>Current value is either equal to or higher/lower than the set alarm limits.</td>
</tr>
<tr>
<td></td>
<td>Low Temp 1 Alarm</td>
<td></td>
</tr>
<tr>
<td>T2 (F/C)</td>
<td>High Temp 2 Alarm</td>
<td>Current value is either equal to or higher/lower than the set alarm limits.</td>
</tr>
<tr>
<td></td>
<td>Low Temp 2 Alarm</td>
<td></td>
</tr>
<tr>
<td>T3 (F/C)</td>
<td>High Temp 3 Alarm</td>
<td>Current value is either equal to or higher/lower than the set alarm limits.</td>
</tr>
<tr>
<td></td>
<td>Low Temp 3 Alarm</td>
<td></td>
</tr>
<tr>
<td>PARAMETER</td>
<td>PANORAMA CENTRAL STATION MESSAGES</td>
<td>REASON</td>
</tr>
<tr>
<td>-----------</td>
<td>---------------------------------</td>
<td>--------</td>
</tr>
<tr>
<td>Delta T (F/C)</td>
<td>High Delta T Alarm Low Delta T Alarm</td>
<td>Current value either equal to or higher/lower than the set alarm limits.</td>
</tr>
<tr>
<td>Temperature Label: ESO, NAS, REC, BLA, AXI, SKI</td>
<td>High ESO Alarm Low ESO Alarm High NAS Alarm Low NAS Alarm High REC Alarm Low REC Alarm High BLA Alarm Low BLA Alarm High AXI Alarm Low AXI Alarm High SKI Alarm Low SKI Alarm</td>
<td>Current value either equal to or higher/lower than the set alarm limits.</td>
</tr>
<tr>
<td>ESO Not Calibrated/Zeroed NAS Not Calibrated/Zeroed REC Not Calibrated/Zeroed BLA Not Calibrated/Zeroed AXI Not Calibrated/Zeroed SKI Not Calibrated/Zeroed</td>
<td>Sensor has not been calibrated or zeroed.</td>
<td></td>
</tr>
<tr>
<td>T Blood (F/C)</td>
<td>Low T Blood Alarm High T Blood Alarm</td>
<td>Current value is either equal to or higher/lower than the set alarm limits.</td>
</tr>
<tr>
<td>ST Single</td>
<td>ST Single Alarm</td>
<td>Current value of one ST lead is either equal to or higher/lower than the set alarm limits. A separate ST Single alarm will be shown for each lead of ST that independently violates the alarm.</td>
</tr>
<tr>
<td>ST Dual</td>
<td>ST Dual Alarm</td>
<td>Current value for two or more ST leads are either equal to or higher/lower than the set alarm limits. A separate ST Dual alarm will be shown for every two leads of ST that independently violate the alarm.</td>
</tr>
<tr>
<td>Apnea</td>
<td>Apnea Alarm</td>
<td>Apneic episode is either equal to or longer than the set alarm limits.</td>
</tr>
<tr>
<td>PVC</td>
<td>High PVC Alarm</td>
<td>Current PVCs/per minute threshold is in violation.</td>
</tr>
<tr>
<td>BIS</td>
<td>High BIS Alarm Low BIS Alarm High BIS L Alarm Low BIS L Alarm High BIS R Alarm Low BIS R Alarm</td>
<td>Current value either equal to or higher/lower than the set alarm limits.</td>
</tr>
</tbody>
</table>
### 13.2 Physiological Alarms and Events: Arrhythmias

This section lists the alarm messages that may appear at the Panorama Central Station when an Arrhythmia alarm occurs.

#### Arrhythmia Alarms

<table>
<thead>
<tr>
<th>EVENT</th>
<th>ALARM PRIORITY</th>
<th>PANORAMA CENTRAL STATION MESSAGES</th>
<th>REASON</th>
</tr>
</thead>
<tbody>
<tr>
<td>V-Tach</td>
<td>1</td>
<td>V-Tach</td>
<td>The configured number of consecutive PVCs is reached at a rate equal to or greater than the V-Tach threshold. The range of the V-Tach heart rate is between 100 to 180 beats per minute. The range of the V-Tach threshold is 3-15 bpm.</td>
</tr>
<tr>
<td>V-Fib</td>
<td>1</td>
<td>V-Fib</td>
<td>There is an absence of recognizable P, QRS or T waves.</td>
</tr>
<tr>
<td>Asystole</td>
<td>1</td>
<td>Asystole</td>
<td>There are no QRS complexes detected for the configured time period in the absence of Ventricular Fibrillation. When using a View 12™ card, the time period range for an Asystole alarm is between three (3) and ten (10) seconds, and between three (3) and eight (8) seconds when using a 3 or 5-wire lead set.</td>
</tr>
<tr>
<td>Bigeminy</td>
<td>2, 3 or Off</td>
<td>Bigeminy</td>
<td>There are three or more cycles of one PVC followed by a normal beat.</td>
</tr>
<tr>
<td>Couplet</td>
<td>2, 3 or Off</td>
<td>Couplet</td>
<td>There are two consecutive PVCs detected between normal beats.</td>
</tr>
<tr>
<td>Irregular HR</td>
<td>2, 3 or Off</td>
<td>Irregular HR</td>
<td>The measured variation in the R-R interval over a period of time exceeds a preset limit established by the arrhythmia algorithm.</td>
</tr>
<tr>
<td>Run</td>
<td>2, 3 or Off</td>
<td>Run</td>
<td>There are a number of consecutive PVCs occurring at a rate that exceeds the configured V-Tach rate.</td>
</tr>
<tr>
<td>Trigeminy</td>
<td>2, 3 or Off</td>
<td>Trigeminy</td>
<td>Three or more cycles of one PVC coupled to two normal beats are detected.</td>
</tr>
<tr>
<td>V-Rhythm</td>
<td>2, 3 or Off</td>
<td>V-Rhythm</td>
<td>The number of consecutive PVCs is greater than two and the ventricular rate is less than the configured V-Tach threshold.</td>
</tr>
<tr>
<td>PVC</td>
<td>2, 3 or Off</td>
<td>High PVC Alarm</td>
<td>The number of PVCs detected per minute exceeds the configured threshold.</td>
</tr>
</tbody>
</table>

1 Indicates alarm is latched.
2 Alarm will only be detected when using a View 12™ card, a Telepack and V 12/V 21 devices.
3 Alarm will not be detected when using a View 12™ card or a Telepack.
4 Alarm is only available for the Telepack, DPM 6/DPM 7, Passport 12M/Passport 17M, Passport 8/Passport 12, T1 monitor and V 12/V 21 devices.
5 Alarm is only available for the Telepack and V 12/V 21 devices.
<table>
<thead>
<tr>
<th>EVENT</th>
<th>ALARM PRIORITY</th>
<th>PANORAMA CENTRAL STATION MESSAGES</th>
<th>REASON</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brady&lt;sup&gt;3&lt;/sup&gt;</td>
<td>1 or Off</td>
<td>Brady</td>
<td>Heart rate fell to a value 10% lower than the configured low heart rate alarm.</td>
</tr>
<tr>
<td></td>
<td>Factory Default: 1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pause&lt;sup&gt;2&lt;/sup&gt;</td>
<td>1, 2, 3 or Off</td>
<td>Pause</td>
<td>No beat is detected during an interval that is greater than the adjustable set value (120% - 200%) R-R and when the next beat is not a PVC.</td>
</tr>
<tr>
<td></td>
<td>Factory Default: 2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Multifocal PVC&lt;sup&gt;4&lt;/sup&gt;</td>
<td>2, 3 or Off</td>
<td>Multifocal PVC</td>
<td>During the last 15 beats, two or more premature ventricular beats with different morphologies were detected.</td>
</tr>
<tr>
<td></td>
<td>Factory Default: 3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>R-On-T&lt;sup&gt;4&lt;/sup&gt;</td>
<td>1, 2 or Off</td>
<td>R-On-T</td>
<td>During early PVC, a beat is detected as a PVC and the previous beat as a non-PVC.</td>
</tr>
<tr>
<td></td>
<td>Factory Default: 2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pause/min&lt;sup&gt;5&lt;/sup&gt;</td>
<td>1, 2, 3 or Off</td>
<td>Pause/Minute Alarm</td>
<td>The number of pauses detected per minute exceeds the configured threshold. The Pauses/min limit can be set to 1 to 15 pauses per minute.</td>
</tr>
<tr>
<td></td>
<td>Factory Default: 2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>QTc&lt;sup&gt;5&lt;/sup&gt;</td>
<td>2 or 3</td>
<td>High QTc Alarm</td>
<td>Current QTc value is equal to or higher than the set alarm limit.</td>
</tr>
<tr>
<td></td>
<td>Factory Default: 3</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

1 Indicates alarm is latched.
2 Alarm will only be detected when using a View 12™ card, a Telepack and V 12/V 21 devices.
3 Alarm will not be detected when using a View 12™ card or a Telepack.
4 Alarm is only available for the Telepack, DPM 6/DPM 7, Passport 12M/Passport 17M, Passport 8/Passport 12, T1 monitor and V 12/V 21 devices.
5 Alarm is only available for the Telepack and V 12/V 21 devices.
13.3 Technical Alarms and Events

The Panorama Central Station uses Technical alarms to indicate that a patient specific technical issue has occurred.

This section provides the technical event messages that may be posted to the Events database and shown in white text (with the exception of the All Alarms Paused, All Alarms Off, All Non-High Alarms Paused, and All Non-High Alarms Off events) in the Patient Status line. When a number is indicated beside the event message, refer to the table footnotes.

Technical Events

<table>
<thead>
<tr>
<th>MESSAGE</th>
<th>REASON</th>
</tr>
</thead>
<tbody>
<tr>
<td>All Alarms Paused 1,2</td>
<td>The All Alarms Pause sidebar button was selected in the Patient Alarms tab.</td>
</tr>
<tr>
<td></td>
<td><strong>NOTE:</strong> This event message will display in red text in the Patient Status line.</td>
</tr>
<tr>
<td>All Alarms Off</td>
<td>The All Alarms Off sidebar button was selected in the Patient Alarms tab.</td>
</tr>
<tr>
<td></td>
<td><strong>NOTE:</strong> This event message will display in red text in the Patient Status line.</td>
</tr>
<tr>
<td>Resume All Alarms 1,2,3</td>
<td>A patient’s alarms are to resume.</td>
</tr>
<tr>
<td>All Non-High Alarms Paused 1,2</td>
<td>The All Non-High Alarms Pause sidebar button is selected in the Patient Alarms tab.</td>
</tr>
<tr>
<td></td>
<td><strong>NOTE:</strong> This event message will display in yellow text in the Patient Status line.</td>
</tr>
<tr>
<td>All Non-High Alarms Off</td>
<td>The All Non-High Alarms Off sidebar button is selected in the Patient Alarms tab.</td>
</tr>
<tr>
<td></td>
<td><strong>NOTE:</strong> This event message will display in yellow text in the Patient Status line.</td>
</tr>
<tr>
<td>Resume All Non-High Alarms 1,2,3</td>
<td>A patient’s non-lethal alarms are set to resume.</td>
</tr>
<tr>
<td>CommLoss - Check Bedside Monitor 4,5</td>
<td>The connection between the Panorama Central Station and the attached monitoring equipment has been interrupted.</td>
</tr>
<tr>
<td>Communications Lost 4,5</td>
<td>The connection between the Panorama Central Station and the Telepack has been interrupted.</td>
</tr>
<tr>
<td>Central Communications Lost 4</td>
<td>The connection between the host central station and the ViewStation/WorkStation has been interrupted.</td>
</tr>
<tr>
<td>Signal Lost-Limited Arrhythmia 4</td>
<td>A Telepack is in use, arrhythmia analysis is enabled, and there are multiple disruptions of data that may affect the performance of the arrhythmia algorithm.</td>
</tr>
</tbody>
</table>

1 Event cannot be deleted from Event database
2 Event will not elicit technical event sound
3 Event will not display in Patient Status line
4 Event will always elicit the technical event sound regardless of the Technical Event sound setting
5 Event will post a Page request when the central station has the paging option enabled and the alarm is configured to send a page
Technical Events (Continued)

<table>
<thead>
<tr>
<th>MESSAGE</th>
<th>REASON</th>
</tr>
</thead>
<tbody>
<tr>
<td>Signal Lost 4</td>
<td>A Telepack is in use, arrhythmia analysis is disabled, and there are multiple disruptions of data.</td>
</tr>
<tr>
<td>Standby 1,2</td>
<td>A patient was placed into Standby mode. In Standby mode, all of the patient’s monitoring activity is suspended.</td>
</tr>
</tbody>
</table>

**NOTE:** Message will not be shown in the Patient Status line. Instead, message and/or Standby location will be shown in the patient’s digital data tile in pink text.

<table>
<thead>
<tr>
<th>Resume Monitoring 1,2</th>
<th>The patient was removed from Standby mode.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low Battery 5</td>
<td>The monitoring equipment battery is low.</td>
</tr>
</tbody>
</table>

**NOTE:** If the Low Battery message appears and the monitoring device is a Telepack, the Battery Percentage digital data tile will flash yellow until the issue is resolved.

<table>
<thead>
<tr>
<th>T1 No Battery</th>
<th>The T1 monitor is not powered by the external power and the battery is removed.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Battery Bad Cell</td>
<td>Bad battery cells are detected at the patient monitor battery. This message is only displayed on the Patient Status line.</td>
</tr>
<tr>
<td>Cooling Fan Failure</td>
<td>The bedside monitor cooling fan is not operational and the monitor is running on AC power. This message is only displayed on the Patient Status line.</td>
</tr>
<tr>
<td>Admit 2,3</td>
<td>A patient is admitted to the Panorama Central Station.</td>
</tr>
<tr>
<td>Discharge</td>
<td>The patient was discharged from the Panorama Central Station.</td>
</tr>
<tr>
<td>Arrhythmia On 1,2,3</td>
<td>The Arrhythmia option is enabled on a patient’s monitoring equipment.</td>
</tr>
<tr>
<td>Arrhythmia Off 1,2,3</td>
<td>The Arrhythmia option is disabled on a patient’s monitoring equipment.</td>
</tr>
<tr>
<td>ST On 1,2,3</td>
<td>The ST option is enabled on a patient’s monitoring equipment.</td>
</tr>
<tr>
<td>ST Off 1,2,3</td>
<td>The ST option is disabled on a patient’s monitoring equipment.</td>
</tr>
<tr>
<td>ST Relearn 2,3</td>
<td>The ST algorithm completes a learning phase.</td>
</tr>
<tr>
<td>Arrhythmia Relearn 2,3</td>
<td>The arrhythmia algorithm completes a learning phase.</td>
</tr>
<tr>
<td>Central User Marked Event 2,3</td>
<td>The user has marked an event at the central station.</td>
</tr>
<tr>
<td>Bedside User Marked Event 2,3</td>
<td>The user has marked an event at the bedside monitor.</td>
</tr>
</tbody>
</table>

1 Event cannot be deleted from Event database  
2 Event will not elicit technical event sound  
3 Event will not display in Patient Status line  
4 Event will always elicit the technical event sound regardless of the Technical Event sound setting  
5 Event will post a Page request when the central station has the paging option enabled and the alarm is configured to send a page
Technical Events (Continued)

<table>
<thead>
<tr>
<th>MESSAGE</th>
<th>REASON</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calipers Marked Event 2,3</td>
<td>The user has marked a caliper event at the Panorama Central Station</td>
</tr>
<tr>
<td>Central Transfer 2,3</td>
<td>A central-to-central transfer has been successfully completed at the Central Station.</td>
</tr>
<tr>
<td>Room Transfer 2,3</td>
<td>A room-to-room transfer has been successfully completed at the central station.</td>
</tr>
<tr>
<td>Break of Data 2,3</td>
<td>Data has not been received for a device for 1 second.</td>
</tr>
<tr>
<td>Auto Sync 2,3</td>
<td>For the Telepack, this is performed once every 4 hours of uninterrupted communications. For 608 radio devices, this is performed once every 20 hours of uninterrupted communications. For DPM 6/DPM 7, Passport 12M/Passport 17M, Passport 8/Passport 12, T1 and Passport V devices, this occurs when there is a 3 second or more difference in the most common disclosure data time and the Panorama clock time. This is needed to re-sync the data from the devices with the Panorama system time. No disclosure data is lost during this re-sync.</td>
</tr>
</tbody>
</table>

Unknown Tech Alarm

This occurs when the technical alarms on the DPM 6/DPM 7, Passport 12M/Passport 17M, Passport 8/Passport 12 and T1 devices are not supported by the Panorama Central Station.

HEART RATE TECHNICAL EVENTS

<table>
<thead>
<tr>
<th>MESSAGE</th>
<th>REASON</th>
</tr>
</thead>
<tbody>
<tr>
<td>Check Lead Connections</td>
<td>The bedside monitor detects a lead off condition.</td>
</tr>
<tr>
<td>RA Lead Off 4,5</td>
<td>The RA lead is not connected to the patient, ECG lead I or II is selected while using a 3-wire lead set, 5-wire lead set or a View 12™ card.</td>
</tr>
<tr>
<td>RL Lead Off 4,5</td>
<td>The RL lead is not connected to the patient while using a 5-wire lead set or a View 12™ card.</td>
</tr>
<tr>
<td>LA Lead Off 4,5</td>
<td>The LA lead is not connected to the patient, ECG lead I or III is selected while using a 3-wire lead set, 5-wire lead set, or a View 12™ card.</td>
</tr>
<tr>
<td>LL Lead Off 4,5</td>
<td>The LL lead is not connected to the patient, ECG lead II or III is selected while using a 3-wire lead, 5-wire lead set, or View 12™ card.</td>
</tr>
<tr>
<td>Leads Off 4,5</td>
<td>Two or more of the ECG leads are not connected to the patient and a 3-wire lead set, 5-wire lead set or a View 12™ card is in use. It will also be shown when using a 3-wire lead set and ECG I, II, or III is shown and the LL, LA, or RA leads are off, respectively. When using a View 12™ card and two or more leads are off, unless one of the disconnected leads is the RL lead. If a 5-wire lead set is in use and the RL lead is off.</td>
</tr>
</tbody>
</table>

1 Event cannot be deleted from Event database
2 Event will not elicit technical event sound
3 Event will not display in Patient Status line
4 Event will always elicit the technical event sound regardless of the Technical Event sound setting
5 Event will post a Page request when the central station has the paging option enabled and the alarm is configured to send a page
Technical Events (Continued)

<table>
<thead>
<tr>
<th>MESSAGE</th>
<th>REASON</th>
</tr>
</thead>
<tbody>
<tr>
<td>V Lead Off</td>
<td>A V lead is not connected to the patient and a 5-wire lead set is in use.</td>
</tr>
<tr>
<td>V1-V6 Lead Off</td>
<td>A V1-6 lead is not connected to the patient and a View 12™ card is in use.</td>
</tr>
<tr>
<td>ECG Noise</td>
<td>Noise is present on one or more leads.</td>
</tr>
<tr>
<td>Acquiring ECG</td>
<td>The algorithm is trying to acquire a heart rate from ECG for a Telepack or V 12/V 21.</td>
</tr>
</tbody>
</table>

**RESPIRATION (ECG THORACIC IMPEDANCE) TECHNICAL EVENTS**

<table>
<thead>
<tr>
<th>MESSAGE</th>
<th>REASON</th>
</tr>
</thead>
<tbody>
<tr>
<td>Respiration CVA Present</td>
<td>A CVA condition is detected for the monitored patient.</td>
</tr>
</tbody>
</table>

**SPO2 (MASIMO SET® AND NELLCOR®) TECHNICAL EVENTS**

<table>
<thead>
<tr>
<th>MESSAGE</th>
<th>REASON</th>
</tr>
</thead>
<tbody>
<tr>
<td>SpO2 System Check</td>
<td>There is an SpO2 system check.</td>
</tr>
<tr>
<td>SpO2 Communication Error</td>
<td>There is interference in the communication between the SpO2 source and the bedside monitor.</td>
</tr>
<tr>
<td>SpO2 Failure</td>
<td>A faulty 5V power supply or a faulty positive or negative 15V power supply is detected on the SpO2 board.</td>
</tr>
<tr>
<td>SpO2 Board Fault</td>
<td>The SpO2 board fails.</td>
</tr>
<tr>
<td>SpO2 Unrecognized Cable</td>
<td>The system detects an unrecognized SpO2 cable.</td>
</tr>
<tr>
<td>SpO2 Pulse Search</td>
<td>The SpO2 sensor is attempting to detect a pulse.</td>
</tr>
<tr>
<td>SpO2 Weak Pulse</td>
<td>The SpO2 sensor detects a weak pulse.</td>
</tr>
</tbody>
</table>

**SPO2 (MASIMO) TECHNICAL EVENTS**

<table>
<thead>
<tr>
<th>MESSAGE</th>
<th>REASON</th>
</tr>
</thead>
<tbody>
<tr>
<td>SpO2 Low Perfusion</td>
<td>The SpO2 Masimo sensor detects low perfusion.</td>
</tr>
<tr>
<td>SpO2 Too Much Light</td>
<td>The SpO2 Masimo sensor detects too much light.</td>
</tr>
<tr>
<td>SpO2 Unrecognized Sensor</td>
<td>An unrecognized SpO2 Masimo sensor is detected by the system.</td>
</tr>
<tr>
<td>SpO2 Sensor Off</td>
<td>The SpO2 Masimo sensor is off.</td>
</tr>
<tr>
<td>SpO2 No Sensor</td>
<td>The SpO2 Masimo sensor is not plugged in.</td>
</tr>
<tr>
<td>SpO2 Sensor Failed</td>
<td>Communication with the SpO2 Masimo sensor failed.</td>
</tr>
<tr>
<td>SpO2 Interference</td>
<td>There is interference in the communication with the SpO2 Masimo sensor.</td>
</tr>
</tbody>
</table>

**SPO2 (NELLCOR) TECHNICAL EVENTS**

<table>
<thead>
<tr>
<th>MESSAGE</th>
<th>REASON</th>
</tr>
</thead>
<tbody>
<tr>
<td>SpO2 Motion</td>
<td>The SpO2 Nellcor sensor detects motion.</td>
</tr>
<tr>
<td>SpO2 Check Sensor</td>
<td>The SpO2 Nellcor sensor detects a problem.</td>
</tr>
<tr>
<td>SpO2 No Pulse</td>
<td>The SpO2 Nellcor sensor does not detect a pulse.</td>
</tr>
</tbody>
</table>

---

1. Event cannot be deleted from Event database
2. Event will not elicit technical event sound
3. Event will not display in Patient Status line
4. Event will always elicit the technical event sound regardless of the Technical Event sound setting
5. Event will post a Page request when the central station has the paging option enabled and the alarm is configured to send a page
Technical Events (Continued)

<table>
<thead>
<tr>
<th>MESSAGE</th>
<th>REASON</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>SPO₂ (NONIN) TECHNICAL EVENTS</strong></td>
<td></td>
</tr>
<tr>
<td>SpO₂ (NONIN) Communication Error</td>
<td>There is interference in the communication between the SpO₂ source and the Telepack.</td>
</tr>
<tr>
<td>SpO₂ Check Sensor</td>
<td>The SpO₂ (Nonin) sensor detects a problem.</td>
</tr>
<tr>
<td>SpO₂ No Sensor</td>
<td>The SpO₂ (Nonin) sensor is not plugged in or the sensor is inoperable.</td>
</tr>
<tr>
<td>SpO₂ Low Perfusion</td>
<td>The SpO₂ (Nonin) sensor detects low perfusion.</td>
</tr>
<tr>
<td>SpO₂ Pulse Search</td>
<td>The SpO₂ (Nonin) sensor is attempting to detect a pulse.</td>
</tr>
<tr>
<td><strong>IBPX TECHNICAL EVENTS</strong></td>
<td></td>
</tr>
<tr>
<td>IBPx (Labels include: Art, UA, LV, PA, AO, BAP, FAP, IAP, CVP, ICP, LA, RA, UAP, UVP, ART2) Not Calibrated/Zeroed</td>
<td>The IBP device is not calibrated or zeroed.</td>
</tr>
<tr>
<td>IBPx (Labels include: Art, UA, LV, PA, AO, BAP, FAP, CVP, ICP, LA, RA) Sensor Off</td>
<td>The IBP x sensor is off.</td>
</tr>
<tr>
<td>IBPx (Labels include: Art, UA, LV, PA, AO, BAP, FAP, CVP, ICP, LA, RA, UVP) No Pulse</td>
<td>The system does not detect a IBPx pulse.</td>
</tr>
<tr>
<td>IBPx (Labels include: Art, UA, LV, PA, AO, BAP, FAP, CVP, ICP, LA, RA, UVP) Search Pulse</td>
<td>The system detects a IBPx pulse search.</td>
</tr>
<tr>
<td>NOTE: x = 1, 2, 3 or 4 (IBP3 and 4 are only available if the Spectrum/Spectrum OR External Parameter Module (EPM) is installed)</td>
<td></td>
</tr>
<tr>
<td><strong>CO₂ TECHNICAL EVENTS</strong></td>
<td></td>
</tr>
<tr>
<td>CO₂ Not Calibrated/Zeroed</td>
<td>The CO₂ device is not calibrated.</td>
</tr>
<tr>
<td>CO₂ Occlusion</td>
<td>The CO₂ device detects a blockage in the filter line.</td>
</tr>
<tr>
<td><strong>O₂ TECHNICAL EVENTS</strong></td>
<td></td>
</tr>
<tr>
<td>O₂ Not Calibrated/Zeroed</td>
<td>The O₂ device is not calibrated.</td>
</tr>
<tr>
<td>O₂ Zeroing</td>
<td>The O₂ device is zeroing.</td>
</tr>
<tr>
<td>O₂ Occlusion</td>
<td>The O₂ device detects a blockage in the filter line.</td>
</tr>
<tr>
<td><strong>N₂O TECHNICAL EVENTS</strong></td>
<td></td>
</tr>
<tr>
<td>N₂O Not Calibrated/Zeroed</td>
<td>The N₂O device is not calibrated.</td>
</tr>
<tr>
<td>N₂O Zeroing</td>
<td>The N₂O device is zeroing.</td>
</tr>
<tr>
<td>N₂O Occlusion</td>
<td>The N₂O device detects a blockage in the filter line.</td>
</tr>
<tr>
<td><strong>AGENT TECHNICAL EVENTS</strong></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>Event cannot be deleted from Event database</td>
</tr>
<tr>
<td>2</td>
<td>Event will not elicit technical event sound</td>
</tr>
<tr>
<td>3</td>
<td>Event will not display in Patient Status line</td>
</tr>
<tr>
<td>4</td>
<td>Event will always elicit the technical event sound regardless of the Technical Event sound setting</td>
</tr>
<tr>
<td>5</td>
<td>Event will post a Page request when the central station has the paging option enabled and the alarm is configured to send a page</td>
</tr>
</tbody>
</table>
Technical Events (Continued)

<table>
<thead>
<tr>
<th>MESSAGE</th>
<th>REASON</th>
</tr>
</thead>
<tbody>
<tr>
<td>Agent Not Calibrated/Zeroed</td>
<td>The Agent device is not calibrated.</td>
</tr>
<tr>
<td>Agent Zeroing</td>
<td>The Agent device is zeroing.</td>
</tr>
</tbody>
</table>

**NIBP TECHNICAL EVENTS**

| NIBP Cuff Overpressure                  | The patient’s NIBP cuff has an overpressure episode.                 |
| NIBP Unable to Measure                  | The patient’s NIBP cuff is unable to take a valid measurement.       |
| NIBP Retry                              | There is an NIBP measurement retry.                                  |
| NIBP Pump Higher                        | The patient’s NIBP cuff needs additional pressure to take a measurement. |
| NIBP Measuring                          | The patient’s NIBP cuff is inflating.                                |
| NIBP Deflating                          | The patient’s NIBP cuff is deflating.                                |

**CARDIAC OUTPUT TECHNICAL EVENTS**

| New CO Measurement 2,3          | The Cardiac Output run was accepted at the bedside monitor.          |

**NOTE:** The CO parameter is only available when the Spectrum/Spectrum OR External Parameter Module (EPM) is installed.

**PAWP TECHNICAL EVENTS**

| New PAWP Measurement 2,3        | The PAWP value was accepted on the bedside monitor.                  |

**NURSE CALL TECHNICAL EVENTS**

| Nurse Call 4                   | The Nurse Call button on the Telepack was selected.                  |

**ATTENDANT PRESENT TECHNICAL EVENTS**

| Attendant Present 2,3           | The Attendant Present buttons on the Telepack were selected.         |

**BEDSIDE PRINTER TECHNICAL EVENTS**

| Printer out of paper            | The bedside monitor local printer is out of paper.                   |
| Printer Door is Open            | The bedside monitor local printer has an open door condition.        |

**CENTRAL COMMUNICATION TECHNICAL EVENTS**

| Central Communications Lost     | The connection between the ViewStation and the host central station was interrupted. |

1  Event cannot be deleted from Event database
2  Event will not elicit technical event sound
3  Event will not display in Patient Status line
4  Event will always elicit the technical event sound regardless of the Technical Event sound setting
5  Event will post a Page request when the central station has the paging option enabled and the alarm is configured to send a page
13.4 System Event Messages and Error Codes

The Panorama Central Station displays system event messages and error codes to indicate system-wide issues at the central station and/or at the ViewStation. They are listed based on the system violation, and are resolved when the system-wide condition no longer exists.

This section provides the system event messages and error codes that may be shown in the System Status line in white text.

System Event Messages and Error Codes

<table>
<thead>
<tr>
<th>MESSAGE</th>
<th>REASON</th>
</tr>
</thead>
<tbody>
<tr>
<td>No Printer Selected or Available</td>
<td>When the print request was made at the central station or remote workstation, no printer was selected, printer was unavailable, or printer was not properly configured.</td>
</tr>
<tr>
<td>Printer Offline</td>
<td>The central station or remote workstation default printer’s Offline button is selected.</td>
</tr>
<tr>
<td></td>
<td><strong>NOTE:</strong> Button may not be labeled &quot;Offline.&quot; Refer to printer user guide for specific button name.</td>
</tr>
<tr>
<td>Printer Paper Jam</td>
<td>The central station or remote workstation default printer paper tray is removed as the print job is starting to print.</td>
</tr>
<tr>
<td>Printer Out of Paper</td>
<td>The central station or remote workstation default printer is out of paper.</td>
</tr>
<tr>
<td>Clock Adjusted</td>
<td>The central station or remote workstation system time or date was adjusted.</td>
</tr>
<tr>
<td>Print Halted...Print Queue Full</td>
<td>The central station or remote workstation default printer has been halted because the print queue is full.</td>
</tr>
<tr>
<td>Printer error - no toner</td>
<td>The central station or remote workstation default printer toner is low.</td>
</tr>
<tr>
<td>Printer in error state</td>
<td>The central station or remote workstation default printer has an error that is not related to any of the previous messages.</td>
</tr>
<tr>
<td>All Audio Paused for xx:xx mins</td>
<td>The Audio Pause menu button was selected at the ViewStation or WorkStation.</td>
</tr>
<tr>
<td>All Audio Off</td>
<td>The All Audio Off menu button was selected at the ViewStation or WorkStation and the Permanent option was selected.</td>
</tr>
<tr>
<td>Touch Screen OFF</td>
<td>The central station or remote touch screen is off.</td>
</tr>
<tr>
<td>System Service required - Contact Mindray or Authorized Mindray Distributor (100)</td>
<td>This message will be displayed after the central station and/or workstation automatically restarted because a system issue was detected.</td>
</tr>
<tr>
<td>System Service required - Contact Mindray or Authorized Mindray Distributor (101)</td>
<td>This message will be displayed if the system performed an auto-reboot due to the GUI process not running.</td>
</tr>
</tbody>
</table>
13.5 Operator Position

The operator of the device should be positioned in front of the Panorama Patient Monitoring Network Display at a distance of no more than 1m.

13.6 Alarm Verification

Alarms can be verified by creating an alarm condition on a connected patient monitor and verifying the alarm is present on the Panorama Patient Monitoring Network Display in a normal operator’s position.
14.0  Network Tab (Optional)

This section outlines how to configure a ViewStation or a WorkStation to display patient tiles from a Panorama Central Station(s) using the Network tab.

- Network Tab
- Troubleshooting
14.1 Network Tab

The **Network** tab (shown in FIGURE 14-1) is where the accessible patient tiles currently monitored on the C-LAN are displayed. These patient tiles may be selected for viewing at the ViewStation or the WorkStation. The **Network** tab can only be seen from a ViewStation or a WorkStation and is not available from the host central station. The ViewStation and WorkStation are independent of each other. For example, if a patient is added to a patient tile at a ViewStation, the WorkStation will not be affected unless the same patient is added to a patient tile at the WorkStation.

**NOTE:** The host central station(s) C-LAN configuration settings and the software installed at the remote location dictates the functionality available for each patient tile. For additional information, refer to "The ViewStation and the WorkStation" on page 2-4.

14.1.1 Accessing the Network Tab

- From the ViewStation or WorkStation menu bar (shown in FIGURE 2-8 on page 2-11), select the **Network** button. The **Network** tab is displayed (shown in FIGURE 14-1).

![FIGURE 14-1 Network Tab](image)

14.1.2 Sorting the Contents of the Network Tab List Box

When the **Network** tab is first accessed, it will automatically be sorted in ascending order, by the numerical tile assignment number. To sort by another column in the list box, select another column heading, and the rows will be resorted based on the selection. Any changes made to the list after it has been sorted, for example a tile assignment is removed, another **Network** tab dialog is opened from another remote location, or the **Network** tab is closed, will cancel the previous sort order.

14.1.3 Assigning Patient Tiles in the Network Tab

The **Add** button assigns a ViewStation or WorkStation tile number to the selected row in the **Network** tab list box. Once a patient in the **Network** tab list box is assigned a tile number, the patient’s data (which is monitored at the host central station) can be seen at the ViewStation or WorkStation.
NOTE: Once a tile is assigned to an associated tile at the ViewStation or the WorkStation, the association to the host patient tile will continue to exist, even if the patient is discharged or the monitoring equipment is removed from the Equipment List at the host central station. Therefore, when a new patient is admitted to the host central station, the patient data will automatically be displayed at the associated ViewStation or WorkStation tile.

NOTE: To determine which host patient tile the ViewStation or the WorkStation tile is associated with, scroll to the Central column in the Network tab.

To assign a patient tile in the Network tab:

1. Select a row in the Network tab list box that does not contain a tile assignment number.
2. Select the Add button. The Tile Number list box is displayed (shown in FIGURE 14-2).

![Figure 14-2: Tile Number List Box](image)

NOTE: The ViewStation or WorkStation single or dual display configuration dictates the maximum number of tiles that can be assigned. Table 14-1 shows how many tiles can be assigned when using a single or dual display configuration.

<table>
<thead>
<tr>
<th>VIEW ONLY WORKSTATION CONFIGURATION</th>
<th>NUMBER OF TILES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single display - 8 tiles</td>
<td>0 - 7</td>
</tr>
<tr>
<td>Single display - 12 tiles</td>
<td>0 - 11</td>
</tr>
<tr>
<td>Dual display - 24 tiles</td>
<td>0 - 23</td>
</tr>
</tbody>
</table>

3. Select a tile from the list box.
4. Accept or reject the selection.
   - Select the Done button to accept the tile assignment and display the patient’s data, from the assigned tile, at the ViewStation or WorkStation within five (5) seconds.
   - Select the Cancel button to reject the tile assignment and close the Tile Number list box.
14.1.4 Removing a Patient Tile Assignment in the Network Tab

The **Remove** button clears the selected rows’ patient tile assignment from the **Network** tab list box and discontinues displaying the selected host patient tile’s data at the ViewStation or WorkStation.

**NOTE:** Removing a patient tile assignment at the ViewStation or WorkStation will eliminate the connection to the associated host central station patient tile. Therefore, when a patient is admitted to the host central station tile, the patient data will not be displayed at the ViewStation or the WorkStation until another tile association has been established.

To remove a single tile assignment from the **Network** tab:

1. Select a row in the **Network** tab multi-column list box that contains a number in the **Tile Assignment** column.
2. Select the **Remove** button. The message **Are you sure?** is displayed.
   - Select the **Yes** button to remove the number displayed in the **Tile Assignment** column, discontinue displaying monitored data for the selected patient tile at the ViewStation or the WorkStation, and close the message box.
   - Select the **No** button to continue displaying monitored data for the selected patient tile at the ViewStation or the WorkStation, cancel the remove request, and close the message box.

14.1.5 Removing All Patient Tile Assignments in the Network Tab

The **Remove All** button clears all previously assigned patient tile numbers from the **Network** tab list box and discontinues displaying all of the patient tiles monitored data at the ViewStation or WorkStation.

To clear all the tile assignments from the **Network** tab:

- From the **Network** tab, select the **Remove All** button. The message **Are you sure?** is displayed.
- Select the **Yes** button to erase all the assigned tile numbers in the **Tile Assignment** column, discontinue displaying monitored data for all the tiles at the ViewStation or the WorkStation, and close the message box.
- Select the **No** button to continue displaying monitored data for all patients at the ViewStation or the WorkStation, cancel the remove all request, and close the message box.

14.1.6 Refreshing the Network Tab List Box

The **Refresh** button updates the **Network** tab list box for any changes that may have occurred while the **Network** tab was open. Once the **Refresh** button is selected, the **Network** tab list box will regenerate if:
Any new equipment is added at the host central station
Any existing equipment is removed from the host central station
Any existing equipment is edited at the host central station
Any patient demographic information (Last Name, First Name, Patient ID, Bed, Doctor) or Central Station name/tile number is edited at the host central station

14.1.7 Troubleshooting

This section lists some of the potential messages and issues that may occur while using the Network tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>The <strong>Add</strong> button disabled.</td>
<td>No row has been selected in the <strong>Network</strong> tab list box.</td>
<td>Select a row in the <strong>Network</strong> tab list box.</td>
</tr>
<tr>
<td></td>
<td>Selected row is already assigned to a tile at the ViewStation or WorkStation.</td>
<td>Select a different row.</td>
</tr>
<tr>
<td>The <strong>Remove</strong> button is disabled.</td>
<td>The selected row(s) in the <strong>Network</strong> tab does not currently have a tile assignment(s) or no tile(s) have been assigned at the ViewStation or WorkStation.</td>
<td>Establish tile assignment(s) at the ViewStation or WorkStation.</td>
</tr>
<tr>
<td>The <strong>Remove All</strong> button is disabled.</td>
<td>No tile(s) have been assigned at the ViewStation or WorkStation.</td>
<td>Establish tile assignment(s) at the ViewStation or WorkStation.</td>
</tr>
<tr>
<td><strong>You have reached the maximum number of channel licenses available.</strong></td>
<td>All the tiles at the ViewStation or WorkStation are currently occupied.</td>
<td>Remove a patient from one of the patient tiles at the ViewStation or WorkStation.</td>
</tr>
<tr>
<td>Patient tile no longer displaying patient data at ViewStation or WorkStation.</td>
<td>Communication between the host central station and the ViewStation or WorkStation has been interrupted (Central Communication Lost state).</td>
<td>Contact a Service Representative.</td>
</tr>
<tr>
<td>A Room to Room Without Device transfer out for the host tile has been completed.</td>
<td></td>
<td>Refresh the <strong>Network</strong> tab list box, and then assign the transferred patient tile to an available tile at the ViewStation or WorkStation.</td>
</tr>
<tr>
<td>A Central to Central Without Device transfer out for the host tile has been accepted.</td>
<td></td>
<td>Refresh the <strong>Network</strong> tab list box, and then assign the transferred patient tile to an available tile at the ViewStation or WorkStation.</td>
</tr>
<tr>
<td>Host tile at central station is in a Discharge state.</td>
<td></td>
<td>Retrieve patient from Discharge list, if possible, then reassign the retrieved patient tile to an available tile at the ViewStation or WorkStation.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>ViewStation or WorkStation tile goes into a Central Communication Lost state</td>
<td>Host tile at central station is in a Lost Communication state.</td>
<td>Reestablish communication at host central station, and if necessary, contact a Service Representative.</td>
</tr>
<tr>
<td>ViewStation or WorkStation tile goes into a Standby state</td>
<td>Host tile at central station is in a Standby state.</td>
<td>Wait for host patient tile to be removed from Standby state.</td>
</tr>
<tr>
<td>Patient tile not available in Network tab list box</td>
<td><strong>Network</strong> tab needs to be refreshed.</td>
<td>Select the <strong>Network</strong> tab <strong>Refresh</strong> button.</td>
</tr>
<tr>
<td></td>
<td>ViewStation or WorkStation is in a Central Communication Lost state.</td>
<td>Contact a Service Representative.</td>
</tr>
<tr>
<td></td>
<td>Patient not currently being monitored on any of the C-LAN central stations.</td>
<td>Admit the patient to a central station on the C-LAN, then assign the patient to an available tile at the ViewStation or WorkStation.</td>
</tr>
<tr>
<td>Host patient tile information is displayed at the host central station but not at the ViewStation or WorkStation</td>
<td>Association between host central station patient tile and ViewStation or WorkStation patient tile has not been established.</td>
<td>Assign the tile in the <strong>Network</strong> tab at the ViewStation or WorkStation.</td>
</tr>
<tr>
<td>Selected tile is currently assigned.</td>
<td>Attempted to assign a patient tile at the ViewStation or WorkStation that is already assigned.</td>
<td>Assign the patient to another tile number, if one is available.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
15.0  Paging Tab (Optional)

This section outlines the functionality of the **Paging** tab and describes how to configure pagers to properly function with the Panorama Central Station(s), when this option has been purchased.

- Paging Password Tab
- Paging Tab
15.1 Paging Password Tab

The Paging password tab (shown in FIGURE 15-1) provides password protected access to the system paging configuration tab. When the correct password is entered, the Paging tab is opened.

The Paging password tab is only shown if the Paging Password Protection option is set to ON in the Unit Choices tab. If Paging Password Protection is OFF, the Paging menu button provides direct access to the Paging tab. For additional information regarding the Unit Choices tab, refer to “Unit Choices Tab” on page 9-50.

The Paging tab is not available at the ViewStation.

15.1.1 Accessing the Paging Tab

From the Menu Bar, select the Paging menu button. The Paging password tab is displayed.

![FIGURE 15-1 Paging Password tab](image)

15.1.2 Entering a Password

To enter a password:

1. Select the Password text box in the Paging password tab. The keyboard dialog box is displayed.
2. Use the keyboard dialog box to enter the password that was created in the Passwords tab (maximum 15 characters). For security purposes asterisks (*) are shown for each typed character. For additional information regarding passwords, refer to the “Passwords Tab” on page 9-29 or contact your System Administrator.

**NOTE:** Passwords are case sensitive.

- The Backspace (<=) button clears a single character in the edit box
- The Caps Lock button enables/disables the caps lock function. A Caps Lock indicator is shown when Caps Lock is enabled.
- The Close button discards the changes
- The Enter button accepts changes
3. The **Paging** tab (shown in FIGURE 15-2) is displayed after the correct password is entered.

### 15.1.3 Sidebar Buttons

The following sidebar button is used to expand the functionality of this tab.

**Normal Screen**

Select the **Normal Screen** sidebar button to return the display to the full screen mode, and close all tabs, message boxes, and dialog boxes.

### 15.1.4 Troubleshooting

This section lists some of the potential messages and issues that may occur while using the **Paging** password tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Incorrect password. Please try again.</td>
<td>The wrong password was entered in the Password text box.</td>
<td>Select the <strong>OK</strong> button to close the dialog box. Enter the correct password in the text box.</td>
</tr>
<tr>
<td></td>
<td>The password was entered in the wrong case.</td>
<td>Re-enter the password turning on/off the keyboard dialog Caps Lock key.</td>
</tr>
<tr>
<td>Incorrect password. This is your last chance. Please try again</td>
<td>The wrong password was entered two consecutive times in the Password text box.</td>
<td>Select the <strong>OK</strong> button to close the dialog box. Enter the correct password in the text box.</td>
</tr>
<tr>
<td>The <strong>Paging</strong> password tab closes and the central station is returned to a full screen monitoring display.</td>
<td>The wrong password was entered three consecutive times in the Password text box.</td>
<td>Reopen the <strong>Paging</strong> password tab and enter the correct password in the text box.</td>
</tr>
</tbody>
</table>

*Messages are shown in all bold text.*
15.2 Paging Tab

The Paging tab allows for the configuration of pager information for a Panorama Central Station that has the paging option enabled. If the central station does not have a Paging button on the system Menu Bar, this is an indication that the workstation does not have the Paging option installed.

15.2.1 Accessing the Paging Tab

1. From the Menu Bar, select the Paging button.
2. If the Paging password tab is displayed, enter the password in the Password text box. The Paging tab is displayed (shown in FIGURE 15-2).

NOTE: The Paging tab list box is displayed in ascending order by pager number.

![FIGURE 15-2 Paging Tab](image)

15.2.2 Adding a Pager

To add a pager:

1. From the Paging tab, select the Add Pager sidebar button. The Pager Details dialog (shown in FIGURE 15-3) is displayed.

NOTE: The Pager Number and Pager Assignment are required fields.

![FIGURE 15-3 Pager Details Dialog](image)

2. Select the Pager Number text box. The keypad dialog box is displayed.
3. Use the numeric keypad to enter the pager number (maximum of 14 alphanumeric characters), then select the OK button. The numeric keypad closes and the Pager Details dialog is displayed.
NOTE: Pager numbers must be unique. If an existing pager number is entered, the message “Pager Number already exists! Please enter a new number.” is displayed.

4. Enter the name of the person who will be responsible for the pager or some other identifying information in the Pager Assignment text box (maximum of 30 alphanumeric characters).

NOTE: Pager assignments must be unique. If an existing pager assignment is entered, the message “This Pager Assignment already exists. Please enter a new assignment.” is displayed.

5. Accept or reject the entry made in the keyboard dialog box.
   - The Done button accepts the entry, closes the dialog box, and updates the Paging tab list box.
   - The Cancel button discards the entry, closes the dialog box, and returns to the Paging tab.

NOTE: The decimal button shall be disabled in the Pager Assignment keyboard.

15.2.3 Assigning a Patient Tile

To assign a patient tile to a pager:

1. From the Paging tab list box, select a pager.

2. Select the Add Tile button. The Tile Number dialog box (shown in FIGURE 15-4) is displayed. The number of tiles that are displayed is dependent on the number of configured central station displays and available patient tiles. For additional information, refer to TABLE 15-1.

FIGURE 15-4 Tile Number dialog box

### TABLE 15-1

<table>
<thead>
<tr>
<th>CENTRAL STATION CONFIGURATION</th>
<th>PATIENT TILE NUMBER</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single display - 8 tiles</td>
<td>0 - 7</td>
</tr>
<tr>
<td>Single display - 12 tiles</td>
<td>0 - 11</td>
</tr>
<tr>
<td>Dual display - 24 tiles</td>
<td>0 - 23</td>
</tr>
</tbody>
</table>
3. Select a tile number.

NOTE: The list box will only include patient tiles that are not currently assigned to the selected pager.

NOTE: A patient tile number can be assigned to a maximum of 10 pagers.

4. Accept or reject the selections made in the list box.
   - The Done button accepts the selection, closes the dialog box, and updates the Paging tab list box.
   - The Cancel button discards the selection, closes the dialog box, and returns to the Paging tab.

5. Repeat steps 2 - 4 to continue adding patient tiles to the selected pager, if necessary.

NOTE: Multiple tile assignments will be separated by a comma (,) in the Tile column.

15.2.4 Adding All Patient Tiles to a Pager
To assign all available patient tiles to a pager:

1. From the Paging tab list box, select a pager.

2. Select the Add All Tiles button. The Are you sure you want to Add All Tiles? question dialog box is displayed.
   - The Yes selection confirms the selection, closes the question dialog box, and updates the Tile column in the Paging tab list box.
   - The No selection cancels the selection, closes the confirmation dialog, and returns to the Paging tab list box.

15.2.5 Removing a Patient Tile Assignment
To remove an existing tile assignment from a pager:

1. From the Paging tab list box, select a pager.

2. Select the Remove Tile button. The Tile Number dialog box (shown in FIGURE 15-4) is displayed. The number of tiles that are displayed is dependent on the number of patient tiles assigned to the selected pager.

3. Select a patient tile.

4. Accept or reject the selections made in the list box.
   - The Done button accepts the selection, closes the dialog box, and returns to the Paging tab where the list box updates are displayed.
   - The Cancel button discards the selection, closes the dialog box, and returns to the Paging tab.

15.2.6 Removing All Patient Tiles from a Pager
To remove all patient tiles assignments from a pager:

1. From the Paging tab list box, select a pager.
2. Select the **Remove All Tiles** button. The **Are you sure you want to Remove All Tiles?** question dialog box is displayed.
   - The **Yes** selection confirms the selection, closes the question dialog box, and updates the Tile column in the **Paging** tab list box.
   - The **No** selection cancels the selection, closes the confirmation dialog, and returns to the **Paging** tab list box.

15.2.7 Editing a Pager

To edit an existing pager number or pager assignment:

1. Select the pager from the **Paging** tab list box.
2. Select the **Edit Pager** sidebar button. The Pager Details dialog box is displayed.

![FIGURE 15-5 Pager Details (Edit Mode)](image)

3. Edit the pager information.

   **NOTE:** The Pager Number and Pager Assignment are required fields.

4. Accept or reject the changes made in the dialog box.
   - The **Done** button accepts the changes, closes the dialog box, and returns to the **Paging** tab where the list box updates are displayed.
   - The **Cancel** button discards the changes, closes the dialog box, and returns to the **Paging** tab.

15.2.8 Removing a Pager

To remove a pager:

1. Follow steps 1 and 2 from the "Editing a Pager" section.
2. Select the **Remove Pager** button in the Pager Details dialog box.
3. Verify the selection. The **Are you sure you want to delete this pager and all its assignments?** question dialog box is displayed.
   - The **Yes** selection confirms the deletion, closes the question dialog box, and removes the pager from the **Paging** tab list box.
   - The **No** selection cancels the deletion, closes the confirmation dialog, and returns to the **Paging** tab list box.
15.2.9 Resending a Failed Page

If an attempt was made to send a page and was unsuccessful, the page will be added to the Failed Pages view. The Failed Pages view contains a list box that can display a maximum of 100 failed pages. The list box orders the failed pages chronologically in descending order.

NOTE: If the system is reset, all failed pages listed in the “Failed Pages” view will be removed.

To resend a failed page:

1. Select the Failed Page sidebar button. The Failed Pages view (shown in FIGURE 15-6) is displayed.

NOTE: The Failed Page sidebar button will only be available if there is failed page information in the Failed Pages view.

FIGURE 15-6 Failed Pages View

2. Select a failed page from the list box using the navigational buttons.

3. Select the Resend sidebar button. The failed page is resent and the row is removed from the list box if the page was successfully resent. If the page was not successfully resent, a new row will be added to the top of the Failed Pages view list box with an updated date and time stamp.

Navigating in the Failed Pages View

To navigate in the Failed Pages view, select any of the following buttons to:

<table>
<thead>
<tr>
<th>BUTTON</th>
<th>DESCRIPTION</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="Image" alt="Button" /></td>
<td>Display the oldest failed pages in the list box.</td>
</tr>
<tr>
<td><img src="Image" alt="Button" /></td>
<td>Scroll down the list one page at a time.</td>
</tr>
<tr>
<td><img src="Image" alt="Button" /></td>
<td>Scroll down one row in the list at a time.</td>
</tr>
<tr>
<td><img src="Image" alt="Button" /></td>
<td>Scroll up one row in the list at a time.</td>
</tr>
</tbody>
</table>
15.2.10 Generating a Paging Report

To review all of the currently assigned pagers, determine who each pager is assigned to, and to which tile the pager is assigned, generate a Paging Report. This report can be generated directly from the Paging tab by selecting the Print button from the Menu Bar.

For additional information regarding the Paging Report, refer to the “Paging Report” on page 8-36.

15.2.11 Troubleshooting

This section lists some of the potential messages and issues that may occur while using the Paging tab.

<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>You have reached the maximum number of pagers that can be assigned to a tile.</td>
<td>A patient tile cannot be assigned to more than 10 pagers.</td>
<td>Remove one pager assignment from the patient tile.</td>
</tr>
<tr>
<td>The Add Tile button is disabled.</td>
<td>There are no pagers listed in the Paging tab’s list box.</td>
<td>Add a pager and then assign it to a tile.</td>
</tr>
<tr>
<td>All the patient tiles at the central station are already assigned to the selected pager in the Paging tab’s list box.</td>
<td>Assign the pager to another patient tile.</td>
<td></td>
</tr>
<tr>
<td>The Remove Tile button is disabled.</td>
<td>There are no pagers listed in the Paging tab’s list box.</td>
<td>Add a pager and then assign it to a tile.</td>
</tr>
<tr>
<td>None of the patient tiles at the central station are assigned to the selected pager in the Paging tab’s list box.</td>
<td>Assign a pager to one of the patient tiles.</td>
<td></td>
</tr>
<tr>
<td>Are you sure you want to Add All Tiles?</td>
<td>The Add All Tiles button was selected in the Paging tab’s list box.</td>
<td>Confirm that this was the intended function, then select the Yes button to confirm or the No button to cancel the function and return to the Paging tab.</td>
</tr>
<tr>
<td>Some tiles could not be assigned to this pager.</td>
<td>The Add All Tiles button was selected.</td>
<td>Some of the patient tiles already contain 10 pager assignments.</td>
</tr>
<tr>
<td>Are you sure you want to Remove All Tiles?</td>
<td>The Remove All Tiles button was selected in the Paging tab’s list box.</td>
<td>Confirm that this was the intended function, then select the Yes or No button.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
<table>
<thead>
<tr>
<th>MESSAGE/ISSUE *</th>
<th>REASON</th>
<th>SOLUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Are you sure that you want to delete the selected event/s?</td>
<td>The Delete or Delete All sidebar button was selected while in the Failed Pages view.</td>
<td>Confirm that this was the intended function, then select the Yes button or the No button.</td>
</tr>
<tr>
<td></td>
<td>The Done button in the Pager Details dialog box is disabled.</td>
<td>Enter a Pager Number and/or Pager Assignment in the Pager Details dialog box.</td>
</tr>
<tr>
<td>Pager Number already exists! Please enter a new number.</td>
<td>Pager numbers must be unique. The pager number entered is a duplicate of an existing pager number.</td>
<td>Enter a unique pager number, then select the OK button.</td>
</tr>
<tr>
<td></td>
<td>No data has been entered into the Pager Number and/or the Pager Assignment text box.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>A maximum of 15 alphanumeric characters can be entered to the Pager Number field.</td>
<td>Verify the Pager Number is correct.</td>
</tr>
<tr>
<td></td>
<td>A maximum of 30 alphanumeric characters can be added to the Pager Assignment field.</td>
<td>Abbreviate the assignee name or identifying title.</td>
</tr>
<tr>
<td>A failed page was removed from the Failed Pages list box.</td>
<td>The failed page was successfully resent which removes the page from the Failed Pages view.</td>
<td>Locate the event in the Events List and verify that a page was sent for the event.</td>
</tr>
<tr>
<td></td>
<td>The list box maximum of 100 records was reached and the failed page was automatically deleted from the database.</td>
<td>Periodically delete or resend the failed pages from the Failed Pages list box to avoid the automatic deletion of failed pages.</td>
</tr>
<tr>
<td></td>
<td>A failed page was removed from the Failed Pages list box.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>If the system is reset, all failed pages listed in the “Failed Pages” view will be removed.</td>
<td></td>
</tr>
<tr>
<td>The selected event has been deleted.</td>
<td>The failed page event has been deleted from the database because it reached its maximum number of records.</td>
<td>Select another failed page from the list box.</td>
</tr>
<tr>
<td>You have reached the maximum number of pagers.</td>
<td>The Add Pager sidebar push button was selected when the the maximum number of pagers has already been assigned at the central station.</td>
<td>Review the list and remove any unused pagers from the list.</td>
</tr>
<tr>
<td>This Pager Assignment already exists. Please enter a new assignment.</td>
<td>The pager assignment entered in the Pager Assignment text box is a duplicate of a pager assignment that already exists.</td>
<td>Enter a different pager assignment.</td>
</tr>
</tbody>
</table>

* Messages are shown in all bold text.
16.0  

Performance Specifications

This section outlines performance specifications for the Panorama Patient Monitoring Network.

- Parameter Performance
- Panorama Central Station and Panorama Telemetry Server
- Keyboard
- Mouse
- Displays
- Network Printer
- Telepack Specifications
- Nonin SpO2 Module Specifications
16.1 Parameter Performance

16.1.1 ECG Performance Requirements

Three Lead Displayable Leads: I, II, III (one vector at a time)

Five Lead Displayable Leads: I, II, III, aVR, aVL, aVF, V

Twelve Lead Displayable Leads: I, II, III, aVR, aVL, aVF, V1, V2, V3, V4, V5, and V6

Standardizing Voltage: The ECG size is indicated by a scale bar displayed on the left of the waveform window with a scale indicating the height of the bar in mV. Printed output to have standardizing pulse printed at beginning of strip

Time base selection (Non-permanent display): 25 mm/s

Time base error (Non-permanent display): ±10%

Impulse Response: 0.1 mV max
0.3 mV/sec max slope

Aspect Ratio: 0.4 ± 0.08 sec/mV (met by 25 mm/s)

Overall System Error: Greater of 5% or 40 µV
ANSI/AAMI EC13-2002/(R)2007, 4.2.7.1 (a)

ECG Input dynamic range: +/− 5mV max

Channel Width: 30 mm. min
<table>
<thead>
<tr>
<th>Performance Specifications</th>
<th>Parameter Performance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Waveform Displays</td>
<td>Each waveform in the All Strips, Event List Zoom-In, Full Disclosure Zoom-In and ST reports will be displayed on a calibrated grid (5mm x 5mm) that conforms to ANSI/AAMI EC13: 2002/(R)2007.4.2.9.7d. Each second of the waveforms in the All Strips, Event List Zoom-In, Full Disclosure Zoom-In and ST reports will be marked with a time annotation marker that conforms to ANSI/AAMI EC13: 2002/(R)2007.4.2.9.7e.</td>
</tr>
<tr>
<td>Heart Rate Meter Accuracy</td>
<td>Accuracy is ± 3 bpm or ± 3% in the range of 20 bpm to 250 bpm, whichever is greater Accuracy is ± 5% in the range of 251 bpm to 300 bpm.</td>
</tr>
<tr>
<td>Heart Rate Averaging Method</td>
<td>In compliance with the requirements in Clause 4.1.2.1 (d) of ANSI/AAMI EC13-2002, the following method is used: The average heart rate is calculated on the basis of the mean RR-interval of the last 16 beats, unless the heart rate calculated using the last 4 beats is less than or equal to 48, then this rate is used. The heart rate value is updated on the display every second.</td>
</tr>
<tr>
<td>Response to Irregular Rhythm</td>
<td>In compliance with the requirements in Clause 4.1.2.1 (e) of ANSI/AAMI EC13:2002/(R)2007, the indicated heart rate after 20 seconds of stabilization period is: • Ventricular bigeminy (3a): 80 bpm • Slow alternating ventricular bigeminy (3b): 60 bpm • Rapid alternating ventricular bigeminy (3c): 120 bpm • Bidirectional systoles (3d): 90 bpm Accuracy for above is ± 3 bpm or ± 3%, whichever is greater.</td>
</tr>
<tr>
<td>Time to Alarm for Tachycardia</td>
<td>The alarm response time for Tachycardia is less than 10 seconds in accordance with ANSI/AAMI EC13-2002/(R)2007 Section 4.1.2.1g.</td>
</tr>
</tbody>
</table>
Pacer Rejection: The heart rate meter rejects all pacer pulses of amplitude signals from ±2mV to ±700mV and duration 0.1ms to 2ms with no tail according to ANSI/AAMI EC13-2002/(R)2007 Section 4.1.4.1 (No tails).

Pacemaker Pulse Rejection Capability: The pacer detector's slew rate is 6.024V/s according to ANSI/AAMI EC13-2002/(R)2007 Section 4.1.4.3.

Tall T-Wave Rejection: Tall T-wave rejection is as per ANSI/AAMI EC13:2002/(R)2007 Section 4.1.2.1 (c). The heart rate meter rejects all T-waves with amplitudes less than 120% of a 1mV, 100 msec QRS, and a T wave duration of 180ms and a Q-T interval of 350 msec.

Response Time of Heart Rate Meter to Change in Heart Rate: The response time of the heart rate meter to change in heart rate is less than 10 seconds for step increase from 80 to 120 BPM and less than 10 seconds for step decrease from 80 to 40 BPM as per ANSI/AAMI EC13:2002/(R)2007 Section 4.1.2.1 (f).

Lead-off Detection Current: Measuring electrode: <0.1 µA
Drive electrode: <1 µA

NOTE: The heart rate meter does not reject Pacemaker pulses with tails.

NOTE: The heart rate meter does not reject Pacemaker with fast ECG signals.

NOTE: There is no Pacemaker pulse auxiliary output.

16.1.2 ECG Derived Heart Rate Meter Performance Requirements

| Range: | 20 to 300 bpm Adult/Pediatric ANSI/AAMI EC13–2002/(R)2007, 4.2.7 |
|        | 20 to 350 bpm Neonatal ANSI/AAMI EC13–2002/(R)2007, 4.2.7 |
| Resolution: | 1 bpm |
16.1.2.1 Alarm Response

High HR Alarm Range
- Off
- 60-250 bpm Adult
- 100-300 bpm Pediatric
- 100-350 bpm Neonatal
(ANSI/AAMI EC13-2002/(R)2007, 4.2.8.1)

Low HR Alarm Range
- Off
- 30-120 bpm Adult
- 30-150 bpm Pediatric
- 30-200 bpm Neonatal
(ANSI/AAMI EC13-2002/(R)2007, 4.2.8.1)

Extreme Bradycardia Alarm Range
- Off
- 25-200 bpm Adult
- 25-200 bpm Pediatric
- 25-200 bpm Neonatal

Extreme Tachycardia Alarm Range
- Off
- 50-300 bpm Adult
- 50-300 bpm Pediatric
- 50-350 bpm Neonatal

Resolution
- 5 bpm max.
(ANSI/AAMI EC13-2002/(R)2007, 4.2.8.2)

Accuracy
- ± 5 bpm or ± 10%
(ANSI/AAMI EC13-2002/(R)2007, 4.2.8.3)

Time to alarm
- Step Change Response Time
- Less than 10 seconds for 60 bpm low limit alarm to sound when stepping from a heart rate of 80 to 40 bpm
(ANSI/AAMI EC13-2002/(R)2007, 4.2.8.4 and 4.2.8.5).
- Less than 10 seconds for 100 bpm high limit alarm to sound when stepping from a heart rate of 80 to 120 bpm
(ANSI/AAMI EC13-2002/(R)2007, 4.2.8.6)

16.1.3 ST Segment Analysis

Enabling:
- Enabled in Adult and Pediatric (from a bedside monitor only) modes only. ST for Telepack units may be enabled at the Panorama Central Station.
Default ST Measurement Point: 80 ms after J point for heart rates ≤120 bpm
60 ms after the J point for heart rates >120 bpm

**User Selectable ST**

Measurement Points: 40, 60 and 80 ms after J point (heart rate independent)
Or 60/80 Heart Rate Dependent

Default ISO Point: Located between the P and Q waves.
User adjustable ISO Point: User adjustable from “R peak” - 10 ms to “R peak” - 200 ms in 8 ms increments.

Default J Point: The end of the QRS complex.
User adjustable J Point: User adjustable from “R peak” - 10 ms to “R peak” - 200 ms in 8 ms increments.

Numerical ST Update Rate: 3/5 Lead Wire: every 10 seconds
View 12 card: every 2 seconds

**16.1.4 Arrhythmia Analysis**

Arrhythmia analysis identifies ventricular arrhythmia.

The following arrhythmias are detectable:

- Asystole
- Bigeminy
- Bradycardia (Brady)
- Couplet
- Irregular Heart Rate
- PVCs per minute
- R-On-T PVC
- Pauses per minute
- QTc (V 12/V 21 and Telepack only)

**16.1.5 NIBP Performance Characteristics**

**Systolic Pressure Measurement**

- Range: 55 to 235 mmHg in Adult mode
- 55 to 160 mmHg in Pediatric mode
- 45 to 120 mmHg in Neonatal mode

- Resolution: 1 mmHg

**Diastolic Pressure Measurement**
16.1.6 Pulse Rate

Range: 35-245 bpm, for Adults/Pediatric
       70–245 bpm, for Neonate

Resolution: 1 bpm

16.1.7 IBP

IBP Pressure Range

Range: -50 to 300 mmHg

IBP Heart Rate Meter

Range: 30 to 300 bpm Adult/Pediatric
       30 to 333 bpm Neonatal

Resolution: 1 bpm

16.1.8 Temperature Performance Requirements

Scale: Selectable Celsius or Fahrenheit

Range: 15 °C to 45 °C
       59 °F to 113 °F

Resolution: 0.1 °C
            0.1 °F

16.1.9 ECG Respiration Performance Requirements

Range: 4 to 199 breaths per minute

Resp. Scale: 1, 2, 3, 4 or 5 with standard ECG cable

16.1.10 SpO2

SpO2 Range: 70% to 100%

Pulse Rate Range: 30 to 235
16.1.10.1 Nonin SpO₂ (Telepack Only)

SpO₂ Range: 0% to 100%
Pulse Rate Range: 18 to 300

* Measurement accuracy verification: The SpO₂ accuracy has been verified in human experiments by comparing with arterial blood sample reference measured with a co-oximeter.

** The testing consisted of 12 healthy consenting subjects who were non-smokers and between approximately nineteen to thirty-five years of age. There was a moderate range of skin tones from light to dark. A pool of seven male and five female subjects were included in the study.

16.1.11 CO₂ Performance Requirements

Range: 0 - 13%:

Respiration Rate Range: 0 - 150 breaths/minute

16.1.11.1 CO₂ Alarm

ET CO₂ High Alarm Range: Off, 20 to 80 mmHg

ET CO₂ Low Alarm Range: Off, 5 to 50 mmHg

Insp. CO₂ High Alarm Range: Off, 5 to 30 mmHg

16.1.12 BIS Performance Requirements

Range: 0 to 100

Resolution: 1
16.2 Panorama Central Station and Panorama Telemetry Server

Panorama Central Station hardware consists of a computer, hard drives, Ethernet, display controllers, case and a power supply.

There is one basic hardware platform for the Panorama Central Station computer system. It contains the following:

- CPU Motherboard and processor
- 1 GB of RAM
- 1280 x 1024 dpi 24-bit (16.7 million colors) display controllers
- 280 GB system hard drive, minimum
- Four channel serial interface for touch screen inputs, Telepack programming, and future features
- 2 on-board Ethernet connectors
- CD-R/W

Panorama Telemetry Server hardware consists of a computer, hard drives, Ethernet, display controllers, case and a power supply. The system is populated with the following:

- CPU Motherboard and processor
- 1 GB of RAM, minimum
- 800 x 600 dpi display controllers
- 34 GB system hard drive, minimum
- 1 channel serial port
- 2 on-board Ethernet connectors
- CD-R/W

16.2.1 Real Time Clock

This is used for various time-related functions in the Panorama Central Station and the Panorama Telemetry Server.
16.2.2 Power Supply

Voltage
- 100-240 VAC (±10%)

Frequency
- 60/50 Hz (±3Hz)

Panorama Central Station Power Consumption Maximum
- 115 VAC @ 1.4 A + 20%
- 230 VAC @ 0.6 A + 20%

16.2.3 Operating Temperature
- +10 to +40 °C

16.2.4 Operating Humidity
- 20% to 80% Relative Humidity Maximum, non-condensing, maximum wet bulb 29 °C

16.2.5 Operating Altitude
- 1060 to 700 hPa (-1250 to 9889 feet ASL) (-380 - 3014 m) (795 to 525 mmHg)

16.2.6 Storage Temperature
- -20 to +60 °C

16.2.7 Storage Humidity
- 5% to 80% Relative Humidity Maximum, non-condensing, maximum wet bulb 29 °C

16.2.8 Storage Altitude
- 1060 to 700 hPa (-1250 to 9889 feet ASL) (-380 - 3014 m) (795 to 525 mm Hg)
16.2.9 Standards
The Panorama Central Station and Panorama Telemetry Server comply with the following standards:

SAFETY
UL 60950-1:2011
CSA-C22.2 60950-1-03
IEC 60601-1-8:2012

SAFETY DESIGNATIONS
(SEE SECTION 16.2.11 (TABLE 16-1, TABLE 16-2, TABLE 16-3, TABLE 16-4)
Mode of Operation Continuous
Protection Against Hazards of Explosion Non Protected (ordinary)
Protection Against Ingress of Liquids IPX0 (ordinary, non-protected)
Degree of Mobility Stationary, Intra Hospital

PERFORMANCE
AAMI/ANSI EC 53:2013 American national standard for ECG cables and leadwires
ANSI/AAMI EC 57:2012 Testing and reporting performance results of cardiac rhythm and ST-segment measurement algorithms
IEC 60601-2-27 Ed.3.0:2011 Particular requirements for the basic safety and essential performance of electrocardiographic monitoring equipment
ISO 80601-2-61 Ed.1.0:2011 Particular requirements for basic safety and essential performance of pulse oximeter equipment
FCC 47CFR, Part 95, Sub Part H, Wireless Medical Telemetry Service (WMTS), for 608 MHz devices and components of the system

SOFTWARE
IEC 62304 Ed.1.0: 2006-05 Software life cycle processes

USABILITY
IIEC 60601-1-6 Ed.3.1:2013 Usability

HAZARD ANALYSIS
ISO 14971:2007/(R) 2010 Application of Risk management to medical devices

ENVIRONMENTAL / EMC
IEC 60601-1-2:2007 Electromagnetic compatibility

SHIPPING
ISTA Test Procedure 2A Shipping Integrity test
16.2.10 Information Display and Control

16.2.10.1 Audio Indicators

Audio alarms are in accordance with IEC60601-1-8.

The measured sound pressure level for the high audio alarm is 45 dB to 85 dB.

The measured sound pressure level for the medium audio alarm is 45 dB to 85 dB.

The measured sound pressure level for the low audio alarm is 45 dB to 85 dB.

16.2.10.2 Recovery from Power Loss

16.2.10.2.1 Alarm Restoration from Power Loss

If the Panorama system experiences a power loss, all alarm settings will be restored upon power reset. For the Telepack, alarm settings will be restored from the Panorama Patient Monitoring Central Station.

**CAUTION:** If the Panorama system experiences a communication loss for more than 60 seconds, all Telepack ST and/or arrhythmia alarms that were active at the time of communication loss will not produce central station alarm notifications once communication is restored. This is due to the automatic relearn function of the ST and/or arrhythmia algorithm after communication is restored.

16.2.10.2.2 Data Logging after Power Loss

All logs will be restored following a power loss of any duration. Logs consist of the event log (service only), error log (service only), and the patient database.

**NOTE:** The Panorama Central Station can store a maximum of 1000 logs. When the logs reaches the maximum amount, the earliest log is cleared, and the new log is added.

16.2.10.3 Alarm Delay

The maximum alarm delay is 2 seconds in the distributed alarm system.
16.2.11 Electromagnetic Compatibility


NOTE: The Panorama Central Station, Panorama Telemetry Server, and Panorama Wireless Transceiver (120V only) needs special precautions regarding Electro Magnetic Compatibility (EMC) and needs to be installed and put into service according to the EMC information provided in this section.

NOTE: Portable and mobile RF communications equipment can affect the Panorama Central Station and Panorama Telemetry. Refer to Table 16-1 on page 16-13, Table 16-2 on page 16-14, and Table 16-4 on page 16-16.

TABLE 16-1
GUIDANCE AND DECLARATION - ELECTROMAGNETIC EMISSIONS

The Panorama Central Station, Telemetry Server, and Panorama Wireless Transceiver (120V only) is intended for use in the electromagnetic environment specified below. The customer or the user of the Panorama Central Station and Telemetry Server should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>EMISSIONS TEST</th>
<th>COMPLIANCE</th>
<th>ELECTROMAGNETIC ENVIRONMENT - GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF emissions CISPR 11</td>
<td>Group 1</td>
<td>The Panorama Central Station, Telemetry Server, and Panorama Wireless Transceiver uses RF energy only for its internal function and external data transmission. However, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.</td>
</tr>
<tr>
<td>Harmonic emissions IEC 61000-3-2 *</td>
<td>Class A</td>
<td>The Panorama Central Station, Telemetry Server, and Panorama Wireless Transceiver is suitable for use in all establishments other than domestic and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.</td>
</tr>
<tr>
<td>Voltage fluctuations/ Flicker emissions IEC 61000-3-3 *</td>
<td>Complies</td>
<td></td>
</tr>
</tbody>
</table>

* Panorama Central Station and Panorama Telemetry Server only
TABLE 16-2

GUIDANCE AND DECLARATION - ELECTROMAGNETIC IMMUNITY

The Panorama Central Station, Telemetry Server, and Panorama Wireless Transceiver is intended for use in the electromagnetic environment specified below. The customer or the user of the Panorama Central Station and Telemetry Server should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>IMMUNITY TEST</th>
<th>IEC 60601 TEST LEVEL</th>
<th>COMPLIANCE LEVEL</th>
<th>ELECTROMAGNETIC ENVIRONMENT - GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrostatic discharge (ESD) IEC 61000-4-2</td>
<td>±6 kV contact ±8 kV air</td>
<td>±6 kV contact ±8 kV air</td>
<td>Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30%.</td>
</tr>
<tr>
<td>Electrical fast transient/burst IEC 61000-4-4</td>
<td>±2 kV for power supply lines ±1 kV for input/output lines</td>
<td>±2 kV for power supply lines ±1 kV for input/output lines</td>
<td>Mains power quality should be typical of a commercial or hospital environment.</td>
</tr>
<tr>
<td>Surge IEC 61000-4-5</td>
<td>±1 kV differential mode ±2 kV common mode</td>
<td>±1 kV differential mode ±2 kV common mode</td>
<td>Mains power quality should be typical of a commercial or hospital environment.</td>
</tr>
<tr>
<td>Voltage dips, short interruptions and voltage variations on power supply input lines IEC 61000-4-11</td>
<td>&lt;5% U_T [&gt;95% dip in U_T] for 0,5 cycle</td>
<td>&lt;5% U_T [&gt;95% dip in U_T] for 0,5 cycle</td>
<td>Mains power quality should be typical of a commercial or hospital environment. If the user requires continued operation during power mains interruptions, it is recommended that the device be powered from an uninterruptible power supply or a battery.</td>
</tr>
<tr>
<td></td>
<td>40% U_T [60% dip in U_T] for 5 cycles</td>
<td>40% U_T [60% dip in U_T] for 5 cycles</td>
<td></td>
</tr>
<tr>
<td></td>
<td>70% U_T [30% dip in U_T] for 25 cycles</td>
<td>70% U_T [30% dip in U_T] for 25 cycles</td>
<td></td>
</tr>
<tr>
<td></td>
<td>&lt; 5% U_T [&gt;95% dip in U_T] for 5 sec</td>
<td>&lt; 5% U_T [&gt;95% dip in U_T] for 5 sec</td>
<td></td>
</tr>
<tr>
<td>Power frequency (50/60 Hz) magnetic field IEC 61000-4-8</td>
<td>3 A/m</td>
<td>3 A/m</td>
<td>Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.</td>
</tr>
</tbody>
</table>

U_T is the a.c. mains voltage prior to application of the test level.

NOTE: Panorama Central Station, Telemetry Server, and Panorama Wireless Transceiver, that are rack mounted, shall be powered down prior to cable and/or Panorama Licensing Key attachment or disconnect.
## TABLE 16-3

**GUIDANCE AND DECLARATION - ELECTROMAGNETIC IMMUNITY**

The Panorama Central Station, Telemetry Server, and Panorama Wireless Transceiver is intended for use in the electromagnetic environment specified below. The customer or the user of the Panorama Central Station and Telemetry Server should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>IMMUNITY TEST</th>
<th>IEC 60601 TEST LEVEL</th>
<th>COMPLIANCE LEVEL</th>
<th>ELECTROMAGNETIC ENVIRONMENT - GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conducted RF IEC 61000-4-6</td>
<td>3 Vrms 150 kHz to 80 MHz</td>
<td>3 Vrms</td>
<td>Portable and mobile RF communications equipment should be used no closer to any part of the Telepack, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter.</td>
</tr>
<tr>
<td>Radiated RF IEC 61000-4-3</td>
<td>3 V/m 80 MHz to 2.5 GHz</td>
<td>3 V/m</td>
<td>Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey should be less than the compliance level in each frequency range.</td>
</tr>
</tbody>
</table>

Where $P$ is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and $d$ is the recommended separation distance in meters (m).

Interference may occur in the vicinity of equipment marked with the following symbol: 

\[ d = 1.2 \times \sqrt{P} \]

\[ d = 2.3 \times \sqrt{P} \]

\[ 800 \text{ MHz to } 2.5 \text{ GHz} \]

\[ P \]

\[ P \]

\[ P \]

**NOTE:** At 80 MHz and 800 MHz, the higher frequency range applies.

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

\[ a \] Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment, due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength where the Telepack is used exceeds the applicable RF compliance level, the Telepack should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as reorienting or relocating the Telepack.

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

**NOTE:** Panorama Central Station equipment should not be used in the presence of high energy amplitude modulated (AM) signals that are operating within the audible range. These signals are not normally found in the hospital environment.
TABLE 16-4

RECOMMENDED SEPARATION DISTANCES BETWEEN PORTABLE AND MOBILE RF COMMUNICATIONS EQUIPMENT AND THE PANORAMA CENTRAL STATION AND TELEMETRY SERVER

The Panorama Central Station, Telemetry Server, and Panorama Wireless Transceiver is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the Panorama Central Station, Telemetry Server, and Panorama Wireless Transceiver can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the Panorama Central Station, Telemetry Server, and Panorama Wireless Transceiver as recommended below, according to the maximum output power of the communications equipment.

<table>
<thead>
<tr>
<th>RATED MAXIMUM OUTPUT POWER OF TRANSMITTER W (WATTS)</th>
<th>SEPARATION DISTANCE ACCORDING TO FREQUENCY OF TRANSMITTER M (METERS)</th>
</tr>
</thead>
<tbody>
<tr>
<td>150 KHZ TO 800 MHZ</td>
<td>800 MHZ TO 2.5 GHZ</td>
</tr>
<tr>
<td>D = 1.2 x \sqrt{P}</td>
<td>D = 2.3 x \sqrt{P}</td>
</tr>
<tr>
<td>150 KHZ TO 800 MHZ</td>
<td>800 MHZ TO 2.5 GHZ</td>
</tr>
<tr>
<td>D = 1.2 x \sqrt{P}</td>
<td>D = 2.3 x \sqrt{P}</td>
</tr>
</tbody>
</table>

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

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

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

16.2.12 Wireless Essential Performance

The Panorama Central Station has the following wireless essential performance:

- Network connection
- Data transmission
- Caution for disconnection
16.2.13  **Physical Characteristics**

**Maximum Size (Rack Mount (2U) CPU Case)**
- Depth: 28.0" (711.2 mm)
- Height: 3.5" (88.9 mm)
- Width: 19.3" (490.2 mm)

**Maximum Weight**
- 36.0 lbs (16.33 kg)

**Maximum Size (Vertical CPU Case)**
- Depth: 17.5" (445.5 mm)
- Height: 17.0" (431.8 mm)
- Width: 7.5" (190.5 mm)

**Maximum Weight**
- 30.0 lbs (13.6 kg)

16.3  **Keyboard**

- Keyboard is supported for system diagnostics only.
- This keyboard is offered in English language only and is not required for normal operation, but will be required for non-application system maintenance.
- The System will not require a keyboard to boot up and enter Panorama Central Station application software.

16.4  **Mouse**

- The mouse is supported for system diagnostics and user interface.
- Supported as an interface control device.
- The mouse operates independent of a touch screen interface.
- The mouse/touch screen is used together with the display to make selections from various menus and screens.

16.5  **Displays**

16.5.1  **21" CRT Display**
(Used with earlier versions of the Panorama Central Station software.)

This is an integrated monitor that includes an internal touch screen controller and has the following features:
16.5.2 20" CRT Display
(Used with earlier versions of the Panorama Central Station software.)

This is an integrated monitor that includes an internal touch screen controller and has the following features:

- SXGA
- 1280 x 1024 dpi capability
- 0.27mm dot pitch
- 19" viewable image measured diagonally

16.5.3 19" Flat Panel
(Used with current versions of the Panorama Central Station software.)

This is an integrated monitor that includes an internal touch screen controller and has the following features:

- SXGA
- 1280 x 1024 dpi capability
- 0.28mm dot pitch
- 19.0" viewable image measured diagonally

16.5.4 18.1" Flat Panel
(Used with earlier versions of the Panorama Central Station software.)

This is an integrated monitor that includes an internal touch screen controller and has the following features:

- SXGA
- 1280 x 1024 dpi capability
- 0.28mm dot pitch
- 18.1" viewable image measured diagonally

**WARNING:** The 18.1" flat panel may tip over, if the display head is inclined to an angle greater than 45° backward tilt. If the user elects to have the display head inclined to an angle greater than 45° backward tilt, the flat panel must be attached to a secure mounting surface via three screw locations on the bottom of the base.
16.5.5 Touch Screen

- A Surface Acoustical Wave (SAW) touch screen is supported as the standard input device on the 18.1", 19", 20", and/or 21" display.
- The touch screen operates independent of a mouse interface.
- The actual touch screen and its controller are integrated into the display.
- The touch screen communication is accepted by Panorama Central Station via a serial port.
- Use of the touch screen does not preclude the use of a mouse.

16.6 Network Printer

- The network printer is the standard hard copy output device for the output of Panorama Central Station waveform strip-charts and reports.
- There is no provision to support a printer mounted within Panorama Central Station CPU enclosure.
- The printer connects to the system via the Panorama Central Network (C-LAN).
- Two network laser printers are supported per Panorama Central Station.

For additional information, refer to the Panorama Printer Configuration Guide (P/N 0070-00-0561).

16.6.1 Requirements

Communications connection: 10/100 Base-TX Ethernet with internal print server
Speed: 17 pages per minute printing at full resolution
Resolution: 600-dpi output
Media: Plain paper sheets
Media sizes: 8.5 in. by 11 in. and A4

**NOTE:** Please refer to the manufacturer's printer manual for specific instructions regarding the printer.
16.7 **Wireless Communication**

The Panorama Central Station can communicate with the monitor with Wi-Fi function. The specifications are as follows.

<table>
<thead>
<tr>
<th>Specification</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Protocol</td>
<td>IEEE 802.11b/g/n</td>
</tr>
<tr>
<td>Modulation mode</td>
<td>DSSS and OFDM</td>
</tr>
<tr>
<td>Operating frequency</td>
<td>2412–2472 MHz</td>
</tr>
<tr>
<td>Channel spacing</td>
<td>5 MHz</td>
</tr>
<tr>
<td>Wireless baud rate (data rate)</td>
<td>IEEE 802.11b: 1–11 Mbps</td>
</tr>
<tr>
<td></td>
<td>IEEE 802.11g: 6–54 Mbps</td>
</tr>
<tr>
<td></td>
<td>IEEE 802.11n: 6.5–150 Mbps</td>
</tr>
<tr>
<td>Output power (transfer power)</td>
<td>&lt; 20 dBm (CE requirement: detection mode – RMS)</td>
</tr>
<tr>
<td></td>
<td>&lt; 30 dBm (FCC requirement: detection mode – peak power)</td>
</tr>
<tr>
<td>Operating mode</td>
<td>Infrastructure</td>
</tr>
<tr>
<td>Data security</td>
<td>WPA2-PSK encryption supported</td>
</tr>
<tr>
<td>Data integrity</td>
<td>Code error (UDP Layer) &lt; 3%</td>
</tr>
<tr>
<td>Data latency</td>
<td>Total delay of the wireless patient monitor transmitting data to the central station &lt;= 2s.</td>
</tr>
<tr>
<td></td>
<td>Total delay of the central station configuring related settings of the wireless patient monitor &lt;= 2s.</td>
</tr>
<tr>
<td>Priority</td>
<td>All communication data types shall have the same priority.</td>
</tr>
<tr>
<td>Transmission distance</td>
<td>Distance of distinct vision from the wireless patient monitor to AP shall be no less than 50 m.</td>
</tr>
<tr>
<td>Roaming</td>
<td>Network switchover is automatically implemented when the wireless patient monitor moves from the coverage area of AP1 to that of AP2.</td>
</tr>
<tr>
<td>System capacity</td>
<td>Number of the wireless patient monitor supported by a single AP: &lt;= 16. Each wireless patient monitor can communicate with the central station. The wireless functions of all wireless monitors are normal at the same time.</td>
</tr>
<tr>
<td>Resistance to wireless interference</td>
<td>When the distance between interfering devices (including wireless devices at the frequency of 2.4GHz, except Wi-Fi, cellular mobile communication networks, microwave ovens, interphones, cordless phones and electro-surgical units) and the wireless patient monitor is farther than 20 cm, the wireless functions of the wireless patient monitor are normal.</td>
</tr>
<tr>
<td>Dynamic networking stability</td>
<td>The wireless functions of the test monitor are normal when the wireless patient monitor is moving at the rate of no more than 3.75 m/s within a 15m non-blocking linear distance.</td>
</tr>
<tr>
<td>Network interruption alarm</td>
<td>When network interruption occurs, the wireless patient monitor screen shows a disconnection icon, and the system initiates the related alarms. When the network is reconnected, wireless connection recovers automatically.</td>
</tr>
<tr>
<td>Wireless Networking stability</td>
<td>The 16 wireless patient monitors connected to one AP do not encounter Communication Lost events on Panorama central stations.</td>
</tr>
</tbody>
</table>
Federal Communications Commission (FCC) Statement

The wireless module has been tested and found to comply with the limits for a class B digital device, pursuant to Part 15 of the FCC rules. These limits are designed to provide reasonable protection against harmful interference in a residential installation. This equipment generates, uses and can radiate radio frequency energy and, if not installed and used in accordance with the instructions, may cause harmful interference to radio communications. However, there is no guarantee that interference will not occur in a particular installation. If this equipment does cause harmful interference to radio or television reception, which can be determined by turning the equipment off and on, the user is encouraged to try to correct the interference by one or more of the following measures:

- Reorient or relocate the receiving antenna.
- Increase the separation between the equipment and receiver.
- Connect the equipment into an outlet on a circuit different from that to which the receiver is connected.
- Consult the dealer or an experienced radio/TV technician for help.

WARNING: This compliance to general radiation exposure limits for an uncontrolled environment, and minimum of 20 cm separation between monitor and human body.

16.8 Panorama Wireless Transceiver/Splitters/Antennas/Cable Repeaters

16.8.1 Panorama Wireless Transceiver

16.8.1.1 Power Supply

Voltage
- 100-120 VAC (± 10%)

Frequency
- 60 Hz (± 3 Hz)

Power Consumption
- 1.25 Amps (maximum)

16.8.1.2 Physical Characteristics

Maximum Size
- Depth: 16.2" (411.48 mm)
- Height: 5.2" (132.08 mm)
- Width: 17" (431.8 mm)

Maximum Weight
• 12.35 lbs (5.6 kg)

16.8.1.3 Frequency Range

Receiver Range
• 608-614 MHz

Transmit Range
• 1395-1400 MHz and 1427-1429.5 MHz

16.8.2 Panorama Antenna

Physical Characteristics

Maximum Size
• Depth: 7.6" diameter (193.04 mm)
• Height: 5.5" (139.7 mm)

Maximum Weight
• 0.88 lbs (0.4 kg)

Frequency Range
• Receiver Range: 608-614 MHz
• Transmit Range: 1395-1400 MHz and 1427-1429.5 MHz

16.8.3 Cable Repeater

Physical Characteristics

Maximum Size
• Depth: 4.75" (120.65 mm)
• Height: 6.75" (171.45 mm)
• Width: 9.75" (247.65 mm)

Maximum Weight
• 3.96 lbs (1.8 kg)

Frequency Range
• Receiver Range: 608-614 MHz
• Transmit Range: 1395-1400 MHz and 1427-1429.5 MHz

16.8.4 Splitter

Physical Characteristics
Maximum Size
- Depth: 2" (50.8 mm)
- Height: 2" (50.8 mm)
- Width: 1.25" (31.75 mm)

Maximum Weight
- 0.22 lbs. (0.1 kg)

16.8.5 Environmental Characteristics for the Panorama Wireless Transceiver/Splitters/Antennas

16.8.5.1 Operating Temperature
- +10 to +40 °C

16.8.5.2 Operating Humidity
- 20 % to 80 % Relative Humidity Maximum, non-condensing, maximum wet bulb 29 °C

16.8.5.3 Operating Altitude
- 1060 to 700 hPa (-1250 to 9889 feet ASL) (795 to 525 mmHg)

16.8.5.4 Storage Temperature
- -20 to +60 °C

16.8.5.5 Storage Humidity
- 5% to 80% Relative Humidity Maximum, non-condensing, maximum wet bulb 35 °C

16.8.5.6 Storage Altitude
- 1060 to 700 hPa (-1250 to 9889 feet ASL) (795 to 525 mmHg)

16.8.5.7 Safety and Electromagnetic Compatibility
Refer to sections 16.2.9 and 16.2.11.

16.8.5.8 FCC Labeling

Panorama Wireless Transceiver
FCC ID: DXXVT0190-00000

This device complies with part 15 of the FCC rules.

NOTE: Operation is subject to the following two conditions: (1) this device may not cause harmful interference, and (2) this device must accept any interference received, including interference that may cause undesired operation of the device.
16.9 Telepack Specifications

The following are the specifications for the Telepack:

### TABLE 16-5

<table>
<thead>
<tr>
<th>SPECIFICATION</th>
<th>DESCRIPTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acquired Leads</td>
<td>ECG 3-Lead: I, II, III (one vector at a time)</td>
</tr>
<tr>
<td></td>
<td>ECG 5-Lead: Three vectors (I, II, III, aVR, aVF, V(n))</td>
</tr>
<tr>
<td>Battery Type and Runtime</td>
<td>40 hrs minimum [Telepack-608]</td>
</tr>
<tr>
<td></td>
<td>24 hrs minimum [Telepack-608 with Nonin SpO2 module]</td>
</tr>
<tr>
<td></td>
<td>Conditions: All Telepack units using two [2] AA alkaline cells with</td>
</tr>
<tr>
<td></td>
<td>ECG at a HR of 60 bpm common to the Panorama Central Station (using battery P/N 0146-00-0077-XX)</td>
</tr>
<tr>
<td>Battery Shelf Life</td>
<td>The battery shelf life is indicated on the battery casing.</td>
</tr>
<tr>
<td>Lead Current</td>
<td>Max lead current: 25nA</td>
</tr>
</tbody>
</table>

16.9.1 Safety Designations

### TABLE 16-6

<table>
<thead>
<tr>
<th>DESIGNATION</th>
<th>DESCRIPTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Degree of protection against electric shock</td>
<td>ECG-Type CF defibrillation protected.</td>
</tr>
<tr>
<td>Supply Connection</td>
<td>3 VDC Internal Battery</td>
</tr>
<tr>
<td>Mode of Operation</td>
<td>Continuous</td>
</tr>
<tr>
<td>Protection Against Hazards of Explosion</td>
<td>Not Protected (Ordinary)</td>
</tr>
<tr>
<td>Degree of Electrical Connection between Equipment and Patient</td>
<td>Equipment designed for direct electrical connection to the patient</td>
</tr>
<tr>
<td>Degree of Mobility</td>
<td>Transportable, Intra Hospital</td>
</tr>
<tr>
<td>Protection against ingress of liquid</td>
<td>IPX4 (with or without SpO2 module attached)</td>
</tr>
<tr>
<td>Protection against splashing water</td>
<td>IPX4 (with or without SpO2 module attached)</td>
</tr>
</tbody>
</table>

16.9.2 Panorama Server Analysis

The bedside monitor collects and analyses patient information and then transfers its analysis to the Panorama Central Station through the Panorama Telemetry Server; therefore, no calculations are performed by the Panorama Telemetry Server.
16.9.3 Environmental Characteristics

The environmental characteristics of the Telepack device are as follows:

**TABLE 16-7**

<table>
<thead>
<tr>
<th>SPECIFICATION</th>
<th>DESCRIPTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maximum Size (Telepack)</td>
<td>Height: 5.10” (129.54 mm)</td>
</tr>
<tr>
<td></td>
<td>Width: 2.95” (74.93 mm)</td>
</tr>
<tr>
<td></td>
<td>Depth: 1.23” (31.24 mm)</td>
</tr>
<tr>
<td>Operating Environment</td>
<td>Temperature: + 5 °C to + 40 °C (+41 °F to +104 °F)</td>
</tr>
<tr>
<td></td>
<td>Altitude: 1060 hPa to 700 hPa</td>
</tr>
<tr>
<td></td>
<td>1250 feet to 9889 feet ASL</td>
</tr>
<tr>
<td></td>
<td>-380 m to 3014 m</td>
</tr>
<tr>
<td></td>
<td>795 mmHg to 525 mmHg</td>
</tr>
<tr>
<td>Operating and Storage</td>
<td>Storage: 10% to 95% maximum, non-condensing</td>
</tr>
<tr>
<td>Humidity</td>
<td>Operating: 15% to 95% maximum, non-condensing</td>
</tr>
<tr>
<td>Storage Temperature</td>
<td>-20 °C to +60 °C (-4 °F to +140 °F)</td>
</tr>
<tr>
<td>Storage Altitude</td>
<td>1,060 hPa to 700 hPa</td>
</tr>
<tr>
<td></td>
<td>1250 feet to 9889 feet ASL</td>
</tr>
<tr>
<td></td>
<td>-380 m to 3,014 m</td>
</tr>
<tr>
<td></td>
<td>795 mmHg to 525 mmHg</td>
</tr>
</tbody>
</table>

16.9.4 Electromagnetic Compatibility

The following section describes the electromagnetic compatibility of the Telepack device.

If the Telepack device is operated within the electromagnetic environment listed in Table 16-9 on page 16-26 and Table 16-10 on page 16-28, the system will remain safe and provide the following essential performances:

- Operating mode
- Accuracy
  - Heart rate (from ECG) accuracy: ±1 bpm
- Function
- Accessories identification
- Alarm
- Detection of the connection

**NOTE:** The Telepack needs special precautions regarding Electro Magnetic Compatibility (EMC) and needs to be installed and put into service according to the EMC information provided in this section.

**NOTE:** Portable and mobile RF communications equipment can affect the Telepack. See Table 16-8 on page 16-26, Table 16-9 on page 16-26, Table 16-10 on page 16-28 and Table 16-11 on page 16-29.
TABLE 16-8

GUIDANCE AND DECLARATION - ELECTROMAGNETIC EMISSIONS

It is important that the Telepack is used in the electromagnetic environment specified in the following table.

<table>
<thead>
<tr>
<th>EMISSIONS TEST</th>
<th>COMPLIANCE</th>
<th>ELECTROMAGNETIC ENVIRONMENT - GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF emissions CISPR 11</td>
<td>Group 1</td>
<td>The Telepack uses RF energy only for its internal function and external data transmission. However, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.</td>
</tr>
<tr>
<td>RF emissions CISPR 11</td>
<td>Class A</td>
<td>The Telepack is suitable for use in all establishments other than domestic establishments and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.</td>
</tr>
<tr>
<td>Harmonic emissions IEC 61000-3-2</td>
<td>N/A</td>
<td></td>
</tr>
<tr>
<td>IEC 61000-3-3 Voltage fluctuations/ Flicker emissions</td>
<td>N/A</td>
<td></td>
</tr>
</tbody>
</table>

TABLE 16-9

GUIDANCE AND DECLARATION - ELECTROMAGNETIC IMMUNITY

It is important that the Telepack is used in the electromagnetic environment specified in the following table.

<table>
<thead>
<tr>
<th>IMMUNITY TEST</th>
<th>IEC 60601 TEST LEVEL</th>
<th>COMPLIANCE LEVEL</th>
<th>ELECTROMAGNETIC ENVIRONMENT - GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrostatic discharge (ESD) IEC 61000-4-2</td>
<td>±6 kV contact ±8 kV air</td>
<td>±6 kV contact ±8 kV air</td>
<td>Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30%.</td>
</tr>
<tr>
<td>Electrical fast transient/burst IEC 61000-4-4</td>
<td>±2 kV for power supply lines ±1 kV for input/ output lines</td>
<td>N/A *</td>
<td>Mains power quality should be typical of a commercial or hospital environment.</td>
</tr>
<tr>
<td>Surge IEC 61000-4-5</td>
<td>±1 kV differential mode ±2 kV common mode</td>
<td>N/A *</td>
<td>Mains power quality should be typical of a commercial or hospital environment.</td>
</tr>
</tbody>
</table>

* Telepack operates only from the internal battery

$U_T$ is the a.c. mains voltage prior to application of the test level.
**GUIDANCE AND DECLARATION - ELECTROMAGNETIC IMMUNITY**

It is important that the Telepack is used in the electromagnetic environment specified in the following table.

<table>
<thead>
<tr>
<th>IMMUNITY TEST</th>
<th>IEC 60601 TEST LEVEL</th>
<th>COMPLIANCE LEVEL</th>
<th>ELECTROMAGNETIC ENVIRONMENT - GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Voltage dips, short interruptions and voltage variations on power supply input lines IEC 61000-4-11</td>
<td>&lt;5% U&lt;sub&gt;T&lt;/sub&gt; [&gt;95% dip in U&lt;sub&gt;T&lt;/sub&gt;] for 0.5 cycle</td>
<td>N/A *</td>
<td>Mains power quality should be typical of a commercial or hospital environment. If the user requires continued operation during power mains interruptions, it is recommended that the device be powered from an uninterruptible power supply or a battery.</td>
</tr>
<tr>
<td></td>
<td>40% U&lt;sub&gt;T&lt;/sub&gt; [60% dip in U&lt;sub&gt;T&lt;/sub&gt;] for 5 cycles</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>70% U&lt;sub&gt;T&lt;/sub&gt; [30% dip in U&lt;sub&gt;T&lt;/sub&gt;] for 25 cycles</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>&lt;5% U&lt;sub&gt;T&lt;/sub&gt; [&gt;95% dip in U&lt;sub&gt;T&lt;/sub&gt;] for 5 sec</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Power frequency (50/60 Hz) magnetic field IEC 61000-4-8</td>
<td>3 A/m</td>
<td>3 A/m</td>
<td>Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.</td>
</tr>
</tbody>
</table>

* <sub>T</sub> is the a.c. mains voltage prior to application of the test level.

* Telepack operates only from the internal battery
TABLE 16-10
GUIDANCE AND DECLARATION - ELECTROMAGNETIC IMMUNITY

It is important that the Telepack is used in the electromagnetic environment specified in the following table.

<table>
<thead>
<tr>
<th>IMMUNITY TEST</th>
<th>IEC 60601 TEST LEVEL</th>
<th>COMPLIANCE LEVEL</th>
<th>ELECTROMAGNETIC ENVIRONMENT - GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conducted RF</td>
<td>IEC 61000-4-6</td>
<td>3 Vrms 150 kHz to 80 MHz</td>
<td>N/A Portable and mobile RF communications equipment should be used no closer to any part of the Telepack, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter.</td>
</tr>
<tr>
<td>Radiated RF</td>
<td>IEC 61000-4-3</td>
<td>3 V/m 80 MHz to 2.5 GHz</td>
<td>3 V/m d = 1.2 x √(P) 80 MHz to 800 MHz</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>d = 2.3 x √(P) 800 MHz to 2.5 GHz</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Where (P) is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and (d) is the recommended separation distance in meters (m).</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey(^a) should be less than the compliance level in each frequency range(^b).</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Interference may occur in the vicinity of equipment marked with the following symbol:</td>
</tr>
</tbody>
</table>

**NOTE:** At 80 MHz and 800 MHz, the higher frequency range applies.

**NOTE:** The EMC guidelines in Table 16-10 on page 16-28 may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

\(^a\) Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment, due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength where the Telepack is used exceeds the applicable RF compliance level, the Telepack should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as reorienting or relocating the Telepack.

\(^b\) Over the frequency range 150 kHz to 80 MHz, field strengths should be less than 3V/m.
### TABLE 16-11

**RECOMMENDED SEPARATION DISTANCES BETWEEN PORTABLE AND MOBILE RF COMMUNICATIONS EQUIPMENT AND THE TELEPACK**

The Telepack is intended for use in an electromagnetic environment where radiated RF disturbances are controlled. Prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the Telepack as recommended in the following table, according to the maximum output power of the communications equipment.

<table>
<thead>
<tr>
<th>RATED MAXIMUM OUTPUT POWER OF TRANSMITTER W (WATTS)</th>
<th>SEPARATION DISTANCE ACCORDING TO FREQUENCY OF TRANSMITTER M (METERS)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>150 KHZ TO 80 MHZ</td>
</tr>
<tr>
<td></td>
<td>( D = 1.2 \times \sqrt{P} )</td>
</tr>
<tr>
<td>0.01</td>
<td>0.12</td>
</tr>
<tr>
<td>0.1</td>
<td>0.38</td>
</tr>
<tr>
<td>1</td>
<td>1.2</td>
</tr>
<tr>
<td>10</td>
<td>3.8</td>
</tr>
<tr>
<td>100</td>
<td>12</td>
</tr>
</tbody>
</table>

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

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

**NOTE:** The separation distances guidelines in Table 16-11 on page 16-29 may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.
16.9.5 FCC Labeling

- Radio Transceiver - The Telepack-608 (FCC ID:DXXWT0191-00000) utilizes the frequency range 608 MHz - 614 MHz. Modulation is 2-level FSK.
- This device complies with part 15 of the FCC rules. Operation is subject to the following two conditions: (1) this device may not cause harmful interference, and (2) this device must accept any interference received, including interference that may cause undesired operation of the device.
- This Class B digital apparatus meets all requirements of the Canadian Interference Causing Equipment Regulations. Operation is subject to the following two conditions: (1) this device may not cause harmful interference, and (2) this device must accept any interference received, including interference that may cause undesired operation.

Cet appareillage numérique de la classe B répond à toutes les exigences de l’interférence canadienne causant des règlements d’équipement. L’opération est sujette aux deux conditions suivantes: (1) ce dispositif peut ne pas causer l’interférence nocive, et (2) ce dispositif doit accepter n’importe quelle interférence reçue, y compris l’interférence qui peut causer l’opération peu désirée.

16.9.6 Compliance
The Telepack complies with the following standards:

**AGENCY AND INDUSTRY STANDARDS**

<table>
<thead>
<tr>
<th>Standard</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>IEC 60601-2-27 Ed.3.0:2011</td>
<td>AAMI/ANSI EC57:2012</td>
</tr>
<tr>
<td>IEC 62304 Ed.1.0:2006-05</td>
<td>IEC 62304 Ed.1.0:2006-05</td>
</tr>
<tr>
<td>ISO 80601-2-61 Ed.1.0:2011</td>
<td>IEC 62304 Ed.1.0:2006-05</td>
</tr>
<tr>
<td>IEC 60601-1-6 Ed.3.1:2013</td>
<td>ISO 80601-2-61 Ed.1.0:2011</td>
</tr>
<tr>
<td>IEEE 802.11 Std 802.11:1999 - Part 11</td>
<td>IEC 60601-1-6 Ed.3.1:2013</td>
</tr>
<tr>
<td>ISTA: Test Procedure 1A</td>
<td>IEEE 802.11 Std 802.11:1999 - Part 11</td>
</tr>
<tr>
<td>FDA Guidance for the Content of Premarket Submissions for Software Contained in Medical Devices, May 11, 2005</td>
<td>ISTA: Test Procedure 1A</td>
</tr>
</tbody>
</table>
16.9.7 Specific Absorption Rate

The Panorama Telepack meets the government’s requirements for exposure to radio waves.

The Telepack is a radio transmitter and receiver. It is designed and manufactured not to exceed limits for exposure to radio frequency (RF) energy set forth by the Federal Communications Commission (FCC) of the U.S. Government and by Industry Canada. These limits are part of comprehensive guidelines and establish permitted levels of RF energy for the general population. These guidelines are based on standards that were developed by independent scientific organizations through periodic and thorough evaluation of scientific studies. These standards include a substantial safety margin designed to assure the safety of all persons, regardless of age or health.

The exposure standard employs a unit of measurement known as the Specific Absorption Rate, or SAR. The SAR limit set by the FCC and Industry Canada is 1.6 W/kg.

The Panorama Telepack has been evaluated for body-worn Specific Absorption Rate (SAR) compliance under FCC Bulletin OET 65C and found to be compliant to the requirements as set forth in CFR 47 Sections 2.1093 and 15.242 addressing RF exposure from radio frequency devices. The highest SAR level measured was 0.002 W/kg.
16.10 Nonin SpO₂ Module Specifications

16.10.1 Environmental Characteristics

The environmental characteristics for the Nonin SpO₂ module device are:

**TABLE 16-12**

<table>
<thead>
<tr>
<th>SPECIFICATION</th>
<th>DESCRIPTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maximum Size of Telepack with Nonin SpO₂ module</td>
<td>Height: 6.25” (158.75 mm) Width: 3.55” (90.17 mm) Depth: 1.75” (44.45 mm)</td>
</tr>
<tr>
<td>Maximum Weight (Nonin SpO₂ module)</td>
<td>4.5 oz. excluding cable and sensor</td>
</tr>
<tr>
<td>Temperature</td>
<td>Operating: + 5 °C to + 40 °C Storage: - 20 °C to + 50 °C</td>
</tr>
<tr>
<td>Humidity</td>
<td>Operating: 15% to 90% maximum, non-condensing Storage: 10% to 95% maximum, non-condensing</td>
</tr>
</tbody>
</table>

16.10.2 Measurement Specifications

The measurement specifications for the Nonin SpO₂ module are:

<table>
<thead>
<tr>
<th>SPECIFICATION</th>
<th>DESCRIPTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oxygen Saturation Range</td>
<td>0 to 100%</td>
</tr>
<tr>
<td>Pulse Rate Range</td>
<td>18 to 300 pulses per minute</td>
</tr>
<tr>
<td>Measurement wavelength red</td>
<td>660 nanometers @3mW nominal</td>
</tr>
<tr>
<td><strong>NOTE:</strong> Information about wavelength range can be especially useful to clinicians.</td>
<td></td>
</tr>
<tr>
<td>Measurement wavelength infrared</td>
<td>910 nanometers @3mW nominal</td>
</tr>
<tr>
<td><strong>NOTE:</strong> Information about wavelength range can be especially useful to clinicians.</td>
<td></td>
</tr>
</tbody>
</table>

**SPO₂ ACCURACY**

<table>
<thead>
<tr>
<th>SpO₂ (70-100%) ± 1 standard deviation</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>No Motion</td>
<td>reusable finger clip ± 2 digits</td>
</tr>
<tr>
<td></td>
<td>disposable ± 3 digits</td>
</tr>
<tr>
<td></td>
<td>low perfusion ± 2 digits</td>
</tr>
<tr>
<td>Motion</td>
<td>reusable finger clip ± 2 digits</td>
</tr>
<tr>
<td></td>
<td>low perfusion ± 2 digits</td>
</tr>
</tbody>
</table>

**PULSE RATE ACCURACY**

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>no motion adults (18-300 bpm)</td>
<td>± 3 digits</td>
</tr>
<tr>
<td>motion adults (40-240 bpm)</td>
<td>± 5 digits</td>
</tr>
<tr>
<td>low perfusion adults (40-240 bpm)</td>
<td>± 3 digits</td>
</tr>
</tbody>
</table>

**DATA UPDATE RATE**

SpO₂ value and heart rate is updated 3 times per second
16.10.3 Electromagnetic Compatibility

The following section describes the electromagnetic compatibility of the Nonin SpO2 device.

The Nonin SpO2 module meets the requirements of:

- IEC 60601-1-2
- ISO 80601-2-61

If the Nonin SpO2 is operated within the electromagnetic environment listed in Table 16-9 on page 16-26 and Table 16-10 on page 16-28, the system will remain safe and provide the following essential performances:

- Accuracy
  - SpO2 accuracy: ±2%
  - Pulse rate accuracy: ±3 bpm
- Function
- Accessories identification
- Detection of the connection
- Alarm limit (When patient size is adult, the high alarm limit for the SpO2 (Nonin) parameter on the ViewStation shall have a range of 80% to 100%)

**TABLE 16-13**

**GUIDANCE AND DECLARATION - ELECTROMAGNETIC EMISSIONS**

It is important that the Nonin SpO2 module is used in the electromagnetic environment specified in the following table.

<table>
<thead>
<tr>
<th>EMISSIONS TEST</th>
<th>COMPLIANCE</th>
<th>ELECTROMAGNETIC ENVIRONMENT - GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF emissions CISPR 11</td>
<td>Group 1</td>
<td>The Nonin SpO2 module uses RF energy only for its internal function and external data transmission. However, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.</td>
</tr>
<tr>
<td>RF emissions CISPR 11</td>
<td>Class B</td>
<td>The Nonin SpO2 module is suitable for use in all establishments other than domestic establishments and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.</td>
</tr>
<tr>
<td>Harmonic emissions IEC 61000-3-2</td>
<td>N/A</td>
<td></td>
</tr>
<tr>
<td>IEC 61000-3-3 Voltage fluctuations/ Flicker emissions</td>
<td>N/A</td>
<td></td>
</tr>
</tbody>
</table>
### TABLE 16-14

**GUIDANCE AND DECLARATION - ELECTROMAGNETIC IMMUNITY**

It is important that the Nonin SpO₂ module is used in the electromagnetic environment specified in the following table.

<table>
<thead>
<tr>
<th>IMMUNITY TEST</th>
<th>IEC 60601 TEST LEVEL</th>
<th>COMPLIANCE LEVEL</th>
<th>ELECTROMAGNETIC ENVIRONMENT - GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrostatic discharge (ESD) IEC 61000-4-2</td>
<td>±6 kV contact ±8 kV air</td>
<td>±6 kV contact ±8 kV air</td>
<td>Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30%.</td>
</tr>
<tr>
<td>Electrical fast transient/burst IEC 61000-4-4</td>
<td>N/A</td>
<td>N/A*</td>
<td>Mains power quality should be typical of a commercial or hospital environment.</td>
</tr>
<tr>
<td>Surge IEC 61000-4-5</td>
<td>N/A</td>
<td>N/A*</td>
<td>Mains power quality should be typical of a commercial or hospital environment. If the user of the device requires continued operation during power mains interruptions, it is recommended that the device be powered from an uninterruptible power supply or battery pack.</td>
</tr>
<tr>
<td>Voltage dips, short interruptions and voltage variations on power supply input lines IEC 61000-4-11</td>
<td>N/A</td>
<td>N/A*</td>
<td>Mains power quality should be typical of a commercial or hospital environment. If the user requires continued operation during power mains interruptions, it is recommended that the device be powered from an uninterruptible power supply or a battery.</td>
</tr>
<tr>
<td>Power frequency (50/60 Hz) magnetic field IEC 61000-4-8</td>
<td>3 A/m</td>
<td>3 A/m</td>
<td>Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.</td>
</tr>
</tbody>
</table>

* The Nonin SpO₂ module receives power from the Telepack which operates only from an internal battery.
TABLE 16-15
GUIDANCE AND DECLARATION - ELECTROMAGNETIC IMMUNITY

It is important that the Nonin SpO₂ module is used in the electromagnetic environment specified in the following table.

Portable and mobile RF communications equipment should be used no closer to any part of the device, including cables, than the recommended separation calculated from the equation applicable to the frequency of the transmitter.

<table>
<thead>
<tr>
<th>IMMUNITY TEST</th>
<th>IEC 60601 TEST LEVEL</th>
<th>COMPLIANCE LEVEL</th>
<th>ELECTROMAGNETIC ENVIRONMENT - GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conducted RF</td>
<td>IEC 61000-4-6 3 Vrms</td>
<td>3 V</td>
<td>Recommended separation distance</td>
</tr>
<tr>
<td></td>
<td>150 kHz to 80 MHz</td>
<td></td>
<td>$d = 1.17 \times \sqrt{P}$</td>
</tr>
<tr>
<td>Radiated RF</td>
<td>IEC 61000-4-3 3 V/m</td>
<td>3 V/m</td>
<td>$d = 1.17 \times \sqrt{P}$ 80 MHz to 800 MHz</td>
</tr>
<tr>
<td></td>
<td>80 MHz to 2.5 GHz</td>
<td></td>
<td>$d = 2.33 \times \sqrt{P}$ 800 MHz to 2.5 GHz</td>
</tr>
</tbody>
</table>

Where $P$ is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and $d$ is the recommended separation distance in meters (m).

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

Interference may occur in the vicinity of equipment marked with the following symbol: 📣

**NOTE:** At 80 MHz and 800 MHz, the higher frequency range applies.

**NOTE:** The EMC guidelines in Table 16-15 on page 16-35 may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

- Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment, due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength where the Nonin SpO₂ module is used exceeds the applicable RF compliance level, the Nonin SpO₂ module should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as reorienting or relocating the Nonin SpO₂ module.

- Over the frequency range 150 kHz to 80 MHz, field strengths should be less than 3 V/m.
### TABLE 16-16

**RECOMMENDED SEPARATION DISTANCES BETWEEN PORTABLE AND MOBILE RF COMMUNICATIONS EQUIPMENT AND THE NONIN SPO2 MODULE**

The Nonin SpO2 module is intended for use in an electromagnetic environment where radiated RF disturbances are controlled. Prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the Nonin SpO2 module as recommended in the following table, according to the maximum output power of the communications equipment.

<table>
<thead>
<tr>
<th>RATED MAXIMUM OUTPUT POWER OF TRANSMITTER W (WATTS)</th>
<th>SEPARATION DISTANCE ACCORDING TO FREQUENCY OF TRANSMITTER M (METERS)</th>
</tr>
</thead>
<tbody>
<tr>
<td>150 KHZ TO 80 MHZ</td>
<td>80 MHZ TO 800 MHZ</td>
</tr>
<tr>
<td>D = 1.17 × \sqrt{P}</td>
<td>D = 1.17 × \sqrt{P}</td>
</tr>
<tr>
<td>0.01</td>
<td>0.12</td>
</tr>
<tr>
<td>0.1</td>
<td>0.37</td>
</tr>
<tr>
<td>1</td>
<td>1.2</td>
</tr>
<tr>
<td>10</td>
<td>3.7</td>
</tr>
<tr>
<td>100</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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

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

**NOTE:** The separation distances guidelines in Table 16-16 on page 16-36 may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.
16.11 Panorama Instrument Radio-608

16.11.1 Approved Accessory
Kit, Instrument Radio-608, Passport 2/Spectrum (P/N 0040-00-0361-01)

16.11.2 FCC Label

- Radio Transceiver - The Panorama Instrument Radio-608 [FCC ID: DXXRA0743-00000] utilizes the frequency range 608 MHz - 614 MHz and 1427MHz -1429.5 MHz. Modulation is 4-level FSK.
- This device complies with part 15 of the FCC Rules. Operation is subject to the following two conditions: 1) This device may not cause harmful interference, and 2) this device must accept any interference received, including interference that may cause undesired operation.
17.0  

**Glossary**

This section provides a glossary of terminology used throughout the Panorama Operating Instructions.

- Terms, Acronyms, and Abbreviations
## 17.1 Terms, Acronyms, and Abbreviations

<table>
<thead>
<tr>
<th>TERMS</th>
<th>DEFINITIONS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Agent</td>
<td>The gas used to anesthetize a patient during surgery</td>
</tr>
<tr>
<td>Arrhythmia</td>
<td>Abnormal ECG rhythms</td>
</tr>
<tr>
<td>Bedside Monitor</td>
<td>A monitoring device that has its own user interface and does not require the Panorama Central Station to be its primary monitoring source</td>
</tr>
<tr>
<td>BIS</td>
<td>Bispectral Index</td>
</tr>
<tr>
<td>BSA</td>
<td>Body Surface Area</td>
</tr>
<tr>
<td>CI</td>
<td>Cardiac Index</td>
</tr>
<tr>
<td>CO</td>
<td>Cardiac Output</td>
</tr>
<tr>
<td>Heart Rate</td>
<td>Number of heartbeats detected during a one minute period</td>
</tr>
<tr>
<td>Internet Protocol (IP)</td>
<td>TCP/IP is the standard protocol that defines the IP datagram as the unit of information passed across an Internet. IP provides the basis for connectionless, best effort packet delivery of services.</td>
</tr>
<tr>
<td>Invasive Blood Pressure (IBP)</td>
<td>Method of obtaining blood pressure using an internal pressure sensor</td>
</tr>
<tr>
<td>Monitoring Device</td>
<td>The instrumentation connected to the patient. The monitoring device acquires the patient data and then transmits it onto the network.</td>
</tr>
<tr>
<td>Non-Invasive Blood Pressure (NIBP)</td>
<td>Method of obtaining blood pressure using an external cuff</td>
</tr>
<tr>
<td>Panorama Central Network</td>
<td>Supports hardwired communication between Panorama Central Stations and laser printers</td>
</tr>
<tr>
<td>Panorama Patient Network</td>
<td>Supports hardwired communication of the real-time patient data collected by monitoring devices attached to the patient</td>
</tr>
<tr>
<td>Panorama Wireless Network</td>
<td>Supports wireless communications between the wireless monitoring devices and the Panorama Server.</td>
</tr>
<tr>
<td>PAWP</td>
<td>Pulmonary Artery Wedge Pressure</td>
</tr>
<tr>
<td>Persistent</td>
<td>Data or settings maintained and remain available until purposely edited or deleted</td>
</tr>
<tr>
<td>Respiration</td>
<td>Number of breaths during a one minute time period</td>
</tr>
<tr>
<td>SpO₂</td>
<td>Level of Oxygen Saturation in the blood</td>
</tr>
<tr>
<td>ST</td>
<td>The ST portion of the ECG waveform</td>
</tr>
<tr>
<td>ACRONYMS/ABBREVIATIONS</td>
<td>DEFINITIONS</td>
</tr>
<tr>
<td>------------------------</td>
<td>-------------</td>
</tr>
<tr>
<td>ADT</td>
<td>Admit Discharge Transfer</td>
</tr>
<tr>
<td>AHA</td>
<td>American Heart Association</td>
</tr>
<tr>
<td></td>
<td>American Hospital Association</td>
</tr>
<tr>
<td>ART</td>
<td>Arterial Line</td>
</tr>
<tr>
<td>AXI</td>
<td>Axillary</td>
</tr>
<tr>
<td>BIS</td>
<td>Bispectral Index</td>
</tr>
<tr>
<td>BLA</td>
<td>Bladder</td>
</tr>
<tr>
<td>bpm</td>
<td>Beats per minute</td>
</tr>
<tr>
<td>C</td>
<td>Current ST</td>
</tr>
<tr>
<td></td>
<td>Chest Lead (IEC)</td>
</tr>
<tr>
<td>CCI</td>
<td>Continuous Cardiac Index</td>
</tr>
<tr>
<td>CCI STAT</td>
<td>Continuous Cardiac Index STAT</td>
</tr>
<tr>
<td>CCO</td>
<td>Continuous Cardiac Output</td>
</tr>
<tr>
<td>CCO STAT</td>
<td>Continuous Cardiac Output STAT</td>
</tr>
<tr>
<td>cm</td>
<td>Centimeters</td>
</tr>
<tr>
<td>CIS</td>
<td>Clinical Information System</td>
</tr>
<tr>
<td>C-LAN</td>
<td>Panorama Central Network</td>
</tr>
<tr>
<td>CO₂</td>
<td>Carbon Dioxide</td>
</tr>
<tr>
<td>CVA</td>
<td>Cardiovascular Artifact</td>
</tr>
<tr>
<td>CVP</td>
<td>Central Venous Pressure</td>
</tr>
<tr>
<td>Des</td>
<td>Desflurane</td>
</tr>
<tr>
<td>DL</td>
<td>Down Link</td>
</tr>
<tr>
<td>ECG</td>
<td>Electrocardiogram</td>
</tr>
<tr>
<td>E-LAN</td>
<td>Panorama Patient Network</td>
</tr>
<tr>
<td>EMC</td>
<td>Electromagnetic Compatibility</td>
</tr>
<tr>
<td>Enf</td>
<td>Enflurane</td>
</tr>
<tr>
<td>EMR</td>
<td>Electronic Medical Records</td>
</tr>
<tr>
<td>EPM</td>
<td>Spectrum/Spectrum OR External Parameter Module</td>
</tr>
<tr>
<td>ESO</td>
<td>Esophageal</td>
</tr>
<tr>
<td>ET</td>
<td>End-Tidal when referencing CO₂/O₂/N₂O/Agent parameters</td>
</tr>
<tr>
<td></td>
<td>Elapsed Time when referencing NIBP</td>
</tr>
<tr>
<td>F</td>
<td>Left Leg when referring to ECG Leads (IEC)</td>
</tr>
<tr>
<td>Hal</td>
<td>Haloflurane</td>
</tr>
<tr>
<td>HIS</td>
<td>Hospital Information System</td>
</tr>
<tr>
<td>ACRONYMS/ABBREVIATIONS</td>
<td>DEFINITIONS</td>
</tr>
<tr>
<td>-------------------------</td>
<td>-------------</td>
</tr>
<tr>
<td>HR</td>
<td>Heart Rate</td>
</tr>
<tr>
<td>IBP</td>
<td>Invasive Blood Pressure</td>
</tr>
<tr>
<td>ICP</td>
<td>Intracranial Pressure</td>
</tr>
<tr>
<td>IEC</td>
<td>International Electro-Technical Commission</td>
</tr>
<tr>
<td>in</td>
<td>Inches</td>
</tr>
<tr>
<td>INSP</td>
<td>Inspired</td>
</tr>
<tr>
<td>IP</td>
<td>Internet Protocol</td>
</tr>
<tr>
<td>ISO</td>
<td>Isoflurane when referencing Agent</td>
</tr>
<tr>
<td></td>
<td>Isoelectric when referencing ST monitoring</td>
</tr>
<tr>
<td>kg</td>
<td>Kilograms</td>
</tr>
<tr>
<td>kPa</td>
<td>Kilopascals Per Atmosphere</td>
</tr>
<tr>
<td>L</td>
<td>Left Arm when referring to ECG Leads (IEC)</td>
</tr>
<tr>
<td>LA</td>
<td>Left Arm when referring to ECG Leads (AHA)</td>
</tr>
<tr>
<td>lbs</td>
<td>Pounds</td>
</tr>
<tr>
<td>LL</td>
<td>Left Leg when referring to ECG Leads (AHA)</td>
</tr>
<tr>
<td>LV</td>
<td>Left Ventricle</td>
</tr>
<tr>
<td>MAP</td>
<td>Mean Arterial Pressure</td>
</tr>
<tr>
<td>MDD</td>
<td>Medical Device Directive</td>
</tr>
<tr>
<td>mmHg</td>
<td>Millimeters of Mercury</td>
</tr>
<tr>
<td>N</td>
<td>Right Leg when referring to ECG Leads (IEC)</td>
</tr>
<tr>
<td>N2O</td>
<td>Nitrous Oxide</td>
</tr>
<tr>
<td>NAS</td>
<td>Nasal</td>
</tr>
<tr>
<td>NIBP</td>
<td>Non-Invasive Blood Pressure</td>
</tr>
<tr>
<td>NSR</td>
<td>Normal Sinus Rhythm</td>
</tr>
<tr>
<td>O2</td>
<td>Oxygen</td>
</tr>
<tr>
<td>PA</td>
<td>Pulmonary Artery</td>
</tr>
<tr>
<td>Pleth</td>
<td>Plethysmograph</td>
</tr>
<tr>
<td>PVC</td>
<td>Premature Ventricular Contraction</td>
</tr>
<tr>
<td>R</td>
<td>Right Arm when referring to ECG Leads (IEC)</td>
</tr>
<tr>
<td></td>
<td>Referenced ST</td>
</tr>
<tr>
<td>RA</td>
<td>Right Atrium when referring to IBP</td>
</tr>
<tr>
<td></td>
<td>Right Arm when referring to ECG Leads (AHA)</td>
</tr>
<tr>
<td>REC</td>
<td>Rectal</td>
</tr>
<tr>
<td>RESP</td>
<td>Respiration Rate</td>
</tr>
<tr>
<td>ACRONYMS/ABBREVIATIONS</td>
<td>DEFINITIONS</td>
</tr>
<tr>
<td>------------------------</td>
<td>-------------</td>
</tr>
<tr>
<td>RL</td>
<td>Right Leg when referring to ECG Leads (AHA)</td>
</tr>
<tr>
<td>rpm</td>
<td>Respirations per minute</td>
</tr>
<tr>
<td>ScvO₂</td>
<td>Central Venous Oxygen Saturation</td>
</tr>
<tr>
<td>Sev</td>
<td>Sevoflurane</td>
</tr>
<tr>
<td>SKI</td>
<td>Skin</td>
</tr>
<tr>
<td>SpO₂</td>
<td>Oxygen Saturation</td>
</tr>
<tr>
<td>SvO₂</td>
<td>Mixed Venous Saturation</td>
</tr>
<tr>
<td>SVR</td>
<td>System Vascular Resistance</td>
</tr>
<tr>
<td>SVRI</td>
<td>System VascularResistance Index</td>
</tr>
<tr>
<td>TEMP</td>
<td>Temperature</td>
</tr>
<tr>
<td>UA</td>
<td>Umbilical Artery</td>
</tr>
<tr>
<td>V</td>
<td>Chest Lead (AHA)</td>
</tr>
</tbody>
</table>
This page intentionally left blank.
18.0  **Anomalies**

This section provides the Panorama Central Station anomalies.

- Anomalies Description
18.1 Anomalies Description

- When scrolling in the Graphic view of the Trends tab using the left or right arrow button up to a break of data, the “Time Out of Range” message will be displayed even though more data remains.

**NOTE:** To correct it, select the Page Left or Page Right button in the Graphic view of the Trends tab to display the remaining data.

- A patient on a Telepack device will automatically re-admit itself to the central station, if a communication is lost and re-connected after the patient discharge.

**NOTE:** To correct it, discharge the patient again after verifying that the Telepack device and the central station communication is stable.

- When entering the Disclosure tab, the most recent record (1 minute) may not be shown if the data is not fully collected.

**NOTE:** To correct it, use the single up-arrow button to scroll the most recent data.

- The Trends tab and Event tab may become disabled and empty if the user frequently switches the two tabs.

**NOTE:** To correct it, exiting and re-entering the tab will display the stored data.

- When the second most recent trend record is selected in the Trends tab, and the most recent record has events associated with it, selecting the Next Event button may display the improper message No more events found.

**NOTE:** To correct it, close the dialog box and re-enter the Trends tab.

- If a patient tile on the central station is assigned to the Network tab on the WorkStation and ViewStation, and then edit the Device ID of the patient tile, two devices with same patient tile are displayed in the Network tab on the WorkStation and ViewStation.

**NOTE:** To correct it, remove the original assigned device and refresh the Network tab on the WorkStation and ViewStation.
• A gray box will display overlaid in the Trends tab or Event tab when a patient is discharged and then re-admitted at the bedside monitor.

NOTE: To correct it, exiting the and re-entering the tab.

• After transferring a patient at the central station from one bedside monitor to another which is of different model, when the patient sizes are configured differently and patient height and weight are configured to the the minimum on the two monitors, the patient’s height and weight may be incorrectly displayed.

NOTE: To correct it, re-enter the height and weight for the patient.

• The Discharge button on the central station may be incorrectly labelled as Discharge Patient after the bedside monitor enters and exits Standby, if the patient’s Discharge tab is open while a Central to Central Transfer Without Device is initiated. The button should display as Panorama and Bedside Discharge.

NOTE: To correct it, exit and re-enter the Discharge tab.

• When the monitoring device is disconnected from the central station, and then the central station restarts, the patient tile of this device may not display the message Waiting for Patient on the central station.

NOTE: To correct it, restore the connectivity between the monitoring device and the central station.

• After the central station restarts, the display and ST configurations of the Telepack tile may gets reset to the system default settings.

NOTE: To correct it, re-configure display settings and ST config as required.

• Selecting the OFF button on the IBP Low Systolic Alarm will not update the Low Systolic alarm slider to OFF.

NOTE: The alarm value is unaffected by this issue. Even though the alarm slider is not indicating OFF, the alarm is off.

NOTE: To correct it, click the single arrow-down button on the Alarm Control to move the slider to the bottom.

• If the start time for the All Strips report is during the time which a break of data occurred, then the report will not print.

CAUTION: To avoid the problem, only turn on the Print On Alarm option of the alarms which are needed to print.
• After adjusting the clock back in time, and scrolling downward past the point in the trend data where the clock was adjusted, you cannot scroll back up. Additionally, when attempting to print the trend data associated with the time change, you will be able to print only the trend data above the change when you are viewing above the change, and all the trend data when you are below the change. This situation will be corrected when the change event expires from the trend database.

**CAUTION:** Make sure that the central station system time is correct before use and don't adjust the clock back during use.

• When changing the system time from PM to AM, a Full Disclosure Report printed at the time of the change may not print all the data available. Scrolling past the change in system time will allow the user to print all the data.

**CAUTION:** Make sure that the central station system time is correct before use and don't adjust the clock back during use.

• When an out of order event is encountered in the Event database, the Event List Report will not include the events that occurred before that event. Selecting the event that occurred before the out of order event and then selecting the Print button will print all of the events.

**CAUTION:** Make sure that the central station system time is correct before use and don't adjust the clock back during use.

• When switching back and forth between Adult size and Neonatal size and pressing the Restore Previous button, the alarm limits for the selected patient tile may display alarm limit data applicable to the previously chosen patient size. Therefore, alarms as well as the Patient Alarm Report may be based on the wrong patient size. To correct, check that alarm limits are set correctly for the patient size.

**CAUTION:** Don't press Restore Previous Settings sidebar button in Patient Alarm Limits tab after changing the patient size.

• Transferring patient data between Panorama Central Stations which have different Full Disclosure options may result in the newest data in the destination disclosure database only being accessible via the Disclosure Zoom In tab. The Disclosure tab’s disclosure data may not be displayed at its correct time and the tab may not be easily maneuvered. This issue can be corrected once the patient tile is discharged.

**CAUTION:** Don't transfer patient between Central Stations with different full disclosure storage volume.
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