



iM50/iM80

Patient Monitor

Version 1.0



About this Manual

P/N: 01.54.455473-10

Release Date: Jan. 2012

© Copyright EDAN INSTRUMENTS, INC. 2012. All rights reserved.

Statement

This manual will help you understand the operation and maintenance of the product better. It is reminded that the product shall be used strictly complying with this manual. User's operation failing to comply with this manual may result in malfunction or accident for which EDAN INSTRUMENTS, INC. (hereinafter called EDAN) can not be held liable.

EDAN owns the copyrights of this manual. Without prior written consent of EDAN, any materials contained in this manual shall not be photocopied, reproduced or translated into other languages.

Materials protected by the copyright law, including but not limited to confidential information such as technical information and patent information are contained in this manual, the user shall not disclose such information to any irrelevant third party.

The user shall understand that nothing in this manual grants him, expressly or implicitly, any right or license to use any of the intellectual properties of EDAN.

EDAN holds the rights to modify, update, and ultimately explain this manual.

Responsibility of the Manufacturer

EDAN only considers itself responsible for any effect on safety, reliability and performance of the equipment if:

Assembly operations, extensions, re-adjustments, modifications or repairs are carried out by persons authorized by EDAN, and

The electrical installation of the relevant room complies with national standards, and

The instrument is used in accordance with the instructions for use.

Upon request, EDAN may provide, with compensation, necessary circuit diagrams, and other information to help qualified technician to maintain and repair some parts, which EDAN may define as user serviceable.

Terms Used in this Manual

This guide is designed to give key concepts on safety precautions.

WARNING

A **WARNING** label advises against certain actions or situations that could result in personal injury or death.

CAUTION

A **CAUTION** label advises against actions or situations that could damage equipment, produce

inaccurate data, or invalidate a procedure.

NOTE

A **NOTE** provides useful information regarding a function or a procedure.

Table of Contents

| | |
|---|-----------|
| Chapter 1 Indication for Use and Safety Guidance | 1 |
| 1.1 Indication for Use..... | 1 |
| 1.2 Safety Guidance | 1 |
| 1.3 Explanation of Symbols on the Monitor | 3 |
| Chapter 2 Installation | 7 |
| 2.1 Initial Inspection..... | 7 |
| 2.2 Mounting the Monitor | 7 |
| 2.2.1 Installing Wall Mount for the Monitor | 7 |
| 2.3 Connecting the Power Cable..... | 7 |
| 2.4 Checking Out the Monitor | 7 |
| 2.5 Checking the Recorder..... | 8 |
| 2.6 Setting the Date and Time | 8 |
| 2.7 Handing Over the Monitor | 8 |
| Chapter 3 Basic Operation | 9 |
| 3.1 Introducing iM50/iM80 Series..... | 9 |
| 3.1.1 iM50 Major Parts and Keys | 9 |
| 3.1.2 iM80 Major Parts and Keys | 12 |
| 3.1.3 iM50/iM80 Configuration..... | 15 |
| 3.2 Operating and Navigating | 15 |
| 3.2.1 Using Keys | 17 |
| 3.3 Operating Mode | 18 |
| 3.3.1 Demo Mode..... | 18 |
| 3.3.2 Standby Mode | 19 |
| 3.4 Changing Monitor Settings | 19 |
| 3.4.1 Adjusting Screen Brightness | 19 |
| 3.4.2 Changing Date and Time..... | 19 |
| 3.5 Adjusting Volume..... | 20 |
| 3.5.1 Adjusting Key Volume | 20 |
| 3.5.2 Adjusting Alarm Volume..... | 20 |
| 3.5.3 Adjusting Beat Volume..... | 20 |
| 3.6 Checking Your Monitor Version | 20 |
| 3.7 Networked Monitoring..... | 20 |
| 3.8 Setting Languages | 20 |
| 3.9 Understanding Screens..... | 21 |
| 3.10 Calibrating Screens | 21 |
| 3.11 Disabling the TouchScreen..... | 21 |
| 3.12 Using the Barcode Scanner | 21 |
| Chapter 4 Alarms | 22 |

| | |
|--|-----------|
| 4.1 Alarm Category | 22 |
| 4.1.1 Physiological alarms | 22 |
| 4.1.2 Technical Alarms | 22 |
| 4.1.3 Prompts | 22 |
| 4.2 Alarm Levels | 22 |
| 4.3 Controlling Alarm | 23 |
| 4.3.1 Switching the Individual Alarm Off | 23 |
| 4.3.2 Temporary Alarm Mute | 23 |
| 4.3.3 Alarm Mute | 24 |
| 4.3.4 Controlling Alarm Volume | 24 |
| 4.3.5 Setting Alarm Limits | 24 |
| 4.4 Latching Alarms | 25 |
| 4.5 Disabling Sensor Off Alarms | 25 |
| 4.6 Testing Alarms | 25 |
| Chapter 5 Alarm Information | 26 |
| 5.1 Physiological Alarm Information | 26 |
| 5.2 Technical Alarm Information | 30 |
| 5.3 Prompts | 41 |
| 5.4 Adjustable Range of Alarm Limits | 42 |
| Chapter 6 Managing Patients | 47 |
| 6.1 Admitting a Patient | 47 |
| 6.1.1 Patient Category and Paced Status | 47 |
| 6.2 Quick Admit | 48 |
| 6.3 Editing Patient Information | 48 |
| 6.4 Updating a Patient | 48 |
| 6.5 Central Monitoring System | 48 |
| Chapter 7 User Interface | 49 |
| 7.1 Setting Interface Style | 49 |
| 7.2 Selecting Display Parameters | 49 |
| 7.3 Changing Waveform Position | 49 |
| 7.4 Changing Interface Layout | 49 |
| 7.5 Viewing Trend Screen | 49 |
| 7.6 Viewing Oxygen Screen | 49 |
| 7.7 Viewing Large Font Screen | 50 |
| 7.8 Changing Parameter and Waveform Colors | 50 |
| 7.9 Transferring a Configuration | 50 |
| 7.10 Default Configuration | 51 |
| Chapter 8 Monitoring ECG | 52 |
| 8.1 Overview | 52 |

| | |
|--|-----------|
| 8.2 ECG Safety Information | 52 |
| 8.3 ECG Display | 53 |
| 8.3.1 Changing the Size of the ECG Wave | 53 |
| 8.3.2 Changing the ECG Filter Settings..... | 54 |
| 8.4 Using ECG Alarms..... | 54 |
| 8.5 Selecting Calculation Lead | 54 |
| 8.6 Monitoring Procedure | 54 |
| 8.6.1 Preparation | 54 |
| 8.6.2 Connecting ECG Cables | 55 |
| 8.7 Selecting Lead Type | 55 |
| 8.8 Installing Electrodes..... | 55 |
| 8.8.1 Electrode Placement for 3-lead | 56 |
| 8.8.2 Electrode Placement for 5-lead | 56 |
| 8.8.3 Electrode Placement for 12-lead | 58 |
| 8.8.4 Recommended ECG Lead Placement for Surgical Patients | 59 |
| 8.9 ECG Menu Setup | 60 |
| 8.9.1 Setting Alarm Source | 60 |
| 8.9.2 Smart Lead Off..... | 60 |
| 8.9.3 Setting Beat Volume..... | 60 |
| 8.9.4 ECG Display | 60 |
| 8.9.5 Setting Pace Status | 61 |
| 8.9.6 ECG Calibration..... | 61 |
| 8.9.7 ECG Waveform Settings | 61 |
| 8.9.8 12 Leads ECG | 61 |
| 8.10 ST Segment Monitoring..... | 62 |
| 8.10.1 Setting ST Analysis | 62 |
| 8.10.2 ST Display..... | 62 |
| 8.10.3 ST Analysis Alarm Setting | 62 |
| 8.10.4 About ST Measurement Points..... | 62 |
| 8.10.5 Adjusting ST and ISO Measurement Points..... | 63 |
| 8.11 Arr. Monitoring..... | 63 |
| 8.11.1 Arrhythmia Analysis..... | 63 |
| 8.11.2 ARR Analysis Menu..... | 64 |
| 8.12 12-Lead ECG Monitoring | 65 |
| 8.12.1 Diagnosis Function | 65 |
| 8.12.2 Measurement and Interpretation | 66 |
| Chapter 9 Monitoring RESP..... | 67 |
| 9.1 Overview | 67 |
| 9.2 RESP Safety Information | 67 |

| | |
|---|-----------|
| 9.3 Resp Display | 67 |
| 9.4 Electrode Placement for Monitoring Resp..... | 68 |
| 9.5 Cardiac Overlay | 68 |
| 9.6 Chest Expansion..... | 68 |
| 9.7 Abdominal Breathing | 68 |
| 9.8 Selecting Resp Lead..... | 68 |
| 9.9 Changing Hold Type | 69 |
| 9.10 Changing the Size of the Respiration Wave..... | 69 |
| 9.11 Using Resp Alarms..... | 69 |
| 9.12 Changing the Apnea Time..... | 69 |
| Chapter 10 Monitoring SpO₂ | 70 |
| 10.1 Overview | 70 |
| 10.2 SpO ₂ Safety Information | 70 |
| 10.3 Measuring SpO ₂ | 71 |
| 10.4 Measurement Procedure..... | 71 |
| 10.5 Understanding SpO ₂ Alarms | 72 |
| 10.6 Adjusting Alarm Limits..... | 72 |
| 10.7 Setting SpO ₂ as Pulse Source..... | 72 |
| 10.8 Setting Pitch Tone | 72 |
| 10.9 Setting Sensitivity | 72 |
| Chapter 11 Monitoring PR..... | 73 |
| 11.1 Overview | 73 |
| 11.2 Setting PR Source..... | 73 |
| 11.3 Setting PR Volume | 73 |
| 11.4 Using Pulse Alarms | 73 |
| 11.5 Selecting the Active Alarm Source..... | 73 |
| Chapter 12 Monitoring NIBP | 74 |
| 12.1 Overview | 74 |
| 12.2 NIBP Safety Information | 74 |
| 12.3 Introducing the Oscillometric NIBP Measurement..... | 75 |
| 12.4 Measurement Limitations | 75 |
| 12.5 Measurement Methods | 75 |
| 12.6 Measurement Procedures | 76 |
| 12.7 Operation Prompts | 77 |
| 12.8 Correcting the Measurement if Limb is not at Heart Level | 78 |
| 12.9 NIBP Alarm..... | 78 |
| 12.10 Resetting NIBP | 78 |
| 12.11 Calibrating NIBP..... | 78 |
| 12.12 Leak Test | 78 |

| | |
|---|-----------|
| 12.12.1 Procedure of Leak Test..... | 78 |
| Chapter 13 Monitoring TEMP..... | 80 |
| 13.1 Overview | 80 |
| 13.2 TEMP Safety Information | 80 |
| 13.3 TEMP Monitoring Setup | 80 |
| 13.4 Calculating Temp Difference | 80 |
| Chapter 14 Monitoring Quick TEMP | 81 |
| 14.1 Overview | 81 |
| 14.2 Quick TEMP Safety Information | 81 |
| 14.3 Measuring Procedure | 82 |
| 14.3.1 Measurement for Oral Temperature | 82 |
| 14.3.2 Measurements for Rectal Temperatures | 83 |
| 14.3.3 Measurements for Axillary Temperatures | 83 |
| 14.4 Changing Temp Unit | 84 |
| Chapter 15 Monitoring IBP | 85 |
| 15.1 Overview | 85 |
| 15.2 IBP Safety Information | 85 |
| 15.3 Monitoring Procedures..... | 85 |
| 15.4 Selecting a Pressure for Monitoring..... | 86 |
| 15.5 Zeroing the Pressure Transducer | 86 |
| 15.6 Zeroing a Pressure Measurement | 86 |
| 15.7 Troubleshooting the Pressure Zeroing (Taking Art for Example)..... | 87 |
| 15.8 IBP Pressure Calibration | 87 |
| 15.9 Troubleshooting the Pressure Calibration | 88 |
| 15.10 IBP Alarm..... | 88 |
| Chapter 16 Monitoring CO₂..... | 89 |
| 16.1 Overview | 89 |
| 16.2 CO ₂ Safety Information | 89 |
| 16.3 Monitoring Procedures..... | 90 |
| 16.3.1 Zeroing the sensor | 90 |
| 16.3.2 LoFlo CO ₂ Module..... | 90 |
| 16.3.3 C5 CO ₂ Module..... | 92 |
| 16.4 Setting CO ₂ Waveform Setup..... | 94 |
| 16.5 Setting CO ₂ Corrections..... | 94 |
| 16.6 Changing CO ₂ Alarms..... | 94 |
| 16.7 Changing Apnea Alarm | 94 |
| Chapter 17 Monitoring C.O..... | 96 |
| 17.1 Overview | 96 |
| 17.2 C.O. Safety Information..... | 96 |

| | |
|--|------------|
| 17.3 C.O. Monitoring Procedures | 96 |
| 17.4 C.O. Measurement Window | 97 |
| 17.5 Measurement Process..... | 99 |
| 17.6 Editing C.O. | 99 |
| 17.7 Blood Temperature Monitoring..... | 100 |
| 17.8 Setting the Computation Constant | 101 |
| 17.9 Recording C.O. Measurements | 101 |
| 17.10 Setting INJ. TEMPSource | 101 |
| Chapter 18 Monitoring AG | 102 |
| 18.1 Overview | 102 |
| 18.2 Safety Information | 102 |
| 18.2.1 Safety Information for ISA Analyzer | 102 |
| 18.2.2 Safety Information for IRMA Module | 104 |
| 18.3 Monitoring Steps..... | 105 |
| 18.3.1 Monitoring Steps for ISA Analyzer..... | 105 |
| 18.3.2 Monitoring Steps for IRMA Module..... | 107 |
| 18.4 Setting Work Mode | 110 |
| 18.5 Setting Alarms..... | 111 |
| 18.6 Setting Apnea Alarm Time | 111 |
| 18.7 Working Status of ISA analyzer | 111 |
| 18.8 Working Status of IRMA Module | 111 |
| 18.9 N ₂ O and O ₂ Compensations | 111 |
| 18.10 Effects of humidity..... | 112 |
| Chapter 19 Freeze | 113 |
| 19.1 Overview | 113 |
| 19.2 Entering/Exiting Freeze Status..... | 113 |
| 19.2.1 Entering Freeze Status..... | 113 |
| 19.2.2 Exiting Freeze Status..... | 113 |
| 19.3 Reviewing Frozen Waveform..... | 114 |
| Chapter 20 Review | 115 |
| 20.1 Trend Graph Review | 115 |
| 20.1.1 Selecting Trend Graph of Specific Parameter..... | 115 |
| 20.1.2 Setting Resolution | 115 |
| 20.1.3 Scrolling Left and Right the Screen | 115 |
| 20.1.4 Switching to the Trend Table | 116 |
| 20.1.5 Record | 116 |
| 20.2 Trend Table Review | 116 |
| 20.2.1 Setting Resolution | 116 |
| 20.2.2 Scrolling the Screen | 116 |

| | |
|---|------------|
| 20.2.3 Switching to Trend Graph | 116 |
| 20.2.4 Recording | 116 |
| 20.3 NIBP Review..... | 116 |
| 20.3.1 Scrolling the Screen | 117 |
| 20.3.2 Recording | 117 |
| 20.4 Alarm Review..... | 117 |
| 20.4.1 Scrolling the Screen | 117 |
| 20.4.2 Selecting Alarm Event of Specific Parameter..... | 117 |
| 20.4.3 Setting Time Index | 117 |
| 20.5 Arr Review | 117 |
| 20.5.1 Scrolling the Screen | 117 |
| 20.6 12-lead Diagnosis Review | 118 |
| 20.6.1 Scrolling the Screen | 118 |
| 20.6.2 Deleting Diagnosis Results | 118 |
| 20.6.3 Switching Between Waveforms and Results..... | 118 |
| 20.6.4 Recording | 118 |
| Chapter 21 Calculation and Titration Table..... | 119 |
| 21.1 Drug Calculation | 119 |
| 21.1.1 Calculation Procedures | 119 |
| 21.1.2 Calculation Unit | 120 |
| 21.2 Titration Table | 120 |
| Chapter 22 Recording..... | 121 |
| 22.1 General Information | 121 |
| 22.2 Performance of the Recorder | 121 |
| 22.3 Recording Type | 122 |
| 22.4 Starting and Stopping Recording | 122 |
| 22.5 Recorder Operations and Status Messages | 123 |
| 22.5.1 Record Paper Requirement | 123 |
| 22.5.2 Proper Operation | 123 |
| 22.5.3 Paper Out..... | 123 |
| 22.5.4 Replacing Paper | 123 |
| 22.5.5 Removing Paper Jam..... | 124 |
| Chapter 23 Other Functions..... | 126 |
| 23.1 Nurse Call | 126 |
| Chapter 24 Using Battery | 127 |
| 24.1 Battery Power Indicator | 127 |
| 24.2 Battery Status on the Main Screen | 127 |
| 24.3 Checking Battery Performance | 127 |
| 24.4 Replacing the Battery..... | 128 |

| | |
|--|------------|
| 24.5 Recycling the Battery | 129 |
| 24.6 Maintaining the Battery..... | 129 |
| Chapter 25 Care and Cleaning | 130 |
| 25.1 General Points | 130 |
| 25.2 Cleaning | 130 |
| 25.2.1 Cleaning the Monitor | 130 |
| 25.2.2 Cleaning the Accessories..... | 131 |
| 25.3 Disinfection | 133 |
| Chapter 26 Maintenance | 134 |
| 26.1 Inspecting | 134 |
| 26.2 Maintenance Task and Test Schedule..... | 134 |
| Chapter 27 Warranty and Service..... | 136 |
| 27.1 Warranty | 136 |
| 27.2 Contact information | 136 |
| Chapter 28 Accessories | 137 |
| 28.1 ECG Accessories | 137 |
| 28.2 SpO ₂ Accessories | 138 |
| 28.3 NIBP Accessories | 139 |
| 28.4 Temp Accessories | 140 |
| 28.5 Quick Temp Accessories | 140 |
| 28.6 IBP Accessories..... | 140 |
| 28.7 CO ₂ Accessories | 140 |
| 28.8 C.O. Accessories | 141 |
| 28.9 AG Accessories | 141 |
| 28.10 Other Accessories..... | 142 |
| A Product Specification | 143 |
| A.1 Classification | 143 |
| A.2 Physical Specifications..... | 143 |
| A.2.1 Size and Weight..... | 143 |
| A.3 Environment Specification..... | 143 |
| A.4 Display | 144 |
| A.5 Battery Specification | 144 |
| A.6 Recorder | 145 |
| A.7 Review..... | 145 |
| A.8 ECG..... | 145 |
| A.8.1 3, 5-lead Monitoring..... | 145 |
| A.8.2 12-lead Monitoring..... | 149 |
| A.9 RESP | 154 |
| A.10 NIBP..... | 154 |

| | |
|---|------------|
| A.10.1 NIBP from EDAN Module..... | 154 |
| A.10.2 NIBP from M3600 Module | 155 |
| A.11 SpO ₂ | 156 |
| A.11.1 SpO ₂ from EDAN Module | 156 |
| A.11.2 SpO ₂ from Nellcor Module | 157 |
| A.12 TEMP | 157 |
| A.13 Quick TEMP | 158 |
| A.14 IBP..... | 158 |
| A.15 CO ₂ | 158 |
| A.16 C.O. | 160 |
| A.17 AG | 161 |
| A.17.1 Phasein Sidestream..... | 161 |
| A.17.2 Phasein Mainstream | 163 |
| A.18 Wirless Network..... | 166 |
| B EMC Information..... | 167 |
| B.1 Electromagnetic Emissions - for all EQUIPMENT and SYSTEMS..... | 167 |
| B.2 Electromagnetic Immunity - for all EQUIPMENT and SYSTEMS | 167 |
| B.3 Electromagnetic Immunity - for EQUIPMENT and SYSTEMS that are not LIFE-SUPPORTING..... | 168 |
| B.4 Recommended Separation Distances | 170 |
| C Default Settings..... | 171 |
| C.1 Patient Information Default Settings | 171 |
| C.2 Alarm Default Settings | 171 |
| C.3 ECG Default Settings | 171 |
| C.4 RESP | 172 |
| C.5 SpO ₂ | 173 |
| C.6 PR | 173 |
| C.7 NIBP | 173 |
| C.8 TEMP | 174 |
| C.9 Quick TEMP..... | 174 |
| C.10 IBP..... | 175 |
| C.11 CO ₂ | 175 |
| C.12 C.O. | 176 |
| C.13 AG | 176 |
| D Abbreviations..... | 178 |

Chapter 1 Indication for Use and Safety Guidance

1.1 Indication for Use

iM80:

The monitor monitors parameters such as ECG (3-lead, 5-lead, 12-lead selectable), Respiration (RESP), Functional arterial oxygen saturation (SpO₂), Invasive or noninvasive blood pressure (2/4 channels IBP NIBP), Cardiac Output (C.O.), Temperature (dual-TEMP), Expired CO₂ and Anesthetic gas (AG). The monitor is equipped with alarms that indicate system faults (such as loose or defective electrodes), physiologic parameters that have exceeded the limits set by the operator, or both.

iM50:

The monitor monitors parameters such as ECG (3-lead, 5-lead selectable), Respiration (RESP), Functional arterial oxygen saturation (SpO₂), Invasive or noninvasive blood pressure (dual-IBP, NIBP), Temperature (dual-TEMP), Expired CO₂ and Quick Temperature (Quick TEMP). The monitor is equipped with alarms that indicate system faults (such as loose or defective electrodes), physiologic parameters that have exceeded the limits set by the operator, or both.

The arrhythmia detection and ST Segment analysis are not intended for neonatal patients.

1.2 Safety Guidance

WARNING

- 1 Before using the device, the equipment, patient cable and electrodes etc. should be checked. Replacement should be taken if there is any evident defect or signs of aging which may impair the safety or performance.
 - 2 Medical technical equipment such as these monitor/monitoring system must only be used by persons who have received adequate training in the use of such equipment and who are capable of applying it properly.
 - 3 EXPLOSION HAZARD-Do not use the device in a flammable atmosphere where concentrations of flammable anesthetics or other materials may occur.
 - 4 SHOCK HAZARD-The power receptacle must be a three-wire grounded outlet. A hospital grade outlet is required. Never adapt the three-prong plug from the monitor to fit a two-slot outlet.
 - 5 Extreme care must be exercised when applying medical electrical equipment. Many parts of the human/machine circuit are conductive, such as the patient, connectors, electrodes, transducers. It is very important that these conductive parts do not come into contact with other grounded, conductive parts when connected to the isolated patient input of the device. Such contact would bridge the patient's isolation and cancel the protection provided by the isolated input. In particular, there must be no contact of the neutral electrode and ground.
-

WARNING

- 6 Magnetic and electrical fields are capable of interfering with the proper performance of the device. For this reason make sure that all external devices operated in the vicinity of the monitor comply with the relevant EMC requirements. X-ray equipment or MRI devices are a possible source of interference as they may emit higher levels of electromagnetic radiation.
 - 7 Route all cables away from patient's throat to avoid possible strangulation.
 - 8 Devices connecting with monitor should be equipotential.
 - 9 Accessory equipment connected to the analog and digital interfaces must be certified according to the respective IEC/EN standards (e.g. IEC/EN 60950 for data processing equipment and IEC/EN 60601-1 for medical equipment). Furthermore all configurations shall comply with the valid version of the standard IEC/EN 60601-1-1. Therefore anybody, who connects additional equipment to the signal input or output connector to configure a medical system, must make sure that it complies with the requirements of the valid version of the system standard IEC/EN60601-1-1. If in doubt, consult our technical service department or your local distributor.
 - 10 Only patient cable and other accessories supplied by EDAN can be used. Or else, the performance and electric shock protection can not be guaranteed, and the patient may be injured.
 - 11 Do not rely exclusively on the audible alarm system for patient monitoring. Adjustment of alarm volume to a low level or off during patient monitoring may result in a hazard to the patient. Remember that the most reliable method of patient monitoring combines close personal surveillance with correct operation of monitoring equipment.
 - 12 When interfacing with other equipment, a test for leakage current must be performed by qualified biomedical engineering personnel before using with patients.
 - 13 During monitoring, if the power supply is off and there is no battery for standby, the monitor will be off, and only the patient information and alarm settings can be saved. After reconnecting the power supply, the user should turn on the monitor for monitoring.
 - 14 Keep away from fire immediately when leakage or foul odor is detected.
 - 15 The device and accessories are to be disposed of according to local regulations after their useful lives. Alternatively, they can be returned to the dealer or the manufacturer for recycling or proper disposal. Batteries are hazardous waste. Do NOT dispose them together with house-hold garbage. At the end of their life hand the batteries over to the applicable collection points for the recycling of waste batteries. For more detailed information about recycling of this product or battery, please contact your local Civic Office, or the shop where you purchased the product.
 - 16 Dispose of the package material, observing the waste control regulations and keeping it out of children's reach.
 - 17 After defibrillation, the screen display recovers within 10 seconds if the correct electrodes are used and applied based on the manufacturers' instructions.
 - 18 This equipment is not intended for family usage.
-
-


CAUTION














- 1 Electromagnetic Interference - Ensure that the environment in which the patient monitor is installed is not subject to any sources of strong electromagnetic interference, such as radio transmitters, mobile telephones, etc.
 - 2 Keep the environment clean. Avoid vibration. Keep it far away from corrosive medicine, dust area, high temperature and humid environment.
 - 3 Do not immerse transducers in liquid. When using solutions, use sterile wipes to avoid pouring fluids directly on the transducer.
 - 4 Do not use autoclave or gas to sterilize the monitor, recorder or any accessories.
 - 5 The device and reusable accessories could be sent back to the manufacturer for recycling or proper disposal after their useful lives.
 - 6 Disposable devices are intended for single use only. They should not be reused as performance could degrade or contamination could occur.
 - 7 Remove a battery whose life cycle has expired from the monitor immediately.
 - 8 Avoid liquid splash on the device. The temperature must be kept between 5°C and 40°C while working. And it should be kept between -20°C and 55°C during transportation and storage.
 - 9 To ensure patient safety, use only parts and accessories manufactured or recommended by EDAN.
 - 10 Federal law restricts this device to sale by or on the order of a physician.
-
-













NOTE:








- 1 Position the device in a location where the operator can easily see the screen and access the operating controls.
- 2 The monitor can only be used on one patient at a time.
- 3 If the monitor gets damp or liquid pours on the monitor, please contact the service personnel of EDAN.
- 4 This monitor is not a device for treatment purposes.
- 5 The pictures and interfaces in this manual are for reference only.
- 6 Regular preventive maintenance should be carried out every two years. You are responsible for any requirements specific to your country.

1.3 Explanation of Symbols on the Monitor

| | |
|---|---|
|  | <p>This symbol indicates that the equipment is IEC/EN60601-1 Type CF equipment. The unit displaying this symbol contains an F-Type isolated (floating) patient applied part providing a high degree of protection against shock, and is suitable for use during defibrillation.</p> |
|---|---|

| | |
|---|--|
|  | This symbol indicates that the instrument is IEC/EN 60601-1 Type BF equipment. The unit displaying this symbol contains an F-Type isolated (floating) patient applied part providing a high degree of protection against shock, and is suitable for use during defibrillation. |
|  | Symbol for “Caution” |
|  | Equipotential grounding terminal |
|  | Alternating Current |
|  | Power Supply switch |
|  | Serial number |
|  | Network port |
|  | USB (Universal Serial Bus) Connection |
|  | Audio alarm is off |
|  | NIBP measurement |
|  | Trend graph |
|  | Freeze |
|  | Record |

| | |
|---|---|
|  | Menu |
|  | VGA output, External Monitor |
|  | RS-232 port |
|  | Nurse call port |
|  | SD Card port |
|  | Signal output port |
|  | Signal output |
|  | The symbol indicates that the device complies with the European Council Directive 93/42/EEC concerning medical devices. |
|  | Authorised representative in the European community |
|  | Date of manufacture |
|  | Manufacturer |
| P/N | Part Number |
|  | Recycle |

| | |
|---|---|
|  | The symbol indicates that the device should be sent to the special agencies according to local regulations for separate collection after its useful life. |
|  | Consult Instructions For Use |
|  | Locked position |
|  | Gas inlet |
|  | Gas outlet (evac) |
|  | ISA equipped to measure CO ₂ only. |
|  | ISA equipped to measure multiple gases. |

Chapter 2 Installation

NOTE:

- 1 The monitor settings must be specified by the authorized hospital personnel.
- 2 To ensure that the monitor works properly, please read the user manual and follow the steps before using the monitor.

2.1 Initial Inspection

Before unpacking, check the packaging and ensure that there are no signs of mishandling or damage. If the shipping cartons are damaged, contact the carrier for compensation and package them again.

Open the package carefully and remove the monitor and accessories. Check that the contents are complete and that the correct options and accessories have been delivered.

If you have any question, please contact your local supplier.

2.2 Mounting the Monitor

If all situations are normal, please place the monitor on a flat, level surface, hung on the bed rail, or mounted on a wall. About how to install the wall mount for the monitor, please refer to the following content.

2.2.1 Installing Wall Mount for the Monitor

For how to install wall mount for the monitor, please refer to *Wall Mounting Bracket Assembly Instruction*.

2.3 Connecting the Power Cable

Connection procedure of the AC power line is listed below:

- 1 Make sure the AC power supply complies with the following specifications: 100V-240V~, 50Hz/60Hz.
- 2 Apply the power line provided with the monitor. Plug the power line to inlet interface of the monitor. Connect the other end of the power line to a grounded 3-phase power output.

NOTE:

Connect the power line to the jack special for hospital usage.

2.4 Checking Out the Monitor

Make sure there is no damage on the measurement accessories and cables. Then turn on the monitor, check whether the monitor can start normally. Make sure all alarm lamps light up and the alarm sound is heard when turning on the monitor.

WARNING

If any sign of damage is detected, or the monitor displays some error messages, do not use it on any patient. Contact Customer Service Center immediately.

NOTE:

- 1 Check all the functions of the monitor and make sure that the monitor is in good status.
- 2 If rechargeable batteries are provided, charge them after using the device every time, to ensure the electric power is enough.
- 3 The interval between double pressing of POWER switch should be longer than 1 minute.
- 4 After continuous 360-hour runtime, please restart the monitor to ensure the monitor's steady performance and long lifespan.

2.5 Checking the Recorder

If your monitor is equipped with a recorder, open the recorder's door to check if paper is properly installed in the slot. If no paper exists, refer to Chapter *Recording* for details.

2.6 Setting the Date and Time

To set the date and time:

1. Select **Menu > System Setup > Date Time Setup**.
2. Adjust the date display format based on the user's habit.
3. Set the correct time of year, month, day, hour, min and sec from the popup menu and press **Exit**.

2.7 Handing Over the Monitor

If you are handing over the monitor to the end-users directly after configuration, make sure that it is in the monitoring mode.

The users must be adequately trained to use the monitor before monitoring a patient. To achieve this, they should have access to, and read, the following documentation delivered with the monitor:

- User Manual (this book) - for full operating instructions.
- Quick Reference Card - for quick reminders during use.

Chapter 3 Basic Operation

This manual is for clinical professionals using the iM50 and iM80 patient monitors. Unless otherwise specified, the information here is valid for all the above products.

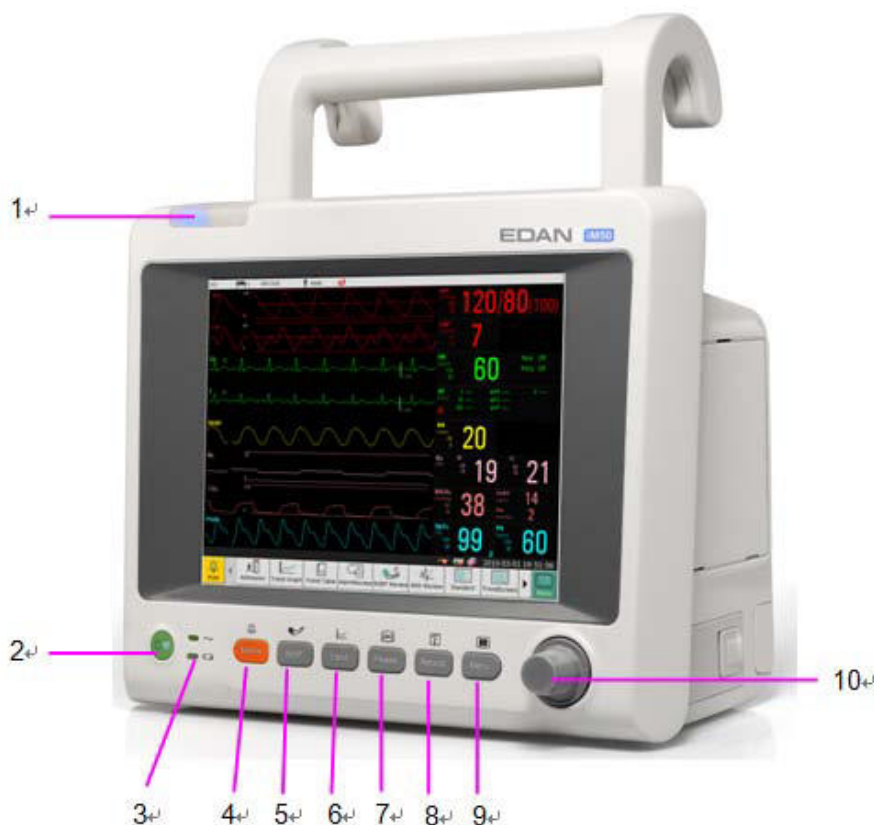
This user manual describes all features and options. Your monitor may not have all of them; they are not all available in all geographies. Your monitor is highly configurable. What you see on the screen, how the menus appear and so forth, depends on the way it has been tailored for your hospital and may not be exactly as shown here.

3.1 Introducing iM50/iM80 Series

The iM50/iM80 series patient monitors offers a monitoring solution optimized for the surgical, cardiac, medical and neonatal care environments. The monitor stores data in trend and event. You can see tabular trends (vital signs) and document them on a local.



3.1.1 iM50 Major Parts and Keys

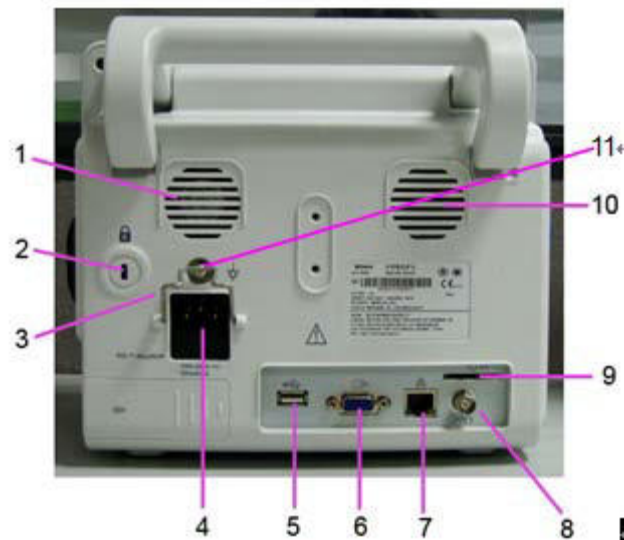
The iM50 patient monitor has an 8.4-inch TFT color flat panel display. Up to 11 waves can be shown on iM50 screens.



iM50 Front View

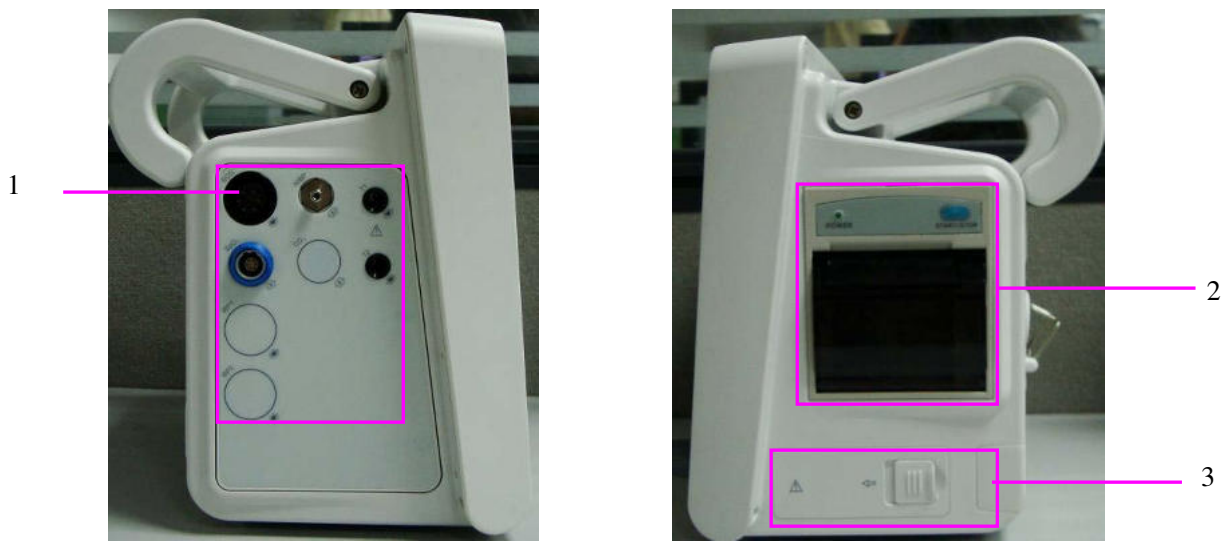
| | |
|---|--|
| 1 | Alarm indicator — when an alarm occurs, the alarm indicator will light or flash. The color of light represents the alarm level. |
| 2 | Power supply switch — when the monitor is connected to the AC power supply, press the key to turn the monitor on. When the monitor is turned |

| | |
|----|---|
| | on, press the key to turn the monitor off. |
| 3 | Battery indicator, refer to Section <i>Battery Indicator</i> for details. |
| 4 | <p>Mute — Press this button to pause the alarm. All the audio alarm will be closed. At the same time, the message of Temporary Alarm Mute **s and the symbol  will be displayed in the information area. When you repress it or the pause time is over, the system will resume the normal monitoring status and the message of Temporary Alarm Mute **s and icon will vanish. Symbol  is shown in the information area. Pressing or holding the button again can resume the alarm.</p> <p>Further Alarm Mute information can be found in the chapter Alarm Mute.</p> |
| 5 | Start / Stop NIBP measurement — Press this button to inflate the cuff and start blood pressure measurement. During the measurement, press the button to stop the measurement. |
| 6 | Trend Key — Press this button to enter trend table review interface. |
| 7 | Freeze /Unfreeze — In normal mode, press this button to freeze all the waveforms on the screen. In Freeze mode, press this button to restore the waveform refreshing. |
| 8 | Start / Stop Recording — Press this button to start a real-time recording. During the recording, press this button again to stop recording. |
| 9 | Menu — Press this button to return to the main interface when there is no menu open. |
| 10 | Rotary Knob (hereinafter called knob) — The user can rotate the knob clockwise or anticlockwise. This operation can make the highlighted item shift up, down, left or right to choose the desired item. Remember, when using the knob, rotate this button to highlight, and press it to select the item. |



iM50 Back View

| | |
|----|--|
| 1 | Fan |
| 2 | Anti-theft lock interface |
| 3 | Security Lock |
| 4 | Power Supply Inlet |
| 5 | USB Interface |
| 6 | VGA Interface |
| 7 | Network Interface |
| 8 | Defibrillator Synchronization/ Analog Output |
| 9 | SD Card |
| 10 | Speaker |
| 11 | Equipotential grounding terminal, if the monitor or other processing unit are used in internal examinations on the heart, ensure that the room incorporates an equipotential grounding system to which the monitor and other processing unit have separate connection. |

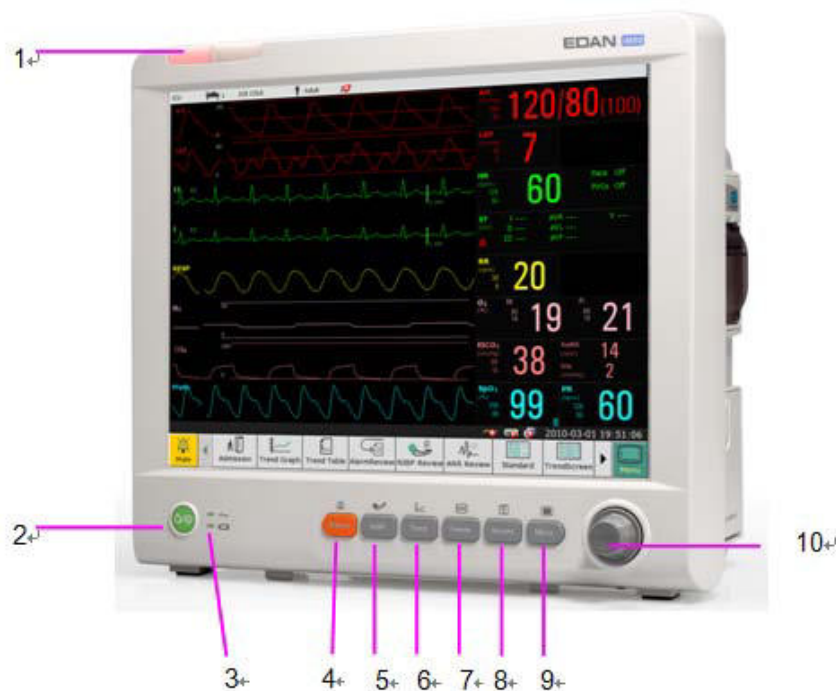


iM50 Side View

| | |
|---|------------------|
| 1 | Sensor interface |
| 2 | Recorder door |
| 3 | Battery door |



3.1.2 iM80 Major Parts and Keys

The iM80 patient monitor has a 15-inch TFT color flat panel display. Up to 13 waves can be shown on iM80 screens.



iM80 Front View

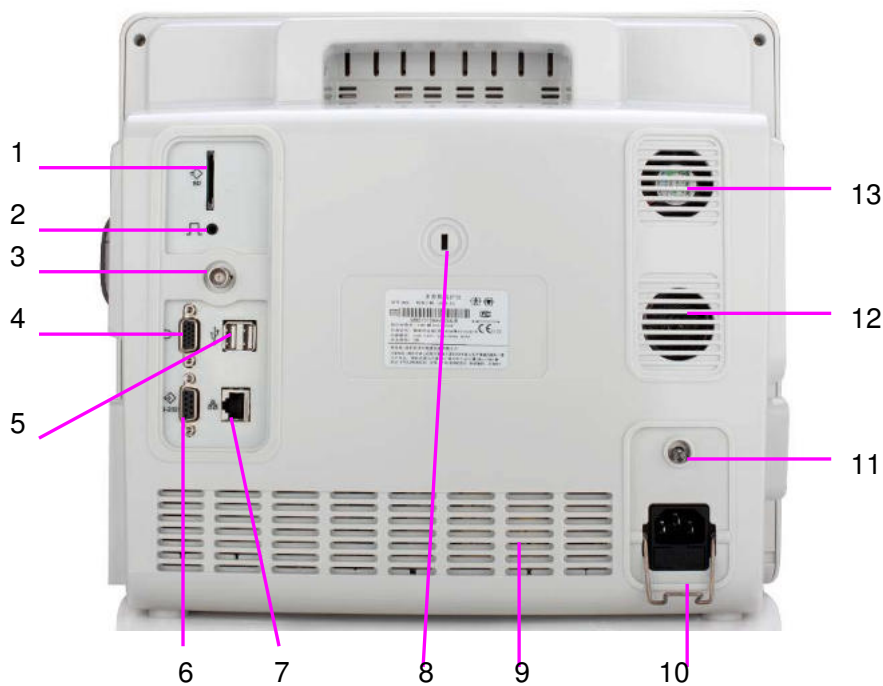
| | |
|---|--|
| 1 | Alarm indicator — when an alarm occurs, the alarm indicator will light |
|---|--|

| | |
|----|---|
| | or flash. The color of light represents the alarm level. |
| 2 | Power supply switch — when the monitor is connected to the AC power supply, press the key to turn the monitor on. When the monitor is turned on, press the key to turn the monitor off. |
| 3 | Battery indicator, refer to Section <i>Battery Indicator</i> for details. |
| 4 | Mute — Press this button to pause the alarm. All the audio alarm will be closed. At the same time, the message of Temporary Alarm Mute **s and the symbol  will be displayed in the information area. When you repress it or the pause time is over, the system will resume the normal monitoring status, and the message of Temporary Alarm Mute **s and icon will vanish. Symbol  is shown in the information area. Pressing or holding the button again can resume the alarm. Further Alarm Mute information can be found in the chapter Alarm Mute. |
| 5 | Start / Stop NIBP measurement — Press this button to inflate the cuff and start blood pressure measurement. During the measurement, press the button to stop the measurement. |
| 6 | Trend Key — Press this button to enter trend table review interface. |
| 7 | Freeze /Unfreeze — In normal mode, press this button to freeze all the waveforms on the screen. In Freeze mode, press this button to restore the waveform refreshing. |
| 8 | Start / Stop Recording — Press this button to start a real-time recording. During the recording, press this button again to stop recording. |
| 9 | Menu — Press this button to return to the main interface when there is no menu open. |
| 10 | Rotary Knob (hereinafter called knob) — The user can rotate the knob clockwise or anticlockwise. This operation can make the highlighted item shift up, down, left or right to choose the desired item. Remember, when using the knob, rotate this button to highlight, and press it to select the item. |



iM80 Side View

| | |
|---|---------------------|
| 1 | Sensor port |
| 2 | AG fixation bracket |
| 3 | Recorder door |
| 4 | Battery door |



iM80 Back View

| | |
|----|--|
| 1 | SD Card |
| 2 | Nurse call port, this port is connected to the call system. When there is an alarm, the monitor outputs nurse call signal to notify the nurse. |
| 3 | Defibrillator synchronization / analog output. When the user selects Analog Output , the monitor outputs the waveform through the auxillary output port. When the user selects Defibrillator , the monitor outputs the defibrillator synchronization signal through the auxillary output port. |
| 4 | VGA output |
| 5 | USB interface, this port is used to connect the USB device. |
| 6 | RS232 interface |
| 7 | Network interface, this port is used to connect to the central monitoring system through the standard network wire. |
| 8 | Anti-theft lock interface |
| 9 | Heat emission hole |
| 10 | Security lock, used to prevent the power supply cord from falling. |
| 11 | Equipotential grounding terminal, if the monitor or other processing unit are used in internal examinations on the heart, ensure that the room incorporates an equipotential grounding system to which the monitor and other processing unit have separate connection. |
| 12 | Speaker |
| 13 | Fan |

3.1.3 iM50/iM80 Configuration

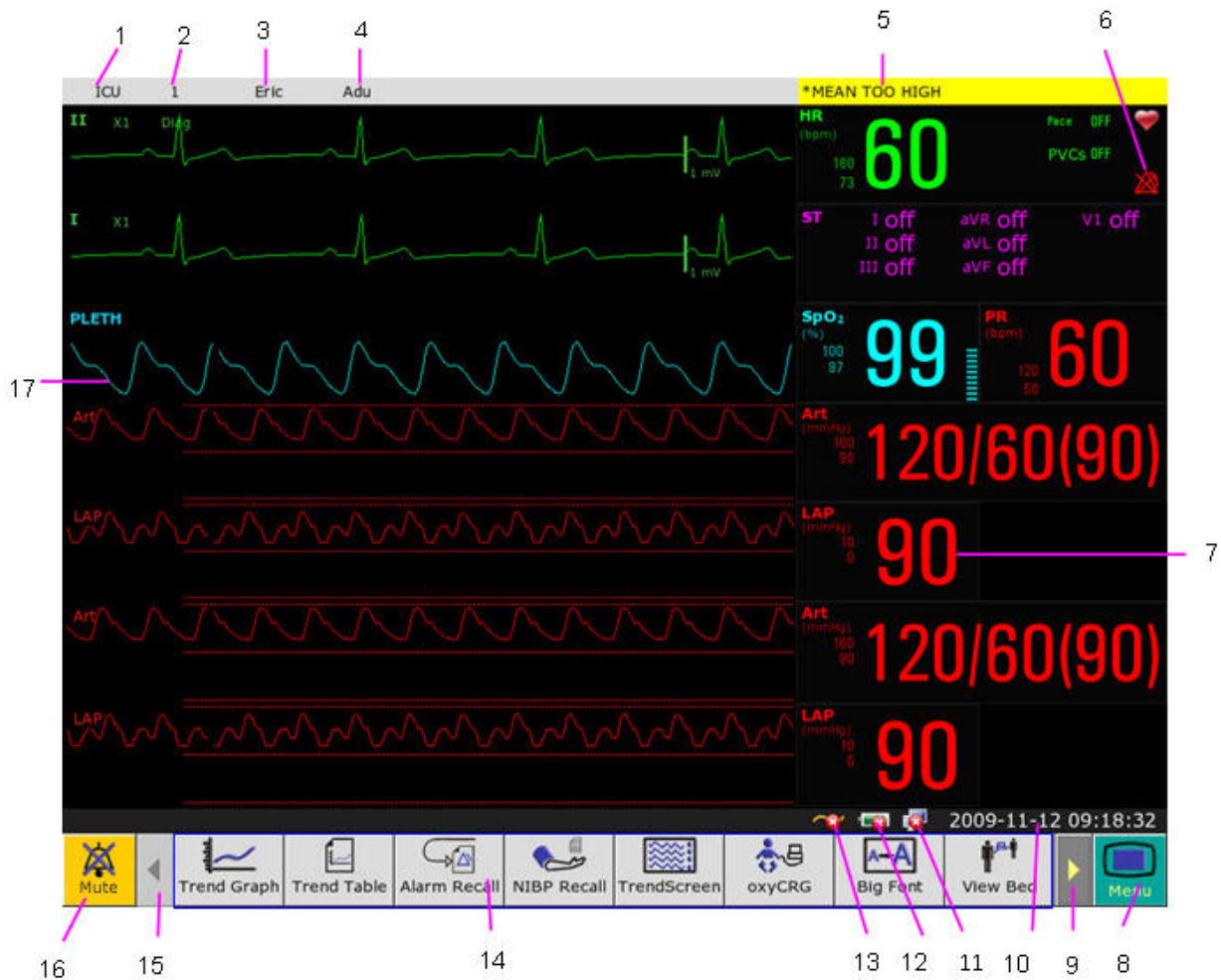
iM50/iM80 configuration is listed as below:

| Model | Size (L×W×H) | Figure | Function Configuration |
|-------|--------------------------------------|--------|--|
| iM50 | 260 mm (L) × 140 mm (W) × 205 mm (H) | Square | ECG, RESP, SpO ₂ , NIBP, IBP, TEMP, Quick TEMP, CO ₂ |
| iM80 | 370 mm (L) × 175 mm (W) × 320 mm (H) | Square | ECG, RESP, SpO ₂ , NIBP, TEMP, IBP, C.O., CO ₂ , AG |

3.2 Operating and Navigating

Everything you need to operate the monitor is contained on its screen. Almost every element on the screen is interactive. Screen elements include measurement numerics, waveforms, screen keys,

information fields, alarms fields and menus. The configurability of the monitor means that often you can access the same element in different ways. For example, you might be able to access an item through its on-screen setup menu, via a hard key, or via a shortcut key. The User Manual always describes how to access items via an on-screen menu. You may use whichever way you find most convenient.



| | |
|---|--|
| 1 | Department |
| 2 | Bed number |
| 3 | Patient name |
| 4 | Patient type |
| 5 | Alarm status area |
| 6 | Alarm off |
| 7 | Measurement value |
| 8 | Menu |
| 9 | Scroll right to display more shortcut keys |

| | |
|----|---|
| 10 | Date and time |
| 11 | Networking symbol |
| 12 | Battery status symbol |
| 13 | AC power supply symbol |
| 14 | Shortcut key area |
| 15 | Scroll left to display more shortcut keys |
| 16 | Mute key |
| 17 | Parameter waveform |

3.2.1 Using Keys

The monitor has four different types of keys:

3.2.1.1 Permanent Keys

A permanent key is a graphical key that remains on the screen all the time to give you fast access to functions.



Menu – enter the main setup menu.


















Mute key – close the audio alarm to switch off the alarm.

3.2.1.2 Shortcut Keys

A shortcut key is a configurable graphical key, located at the bottom of the main screen. It gives you fast access to functions. The selection of shortcut keys available on your monitor depends on your monitor configuration and on the options purchased.

| | |
|--|---|
| | Directly select this item on the screen to quickly admit a patient |
| | Directly select this item on the screen to enter trend graph review interface |
| | Directly select this item on the screen to enter trend table review interface |
| | Directly select this item on the screen to enter the alarm event review interface |
| | Directly select this item on the screen to enter the NIBP review interface |
| | Directly select this item on the screen to enter the ARR review interface |

| | |
|---|--|
|  | Directly select this item on the screen to enter the TrendScreen interface |
|  | Directly select this item on the screen to enter the standard interface |
|  | Directly select this item on the screen to enter the OxyCRG interface |
|  | Directly select this item on the screen to enter the Large Font interface |
|  | Directly select this item on the screen to enter the module switch interface |
|  | Directly select this item on the screen to change the key volume |
|  | Directly select this item on the screen to adjust the screen brightness |
|  | Directly select this item on the screen to calibrate the touch screen |
|  | Directly select this item on the screen to zero the IBP sensor |
|  | Directly select this item on the screen to enter the alarm setup interface |
|  | Directly select this item on the screen to change the beat volume |
|  | Directly select this item on the screen to enter the menu |
|  | Directly select this item on the screen to mute the alarm |
|  | Directly select this item on the screen to enter standby mode |
|  | Select this item by the knob to enable the touchscreen operation |

3.2.1.3 Hardkeys

A hardkey is a physical key on a monitoring device, such as the recording key on the front panel.

3.2.1.4 Pop-up keys

Pop-up keys are task-related graphical keys that appear automatically on the screen when required. For example, the confirmation pop-up key appears only when you need to confirm a change.

3.3 Operating Mode

3.3.1 Demo Mode

To change the operating mode into the demo mode, please refer to the following procedure:

1. Select the **Demo** key on the screen directly or.

2. Select **Menu > Common Function**, then choose **Demo Mode** from the popup interface and input password **3045**.

After entering **Demo Mode**, the monitor will perform the followings:


- Storing no data of new patient.
- Pausing to give all types of alarm.
- Pausing to transmit patient data to CMS and other network devices.
- Pausing to store the currently recorded data, and clearing the memory used to store recording and printing data.
- Real data: the parameter measurement value and real waveform displayed on the screen are from the predefined analog data, not the truly monitoring patients' data.
- History data: the monitor will store the analog real-time data in Demo mode, including trend data, patient information, alarm event, waveform and setting.

To exit **Demo Mode**, select **Menu > Common Function > Demo Mode** to exit.

WARNING

This is for demonstration purposes only. You must not change into Demo Mode during monitoring. In Demo Mode, all stored trend information is deleted from the monitor's memory.

3.3.2 Standby Mode

Standby mode can be used when you want to temporarily interrupt monitoring. To enter standby mode, please press the shortcut key  on the screen directly. To resume monitoring, select anything on the screen or press any key.

3.4 Changing Monitor Settings

3.4.1 Adjusting Screen Brightness

To change the screen brightness, please:

1. Press the **Brightness** key on the screen directly or.
2. Select **Menu > Common Function > Brightness**, and select the appropriate setting for the screen brightness. **10** is the brightest, **1** is the least bright.

Your monitor may be configured with lower brightness in standby mode and also for transport to conserve battery power.

3.4.2 Changing Date and Time

To change the date and time, please refer to Section *Setting Date and Time*.

WARNING

Change to date and time will influence the storage of trend data.

3.5 Adjusting Volume

3.5.1 Adjusting Key Volume

The key volume is the volume you hear when you select any field on the monitor screen or when you turn the knob. To adjust the key volume, please:

1. Select the **Key Volume** key on the screen directly or.
2. Select **Menu > System Setup > Key Volume**, then select the appropriate setting for the key volume: **Five** is the loudest and **Zero** is the quietest.

3.5.2 Adjusting Alarm Volume

To change the alarm volume, please

1. Press the **Alarm Volume** key on the screen directly or.
2. Select **Menu > Alarm Setup > Alarm Volume**, and select the desired setting from the popup interface. For detailed information, please refer to Section *Controlling Alarm Volume*.

3.5.3 Adjusting Beat Volume

To change the beat volume, please press the **Beat Volume** key on the screen directly or refer to Section *Setting Beat Volume*.

3.6 Checking Your Monitor Version

To check the monitor version, please select **Menu > Common Function > About** to check the monitor software revision.

3.7 Networked Monitoring

Your monitor can be connected to the wired network. If the monitor is networked, a network symbol is displayed on the screen.

3.8 Setting Languages

To change the language, please:



1. Select **Menu > Maintenance > User Maintain**, then type the correct password **ABC** into the displayed interface.
2. Select the **Language** option on the popup interface to open the language list.
3. Select the desired language from the list. To make the change validate, please restart the monitor.

3.9 Understanding Screens



Your monitor comes with a set of preconfigured screens, optimized for common monitoring scenarios such as OR adult, or ICU neonatal. A screen defines the overall selection, size and position of waves, numerics and shortcut keys on the monitor screen when you switch on the monitor. You can easily switch between different screens during monitoring. Screens do NOT affect alarm settings, patient category and so forth. When you switch from a complex to a less complex screen layout, some measurements may not be visible but are still monitored in the background. For detailed information, please refer to *Chapter User Interface*.

3.10 Calibrating Screens

To calibrate the screen, please refer to the following steps:

1. Select the **Touch Calib** shortcut key on the screen directly or select **Menu > Maintenance > User Maintain**, then input maintenance password **ABC**, then select **TouchScr Calibration** from the popup interface.
2. The symbol  appears on the screen.
3. Click on the central point of the symbol .
4. After successful calibration, the message of **Screen Calibration Completed** appears on the screen. Then select **Exit** to finish the calibration.

3.11 Disabling the TouchScreen

The user can disable touchscreen operation by pressing and holding the **Menu** shortcut key for 3 seconds. A message of **Screen Locked** and the symbol  are displayed at the bottom of screen if the touchscreen is disabled. To enable the touchscreen operation, turn the knob to select the symbol  and press it.

3.12 Using the Barcode Scanner

To enter the barcode setup menu, please select **Menu > Maintenance > User Maintain**, after entering the required password **ABC**, select **Other Setup > BarCode Setup**. Then the user can set serial No, last Name, first Name and so on.

If **Auto Update** is set to **On**, the patient information is updated automatically by using a bar code scanner. If **Auto Update** is set to **Off**, the user needs to update the patient information manually.

Chapter 4 Alarms

The alarm information here applies to all measurements. Measurement-specific alarm information is discussed in the sections of individual measurements.

WARNING

A potential hazard can exist if different alarm presets are used for the same or similar equipment in any single area, e.g. an intensive care unit or cardiac operating room.

4.1 Alarm Category

The monitor provides three types of alarm: physiological alarms, technical alarms and prompts.

4.1.1 Physiological alarms

If one or several physiological parameters of the currently monitored patient exceed the predefined alarm limit, for example: SpO₂ values exceed the alarm limit, the monitor will give an alarm, and this type of alarm is called physiological alarms. About the detailed alarm information, please refer to the Section *physiological alarm information*.

4.1.2 Technical Alarms

If one or several technical status of the device is in abnormal status, such as: lead off or low battery and so on, the monitor will give an alarm. And this type of alarm is called technical alarms. About the detailed alarm information, please refer to Section *technical alarm information*.

4.1.3 Prompts

The monitor can give the character indication of monitoring process or other functions, such as: ARR Relearning and so on. And this character is called prompts. About the detailed alarm information, please refer to Section *Prompts*.

4.2 Alarm Levels

In terms of severity, the device's alarm levels can be classified into three categories: high level alarms, medium level alarms and low level alarms.

1. High level alarms

Indicating that the patient is in a life threatening situation and an emergency treatment is demanded.

2. Medium level alarms

The patient's vital signs appear abnormally or the device system status is abnormal, indicating that prompt operator response is required.

3. Low level alarms

The patient's vital signs appear abnormal or the device system status appears abnormally, indicating that operator awareness is required.

The high/medium/low-level alarms are indicated by the system in following different ways:


| Alarm level | Prompt | Physiological alarms | Technical alarms |
|-------------|--|---|---|
| High | Mode is “DO-DO-DO-----DO-DO, DO-DO-DO-----DO-DO”, which is triggered once every 5 seconds. | The alarm indicator flashes in red, with a frequency of 1.4Hz ~ 2.8Hz. The alarm message flashes with red background, and the symbol *** is displayed at the alarm area. | The alarm indicator flashes in red. The alarm message flashes with red background, and the symbol *** is displayed at the alarm area. |
| Medium | Mode is “DO-DO-DO”, which is triggered once every 25 seconds. | The alarm indicator flashes in yellow, with a frequency of 0.4Hz ~ 0.8Hz. The alarm message flashes with yellow background, and the symbol ** is displayed at the alarm area. | No definition |
| Low | Mode is “DO-”, which is triggered once every 30 seconds. | The alarm indicator flashes in yellow. The alarm message flashes with yellow background, and the symbol * is displayed at the alarm area. | The alarm indicator flashes in blue. The alarm message flashes with yellow background, and the symbol * is displayed at the alarm area. |

The sound pressure range for audible alarm signals is from 45 dB to 85 dB.

4.3 Controlling Alarm


4.3.1 Switching the Individual Alarm Off

To switch the alarm off, please select **XX Setup > Alarm Setup** (XX stands for parameter name)

and set **Switch** to **Off** from the popup list. Alarm Off symbol  is displayed on the parameter area if the parameter alarm is switched to off.

4.3.2 Temporary Alarm Mute


The monitor will give an audible alarm when there is an alarm during the audio alarm pause and the alarm indicator and screen flash indicating there is an alarm. The top of monitor displays the following:

1. Alarm pause symbol .
2. The remaining pause time is displayed in text and the word background is red.

The user can set the audio alarm pause to **60 s**, **120 s**, or **180 s** based on the requirement.

4.3.3 Alarm Mute

To mute the alarm, please select **Menu > Maintenance > User Maintain > Alarm Setup**, and set

Mute to On, then press the  key on the front panel for more than three seconds or press the **Silence** shortcut key on the screen directly.

And alarm reminder tone DO is heard every three minutes.

4.3.4 Controlling Alarm Volume

The monitor provides five levels of alarm volume: **1**, **2**, **3**, **4** and **5**. For adjusting the alarm volume, please refer to Section *Adjusting Alarm Volume*.

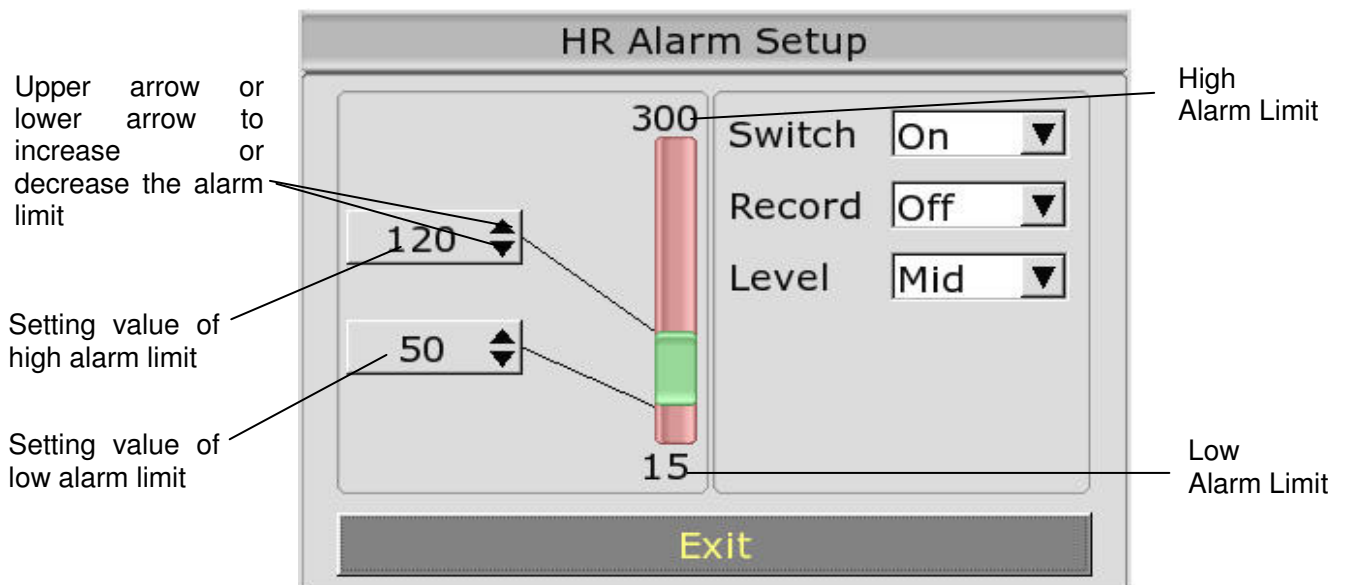
4.3.5 Setting Alarm Limits

WARNING

1. Prior to monitoring, make sure that the alarm limit settings are appropriate for your patient.
2. Setting alarm limits to extreme values may cause the alarm system to become ineffective.

To change individual measurement alarm limits, take HR alarm for example, please refer to the following steps:

1. Select the **HR Parameter** area.
2. Select **HR Setup > Alarm Setup**. Then set the alarm limit to the desired value from the popup interface. For how to set the alarm limit, please refer to the following figure:



4.4 Latching Alarms

To set the alarm latch function, please select **Menu > Maintenance > User Maintain > Alarm Setup** and set **Alarm Latch** from the pull-down list. If it is set to **On**, when an alarm occurs, the monitor will display the alarm message of the parameter in the alarm status area. If the parameter resumes to normal, the alarm information of this parameter still displays in the alarm display area and the alarm time is also displayed. If many parameters appear to be latching alarms, the alarm messages are displayed in the physiological alarm message area in turn.

To deselect the alarm latch, please set **Alarm Latch** to **Off**. When **Alarm Latch** is set to **Off**, the latch function is invalid.

4.5 Disabling Sensor Off Alarms

To set sensor off alarm, please select **Menu > Maintenance > User Maintain** and enter the required password **ABC**. Then select **Alarm Setup** and set **Sensor Off Alm** from the pull-down list. If it is set to **On**, and a sensor off alarm occurs, the user can press the Mute key on the front panel to disable the alarm signal. Then the alarm indicator stops flashing and the monitor is in temporary mute alarm status. If the user presses the Mute key again or the temporary mute time ends, the monitor resumes audible alarm. And the monitor will display prompt message of sensor off alarm.

4.6 Testing Alarms

When you switch the monitor on, a selftest is started. You must check that the alarm indicator lights and that you hear a single tone. This indicates that the visible and audible alarm indicators are functioning correctly. For further testing of individual measurement alarms, perform the measurement on yourself or use a simulator. Adjust alarm limits and check that appropriate alarm behavior is observed.

Chapter 5 Alarm Information

5.1 Physiological Alarm Information

| Message | Cause | Alarm level |
|--------------|---|-----------------|
| ECG Too Weak | Can not detect the signal in designated time period. | High |
| ST-X High | ST measuring value is above the upper alarm limit. (X stands for I, II, III, aVR, aVL, aVF, V, V1, V2, V3, V4, V5 or V6) | User-selectable |
| ST-X Low | ST measuring value is below the lower alarm limit.(X stands for I, II, III, aVR, aVL, aVF, V, V1, V2, V3, V4, V5 or V6) | User-selectable |
| ASYSTOLE | No QRS is detected for 4 consecutive seconds | User-selectable |
| VFIB/VTAC | Ventricular tachycardia: The fibrillation wave lasts for 4 consecutive seconds; or the number of continuous Vent beats is larger than the upper limit of cluster Vent beats (≥ 5). The RR interval is less than 600ms. | User-selectable |
| VT>2 | $3 \leq$ the number of cluster PVCs < 5 | User-selectable |
| COUPLET | 2 consecutive PVCs | User-selectable |
| BIGEMINY | Vent Bigeminy | User-selectable |
| TRIGEMINY | Vent Trigeminy | User-selectable |
| R ON T | A type of single PVC under the condition that HR<100, R-R interval is less than 1/3 the average interval, followed by a compensating pause of 1.25X the average R-R interval (the next R wave advances onto the previous T wave). | User-selectable |
| PVC | Single PVCs not belonging to the type of above mentioned PVCs. | User-selectable |
| TACHY | 5 consecutive QRS complex, RR interval is less than 0.5s. | User-selectable |
| BRADY | 5 consecutive QRS complex, RR interval is longer than 1.5s. | User-selectable |
| MISSED BEATS | When HR is less than 100 beats/min., no heart beat is tested during the period 1.75 times of the average RR interval; or When HR is higher than 100 beat/min, no beat is tested within 1 second. | User-selectable |
| IRR | IRREGULAR RHYTHM: The patient has irregular heart rate, check patient's condition, electrodes, cables and leads. | User-selectable |

| | | |
|---------------------------|--|-----------------|
| PNC | PACE NOT CAPTURE: After the pacemaker is paced, QRS complex can not be detected during 300ms. | User-selectable |
| PNP | PACER NOT PACED: After the QRS complex, no pace is detected during 1.75 times of RR interval. | User-selectable |
| VBRADY | VENTRICULAR BRADYCARDIA: The patient has irregular HR, and his average HR is less than 60bpm. Check his condition, electrodes, cables and leads. | User-selectable |
| VENT | VENTRICULAR RHYTHM: The patient has irregular heart rate, check patient's condition, electrodes, cables and leads. | User-selectable |
| RESP APNEA | RESP can not be measured within specific time interval. | High |
| RR High | RESP measuring value is above upper alarm limit. | User-selectable |
| RR Low | RESP measuring value is below lower alarm limit. | User-selectable |
| HR High | HR measuring value is above the upper alarm limit. | User-selectable |
| HR Low | HR measuring value is below the lower alarm limit. | User-selectable |
| SpO ₂ High | SpO ₂ measuring value is above upper alarm limit. | User-selectable |
| SpO ₂ Low | SpO ₂ measuring value is below lower alarm limit. | User-selectable |
| SpO ₂ No Pulse | The signal of the measurement site is too weak, so the monitor can't detect the pulse signal. | High |
| PR High | PR measuring value is above upper alarm limit. | User-selectable |
| PR Low | PR measuring value is below lower alarm limit. | User-selectable |
| T1 High | Measuring value of T1 channel is above upper alarm limit. | User-selectable |
| T1 Low | Measuring value of T1 channel is below lower alarm limit. | User-selectable |
| T2 High | Measuring value of T2 channel is above upper alarm limit. | User-selectable |
| T2 Low | Measuring value of T2 channel is below lower alarm limit. | User-selectable |
| TD High | Measuring value of TD channel is above upper alarm limit. | User-selectable |
| TEMP High | Measuring value of TEMP is above upper alarm limit. | User-selectable |
| TEMP Low | Measuring value of TEMP is below lower alarm limit. | User-selectable |
| SYS High | SYS measuring value is above upper alarm limit. | User-selectable |
| SYS Low | SYS measuring value is below lower alarm limit. | User-selectable |
| DIA High | DIA measuring value is above upper alarm limit. | User-selectable |

| | | |
|--------------|---|-----------------|
| DIA Low | DIA measuring value is below lower alarm limit. | User-selectable |
| MAP High | MAP measuring value is above upper alarm limit. | User-selectable |
| MAP Low | MAP measuring value is below lower alarm limit. | User-selectable |
| Art SYS High | Art SYS measuring value is above upper alarm limit. | User-selectable |
| Art SYS Low | Art SYS measuring value is below lower alarm limit. | User-selectable |
| Art DIA High | Art DIA measuring value is above upper alarm limit. | User-selectable |
| Art DIA Low | Art DIA measuring value is below lower alarm limit. | User-selectable |
| Art MAP High | Art MAP measuring value is above upper alarm limit. | User-selectable |
| Art MAP Low | Art MAP measuring value is below lower alarm limit. | User-selectable |
| PA SYS High | PA SYS measuring value is above upper alarm limit. | User-selectable |
| PA SYS Low | PA SYS measuring value is below lower alarm limit. | User-selectable |
| PA DIA High | PA DIA measuring value is above upper alarm limit. | User-selectable |
| PA DIA Low | PA DIA measuring value is below lower alarm limit. | User-selectable |
| PA MAP High | PA MAP measuring value is above upper alarm limit. | User-selectable |
| PA MAP Low | PA MAP measuring value is below lower alarm limit. | User-selectable |
| CVP MAP High | CVP MAP measuring value is above upper alarm limit. | User-selectable |
| CVP MAP Low | CVP MAP measuring value is below lower alarm limit. | User-selectable |
| ICP MAP High | ICP MAP measuring value is above upper alarm limit. | User-selectable |
| ICP MAP Low | ICP MAP measuring value is below lower alarm limit. | User-selectable |
| LAP MAP High | LAP MAP measuring value is above upper alarm limit. | User-selectable |
| LAP MAP Low | LAP MAP measuring value is below lower alarm limit. | User-selectable |
| RAP MAP High | RAP MAP measuring value is above upper alarm limit. | User-selectable |
| RAP MAP Low | RAP MAP measuring value is below lower alarm limit. | User-selectable |
| P1 SYS High | P1 SYS measuring value is above upper alarm limit. | User-selectable |
| P1 SYS Low | P1 SYS measuring value is below lower alarm limit. | User-selectable |
| P1 DIA High | P1 DIA measuring value is above upper alarm limit. | User-selectable |
| P1 DIA Low | P1 DIA measuring value is below lower alarm limit. | User-selectable |
| P1 MAP High | P1 MAP measuring value is above upper alarm limit. | User-selectable |
| P1 MAP Low | P1 MAP measuring value is below lower alarm limit. | User-selectable |
| P2 SYS High | P2 SYS measuring value is above upper alarm limit. | User-selectable |
| P2 SYS Low | P2 SYS measuring value is below lower alarm limit. | User-selectable |
| P2 DIA High | P2 DIA measuring value is above upper alarm limit. | User-selectable |
| P2 DIA Low | P2 DIA measuring value is below lower alarm limit. | User-selectable |

| | | |
|-------------------------|--|-----------------|
| P2 MAP High | P2 MAP measuring value is above upper alarm limit. | User-selectable |
| P2 MAP Low | P2 MAP measuring value is below lower alarm limit. | User-selectable |
| EtCO ₂ High | EtCO ₂ measuring value is above upper alarm limit. | User-selectable |
| EtCO ₂ Low | EtCO ₂ measuring value is below lower alarm limit. | User-selectable |
| FiCO ₂ High | FiCO ₂ measuring value is above alarm limits. | User-selectable |
| CO ₂ APNEA | In a specific time interval, no RESP can be detected using CO ₂ module. | High |
| AwRR High | AwRR measuring value is above upper alarm limit. | User-selectable |
| AwRR Low | AwRR measuring value is below lower alarm limit. | User-selectable |
| EtO ₂ High | EtO ₂ measuring value is above upper alarm limit. | User-selectable |
| EtO ₂ Low | EtO ₂ measuring value is below lower alarm limit. | User-selectable |
| FiO ₂ High | FiO ₂ measuring value is above upper alarm limit. | User-selectable |
| FiO ₂ Low | FiO ₂ measuring value is below lower alarm limit. | User-selectable |
| EtN ₂ O High | EtN ₂ O measuring value is above upper alarm limit. | User-selectable |
| FiN ₂ O Low | FiN ₂ O measuring value is below lower alarm limit. | User-selectable |
| EtHAL High | EtHAL measuring value is above upper alarm limit. | User-selectable |
| EtHAL Low | EtHAL measuring value is below lower alarm limit. | User-selectable |
| FiHAL High | FiHAL measuring value is above upper alarm limit. | User-selectable |
| FiHAL Low | FiHAL measuring value is below lower alarm limit. | User-selectable |
| EtENF High | EtENF measuring value is above upper alarm limit. | User-selectable |
| EtENF Low | EtENF measuring value is below lower alarm limit. | User-selectable |
| FiENF High | FiENF measuring value is above upper alarm limit. | User-selectable |
| FiENF Low | FiENF measuring value is below lower alarm limit. | User-selectable |
| EtISO High | EtISO measuring value is above upper alarm limit. | User-selectable |
| EtISO Low | EtISO measuring value is below lower alarm limit. | User-selectable |
| FiISO High | FiISO measuring value is above upper alarm limit. | User-selectable |
| FiISO Low | FiISO measuring value is below lower alarm limit. | User-selectable |
| EtSEV High | EtSEV measuring value is above upper alarm limit. | User-selectable |
| EtSEV Low | EtSEV measuring value is below lower alarm limit. | User-selectable |
| FiSEV High | FiSEV measuring value is above upper alarm limit. | User-selectable |
| FiSEV Low | FiSEV measuring value is below lower alarm limit. | User-selectable |
| EtDES High | EtDES measuring value is above upper alarm limit. | User-selectable |
| EtDES Low | EtDES measuring value is below lower alarm limit. | User-selectable |
| FiDES High | FiDES measuring value is above upper alarm limit. | User-selectable |

| | | |
|-------------------------|---|-----------------|
| FiDES Low | FiDES measuring value is below lower alarm limit. | User-selectable |
| AG FiO ₂ Low | FiO ₂ measurement value is too low. | High |
| AG APNEA | In a specific time interval, no RESP can be detected using AG module. | High |
| TB High | TB measuring value is above upper alarm. | User-selectable |
| TB Low | TB measuring value is below lower alarm. | User-selectable |

5.2 Technical Alarm Information

NOTE:

The ECG alarm information listed in the below table describes the lead names in America. For the corresponding lead names in Europe, please refer to Section *Installing Electrodes*.

| Message | Cause | Alarm Level | Action Taken |
|-------------------|--|-------------|---|
| ECG Lead Off | More than one ECG electrode falls off the skin or ECG cables fall off the monitor. | Low | Make sure that all electrodes, leads and patient cables are properly connected. |
| ECG V Lead Off | ECG electrode V falls off the skin or ECG cables fall off. | Low | |
| ECG LL Lead Off | ECG electrode LL falls off the skin or ECG cables fall off the monitor. | Low | |
| ECG LA Lead Off | ECG electrode LA falls off the skin or ECG cables fall off the monitor. | Low | |
| ECG RA Lead Off | ECG electrode RA falls off the skin or ECG cables fall off the monitor. | Low | |
| ECG Signal Exceed | ECG measuring value is beyond measuring range. | High | |

| | | | |
|---------------------|---|------|---|
| ECG Signal Overflow | The amplitude of ECG signal is too wide. | Low | Please modify the ECG gain. |
| ECG Comm Fail | ECG module failure or communication failure | High | Stop measuring function of ECG module, and notify biomedical engineer or manufacturer's service staff. |
| ECG Noise | ECG measuring signal is greatly interrupted. | Low | Check lead connection and patient condition |
| ECG V1 Lead Off | ECG electrode V1 falls off the skin or ECG cables fall off. | Low | Make sure that all electrodes, leads and patient cables are properly connected. |
| ECG V2 Lead Off | ECG electrode V2 falls off the skin or ECG cables fall off. | Low | |
| ECG V3 Lead Off | ECG electrode V3 falls off the skin or ECG cables fall off. | Low | |
| ECG V4 Lead Off | ECG electrode V4 falls off the skin or ECG cables fall off. | Low | |
| ECG V5 Lead Off | ECG electrode V5 falls off the skin or ECG cables fall off. | Low | |
| ECG V6 Lead Off | ECG electrode V6 falls off the skin or ECG cables fall off. | Low | |
| RESP Comm Fail | RESP module failure or communication failure | High | Stop measuring function of RESP module, and notify biomedical engineer or the manufacturer's service staff. |

| | | | |
|--------------------------------|---|------|--|
| SpO ₂ Sensor Off | SpO ₂ sensor may be disconnected from the patient or the monitor. | Low | Make sure the sensor is well connected to the patient's finger or other parts. |
| SpO ₂ No Sensor | SpO ₂ sensor was not connected well or connected to the monitor, or the connection is loose. | Low | Make sure the monitor and sensor is well connected, reconnect the sensor. |
| SpO ₂ Comm Fail | SpO ₂ module failure or communication failure | High | Stop using measuring function of SpO ₂ module, and notify biomedical engineer or manufacturer's service staff. |
| SpO ₂ Low Perfusion | The pulse signal is too weak or the perfusion of the measurement site is too low | Low | Reconnect the SpO ₂ sensor and change the measurement site. If problem exists, please notify biomedical engineer or manufacturer's service staff. |
| NIBP Comm Fail | NIBP module failure or communication failure | High | Stop using measuring function of NIBP module, and notify biomedical engineer or manufacturer's service staff. |
| NIBP Loose Cuff | Cuff is no properly wrapped or no cuff exists. | Low | Properly wrap the cuff. |
| NIBP Excessive Pressure | Pressure has exceeded the specified upper safety limit. | Low | Measure again, if failure persists, stop measuring function of NIBP module and notify biomedical engineer or manufacturer's service staff. |

| | | | |
|-------------------------|--|------|---|
| NIBP Signal Sturated | Signal amplitude too strong | Low | Stop the patient from moving. |
| NIBP Init Pressure High | The initial pressure is too high during measuring | High | Measure again, if failure persists, stop measuring function of NIBP module and notify biomedical engineer or manufacturer's service staff. |
| NIBP Invalid Reset | The hardware pressure is too high | Low | Measure again, if failure persists, stop using measuring function of NIBP module and notify biomedical engineer or manufacturer's service staff. |
| NIBP Time Out | Measuring time has exceeded the specified time. | Low | Measure again or use other measuring method. |
| NIBP Tube Leak | NIBP cuff or pump has a leakage. | Low | Check the NIBP cuff and pump for leakages. |
| NIBP Cuff Type Error | The cuff type used isn't consistent with the patient type. | Low | Confirm the patient type and change the cuff. |
| Air Pressure Error | Environment atmospheric pressure abnormal or system pressure abnormal. | Low | Check whether the airway is occluded or pressure sensor works properly in pressure meter mode. If the problem still exists, contact your service personnel. |
| NIBP Self Test Error | Sensor or other hardware errors. | High | Contact your service personnel. |

| | | | |
|-----------------------|---|------|---|
| NIBP Pneumatic Leak | NIBP cuff or pump has a leakage. | Low | Check the NIBP cuff and pump for leakages. |
| NIBP System Failure | Hardware abnormal | High | Contact your service personnel. |
| NIBP Cuff Leak | Cuff, pump or airway has a leakage. | Low | NIBP cuff isn't properly connected, or there is a leak in the airway. |
| NIBP Leak Test Error | Hardware abnormal | High | Check whether the airway is occluded or pressure sensor works properly in pressure meter mode. If the problem still exists, contact your service personnel. |
| NIBP Weak Signal | Cuff is too loose or patient pulse is too weak. | Low | Use other method to measure blood pressure. |
| NIBP Excessive Motion | Due to arm motion, signal noise is too large or pulse rate is not regular. | Low | Make sure that the patient under monitoring is motionless. |
| NIBP Range Exceeded | Maybe the patient blood pressure value is beyond the measurement range. | Low | Maybe the patient blood pressure value is beyond the measurement range. |
| NIBP Air Leak | The cuff pressure can't reach the set value within 60sec and 20sec in Neo mode. | Low | Check the connections and the wrapped cuff to see whether they are all prepared well. |
| NIBP Pressure Low | The module isn't able to detect the SYSTOLIC. Apply pressure again. | Low | Inflate again and retry thrice. Check whether the patient has an over high blood pressure or it is interfered by movement. |

| | | | |
|--|--|------|---|
| NIBP Pulse Abnormal | Abnormal oscillometric waveform | Low | Retry twice. Check for hyperkinesia or arrhythmia. |
| NIBP Pulse Signal Weak | Pulse is too low to measure. | Low | Check the patient's condition or the wrapped cuff. |
| TEMP T1 Sensor Off | Temperature cable of TEMP channel1 may be disconnected from the monitor. | Low | Make sure that the cable is properly connected |
| TEMP T2 Sensor Off | Temperature cable of TEMP channe2 may be disconnected from the monitor. | Low | Make sure that the cable is properly connected. |
| Excessive T1 | TEMP1 measuring value is beyond measuring range. | High | Check sensor connection and patient condition |
| Excessive T2 | TEMP2 measuring value is beyond measuring range. | High | Check sensor connection and patient condition |
| TEMP Comm Fail | TEMP module failure or communication failure. | High | Stop measuring function of TEMP module, and notify biomedical engineer or Manufacturer's service staff. |
| YY Sensor Off (YY stands for the IBP label name) | IBP sensor falls off. | Low | Check the sensor connection and reconnect the sensor. |
| YY Comm Fail (YY stands for the label name) | IBP module failure or communication failure | High | Stop measuring function of IBP module, and notify biomedical engineer or Manufacturer's service staff. |

| | | | |
|----------------------------------|---|--------|---|
| C.O. Comm Fail | C.O. module failure or communication failure | High | Stop measuring of C.O. module, or notify biomedical engineer or Manufacturer's service staff. |
| C.O. TI No Sensor | C.O. TI sensor not connected | Low | Insert injectate temperature sensor. |
| C.O. TB No Sensor | C.O. TB sensor not connected | Low | Insert TB sensor. |
| TEMP Out Of Range | TB measuring value is above measuring range. | High | Please check TB sensor. |
| C.O. Lack Param | C.O. measuring needs parameters | High | Please input patient's height and weight. |
| AA Out Of Range | The AA concentration exceeds the accuracy range of AG module. | High | Reduce AA concentration. |
| O ₂ Sensor Error | The oxygen sensor of the sidestream AG module has a failure. | Medium | Stop measuring of AG module, and notify biomedical engineer or Manufacturer's service staff. |
| AG Comm Fail | AG module failure or communication failure. | High | Stop measuring function of AG module, and notify biomedical engineer or Manufacturer's service staff. |
| AG Zero Required | AG module requires zero. | Low | Please perform zeroing. |
| AG Self-Testing | AG module is self testing. | Low | Please wait the self testing finishing. |
| AG Replace O ₂ Sensor | O ₂ sensor needs to be replaced. | High | Please replace the O ₂ sensor. |
| AG Check Adapter | AG module checks adapter. | Low | Please wait check finishing. |
| O ₂ Cali Required | O ₂ needs to be calibrated. | Low | Please calibrate O ₂ . |

| | | | |
|-----------------------------|---|--------|--|
| AG Software Error | AG module software abnormal | High | Please replace software revision. |
| AG Hardware Error | AG module has hardware failure. | High | Please check whether the hardwares work properly. |
| AG Motor Error | AG module motor abnormal | High | Please check whether the motor works properly. |
| AG Uncalibrated | AG module uncalibrated | Low | Please calibrate the AG module. |
| AG Replace Adapter | AG module needs to change adapter. | High | Please replace the adapter. |
| O ₂ Out Of Range | O ₂ is out of range. | High | Please make O ₂ range resume normal. |
| AG TEMP Out Of Range | AG module temperature out of range | High | Please make the temperature resume normal. |
| AG Baro Press Out Of Range | AG module baro pressure out of pressure | High | Please make the baro pressure value resume normal. |
| AG AA Id Unreliable | AG module can't identify the AG agent. | Medium | Reduce gas agent type. |
| AG Span Calib In Progress | AG module is calibrating. | Low | Please wait calibration finishing. |
| AG Calibration Fail | AG module calibration failure | Medium | Please check whether the module works properly. |
| Unable To Calibrate | AG module can't be calibrated. | Medium | Please check whether the module works properly. |
| AG Zero In Progress | AG module is zeroing. | Low | Please wait zeroing. |
| AG Occlusion | The sampling line is clogged. | Medium | Replace the sampling line. |
| AG Init Fail | AG module has a failure. | High | AG module works improperly. |
| AG Data Limit Error | AG module has a failure. | High | AG module works improperly. |

| | | | |
|----------------------------------|--|--------|--|
| AG Usa Error | AG module has a failure. | High | AG module works improperly. |
| AG Cal Fail | AG module fails to calibrate. | High | AG module works improperly. |
| AG Zref Fail | AG module fails to zero. | High | AG module works improperly. |
| AG Change Oxygen Sensor | Replace oxygen sensor of AG module. | Low | Please wait changing finishing. |
| AG No Oxygen Sensor | The oxygen sensor falls off from the AG module. | High | Connect the sensor again. |
| AG Mixed Agents | AG module detects mixture gas agent. | Medium | Close the subsidiary gas agent. |
| CO ₂ Occlude | Water trap of SideStream is occluded. | Low | Make sure the gas exhaust works well |
| CO ₂ Out Of Range | The CO ₂ concentration exceeds the accuracy range of AG module. | High | Reduce CO ₂ concentration. |
| CO ₂ Sensor Faulty | CO ₂ module failure | High | Stop measuring function of CO ₂ module, notify biomedical engineer. |
| CO ₂ Sensor Over Temp | | | |
| CO ₂ Comm Fail | CO ₂ module failure or communication failure | High | Stop using CO ₂ alarm function, and notify biomedical engineer or Manufacturer's service staff. |
| CO ₂ Ram Error | CO ₂ module failure | High | |
| CO ₂ Rom Error | CO ₂ module failure | High | |
| CO ₂ Zero Required | Zero calibration failure | Low | |

| | | | |
|-------------------------------|---|--------|---|
| CO ₂ Check Adapter | The cannula is off or disconnected. | Low | |
| AA Out Range | The AA concentration exceeds the accuracy range of AG module. | High | Reduce AA concentration. |
| O ₂ Sensor Error | The oxygen sensor of the sidestream AG module has a failure. | Medium | Stop measuring of AG module, and notify biomedical engineer or Manufacturer's service staff. |
| QuickTemp Comm Fail | TEMP module failure or communication failure. | High | Stop measuring function of TEMP module, and notify biomedical engineer or manufacturer's service staff. |
| Temp exceed limit | The TEMP value is beyond the range of +25°C ~ +45°C. | Med | Put the sensor into the sensor bracket, take it out and measure again. |
| No Temp Sensor | TEMP sensor is not connected to the TEMP module. | Low | Connect the sensor and the monitor well, and measure again. |
| Ambient temp too high | The Sensor temperature is higher than +40°C. | Low | Put the sensor into the sensor bracket, measure again after the ambient temperature reaches normal value. |
| Ambient temp too low | The Sensor temperature is lower than +10°C. | Low | Put the sensor into the sensor bracket, measure again after the ambient temperature reaches normal value. |

| | | | |
|-----------------------|--|------|---|
| Probe data error | Offline: NTC resistance $>R$ 0 °C; Short: NTC resistance $<R+100$ °C. | Med | Put the sensor into the sensor bracket, take it out and measure again. If the problem persists, stop using measuring function of TEMP module, and notify biomedical engineer or manufacturer's service staff. |
| Probe heater error | Single failure | Med | Put the sensor into the sensor bracket, take it out and measure again. If the problem persists, stop using measuring function of TEMP module, or notify biomedical engineer or manufacturer's service staff. |
| Probe temp too high | The original temperature of sensor $>+33$ °C & $\leq+40$ °C. | Low | Put the sensor into the sensor bracket, measure again after the sensor temperature reaches normal value. |
| Temp Sensor Off | After the sensor temperature reaches Predict value, it descends to the value lower than Predict value. | Med | Reconnect the sensor and make sure that the cable is properly connected. |
| Battery Low | Battery Low | High | Please change the battery or charging. |
| Recorder Out Of Paper | Recorder Out Of Paper | Low | Please install the paper |
| Recorder setup needed | The user presses the RECORD button when the monitor is not installed with a recorder. | Low | Notify the manufacturer's service staff to install and set the recorder. |

5.3 Prompts

| Message | Cause |
|--------------------------------|---|
| ECG Arr Learning | The QRS template building required for Arr. Analysis is in process. |
| SpO ₂ Search Pulse | SpO ₂ module is analyzing the patient signal and searching for the pulse to compute the saturation, when sensor is connected with patient. |
| Manual Measuring | In manual measuring mode |
| Continual Measuring | In continuous measuring mode |
| Auto Measuring | In automatic measuring mode |
| Measure Abort | Measurement over |
| Calibrating | During calibrating |
| Calibrate Abort | Calibration over |
| Leak Testing | During pneumatic test |
| Leak Test Ok | NIBP module has passed leak test. |
| Leak Test Abort | Pneumatic test over |
| Resetting | NIBP module in resetting |
| Please Start | NIBP module is in idle status. |
| Done | NIBP measurement successfully done |
| Continual Measuring | NIBP module performs continual measuring |
| Stat Measuring | NIBP module performs STAT function. |
| Please Switch To Maintain Mode | NIBP module is in normal mode, the user can't start leak test and pressure calibration. Please enter User Maintain > NIBP Maintain and switch to Maintain Mode to perform leak test or pressure calibration. |
| Please Switch To Normal Mode | NIBP module is in maintain mode, the user can't start blood pressure measurement. Please enter User Maintain > NIBP Maintain and switch to Normal Mode to perform blood pressure measurement. |
| Quick TEMP Is Warming Up | Quick TEMP Is Warming Up. |
| Place Probe On Measure Place | Probe isn't placed on the measurement site. |
| CO ₂ Standby | Turn from measuring mode to standby mode, making the module in energy-saving status. |
| CO ₂ Sensor Warm Up | The CO ₂ module is in warm-up state. |
| Excessive C.O. Temp | TB measuring value is beyond measuring range. |

| | |
|---|---|
| Excessive Temp, C.O. Measurement Fail. | C.O. measuring needs parameters. |
| C.O. measure need param | HEMOD calculation needs parameters. |
| Insufficient factors for Hemod Dynamics | HEMO Dynamics calculation needs parameters. |
| No Sensor, C.O. measurement fail | No Sensor,C.O. measurement fail |
| Measuring... | The C.O. module is performing measuring. |
| Ready for new measurement | C.O. module is ready for new measurement. |
| Invalid C.O. result | C.O. measurement result is invalid. |
| C.O. Measurement Complete | C.O. Measurement is completed. |
| C.O. Measurement Abort | C.O. Measurement is aborted. |
| Warm-up over | The monitor displays this message after taking the sensor out of the bracket and warm-up is over. |
| Measure over | After the Predict measuring is over, the data and message display on the interface. |
| Measure time out | No measuring result after the module entering Predict state for 30s. |
| AG Standby | AG module is operating in the standby status. |

5.4 Adjustable Range of Alarm Limits

ECG alarm limits are listed as follows: unit (bpm)

| | Patient Type | ALM HI | ALM LO |
|----|--------------|--------|--------|
| HR | ADU | 300 | 15 |
| | PED | 350 | 15 |
| | NEO | 350 | 15 |

ST analysis alarm limits are listed as follows: unit (mV)

| | ALM HI | ALM LO |
|----|--------|--------|
| ST | 2.0 | -2.0 |

PVCs alarm upper limits are listed as follows:

| | ALM HI | ALM LO |
|------|--------|--------|
| PVCs | 10 | |

RESP alarm limits are listed as follows: unit (rpm)

| Patient Type | ALM HI | ALM LO |
|--------------|--------|--------|
| ADU | 120 | 6 |

| | | |
|-----|-----|---|
| PED | 150 | 6 |
| NEO | 150 | 6 |

SpO₂ alarm limits are listed as follows (unit %):

| | | |
|------------------|--------|--------|
| | ALM HI | ALM LO |
| SpO ₂ | 100 | 0 |

PR alarm limits is listed as follows: unit (bpm)

| | | |
|----|--------|--------|
| | ALM HI | ALM LO |
| PR | 300 | 30 |

NIBP alarm limits are listed as follows (EDAN module): unit (mmHg)

| Patient Type | | ALM HI | ALM LO |
|--------------|-----|--------|--------|
| ADU | SYS | 270 | 40 |
| | DIA | 215 | 10 |
| | MAP | 235 | 20 |
| PED | SYS | 200 | 40 |
| | DIA | 150 | 10 |
| | MAP | 165 | 20 |
| NEO | SYS | 135 | 40 |
| | DIA | 100 | 10 |
| | MAP | 110 | 20 |

NIBP alarm limits are listed as follows (M3600 module): unit (mmHg)

| Patient Type | | ALM HI | ALM LO |
|--------------|-----|--------|--------|
| ADU(PED) | SYS | 250 | 60 |
| | DIA | 200 | 40 |
| | MAP | 235 | 45 |
| NEO | SYS | 120 | 40 |
| | DIA | 90 | 20 |
| | MAP | 100 | 30 |

TEMP alarm limits are listed as follows:

| | | |
|----|----------------|--------------|
| | ALM HI | ALM LO |
| T1 | 50°C (122 ° F) | 0°C (32 ° F) |
| T2 | 50°C (122 ° F) | 0°C (32 ° F) |
| TD | 50°C (90 ° F) | 0°C (0 ° F) |

IBP alarm limits are listed as follows: unit (mmHg)

| | ALM HI | ALM LO |
|-----|--------|--------|
| Art | 300 | 0 |
| RAP | 40 | -10 |
| LAP | 40 | -10 |
| CVP | 40 | -10 |
| PA | 120 | -6 |
| ICP | 40 | -10 |
| P1 | 300 | -50 |
| P2 | 300 | -50 |

CO₂ alarm limits are listed as follows:

| | ALM HI | ALM LO |
|-------------------|----------|--------|
| EtCO ₂ | 150 mmHg | 0 |
| FiCO ₂ | 50 mmHg | 3 |
| AwRR | 150 rpm | 2 rpm |

C.O. alarm limits are listed as follows:

| | ALM HI | ALM LO |
|----|----------------|--------------|
| TB | 43°C (109.4°F) | 23°C(73.4°F) |

Quick Temp alarm limits are listed as follows:

| Patient Type | ALM HI | ALM LO |
|--------------|----------------|-----------------|
| ADU | 42°C (107.6°F) | 35.5°C (95.9°F) |
| PED | 42°C (107.6°F) | 35.5°C (95.9°F) |

AG alarm limits are listed as follows:

| Patient Type | | ALM HI | ALM LO |
|--------------|--------------------|--------|--------|
| ADU | FiCO ₂ | 25.0% | 0.0% |
| | EtCO ₂ | 25.0% | 0.0% |
| | FiO ₂ | 88.0% | 18.0% |
| | EtO ₂ | 90.0% | 18.0% |
| | FiN ₂ O | 100.0% | 0.0% |
| | EtN ₂ O | 100.0% | 0.0% |
| | EtDes | 18.0% | 0% |
| | FiDes | 18.0% | 0% |

| | | | |
|-----|--------------------|---------|-------|
| | EtIso | 5.0% | 0% |
| | FiIso | 5.0% | 0% |
| | EtHal | 5.0% | 0% |
| | FiHal | 5.0% | 0% |
| | EtSev | 8.0% | 0% |
| | FiSev | 8.0% | 0% |
| | EtEnf | 5.0% | 0% |
| | FiEnf | 5.0% | 0% |
| | awRR | 150 rpm | 0 rpm |
| | Apnea Time | 40s | 20s |
| PED | FiCO ₂ | 25.0% | 0.0% |
| | EtCO ₂ | 25.0% | 0.0% |
| | FiO ₂ | 88.0% | 18.0% |
| | EtO ₂ | 90.0% | 18.0% |
| | FiN ₂ O | 100.0% | 0.0% |
| | EtN ₂ O | 100.0% | 0.0% |
| | EtDes | 18.0% | 0% |
| | FiDes | 18.0% | 0% |
| | EtIso | 5.0% | 0% |
| | FiIso | 5.0% | 0% |
| | EtHal | 5.0% | 0% |
| | FiHal | 5.0% | 0% |
| | EtSev | 8.0% | 0% |
| | FiSev | 8.0% | 0% |
| | EtEnf | 5.0% | 0% |
| | FiEnf | 5.0% | 0% |
| | awRR | 150 rpm | 0 rpm |
| | Apnea Time | 40s | 20s |
| NEO | FiCO ₂ | 25.0% | 0.0% |
| | EtCO ₂ | 25.0% | 0.0% |
| | FiO ₂ | 88.0% | 18.0% |
| | EtO ₂ | 90.0% | 18.0% |

| | | | |
|--|--------------------|---------|-------|
| | FiN ₂ O | 100.0% | 0.0% |
| | EtN ₂ O | 100.0% | 0.0% |
| | EtDes | 18.0% | 0% |
| | FiDes | 18.0% | 0% |
| | EtIso | 5.0% | 0% |
| | FiIso | 5.0% | 0% |
| | EtHal | 5.0% | 0% |
| | FiHal | 5.0% | 0% |
| | EtSev | 8.0% | 0% |
| | FiSev | 8.0% | 0% |
| | EtEnf | 5.0% | 0% |
| | FiEnf | 5.0% | 0% |
| | awRR | 150 rpm | 0 rpm |
| | Apnea Time | 40 s | 20 s |

Chapter 6 Managing Patients

6.1 Admitting a Patient

The monitor displays physiological data and stores it in the trends as soon as a patient is connected. This allows you monitor a patient who is not yet admitted. It is however important to admit patients properly so that you can identify your patient on recordings, reports, and networked devices.

During admission you enter data that the monitor needs for safe and accurate operation. For example, the patient category setting determines the algorithm the monitor uses to process and calculate some measurements, the safety limits that are applied for some measurements, and the alarm limit ranges.

To admit a patient, please:

1. Select the **Admission** key on the screen or.
2. Select **Menu > Patient Setup > New Patient**, then a message is displayed to ask the user to confirm to update patient.
3. Click on **No** to cancel this operation; click on **Yes**, the **Patient Info** window is displayed.
4. Enter the patient information:
 - **Serial No:** Enter the patient's medical record number (MRN), for example 12345678.
 - **Last name:** Enter the patient's last name (family name), for example Smith.
 - **First name:** Enter the patient's first name, for example Joseph.
 - **Gender:** Choose **Male** or **Female**.
 - **Type:** Choose the patient type, either **Adult**, **Pediat**, or **Neonat**.
 - **BloodType:** N/A, A, B, AB and O.
 - **Pace:** Choose **On** or **Off** (You must select **On** if your patient has a pacemaker).
 - **Date of Birth:** Enter the patient's date of birth.
 - **Date of Admission:** Enter the patient's date of admission.
 - **Height:** Enter the patient's height.
 - **Weight:** Enter the patient's weight.
 - **Doctor:** Enter any extra information about the patient or treatment.
5. Select **Exit**.

6.1.1 Patient Category and Paced Status

The patient category setting determines the algorithm the monitor uses to process and calculate some measurements, the safety limits that are applied for some measurements, and the alarm limit ranges.

The paced setting determines whether the monitor shows pacemaker pulses or not. When **Pace** is set to **Off**, pace pulses are filtered and therefore do not show in the ECG wave.

WARNING

- 1 Changing the patient category may change the arrhythmia and NIBP alarm limits. Always check alarm limits to make sure that they are appropriate for your patient.
 - 2 For paced patients, you must set Paced to On. If it is incorrectly set to Off, the monitor could mistake a pace pulse for a QRS and fail to give an alarm during asystole.
-
-

6.2 Quick Admit

If you do not have the time or information to fully admit a patient. Complete the rest of the patient information later. To quickly admit a patient, please:

1. Select **Menu > Patient Setup > Quick Admit**, then a message is displayed to ask the user to confirm to update patient.
2. Click on **No** to cancel this operation; click on **Yes** to continue and the **Patient Info** window is displayed, choose **Type** and **Pace** and set them to the correct mode.
3. Select **Exit**.

6.3 Editing Patient Information

To edit the patient information after a patient has been admitted, select **Menu > Patient Setup > Patient Info.**, and make the required changes on the popup interface.

6.4 Updating a Patient

You should always perform an update before starting monitoring for a new patient. When you select **Menu > Patient Setup > Quick Admit**, or **Menu > Patient Setup > New Patient**, a message of **Press 'Yes' to create new patient profile by clearing all current patient data...** is displayed.

- If the user selects **Yes**, the monitor will update the patient information.
- If the user selects **No**, the monitor won't update the patient information and returns to patient setup interface.

NOTE:

Discharging patient will clear the history data in the monitor associated with the patient.

6.5 Central Monitoring System

The monitor can be connected to the central monitoring system. Through the network:

1. The monitor sends patient information, real-time monitoring or measurement data to the central monitoring system.
2. The real-time monitoring information is displayed on the central monitoring system as the same to the monitor, and the central monitoring system can perform some bilateral control. For example: changing patient information, alarm limit and so forth.

For detailed information, please refer to *MFMS-CMS Central Monitoring System User Manual* and *CMS Central Monitoring System User Manual*.

And the monitor supports HL 7 protocol.

Chapter 7 User Interface

7.1 Setting Interface Style

The user can set the interface based on the requirement, and the set options include the following:

- Sweep of the waveform.
- Parameters needing to be monitored.

Change to some settings may have the risk, so only the authorized person can change them. After changing the settings, please notify the operator.

7.2 Selecting Display Parameters

The user can select the display parameters based on the monitoring and measurement requirements. To select the parameter, please:

1. Select **Menu > System Setup > Module Switch**.
2. Select the required parameters from the popup interface.
3. Press **Exit** to exit the menu and the screen will adjust the parameters automatically.

7.3 Changing Waveform Position

The user can exchange the waveform positions of parameter A and parameter B, please refer to the following steps to do so:

1. Select waveform A and open the setup menu of waveform A.
2. Select **Change** from the popup menu and select the desired label name of waveform B from the pull-down list.

7.4 Changing Interface Layout

To change the interface layout, please refer to the following steps:

1. Select **Menu > Display Setting**.
2. Select one interface from the popup menu.
3. The user can implement one kind of function screen based on the requirements. If the user selects the **Large Font** option, there is no function screen to be selected.

7.5 Viewing Trend Screen

To view the short trend screen, the user can press the **Trend Screen** key on the screen directly or select **Menu > Display Setting > View Selection > TrendScreen**.

7.6 Viewing Oxygen Screen

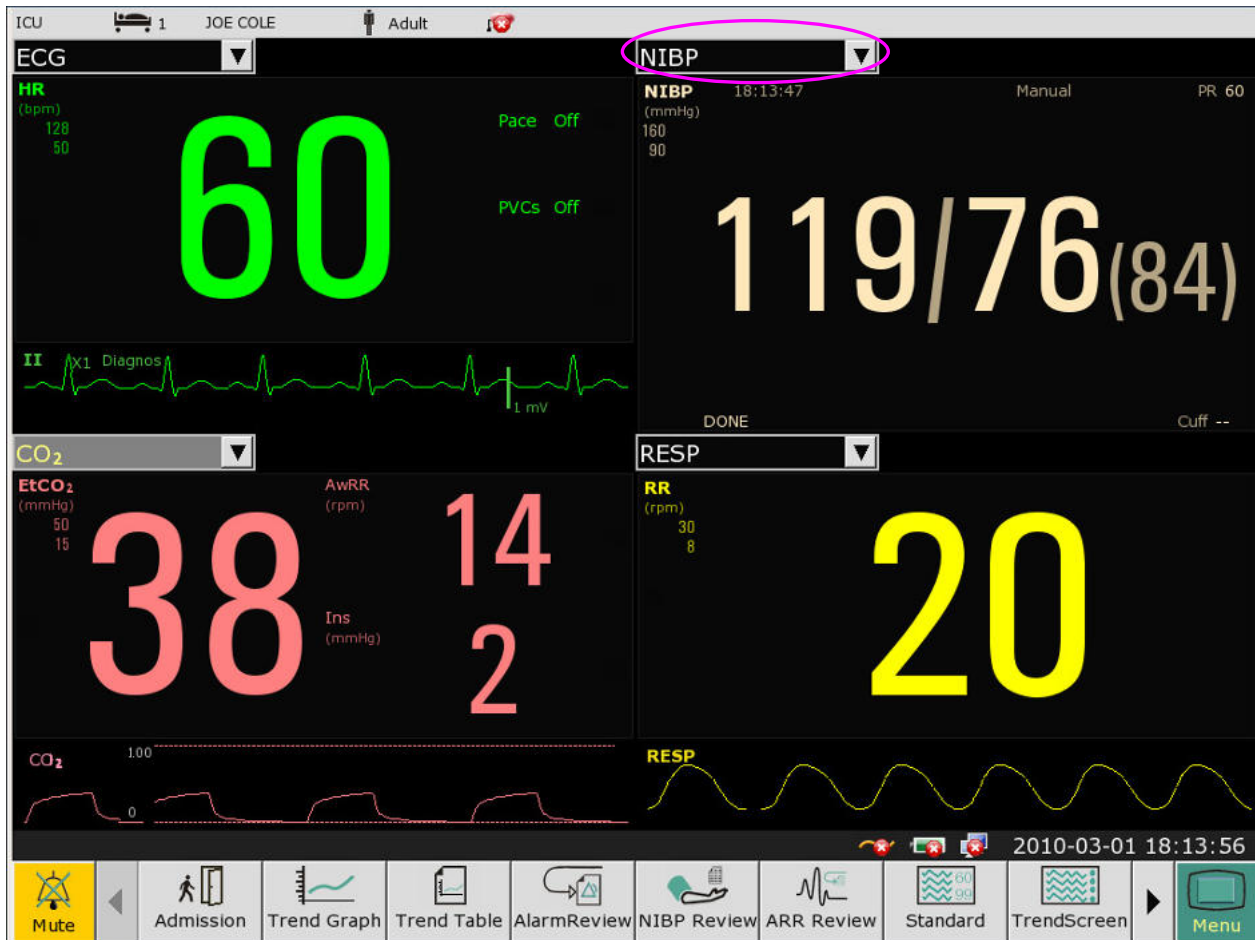
To view the oxygen screen, the user can press the **oxyCRG** key on the screen or select **Menu > Display Setting > View Selection > oxyCRG**. This interface is always used in NICU because the SpO₂, HR and Resp of the neonate are different from those of adults.

7.7 Viewing Large Font Screen

To open the large font screen, please refer to the following steps:

1. Select the **Large Font** key on the screen directly or.
2. Select **Menu > Display Setting > View Selection > Large Font** to open this interface.

To view the large font interface of specific parameter, please select the parameter pull-down dialog on the interface (the red circle shown in the following figure).



7.8 Changing Parameter and Waveform Colors

The user can set the display colors of parameter and waveform as desired. To change the display color, please select **Menu > Maintenance > User Maintain**, enter the required password **ABC**. Then select **Color Select** to make color changes on parameter and color.

NOTE:

To make the color changes effective, please restart the monitor after changing the colors.

7.9 Transferring a Configuration

When installing several monitors with identical user configuration, it is not necessary to set each device separately. An USB drive may be used to transfer the configuration from monitor to monitor.

To export the current monitor's configuration:

1. Connect USB device to the monitor's USB port.
2. Select **USB Configure** in the **Menu > Maintenance > User Maintain**, enter the required password **ABC**.
3. In the **USB Configure** menu, select **Export** to export configuration. A status message will display after the operation.

To import the configuration on the USB drive to the monitor:

1. Connect the USB drive to the monitor's USB port.
2. Select **USB Configure** in the **Menu > Maintenance > User Maintain**, enter the required password **ABC**.
3. In the **USB Configure** menu, select **Import** to import configuration. A status message will display after the operation.

NOTE:

1. The U disk only supports two kinds of format: FAT and FAT 32.
2. The exported configuration files are saved into the folder named USERCONFIG, and the user can't modify files the in the folder.
3. Up to three configuration files can be saved and recognized.

7.10 Default Configuration

To set default configuration, please select **Menu > Default** and choose a configuration (adult, pediatric or neonate) based on your patient category. This configuration is factory configuration.

Chapter 8 Monitoring ECG

8.1 Overview

The electrocardiogram (ECG) measures the electrical activity of the heart and displays it on the monitor as a waveform and a numeric. This chapter also tells you about arrhythmia monitoring and ST monitoring.

8.2 ECG Safety Information

WARNING

- 1 Do not come into contact with the patient, table, or the monitor during defibrillation.
 - 2 Only use the ECG leads supplied by the manufacturer when using the monitor for ECG monitoring.
 - 3 When connecting the cables and electrodes, make sure no conductive part is in contact with the ground. Verify that all ECG electrodes, including neutral electrodes, are securely attached to the patient but not the conductive part or ground.
 - 4 Check every day whether there is skin irritation resulted from the ECG electrodes. If yes, replace electrodes every 24 hours or change their sites.
 - 5 Place the electrode carefully and ensure a good contact.
 - 6 Check if the lead connection is correct before monitoring. If you unplug the ECG cable from the socket, the screen will display the error message "ECG LEAD OFF" and the audible alarm is activated.
 - 7 When using the monitor with the defibrillator or other high-frequency equipment, please use defibrillator-proof ECG lead to avoid burn.
 - 8 In order to avoid being burnt, please keep the electrodes far away from the radio knife while using electrosurgical equipment.
 - 9 When using Electrosurgery (ES) equipment, do not place an electrode near the grounding plate of the Electrosurgery device: otherwise there will be a great deal of interference with the ECG signal.
 - 10 For patients with pacemakers, the pacing impulse analysis function must be switched ON. Otherwise, the pacing impulse may be counted as normal QRS complex, which results in failure of ECG LOST error detection.
 - 11 The electrodes should be made of the same metal materials.
 - 12 ECG cables can be damaged when connected to a patient during defibrillation. Check cables for functionality before using them again.
-
-

NOTE:

- 1 Interference from a non-grounded instrument near the patient and ESU interference can cause inaccuracy of the waveform.

- 2 IEC/EN60601-1-2 (protection against radiation is 3v/m) specifies that the electrical field density exceeding 1v/m may cause measurement error in various frequencies. It is accordingly suggested that do not use equipment generating electrical radiation near ECG/RESP monitoring devices.
- 3 The simultaneous use of cardiac pacemaker and other patient-connected equipment may cause safety hazard.
- 4 If the pacemaker signals are beyond the claimed range, the heart rate may be calculated incorrectly.
- 5 In the default settings of the monitor, the ECG waveforms are the first two waveforms from top in the waveform area.
- 6 For measurements in or near the heart we recommend connecting the monitor to the potential equalization system.
- 7 For protecting environment, the used electrodes must be recycled or disposed of properly.

8.3 ECG Display

The figure below is for reference only.



The symbol “①” indicates lead name of display waveform: there are other leads for selection, such as I, II, III, aVR, aVF, aVL, V. If you want to change the lead, please refer to section *Selecting Calculation Lead*.

The symbol “②” indicates waveform gain: there are several options, such as X0.125, X0.25, X0.5, X1, X2 and Auto. If you want to change it, please refer to section *Changing the size of the ECG Wave*.

The symbol “③” indicates Filter setting, there are three options: monitoring, surgery, diagnos. If you want to change it, please refer to section *Changing the ECG Filter Setting*.

8.3.1 Changing the Size of the ECG Wave

If any of the displayed ECG waveform is too small or clipped, you can change the size of it on the screen. First select **ECG Wave Setup > ECG Gain**, then select an appropriate factor from the pop-up box to adjust the ECG waveform.

X0.125 to make strength of ECG signal waveform of 1mV become 1.25mm;

X0.25 to make strength of ECG signal waveform of 1mV become 2.5mm;

X0.5 to make strength of ECG signal waveform of 1mV become 5mm;

X1 to make strength of ECG signal waveform of 1mV become 10mm;

X2 to make strength of ECG signal waveform of 1mV become 20mm;

Auto let the monitor choose the optimal adjustment factor for all the ECG waves.

8.3.2 Changing the ECG Filter Settings

The ECG filter setting defines how ECG waves are smoothed. An abbreviation indicating the filter type is shown underneath the lead label on the monitor display. Filter settings do not affect ST measurement.

To change the filter setting, in the **ECG Setup** menu, select **Filter** and then select the appropriate setting.

- **Monitor**: Use this mode under normal measurement conditions.
- **Surgery**: The filter reduces interference to the signal. It should be used if the signal is distorted by high frequency or low frequency interference. High frequency interference usually results in large amplitude spikes making the ECG signal look irregular. Low frequency interference usually leads to a wandering or rough baseline. In the operating room, the Filter reduces artifacts and interference from electro-surgical units. Under normal measurement conditions, selecting **Surgery** may suppress the QRS complexes too much and thus interfere with the clinical evaluation of the ECG displayed on the monitor.
- **Diagnos**: Use when diagnostic quality is required. The unfiltered ECG wave is displayed so that changes such as R-wave notching or discrete elevation or depression of the ST segments are visible.

8.4 Using ECG Alarms

ECG alarms can be switched on and off and the changes to high and low alarm limits are just like other measurement alarms, which are described in the Alarms section. Special alarm features applying only to ECG are described here.

8.5 Selecting Calculation Lead

On the **Normal** interface, the users can select either **3 LEADS** or **5 LEADS** for this item. Normal QRS complex should be:

- The normal QRS should be either completely above or below the baseline and it should not be biphasic. For paced patients, the QRS complexes should be at least twice the height of pace pulses.
- The QRS should be tall and narrow.
- The P-waves and the T-waves should be less than 0.2 mV.

8.6 Monitoring Procedure

8.6.1 Preparation

The skin is a poor conductor of electricity, therefore preparation of the patient's skin is important to facilitate good electrode contact to skin.

- Select sites with intact skin, without impairment of any kind.
- Shave hair from sites, if necessary.
- Wash sites thoroughly with soap and water. (Never use ether or pure alcohol, because this increases skin impedance).
- Rub the skin briskly to increase capillary blood flow in the tissues and remove skin scurf and grease.

8.6.2 Connecting ECG Cables

1. Attach clip or snap to electrodes prior to placement.
2. Put the electrodes on the patient. Before attaching, apply some conductive jelly on the electrodes if the electrodes are not electrolyte self-supplied.
3. Connect the electrode lead to the patient's cable.

CAUTION

To protect the monitor from damage during defibrillation, for accurate ECG information and to protect against noise and other interference, use only ECG electrodes and cables specified by EDAN.

8.7 Selecting Lead Type

To change the lead type, please:

1. Select the ECG parameter area, open the **ECG Setup** menu;
2. Set **Lead Type** to **3 Leads**, **5 Leads** or **12 Leads** based on the lead used.

8.8 Installing Electrodes

NOTE:

The following table gives the corresponding lead names used in Europe and America respectively. (Lead names are represented by R, L, F, N, C, C1-C6 in Europe, whose corresponding lead names in America are RA, LA, LL, RL, V, V1-V6.)

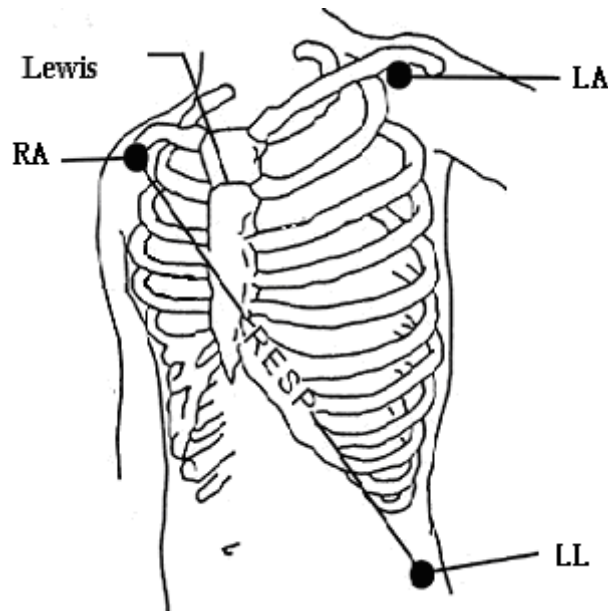
| AHA (American Standard) | | IEC (Europe Standard) | |
|-------------------------|-------|-----------------------|--------|
| Electrode Labels | Color | Electrode Labels | Color |
| RA | White | R | Red |
| LA | Black | L | Yellow |
| LL | Red | F | Green |
| RL | Green | N | Black |
| V | Brown | C | White |

| | | | |
|----|---------------|----|---------------|
| V1 | Brown/ Red | C1 | White/ Red |
| V2 | Brown/ Yellow | C2 | White/ Yellow |
| V3 | Brown/ Green | C3 | White/ Green |
| V4 | Brown/Blue | C4 | White/ Brown |
| V5 | Brown/Orange | C5 | White/ Black |
| V6 | Brown/Purple | C6 | White/ Purple |

8.8.1 Electrode Placement for 3-lead

Take the American standard for example, see the following figure:

- RA placement - directly below the clavicle and near the right shoulder.
- LA placement: directly below the clavicle and near the left shoulder.
- LL placement - on the left hypogastrium.

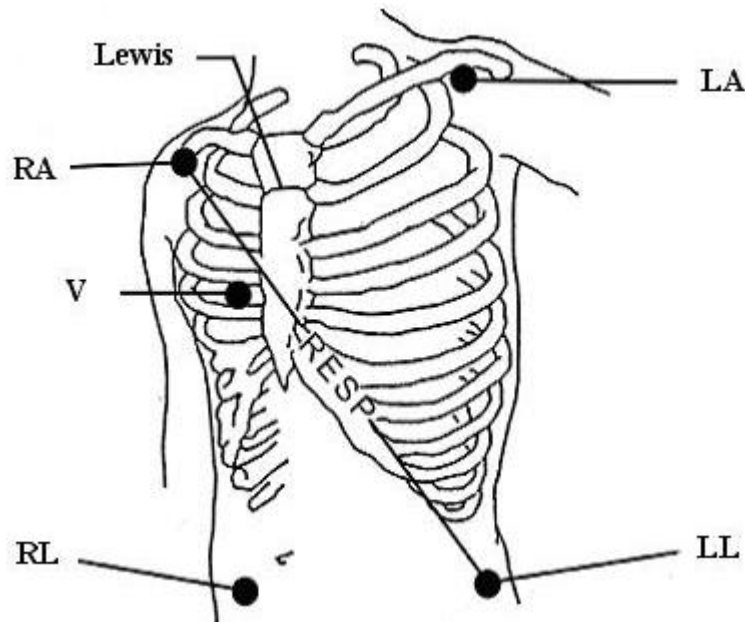


Electrode Placement for 3-lead

8.8.2 Electrode Placement for 5-lead

Take the American standard for example, see the following figure:

- RA placement: directly below the clavicle and near the right shoulder.
- LA placement: directly below the clavicle and near the left shoulder.
- RL placement: on the right hypogastrium.
- LL placement: on the left hypogastrium.
- V placement: on the chest, the position depends on your required lead selection.



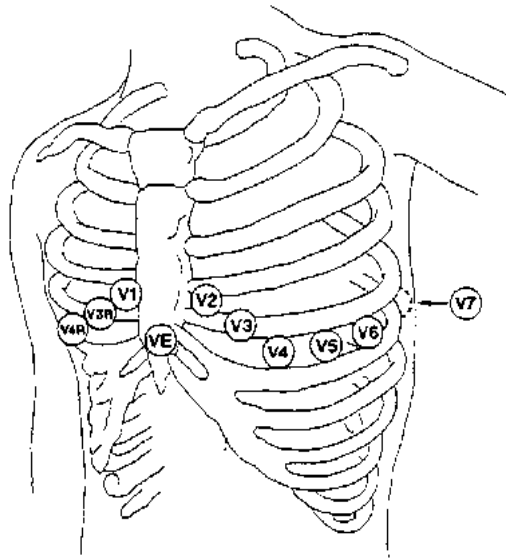
Electrode Placement for 5-lead

NOTE:

To ensure the patient safety, all leads must be attached to the patient.

For 5-lead, attach the V electrode to one of the indicated positions as below:

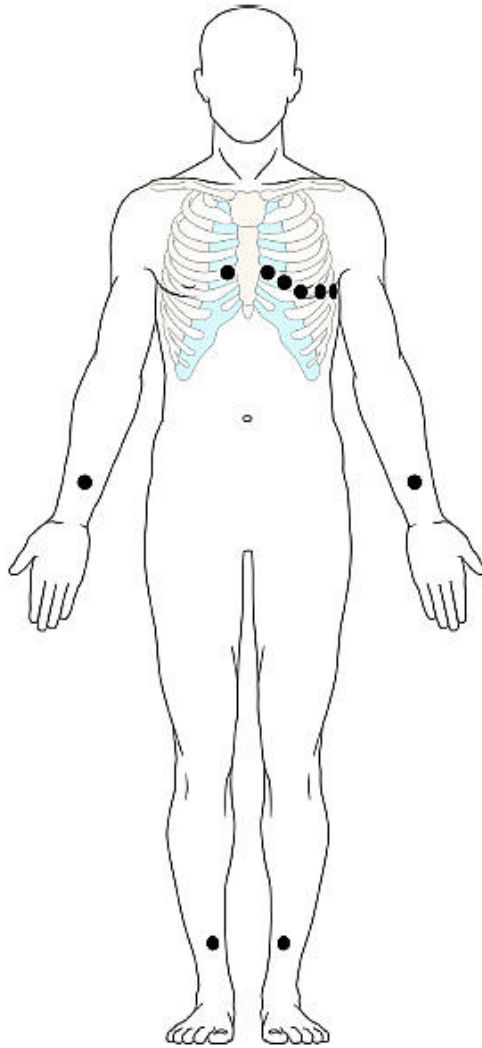
- V1 On the 4th intercostal space at the right sterna margin.
- V2 On the 4th intercostal space at the left sterna margin.
- V3 Midway between V2 and V4 electrodes.
- V4 On the 5th intercostal space at the left clavicular line.
- V5 On the left anterior axillary line, horizontal with V4 electrode.
- V6 On the left middle axillary line, horizontal with V4 electrode.
- V3R-V6R On the right side of the chest in positions corresponding to those on the left.
- VE Over the xiphoid position.
- V7 On the 5th intercostal space at the left posterior axillary line of back.
- V7R On the 5th intercostal space at the right posterior axillary line of back.



V-Electrode Placement for 5-lead

8.8.3 Electrode Placement for 12-lead

Take the American standard for example. The 12-lead electrodes should be placed on extremities and chest. The electrodes for extremities should be placed on the skin of legs or arms, the electrodes placed on chest should follow the doctor's advice. Please see the following figure.



Electrode Placement for 12-lead

8.8.4 Recommended ECG Lead Placement for Surgical Patients

WARNING

When using Electrosurgery (ES) equipment, leads should be placed in a position in equal distance from Electrosurgery electrotome and the ES grounding plate to avoid cautery. Electrosurgery equipment wire and ECG cable must not be tangled up.

Monitoring ECG leads are mainly used for monitoring the patient's vital signs. When using the patient monitor with other electrosurgery equipment, it is advised to use the counteracting defibrillation ECG lead.

The placement of the ECG leads will depend on the type of surgery that is being performed. For example, in an open chest surgery the electrodes may be placed laterally on the chest or on the back. In the operating room, artifacts may affect the ECG waveform due to the use of ES (Electrosurgery) equipment. To help reduce this you can place the electrodes on the right and left shoulders, the right and left sides near the abdomen, and the chest lead on the left side at mid-chest. Avoid placing the electrodes on the upper arms. Otherwise the ECG waveform will be too small.

NOTE:

- 1 If an ECG waveform is not accurate, while the electrodes are tightly attached, try to change the lead.
- 2 Interference from a non-grounded instrument near the patient and ESU interference can cause inaccuracy of the waveform.

8.9 ECG Menu Setup

8.9.1 Setting Alarm Source

To change the alarm source, please select **ECG Setup > Alarm Source**, then a pop-up box is displayed:

HR: the monitor considers the HR as HR/PR alarm source;

PR: the monitor considers the PR as HR/PR alarm source;

AUTO: If the Alarm Source is set to **Auto**, the monitor will use the heart rate from the ECG measurement as the alarm source whenever the ECG measurement is switched on and at least one ECG lead can be measured without a technical condition. The monitor will automatically switch to Pulse as the alarm source if:

- a valid ECG lead can no longer be measured and
- a pulse source is switched on and available.

The monitor then uses the pulse rate from the measurement currently active as system pulse. While Pulse is the alarm source, all arrhythmia and ECG HR alarms are switched off. If an ECG lead becomes available again, the monitor automatically uses HR as alarm source.

8.9.2 Smart Lead Off

In **5 LEADS, 12 LEADS** mode, if **CH1** and **CH2** can not be measured because of the lead off or other reasons, it can shift to other modes to collect an ECG waveform.

To change smart lead off setting, please select **ECG Setup > Smart Leadoff**, then a pop-up menu is displayed.

8.9.3 Setting Beat Volume

Beat volume is from HR or PR, depending on your HR alarm setting. Five selections are available: **0, 1, 2, 3, 4, 5**. **5** indicates the maximum volume. **0** indicates the minimum volume.

To change the beat volume, first select **ECG Setup > Beat Volume**, then select an appropriate volume from the pop-up list.

8.9.4 ECG Display

It varies with **Lead Type**. When **Lead Type** is set to **3 Leads**, **Display** can be set to **Normal**, and it can display one ECG waveform on the main screen.

When **Lead Type** is set to **5 Leads**, **Display** can be set to **Normal, Full-Scr** and **Half-Scr**. Select **Normal** to display two ECG waveforms on the main screen; select **Full-Scr** to display seven ECG waveforms which occupy the area of seven waveforms on the main screen; Select **Half-Scr**

to display seven ECG waveforms on the screen, occupying the area of four waveforms.


NOTE:

If **3 Leads** is selected in the **ECG Setup** menu, only **Normal** can be selected for **Display** in the sub-menu.

8.9.5 Setting Pace Status

It is important to set the paced status correctly when you start monitoring ECG. To change the paced status in the ECG Setup menu, select **Pace** to toggle between **On** or **Off**. When **Pace** is set to **On**:

- Pace Pulse Rejection is switched on. This means that pacemaker pulses are not counted as extra QRS complexes.

- Paced symbol is displayed as  on the main screen.

NOTE:

- 1 When monitoring a patient with a pacemaker, set **Pace** to **On**. If monitoring a patient without a pacemaker, set **Pace** to **Off**.
- 2 If **Pace** is set to **On**, the system will not perform some types of ARR analysis.

WARNING

Some pace pulses can be difficult to reject. When this happens, the pulses are counted as a QRS complex, and could result in an incorrect HR and failure to detect cardiac arrest or some arrhythmias. Keep pacemaker patients under close observation.

8.9.6 ECG Calibration

This item is used to calibrate ECG waveform. When you select this item from ECG Setup menu again, the ECG waveform calibration ends.

NOTE:

The device can't be monitored during ECG calibration.

8.9.7 ECG Waveform Settings

To change this speed, select **ECG Wave Setup > Sweep**, then select an appropriate setting from the pop-up list. The bigger the value is, the wider the waveform is.

8.9.8 12 Leads ECG

When the monitor is installed with 12-lead, it can provide 3-lead, 5-lead and 12-lead ECG monitoring functions.

8.10 ST Segment Monitoring

The monitor performs ST segment analysis on normal and atrially paced beats and calculates ST segment elevations and depressions. This information can be displayed in the form of ST numerics and snippets on the monitor.

ST segment monitoring function is shut off by default. You can switch it to **On** when necessary. When using the ST analysis function, the ST analysis results will be displayed on the right of the main screen, please refer to the following figure.

NOTE:

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.

8.10.1 Setting ST Analysis

To change ST analysis, please select **ECG Setup > ST Analysis**, then select **On** or **Off** from the pop-up list.

8.10.2 ST Display

Your monitor screen may be configured to look slightly different from the illustrations.

| | | | | | | |
|-----------|-----|------|-----|-------|---|------|
| ST | I | 0.08 | aVR | -0.09 | V | 0.04 |
| | II | 0.10 | aVL | 0.03 | | |
| | III | 0.02 | aVF | 0.06 | | |

NOTE:

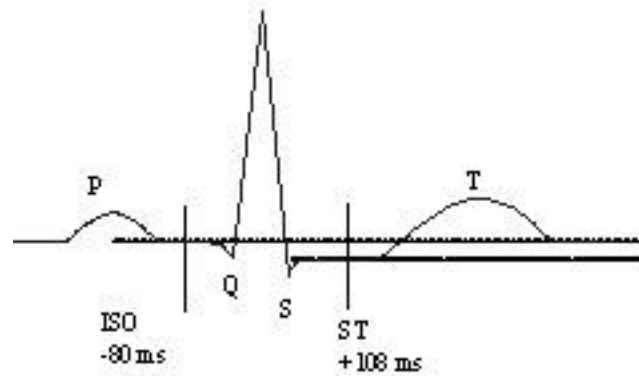
- 1 **ST Analysis** only can be used only in **Adu** mode.
- 2 When setting **ST Analysis** to on, the monitor should be in **Diagnos** mode.
- 3 ECG monitoring should be in **Diagnos** mode.

8.10.3 ST Analysis Alarm Setting

The user can select **ECG Setup > ST Analysis > Alarm Setup** to set the upper alarm limit and lower alarm limit. **ALM HI** can be set to 0.2 mV ~ 2.0 mV, and **ALM LO** can be set to -2.0 mV ~ 0.2 mV. **ALM HI** should be higher than **ALM LO**.

8.10.4 About ST Measurement Points

The ST value for each beat complex is the vertical difference between the ISO point and the ST point, as shown in the diagram below. The isoelectric (ISO) point provides the baseline, and the ST point is at the midpoint of the ST segment. The J point is where the QRS complex changes its slope; as it is a fixed distance away from the ST point, it can be useful to help you position the ST point correctly.



DEF POINT

The ST and ISO measurement points need to be adjusted when you start monitoring, and if the patient's heart rate or ECG morphology changes significantly. Always ensure that ST measurement points are appropriate for your patient. Abnormal QRS complex is not considered in ST segment analysis.

8.10.5 Adjusting ST and ISO Measurement Points

Depending on your monitor's configuration, the ST point can be positioned, too.

These two points can be adjusted by turning the knob. When adjusting ST measurement point, the system will show the ST Measurement Point Window. The system displays the QRS complex template in the window. It is adjustable for the highlight bar in the window. You may select ISO or ST, switch the knob left or right to move the cursor line. When the cursor is at the required position, you may select the base point or the measurement point.

8.11 Arr. Monitoring

8.11.1 Arrhythmia Analysis

The arrhythmia algorithm is used to monitor ECG of neonatal and adult patients in clinics, and detect the changes of heart rate and ventricular rhythm, and also save arrhythmia events and generate alarming information. Arrhythmia algorithm can monitor paced and non-paced patients. Qualified personnel can use arrhythmia analysis to evaluate patient's condition (such as heart rate, PVCs frequency, rhythm and ectopic beat) and decide the treatment. Besides detecting change of ECG, arrhythmia algorithm can also monitor patients and give proper alarm for arrhythmia.

The monitor can support up to 16 different arrhythmia analyses.

| ARR Types | Occurring Condition |
|-----------|--|
| ASYSTOLE | No QRS is detected for 4 seconds |
| VFIB/VTAC | Ventricular tachycardia: The fibrillation wave lasts for 4 consecutive seconds; or the number of continuous Vent beats is larger than the upper limit of cluster Vent beats (≥ 5). The RR interval is less than 600ms. |
| VT>2 | $3 \leq$ the number of cluster PVCs < 5 |


| | |
|--------------|---|
| COUPLET | 2 consecutive PVCs |
| BIGEMINY | Vent Bigeminy |
| TRIGEMINY | Vent Trigeminy |
| R ON T | A type of single PVC under the condition that HR<100, R-R interval is less than 1/3 the average interval, followed by a compensating pause of 1.25X the average R-R interval (the next R wave advances onto the previous T wave). |
| PVC | Single PVCs not belonging to the type of above mentioned PVCs. |
| TACHY | 5 consecutive QRS complex, RR interval is less than 0.5s. |
| BRADY | 5 consecutive QRS complex, RR interval is longer than 1.5s. |
| MISSED BEATS | When HR is less than 100 beats/min., no heart beat is tested during the period 1.75 times of the average RR interval; or When HR is higher than 100beat/min, no beat is tested within 1 second. |
| IRR | The patient has irregular heart rate, check patient's condition, electrodes, cables and leads |
| PNC | After the pacemaker is paced, QRS complex can not be detected during 300ms. |
| PNP | After the QRS complex, no pace is detected during 1.75 times of RR interval. |
| VBRADY | The patient has irregular HR, and his average HR is less than 60bpm. Check his condition, electrodes, cables and leads. |
| VENT | The patient has irregular heart rate, check patient's condition, electrodes, cables and leads. |

8.11.2 ARR Analysis Menu

8.11.2.1 Switching ARR Analysis On and Off

To switch ARR Analysis on or off, in the **ECG Setup** menu, select **ARR Analysis** to toggle between **On** and **Off** from the popup interface.

8.11.2.2 PVCs Alarm

Select **On** in the menu to enable prompt message when an alarm occurs; select **Off** to disable the alarm function, and there will be a symbol  beside **PVCs**.

WARNING

When the PVCs Alarm is set to OFF, the monitor won't give an alarm prompt even if an alarm occurs. In order to avoid endangering the patient's life, the user should use this function cautiously.

8.11.2.3 ARR Relearning

Pick this item to start a learning procedure, and **ECG ARR LEARNING** is displayed on the screen. The ECG ARR LEARNING will start automatically in the following status:

- Changing lead type;
- Connectiong leads;
- Updating the patients;
- Starting ARR learning manually;
- After the ARR analysis is switched on;
- The module is set to on;
- Calibration mode is changed to nomoral measurement mode;
- Exiting the Demo mode;
- Exiting the standby mode;

8.11.2.4 ARR Alarm

The users can switch on or off all arrhythmia alarms by selecting **ECG Setup > ARR Analysis > ARR Alarm**. And some arrhythmia alarms can be individually switched on or off. They are: **ASYSTOLE, VFIB/VTAC, R-ON-T, VT>2, COUPLET, PVC, BIGEMINY, TRIGEMINY, TACHY, BRADY, MISSED BEATS, IRR, PNC, PNP, VBRADY** and **VENT**.

To switch individual alarm on or off, select **ECG Setup > ARR Analysis > ARR Alarm**. The user can set the individual ARR alarm from the popup interface.

8.12 12-Lead ECG Monitoring

8.12.1 Diagnosis Function

If iM80 is equipped with 12 leads monitoring, the device has automatical diagnosis function. To perform this function, please:

- 1 Set **Lead Type** in the **ECG Setup** menu to **12 Leads** and set **Display** in the **ECG Setup** menu to **12 Leads**.
- 2 Select the **ECG Analysis** shortcut key on the screen.
- 3 The **Diagnosis Review** interface is displayed, shown in figure a. And the diagnosis result is displayed on the interface after approximately 10 seconds, shown in figure b.

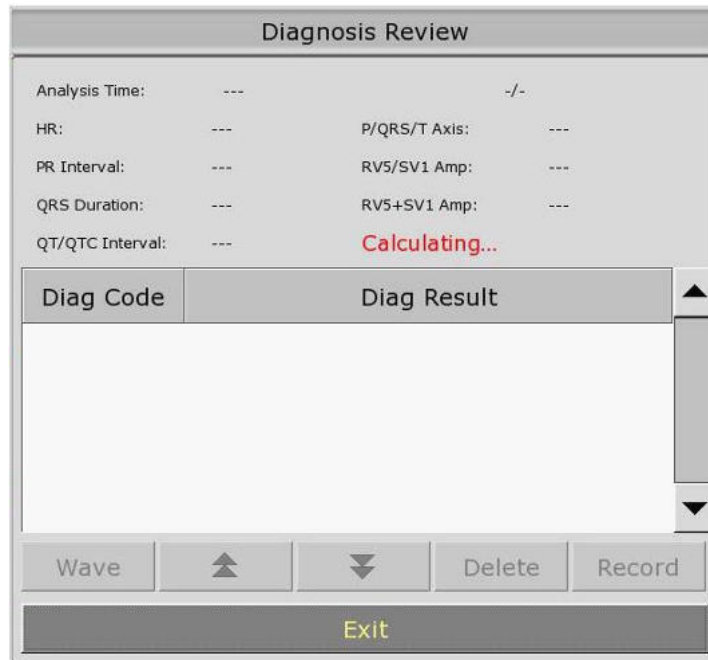


Figure a



Figure b

Figure b shows analysis time, HR (heart rate), P/QRS/T Axis, RR Interval, RVS/SV1 Amp (RVS/SV1 amplitude), QRS Duration, RV5+SV1 Amp (RV5+SV1 amplitude), QT/QTC Interval and Diagnosis code.

More information about diagnosis review, please refer to Section *12-lead Diagnosis Review*.

8.12.2 Measurement and Interpretation

The measurement function provides the automatic measurement of these common parameters, such as heart rate, PR interval, QRS complex duration, QT interval, P/QRS/T axis, RV5/SV1 amplitude etc. The interpretation function provides the automatic diagnosis of hundreds of abnormal cases, such as Arrhythmia, AV block, ventricular conduction block, myocardial infarction, ventricular hypertrophy and atria enlargement, ST-T abnormality and electrical axes deviation.

Chapter 9 Monitoring RESP

9.1 Overview

The monitor measures respiration from the amount of thoracic impedance between two ECG electrodes. The change of impedance between the two electrodes, (due to the thoracic movement), produces a respiratory waveform on the screen.

9.2 RESP Safety Information

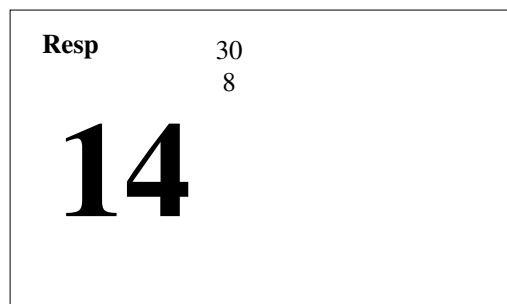
WARNING

- 1 If you do not set the detection level for the respiration correctly in manual detection mode, it may not be possible for the monitor to detect apnea. If you set the detection level too low, the monitor is more likely to detect cardiac activity, and to falsely interpret cardiac activity as respiratory activity in the case of apnea.
 - 2 The respiration measurement does not recognize obstructive and mixed apneas - it only indicates an alarm when a pre-adjusted time has elapsed since the last detected breath.
 - 3 If operating under conditions according to the EMC Standard EN 60601-1-2 (Radiated Immunity 3V/m), field strengths above 1V/m may cause erroneous measurements at various frequencies. Therefore it is recommended to avoid the use of electrically radiating equipment in close proximity to the respiration measurement unit.
 - 4 Cardiogenic artifact in impedance respiration monitoring may make it difficult to detect breaths or may otherwise be counted as breaths. In some instances, the breath rate may also correspond to the heart rate making it difficult to determine if the signal is due to breathing or the cardiac cycle. Do not rely on RESP monitoring as the sole method for detecting cessation of breathing. Follow hospital guidelines and best clinical practices for apnea detection including monitoring additional parameters that indicate the patient's oxygenation status, such as etCO₂ and SpO₂.
-
-

NOTE:

The RESP monitoring is not recommended to be used on patients who are very active, as this can cause false alarms.

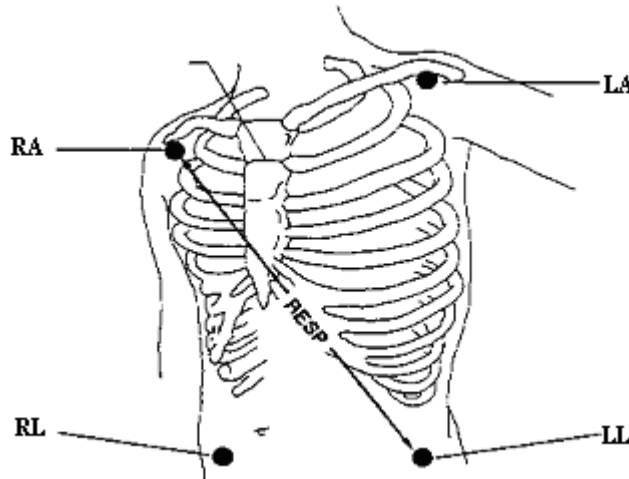
9.3 Resp Display



9.4 Electrode Placement for Monitoring Resp

Correct patient skin preparation techniques for electrode placement are important for Resp measurement: you will find this information in the chapter on ECG.

The Resp signal is always measured between two of the ECG electrodes. There are two standard ECG leads for selection: I lead (RA and LA) and II lead (RA and LL).



Electrodes Placement for 5-lead

9.5 Cardiac Overlay

Cardiac activity that affects the Resp waveform is called cardiac overlay. It happens when the Resp electrodes pick up impedance changes caused by the rhythmic blood flow. Correct electrode placement can help to reduce cardiac overlay: avoid the liver area and the ventricles of the heart in the line between the respiratory electrodes. This is particularly important for neonates.

9.6 Chest Expansion

Some patients, especially neonates, expand their chests laterally. In these cases it is best to place the two respiratory electrodes in the right midaxillary and left lateral chest areas at the patient's maximum point of breathing movement to optimize the respiratory wave.

9.7 Abdominal Breathing

Some patients with restricted chest movement breathe mainly abdominally. In these cases, you may need to place the left leg electrode on the left abdomen at the point of maximum abdominal expansion to optimize the respiratory wave.

NOTE:

Place the red and green electrodes diagonally to optimize the respiration waveform. Avoid the liver area and the ventricles of the heart in the line between the RESP electrodes so as to avoid cardiac overlay or artifacts from pulsating blood flow. This is particularly important for neonates.

9.8 Selecting Resp Lead

To change Resp lead, in the **Resp Setup** menu, select **Resp Lead** to pick up the appropriate lead

from the pop-up list.

9.9 Changing Hold Type

To change the calculation mode, in the **Resp Setup** menu, set **Hold Type** to **Manual** or **Auto**. When it is set to the **AUTO** mode, **Hold High** and **Hold Low** are unavailable, and the monitor can calculate the respiration rate automatically. When it is set to the **Manual** mode, you can adjust the broken lines in RESP area by the **Hold High** and **Hold Low** items.

9.10 Changing the Size of the Respiration Wave

Select the Resp waveform area to open the **Resp Wave Setup** menu:

- Select **AMP**, then choose an appropriate value. The value is bigger, the waveform amplitude is higher.
- Select **Sweep**: select an appropriate setting from the pop-up list.

9.11 Using Resp Alarms

Resp alarms can be switched on and off and the high and low alarm limits can be changed just like other measurement alarms, as described in the Alarms chapter.

9.12 Changing the Apnea Time

The apnea alarm is a high priority red alarm used to detect apneas. The apnea alarm delay time defines the time period between the point where the monitor cannot detect any respiration activity and the indication of the apnea alarm.

1. In the **Resp Setup** menu, select **Apnea Alm**.
2. Select the appropriate setting from the popup list.

Chapter 10 Monitoring SpO₂

10.1 Overview

SpO₂ is based on the absorption of pulse blood oxygen to red and infrared light by means of finger sensor and SpO₂ measuring unit. SpO₂ Plethysmogram measurement is employed to determine the oxygen saturation of hemoglobin in the arterial blood. If, for example, 97% of the hemoglobin molecules in the red blood cells of the arterial blood combine with oxygen, then the blood has a SpO₂ oxygen saturation of 97%. The SpO₂ numeric on the monitor will read 97%. The SpO₂ numeric shows the percentage of hemoglobin molecules which have combined with oxygen molecules to form oxyhemoglobin. The SpO₂/PLETH parameter can also provide a pulse rate signal and a plethysmogram wave.

10.2 SpO₂ Safety Information

WARNING

- 1 If the SpO₂ sensor can not work properly, please reconnect the sensor or change a new one.
 - 2 Do not use the sterile supplied SpO₂ sensors if the packaging or the sensor is damaged and return them to the vendor.
 - 3 Prolonged and continuous monitoring may increase the risk of unexpected change of dermal condition such as abnormal sensitivity, rubescence, vesicle, repressive putrescence, and so on. It is especially important to check the sensor placement of neonate and patient of poor perfusion or immature dermogram by light collimation and proper attaching strictly according to changes of the skin. More frequent examinations may be required for different patients.
 - 4 Tissue damage may be caused by incorrect application or prolonged measurement duration using the sensor (more than 4 hours). Inspect the sensor periodically according to the sensor user manual.
 - 5 Neonate SpO₂ sensor can only be used when required, no more than 20 min at a time.
 - 6 Use only EDAN permitted sensors and extension cables with the monitor. Other sensors or extension cables may cause improper monitor performance and/or minor personal injury.
-
-

NOTE:

- 1 Make sure the nail covers the light window. The wire should be on the backside of the hand.
- 2 SpO₂ waveform is not proportional to the pulse volume.
- 3 Avoid placing the sensor on extremities with an arterial catheter, or intravascular venous infusion line.

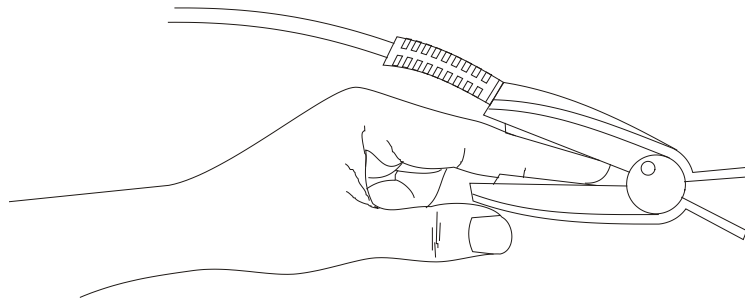
- 4 Don't use the functional tester to access the SpO₂ accuracy.
- 5 The device is calibrated to display functional oxygen saturation.
- 6 The materials with which the patient or any other person can come into contact conform with the standard of ISO10993.

10.3 Measuring SpO₂

1. Select the correct patient category setting (adult/pediatric and neonatal), as this is used to optimize the calculation of the SpO₂ and pulse numerics.
2. During measurement, ensure that the application site:
 - has a pulsatile flow, ideally with a good circulation perfusion.
 - has not changed in its thickness, causing an improper fit of the sensor.

10.4 Measurement Procedure

1. Switch on the monitor.
2. Attach the sensor to the appropriate site of the patient finger.
3. Plug the connector of the sensor extension cable into the SpO₂ socket on the SpO₂ module.



Mounting of the Sensor

WARNING

Inspect the application site every two to three hours to ensure skin quality and correct optical alignment. If the skin quality changes, move the sensor to another site. Change the application site at least every four hours.

NOTE:

Injected dyes such as methylene blue or intravascular dyshemoglobins such as methemoglobin and carboxyhemoglobin may lead to inaccurate measurements.

Interference can be caused by:

- High levels of ambient light or strobe lights or flashing lights (such as fire alarm lamps). (Hint: cover application site with opaque material.)
- High-frequency electrical noise, including electro-surgical apparatus and defibrillators

- Intravascular dye injections
- Significant concentrations of dysfunctional hemoglobin, such as carboxyhemoglobin and methemoglobin
- Excessive patient movement and vibration
- Improper sensor application
- Low perfusion or high signal attenuation
- Venous pulsation
- Placement of the sensor on an extremity that has a blood pressure cuff, arterial catheter, or intravascular line

10.5 Understanding SpO₂ Alarms

This refers to SpO₂ specific alarms. See the Alarms section for general alarm information. SpO₂ offers high and low limit alarms, and the users can adjust them.

10.6 Adjusting Alarm Limits

In the **SpO₂ Setup** menu, select **Alarm Setup**:

- Set the SpO₂ High Alarm Limit to an appropriate value from the popup interface.
- Set the SpO₂ Low Alarm Limit to an appropriate value from the popup interface.

WARNING

High oxygen levels may predispose a premature infant to retrolental fibroplasia. If this is a consideration do NOT set the high alarm limit to 100%, which is equivalent to switching the alarm off.

10.7 Setting SpO₂ as Pulse Source

1. In the **PR Setup** menu, select **PR Source**;
2. Select **SpO₂** from the pop-up list.

10.8 Setting Pitch Tone

If tone modulation is on, the PR sound lowers when the SpO₂ level drops. In the **SpO₂ Setup** menu, select pitch tone to toggle between **On** and **Off**.

10.9 Setting Sensitivity

The different sensitivity indicates different refresh frequency. **High** indicates the refresh frequency of SpO₂ value is the most frequent. To change the sensitivity, please follow the steps:

- 1 Select the **SpO₂ Setup** menu;
- 2 Select **Sensitivity** on the interface and select the desired sensitivity from the popup list.

Chapter 11 Monitoring PR

11.1 Overview

The pulse numeric counts the arterial pulsations that result from the mechanical activity of the heart in beats per minute (bpm). You can display a pulse from any measured SpO₂ signal or any arterial pressure.

11.2 Setting PR Source

The monitor provides PR sources options, but currently only SpO₂ is supported. If the parameter as PR source is switched off, the monitor will switch based on priority. If all parameters producing SpO₂ are switched off, the PR parameter will be switched off.

11.3 Setting PR Volume

Six selections are available: **0**, **1**, **2**, **3**, **4**, and **5**. **5** indicates the maximum volume. **0** indicates no sound. You can change **PR Volume** in the **PR Setup** menu.

11.4 Using Pulse Alarms

You can change pulse rate alarm limits in the **PR Setup** menu by selecting **Alarm Setup**. Pulse alarms are only generated when the active alarm source is set to **PR**, a pulse source is set as system pulse and pulse alarms are switched on.

11.5 Selecting the Active Alarm Source

In most cases, the HR and Pulse numerics are identical. In order to avoid simultaneous alarms on HR and Pulse, the monitor uses either ECG or Pulse as its active alarm source. To change the alarm source, select Alarm Source in the ECG/Pulse Alarms menu, then select

- **HR**: if you want HR to be the alarm source for HR/Pulse.
- **PR**: if you select Pulse as the active alarm source, the monitor will prompt you to confirm your choice. Be aware that if you select Pulse as the alarm source, all arrhythmia and ECG HR alarms are switched off.
- **AUTO**: if the Alarm Source is set to Auto, the monitor will use the heart rate from the ECG measurement as the alarm source whenever the ECG measurement is switched on and at least one ECG lead can be measured without an technical alarm condition. The monitor will automatically switch to Pulse as the alarm source.

Chapter 12 Monitoring NIBP

12.1 Overview

This monitor uses the oscillometric method for measuring NIBP. It can be used for adult, pediatric and neonatal patients.

Oscillometric devices measure the amplitude of pressure changes in the occluding cuff as the cuff deflates from above systolic pressure. The amplitude suddenly increases as the pulse breaks through the occlusion in the artery. As the cuff pressure decreases further, the pulsations increase in amplitude, reach a maximum (which approximates to the mean pressure), and then diminish.

In adult and pediatric mode, the blood pressure measurements determined with this device comply with the American National Standard for Electronic or Automated Sphygmomanometers (ANSI/AAMI SP10-1992) in relation to mean error and standard deviation, when compared to auscultatory measurements in a representative patient population. For the auscultatory reference, the fifth Korotkoff sound was used to determine the diastolic pressure.

In neonatal mode, the blood pressure measurements determined with this device comply with the American National Standard for Electronic or Automated Sphygmomanometers (ANSI/AAMI SP10-1992) in relation to mean error and standard deviation, when compared to intra-arterial measurements in a representative patient population.

12.2 NIBP Safety Information

WARNING

- 1 Do not measure NIBP on patients with sickle-cell disease or any condition where skin damage has occurred or is expected.
 - 2 Use clinical judgement to decide whether to perform frequent unattended blood pressure measurements on patients with severe blood clotting disorders because of the risk of hematoma in the limb fitted with the cuff.
 - 3 Ensure that the correct setting is selected when performing measurements. It may be dangerous for the children to use an over pressure level.
 - 4 The equipment is suitable for use in the presence of electrosurgery.
 - 5 The equipment can provide protective means to prevent the patient from being burned when used with HF SURGICAL EQUIPMENT. The equipment can protect against the effects of the discharge of a defibrillator.
 - 6 Before starting a measurement, verify that you have selected a setting appropriate for your patient (adult, pediatric or neonatal.)
 - 7 Do not apply the cuff to a limb that has an intravenous infusion or catheter in place. This could cause tissue damage around the catheter when infusion is slowed or blocked during cuff inflation.
-
-

WARNING

- 8 Make sure that the air conduit connecting the blood pressure cuff and the monitor is neither blocked nor tangled.
 - 9 Do not attach the cuff to a limb being used for IV infusions as the cuff inflation can block the infusion, potentially causing harm to patient.
-
-

NOTE:

- 1 It is suggested that the user should not start NIBP measuring when the low battery displays, or the monitor may be turned off automatically.
- 2 If an alarm occurs or measurement fails, please discontinue the measurement.
- 3 If you spill liquid onto the equipment or accessories, particularly if there is a chance that it can get inside the tubing or the measurement device, contact your service personnel.
- 4 Continuous use of the automatic measuring mode for short intervals may lead to the discomfort of patient.

12.3 Introducing the Oscillometric NIBP Measurement

Oscillometric devices measure the amplitude of pressure changes in the occluding cuff as the cuff deflates from above systolic pressure. The amplitude suddenly increases as the pulse breaks through the occlusion in the artery. As the cuff pressure decreases further, the pulsations increase in amplitude, reach a maximum (which approximates to the mean pressure), and then diminish.

12.4 Measurement Limitations

Measurements are impossible with heart rate extremes of less than 40 bpm or greater than 240 bpm, or if the patient is on a heart-lung machine.

The measurement may be inaccurate or impossible in the following situations:

- A regular arterial pressure pulse is hard to detect.
- Patients with cardiac arrhythmias.
- Patients with excessive and continuous movement such as shivering or convulsions.
- Patients with rapid blood pressure changes.
- Patients with severe shock or hypothermia that reduces blood flow to the peripheries.
- Patients with obesity, where a thick layer of fat surrounding a limb dampens the oscillations coming from the artery.
- Patients on an edematous extremity.

12.5 Measurement Methods

There are three methods of measuring NIBP:

- Manual - measurement on demand.
- Auto - continually repeated measurements (between 1 and 480 minute adjustable interval).
- Sequence - the measurement will run consecutively in five minutes, then the monitor enters manual mode.

WARNING

Prolonged non-invasive blood pressure measurements in Auto mode may be associated with purport, ischemia and neuropathy in the limb wearing the cuff. When monitoring a patient, examine the extremities of the limb frequently for normal color, warmth and sensitivity. If any abnormality is observed, stop the blood pressure measurements.

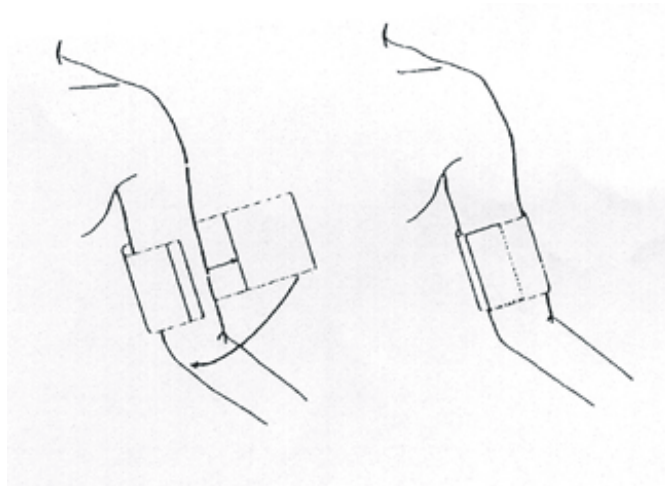
12.6 Measurement Procedures

1. Connect the air hose and switch on the monitor.
2. Apply the blood pressure cuff to the patient's arm or leg and follow the instructions below. Ensure that the cuff is completely deflated.


Apply the appropriate size cuff to the patient (About the cuff size selection, please refer to Section *NIBP accessories*), and make sure that the symbol "Φ" is over the artery. Ensure that the cuff is not wrapped too tightly around the limb. Excessive tightness may cause discoloration and eventual ischemia of the extremity.

NOTE:

The width of the cuff should be either 40% of the limb circumference (50% for neonates) or 2/3 of the upper arm length. The inflatable part of the cuff should be long enough to encircle 50-80% of the limb. The wrong size of cuff can cause erroneous readings. If the cuff size is in question, use a larger cuff.






Cuff Usage

3. Connect the cuff to the air tubing.
4. Check whether the patient mode is appropriately selected. Access the **Patient Setup** menu from **Menu**. Turn the knob to select the required patient **Type** in the **Patient Info.** menu.
5. Select a measurement mode in the **NIBP Setup** menu. Refer to section **Operation Prompts** for details.
6. Press the  button on the front panel to start a measurement.


12.7 Operation Prompts

1. Manual Measuring

Access the **NIBP Setup** menu and set the **Measure Mode** item to **Manual**. Then press the  button on the front panel to start a manual measurement.

During the idle period of measurement process, press the  button on the front panel at any time to start a manual measurement. Then press the  button on the front panel to stop manual measurement and the system continues to execute auto measurement program according to the selected time interval.

2. Automatic Measurement

Access the **NIBP Setup** menu and set the **Measure Mode** item to **Auto**, then press the  button on the front panel to start the automatic measurement according to the selected time interval.

3. Continuous measurement

Access the **NIBP Setup** menu and pick the **Continual** item to start a continuous measurement. The continuous measurement will last 5 minutes.

4. Stopping continuous measurement

During continuous measurement, press the  button on the front panel at any time to stop

continuous measurement.

12.8 Correcting the Measurement if Limb is not at Heart Level

To correct the measurement if the limb is not at heart level to the displayed value:

| | |
|--|--|
| Add 0.75mmHg (0.10kPa) for each centimeter higher or | Deduct 0.75mmHg (0.10kPa) for each centimeter lower or |
| Add 1.9mmHg (0.25kPa) for each inch higher | Deduct 1.9mmHg (0.25kPa) for each inch lower |

12.9 NIBP Alarm

When **NIBP Alarm** is set to **On**, the physiology alarm occurs if any measurement value of Systolic pressure, Mean pressure, Diastolic pressure exceeds alarm limit. The users can adjust the alarm limit by accessing **NIBP Setup > Alarm Setup > Sys Alarm/Map Alarm/Dia Alarm**.

12.10 Resetting NIBP

When the pressure does not work properly and the system fails to give a message for the problem, pick **Reset** in the **User Maintain > NIBP Maintain** menu to activate self-test procedure, and thus restore the system from abnormal performance.

12.11 Calibrating NIBP

NIBP is not user-calibrated. Cuff-pressure transducers must be verified and calibrated on a yearly interval by a qualified service professional. See the Service Manual for details.

12.12 Leak Test

This item is used for leak test. Turn the knob to pick the **Leak Test** item in the **User Maintain > NIBP Maintain** menu to start the air leakage test. When the item is selected, it will change into **Stop**. If this item is selected again, the system will stop air leakage test. And the item returns to **Leak Test**.

WARNING

This pneumatic test other than being specified in the EN 1060-1 standard is to be used by the user to simply determine whether there are air leaks in the NIBP airway. If at the end of the test the system gives the prompt that the NIBP airway has air leaks, please contact the manufacturer for repair.

12.12.1 Procedure of Leak Test

- Connect the cuff securely with the socket for NIBP air hole.
- Wrap the cuff around the cylinder of an appropriate size.
- Access **User Maintain > NIBP Maintain**.
- Turn the knob to the **Leak Test** item and press the item. Then the prompt of **Leak Testing** will appear indicating that the system has started performing leak test.

- The system will automatically inflate the pneumatic system to about 180 mmHg.
- After 20 seconds, the system will automatically open the deflating valve, which marks the completion of a pneumatic measurement.
- If the prompt of **Leak Test OK** appears, it indicates that the airway is in good situation and no air leaks exist. However if the alarm information of **NIBP Cuff Leak** appears, it indicates that the airway may have air leaks. In this case, the user should check for loose connection. After confirming secure connections, the user should re-perform the pneumatic test. If the failure prompt still appears, please contact the manufacturer for repair.

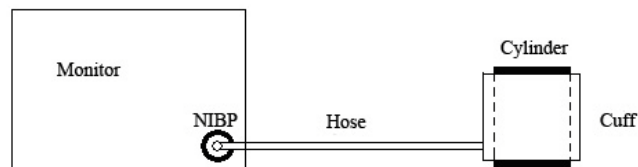


Diagram of NIBP Air Leakage Test

Chapter 13 Monitoring TEMP

13.1 Overview

Body temperature is measured by means of a thermistor probe (a semiconductor whose resistance changes with temperature) that is inserted in the rectum.

Two TEMP probes can be used simultaneously to measure two TEMP values, and get the temperature difference. The standard configuration is axilla sensor for adult.

13.2 TEMP Safety Information

WARNING

- 1 Verify probe cables fault detection before the beginning of monitoring phase. Unplug the temperature probe cable of the channel from the socket, and then the screen will display the error message **TEMP T1 Sensor Off** and the audible alarm is activated. It is the same to the other channel.
 - 2 Take the TEMP probe and cable carefully. When they are not in use, you should coil up the probe and cable into a loose circle. If the wire inside the cable is tensely pulled, it may cause mechanical damage to the probe and the cable.
-

NOTE:

Disposable TEMP probe can only be used once for one patient.

13.3 TEMP Monitoring Setup

- If you are using disposable TEMP probes you need to plug the TEMP cable into the monitor and then connect the probe to the cable. With a reusable TEMP probe you can plug the probe directly into the monitor.
- Apply the TEMP probes securely to the patient.
- Switch on the monitor

It takes 2 min ~ 3 min for the body temperature to stabilize.

13.4 Calculating Temp Difference

The monitor can calculate and display the difference between two temperature values by subtracting the second value from the first. The difference is labeled TD.

Chapter 14 Monitoring Quick TEMP

14.1 Overview

Quick temperature measurement is to establish thermal balance between probe and human body. When the probe is placed on the measurement site until a steady reading is available - after approximately three minutes for oral and rectal measurements and five minutes for axillary measurements. And the measurement temperature is the monitoring temperature. The temperature curve in this process has a certain discipline. The approximately actual temperature curve can be simulated by temperature data sampled earlier. The temperature at thermal balance is calculated through the specific algorithm based on the curve.

The monitor can only measure temperature of adult and pediatric patients. If the user measures temperature of neonatal patient, the monitor will not display data. The Oral/Axillary sensor and Rectal sensor are of standard configuration.

14.2 Quick TEMP Safety Information

WARNING

- 1 To ensure optimal accuracy, always confirm that the correct mode and alarm limit are selected. Changing the measure position may lead to the change of alarm limit.
 - 2 Verify probe cables fault detection before the beginning of monitoring phase. Unplug the temperature probe cable from the socket, and then the screen will display the error message TEMP SENSOR OFF and the audible alarm is activated.
 - 3 Take the TEMP probe and cable carefully. When they are not in use, you should coil up the probe and cable into a loose circle. If the wire inside the cable is tensely pulled, it may cause mechanical damage to the probe and the cable.
 - 4 The calibration of the temperature module is necessary every two years (or as frequently as dictated by your Hospital Procedures Policy). When you need to calibrate the temperature measurement, please contact the manufacturer.
 - 5 Patient actions may interfere with accurate oral temperature readings. Ingesting hot or cold liquids, eating food, chewing gum or mints, brushing teeth, smoking or performing strenuous activity may affect temperature readings for up to 20min after activity has ended.
 - 6 Do not take an axillary temperature through patient's clothing. Direct probe cover to skin contact is required.
 - 7 Biting the sensor tip while taking a temperature may result in damage to the sensor.
-
-

WARNING

- 8 Make sure disposable TEMP sensor covers are used to limit patient cross-contamination. The use of any other probe cover may produce temperature measurement errors or result in inaccurate readings.
- 9 Quick Temp measurement isn't suitable for use during defibrillation.

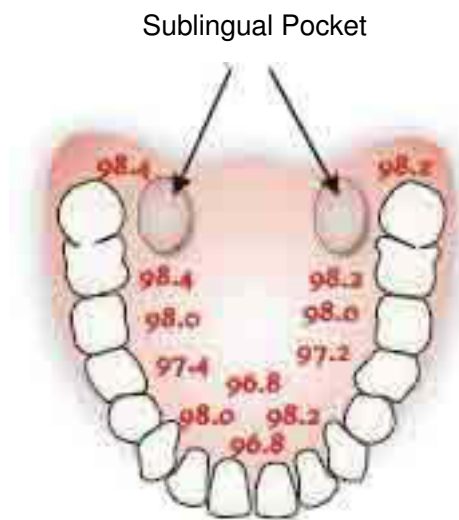
14.3 Measuring Procedure

14.3.1 Measurement for Oral Temperature

1. Ensure the oral probe (white probe) and probe well are installed.
2. Remove the probe from the probe well.
3. Observe the oral mode indicator on the screen (flashing head icon).

If this icon is not flashing, press the **Measure Pos** button and set it to **Oral** until the head icon appears.

4. Load the probe cover.
5. Place the probe tip deep into the patient's sublingual pocket as shown in the following figure.



Measuring Position in Mouth

6. Do not hand the probe to the patient to place in his or her own mouth.
7. Always hold the probe in place, maintaining tissue contact until temperature is complete.

If necessary, repeat the measurement procedure shown above.

NOTE:

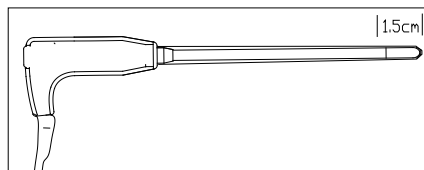
- 1 After one measurement, the user should put the sensor well to the sensor bracket

and then take it out for starting a new measurement.

- 2 To ensure optimal accuracy, always confirm that the correct measurement position is selected.

14.3.2 Measurements for Rectal Temperatures

1. Ensure that the rectal probe (red probe) and probe well are installed.
2. Remove the red probe from the probe well.
3. Observe the Rectal Mode indicator on the display (flashing lower body icon).
4. Load a probe cover. Apply lubricant if desired.
5. Separate the buttocks and gently insert the probe only 1.5cm (5/8 inch), less for infants and children.



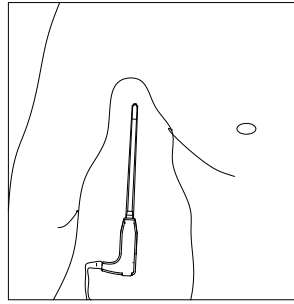
Measuring Position in Rectal

WARNING

Incorrect insertion can cause bowel perforation.

14.3.3 Measurements for Axillary Temperatures

1. Ensure that the white probe and probe well are installed.
2. Remove the probe from the probe well.
3. Press **Quick TEMP Setup > Measure Pos** and select the Axillary mode.
4. Observe the axillary mode indicator on the display (flashing axillary icon).
5. Load a probe cover.
6. Adjust clothing to visualize the axilla.
7. Avoid folds in the axilla and place the probe tip vertically as high as you can as shown.
8. Place the arm at the patient's side. Hold in this position without movement of the arm or probe during the measurement cycle.



Measuring Position in Axillary

NOTE:

Do not take an axillary temperature through the patient's clothing. Direct contact between the patient's skin and the probe is required.

To obtain accurate rectal temperature, use the white temperature probe.

14.4 Changing Temp Unit

To change **Temp Unit**, please:

- 1 Select the **Quick Temp Setup** menu to open it and select **Unit** on the interface.
- 2 Select the appropriate unit from the popup list.

Chapter 15 Monitoring IBP

15.1 Overview

IBP is measured by means of a catheter inserted directly into the circulatory system. A pressure transducer connected to the catheter converts the mechanical force exerted by the blood into an electrical signal, which is displayed graphically as pressure versus time on a monitor screen or numerically on digital display.

The monitor measures direct blood pressure of one selected blood vessel through two channels or four channels, and displays waveforms and pressure of measured direct blood pressure (SYS, DIA and MAP).

15.2 IBP Safety Information

WARNING

- 1 The operator should avoid contact with the conductive parts of the appurtenance when it is connected or applied.
 - 2 When the monitor is used with HF surgical equipment, the transducer and the cables must be avoided from conductive connection to the HF equipment. This is to protect against burns to the patient.
 - 3 Disposable IBP transducer or domes should not be reused.
 - 4 If any kind of liquid, other than solution to be infused in pressure line or transducer, is splashed on the equipment or its accessories, or enters the transducer or the monitor, contact the Hospital Service Center immediately.
-
-

NOTE:

- 1 Use only the pressure transducer listed in the IBP Accessories
- 2 Calibrate the instrument either whenever a new transducer is used, or as frequently as dictated by your Hospital Procedures Policy.

15.3 Monitoring Procedures

Preparatory steps for IBP measurement:

1. Plug the pressure cable into the corresponding socket and switch on the monitor.
2. Flushing through the system with normal saline solution. Ensure that the system is free of air bubbles.
3. Connect the patient catheter to the pressure line, making sure that there is no air present in the catheter or pressure line.
4. Position the transducer so that it is at the same level with the patient's heart, approximately mid-axillary line.
5. For the label name selection, please refer to Selecting a Pressure for Monitoring.

6. To zero the transducer, please refer to *Zeroing the Pressure Transducer*.

WARNING

If there are air bubbles in the pressure line or the transducer, you should flush the system with the solution to be infused.

15.4 Selecting a Pressure for Monitoring

Tell the monitor which pressure you want to monitor by selecting its pressure label. The label is a unique identifier for each type of pressure. When you choose a label, the monitor uses that label's stored settings, for example color, wave scale and alarm settings. The label also determines which algorithm is used to process the pressure signal, so an incorrect label can lead to incorrect pressure values. To select the label, please refer to the following table:

| Label | Description |
|-------|--|
| ART | Arterial blood pressure |
| PA | Pulmonary artery pressure |
| CVP | Central venous pressure |
| ICP | Intracranial pressure |
| LAP | Left atrial pressure |
| RAP | Right atrial pressure |
| P1-P2 | Alternative non-specific pressure labels |

15.5 Zeroing the Pressure Transducer

To avoid inaccurate pressure readings, the monitor requires a valid zero. Zero the transducer in accordance with your hospital policy (at least once per day). You must perform a zero:

- When you use a new transducer or tubing
- Every time you reconnect the transducer cable to the monitor;
- If you think the monitor's pressure readings are not correct.

When using a pressure module, the zero information is stored in the module.

15.6 Zeroing a Pressure Measurement

The zeroing procedure is listed as below:

1. Turn off the stopcock to the patient.
2. Vent the transducer to atmospheric pressure, to compensate for the static and atmospheric pressure exerted on the transducer.
3. In the setup menu for the pressure, select **Zero**.

- When you see the message **Zero Ok**, please close the stopcock to atmospheric pressure, and open the stopcock to the patient.

15.7 Troubleshooting the Pressure Zeroing (Taking Art for Example)

The status message lists the probable cause of an unsuccessful calibration.

| Message | Corrective Action |
|------------------------------|---|
| Art ZERO FAIL | Make sure that the transducer is not attached to the patient |
| Art SENSOR OFF, FAIL | Make sure that transducer is not off, and then proceed zeroing |
| IN DEMO, FAIL | Make sure that the monitor is not in DEMO mode. Contact service technician if necessary |
| PRESSURE OVER RANGE, FAIL | Make sure that the stopcock is vented to atmosphere. If the problem persists, please contact service technician |
| PULSATILE PRESSURE ZERO FAIL | Make sure that the transducer is vented to air, not connected to a patient, and try again. |

15.8 IBP Pressure Calibration

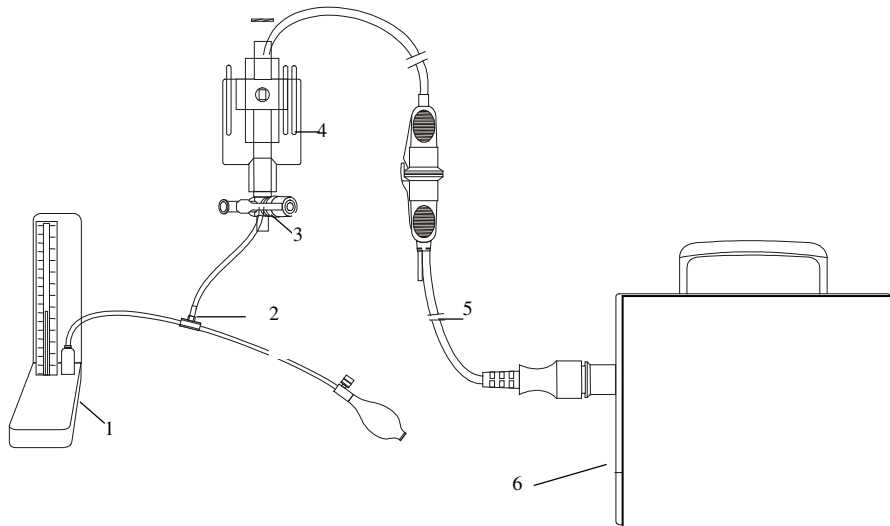
- Mercury calibration should be performed by the biomedical engineering department either whenever a new transducer is used, or as frequently as dictated by your Hospital Procedures Policy.
- The purpose of the calibration is to ensure that the system gives you accurate measurements.
- Before starting a mercury calibration, a zero procedure must be performed.

If you need to perform this procedure yourself, you will need the following equipment: Standard sphygmomanometer, 3-way stopcock and Tubing (approximately 25 cm long).

The calibration procedure is listed below:

- Close the stopcock that was open to atmospheric pressure for the zero calibration.
- Attach the tubing to the sphygmomanometer.
- Ensure that connection to patient is off.
- Connect the 3-way connector to the 3-way stopcock that is not connected to the patient catheter.
- Open the port of the 3-way stopcock to the sphygmomanometer.
- Select the channel to be calibrated in the menu and select the pressure value to which the IBP is to be adjusted.
- Inflate to make the mercury bar rise to the setup pressure value.
- Adjust repeatedly until the value in the menu is equal to the pressure value shown by the mercury calibration.

9. Press the Start button, the device will begin calibrating.
10. Wait for the calibrated result. You should take corresponding measures based on the prompt information.
11. After calibration, disassemble the blood pressure tubing and the attached 3-way valve.



1: Hydrargyrum pressure meter; 2: 3-way connector; 3: 3-way stopcock; 4: Pressure transducer; 5: Pressure transducer interface cable; 6: Monitor

IBP Calibration

15.9 Troubleshooting the Pressure Calibration

The status line lists the probable reasons of an unsuccessful calibration.

| Message | Corrective Action |
|-------------------------------------|--|
| Sensor Off, Fail! | Make sure that sensor is not off, then start the calibration. Contact service technician if necessary. |
| Unable to calibrate in Demo Mode. | Make sure that the monitor is not in DEMO mode. Contact service technician if necessary. |
| Pressure out of normal range, fail. | Make sure that you have selected transducer value in IBP CAL, then start the calibration. Contact service technician if necessary. |
| Pulsate Pressure Calibration Fail | Make sure that pressure value shown by hydrargyrum pressure meter is changeless. Contact service technician if necessary. |

15.10 IBP Alarm

When **Alarm Switch** is set to On, the physiology alarm occurs if any measurement value of Systolic pressure, Mean pressure, or Diastolic pressure exceeds alarm limit. Users can adjust the alarm limit by accessing **XX Options (XX stands for the label name) > Setup > SYS Alarm / MAP Alarm / DIA Alarm**.

Chapter 16 Monitoring CO₂

16.1 Overview

The monitor provides the SideStream and MainStream methods for CO₂ monitoring. LoFlo CO₂ module is used for SideStream measuring, and Capnostat 5 CO₂ module (C5) is used for mainStream measuring.

The principle of CO₂ measurement is primarily based on the fact that CO₂ molecule can absorb 4.3µm infrared ray. Absorption intensity is proportional to CO₂ concentration of patient sample, the CO₂ concentration will compute according to the detecting CO₂ absorption intensity of patient sample.

- SideStream measurement takes a sample of the respiratory gas with a constant sample flow from the patient's airway and analyzes it with a remote CO₂ sensor. You can measure SideStream CO₂ using the monitor's built-in CO₂ measurement.
- MainStream measurement uses a CO₂ sensor attached to an airway adapter directly inserted into the patient's breathing system.

16.2 CO₂ Safety Information

WARNING

- 1 Do not use the device in the environment with flammable anesthetic gas.
 - 2 The device should be used by trained and qualified medical personnel authorized by EDAN.
 - 3 Nitrous oxide, elevated levels of oxygen, helium, xenon, halogenated hydrocarbons, and barometric pressure can influence the CO₂ measurement.
 - 4 The monitor will be damaged if any pipeline from the CO₂ module is disconnected, or the air tube /the air inlet /the air outlet is plugged by water or other materials.
 - 5 The accuracy of the CO₂ measurement will be affected by the following reasons: the airway was highly obstructed; the leakage of air way connection or quick variation of environment temperature.
 - 6 Follow precautions for electrostatic discharge (ESD) and electromagnetic interference (EMI) to and from other equipment.
 - 7 In the presence of electromagnetic devices (i.e., electrocautery), patient monitoring may be interrupted due to electromagnetic interference. Electromagnetic fields up to 20V/m will not adversely affect module performance.
 - 8 Do not place the sensor cables or tubing in any manner that may cause entanglement or strangulation.
 - 9 Do not store the CO₂ Module at temperatures less than -40° F (-40° C) or greater than 158° F (70° C). Do not operate the CO₂ Module at temperatures less than 32° F (0° C) or greater than 104° F (40° C).
-
-

NOTE:

After the low battery alarm appears, please do not start the CO₂ measurement, or the monitor may turn off for the low battery.

16.3 Monitoring Procedures

16.3.1 Zeroing the sensor

You must perform zeroing following the steps when using the new airway adapter.

1. Expose the sensor to room air and keep it away from all sources of CO₂ including the ventilator, the patient's breath and the operator's.
2. In the **CO₂ Setup** menu, please set the **Work Mode** to **Measure**.
3. In the **CO₂ Setup** menu, select **Zero Calibration**.
4. If the system briefly displays **Zero In Progress**, the process is successful. After the zeroing calibration is finished, you can start CO₂ Monitoring. If the system displays **Breath detected** or **Zero required**, zeroing has failed. Zero calibration must be performed again.

16.3.2 LoFlo CO₂ Module

NOTE:

You must perform a zero calibration as described in this procedure each time the ambient temperature changes more than 10°C (for example during transport).



LoFlo CO₂ module

16.3.2.1 Measurement Steps

- 1 Plug the sensor cable into the monitor's CO₂ input connector. Allow the sensor two minutes

for warm-up.

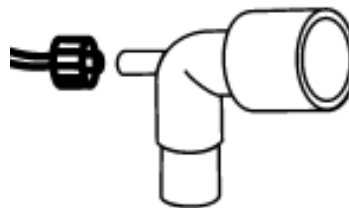
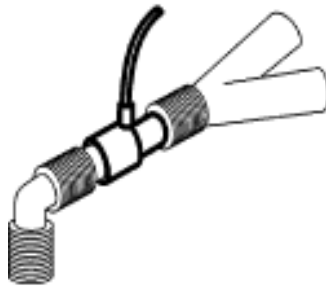
2 Connect the cannula, airway adapter, or sample line as appropriate, to the sensor. It will click into place when seated correctly.



Connecting LoFlo module

3 To zero the sensor, please refer to zeroing the sensor.

4 For intubated patients, an airway adapter is required;



Air adapter

For non-intubated patients: Place the nasal cannula onto the patient.



Place the nasal cannula

NOTE:

- 1 Always connect the airway adapter to the sensor before inserting the airway adapter into the breathing circuit. In reverse, always remove the airway adapter from the breathing circuit before removing the sensor.
- 2 Always disconnect the cannula, airway adapter or sample line from the sensor when the sensor is not in use.

16.3.2.2 Removing Exhaust Gases from the System

WARNING

Anesthetics: When using the sidestream CO₂ measurement on patients who are receiving or have recently received anesthetics, connect the outlet to a scavenging system, to avoid exposing medical staff to anesthetics.

Use an exhaust tube to remove the sample gas to a scavenging system. Attach it to the sidestream sensor at the outlet connector.

16.3.3 C5 CO₂ Module

NOTE:

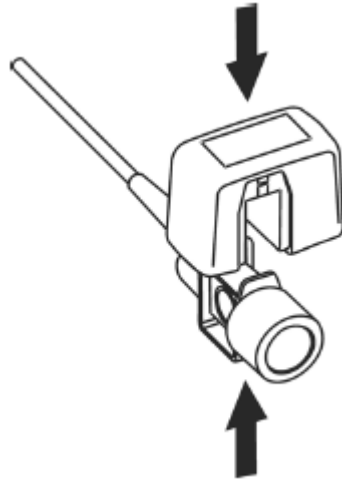
You must perform a zero calibration as described in this procedure each time you use a new airway adapter.



C5 CO₂ module

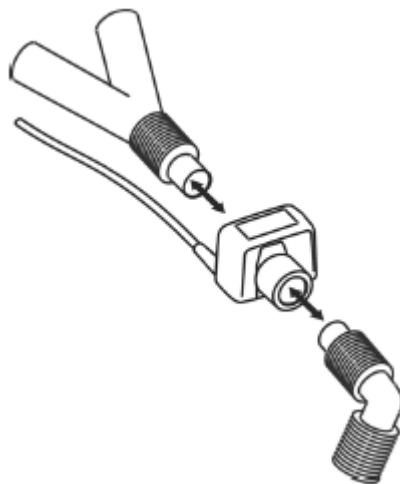
16.3.3.1 Measurement Steps

- 1 Attach the sensor connector to the CO₂ connector on the monitor.
- 2 Wait two minutes, allowing the sensor to reach its operating temperature and a stable thermal condition.
- 3 Choose the appropriate airway adapter and connect it to the sensor head. The airway adapter clicks into place when seated correctly.



Connecting Sensor

- 4 To zero the sensor, please refer to zeroing the sensor.
- 5 Install the airway adapter at the proximal end of the circuit between the elbow and the ventilator Y-section.



Connecting Airway Adapter

WARNING

- 1 No routine user calibration required.
 - 2 Accuracy is affected by temperature and barometric pressure.
 - 3 It is forbidden to insert or draw out the module when the monitor is working, for it can cause instability of the system. If you do it inadvertently, please turn off the module in menu immediately. The module enters STANDBY mode if you reconnect it to monitor which it is powered on. If the readings are inaccurate, you should do calibration.
-

NOTE:

- 1 Replace the airway adapter, if excessive moisture or secretions are observed in the tubing or if the CO₂ waveform changes unexpectedly without a change in patient

status.

- 2 To avoid infection, use only sterilized, disinfected or disposable airway adapters.
- 3 Inspect the airway adapters prior to use. Do not use it if airway adapter appears damaged or broken. Observe airway adapter color coding for patient population.
- 4 Periodically check the flow sensor and tubing for excessive moisture or secretion buildup.

16.3.3.2 Removing Exhaust Gases from the System

WARNING

Anesthetics: when using the mainstream CO₂ measurement on patients who are receiving or have recently received anesthetics, connect the outlet to a scavenging system, to avoid exposing medical staff to anesthetics.

Use an exhaust tube to remove the sample gas to a scavenging system. Attach it to the mainstream sensor at the outlet connector.

16.4 Setting CO₂ Waveform Setup

Select the CO₂ waveform area to open the CO₂ waveform menu:

- Set **Mode** to **Curve** or **Filled** as your desire.
- Set **Sweep** to an appropriate value from the pop-up list. The bigger the value is, the quicker the speed is.

16.5 Setting CO₂ Corrections

Temperature, water vapor in the patient's breath, barometric pressure, and the proportions of O₂, N₂O and Helium in the mixture all influence CO₂ absorption. If values seem inaccurately high or low, check that the monitor is using the appropriate corrections. There are **Baro Press**, **O₂ Compens**, **Balance Gas** and **Anes Agent** in the **Other Setup** menu of the **CO₂ Setup** menu and the user can select the desired item.

16.6 Changing CO₂ Alarms

This refers to CO₂ specific alarms. See the Alarms section for general alarm information. To change the alarm, please refer to the following steps:

1. Select the **CO₂ Setup** menu;
2. Select **EtCO₂ Alarm Setup**, **FiCO₂ Alarm Setup** or **AwRR Alarm Setup** to adjust the alarm limit. About how to adjust the alarm limit, please refer to section Setting alarm limits.

16.7 Changing Apnea Alarm

This determines the time limit after which the monitor gives an alarm if the patient stops breathing.

1. Select the **CO₂ Setup** menu to open it;

2. Select **Apnea Alm** from the menu;
3. Choose the apnea alarm time from the pop-up list.

WARNING

Safety and effectiveness of the respiration measurement method in the detection of apnea, particularly the apnea of prematurity and apnea of infancy, has not been established.

Chapter 17 Monitoring C.O.

17.1 Overview

The Cardiac Output (C.O.) measurement is performed by using Thermodilution method. The monitor can determine blood temperature, measure cardiac output. You can have iced injecta using either the flow through system or individual syringes of injecta. You can perform up to 6 measurements before editing the average Cardiac Output. The prompt message on the screen will tell you when to inject.

17.2 C.O. Safety Information

WARNING

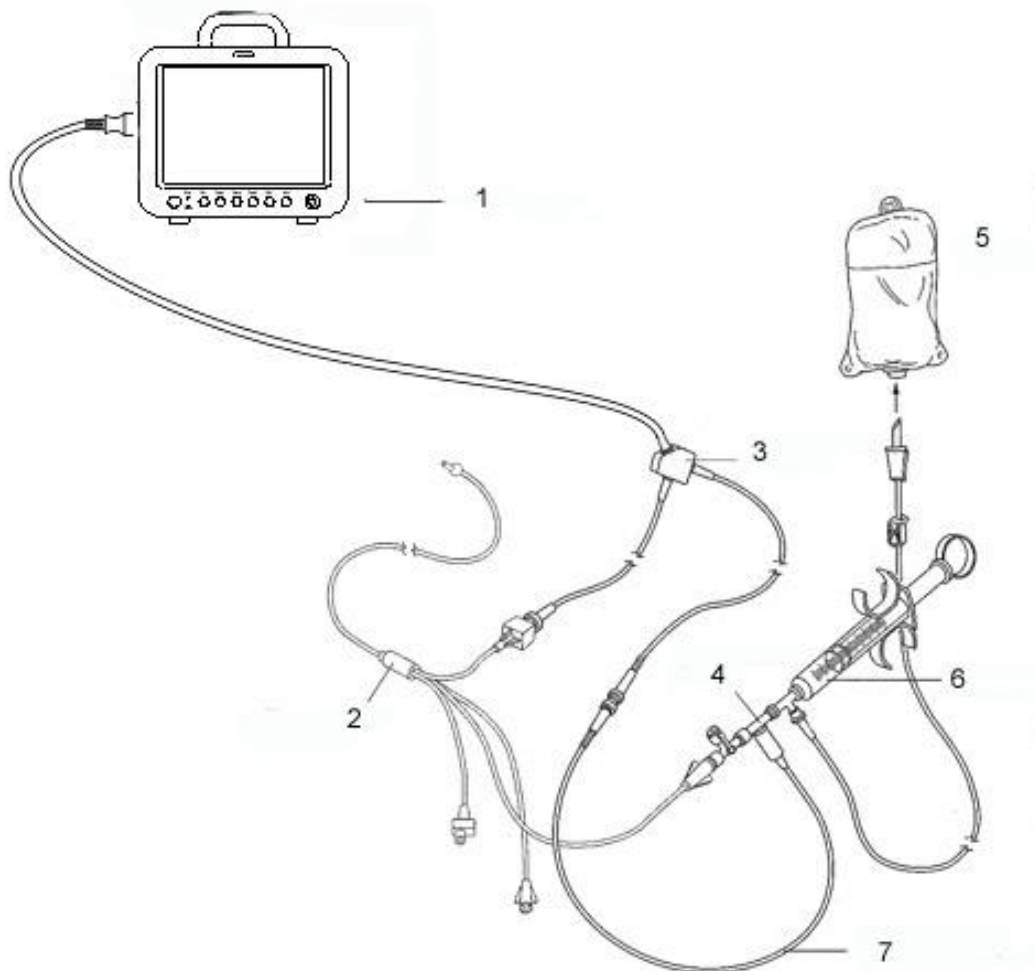
- 1 Make sure that appurtenance applied is in conformity with relevant Medical Device Safety Requirements.
 - 2 Appurtenance should be avoided from contact with conductive metal body when being connected or applied.
-
-

NOTE:

To replace the catheter thermistor, please enter the catheter computation coefficient into the **Constant** item according to the instruction.

17.3 C.O. Monitoring Procedures

1. Plug the C.O. interface cable into the C.O. socket and turn on the monitor.
2. Attach the injectate probe connector and catheter thermistor connector to the appropriate parts of the cardiac output interface cable. And open the patient information window to confirm the patient's height and weight.
3. Pick the **C.O. Measure** item in the **C.O. Option** menu.
4. You can perform more than one measurement as required.
5. After the completion of the measurement, access the **C.O. Measure** window for **Review** to edit the measured data.



1: Monitor; 2: Thermodilution Catheter; 3: Cardiac Output Cable; 4: Injectate Sensor Housing; 5: Injectate; 6: Delivery System; 7: In-line injectate Temperature probe.

C.O. Sensor Connection

WARNING

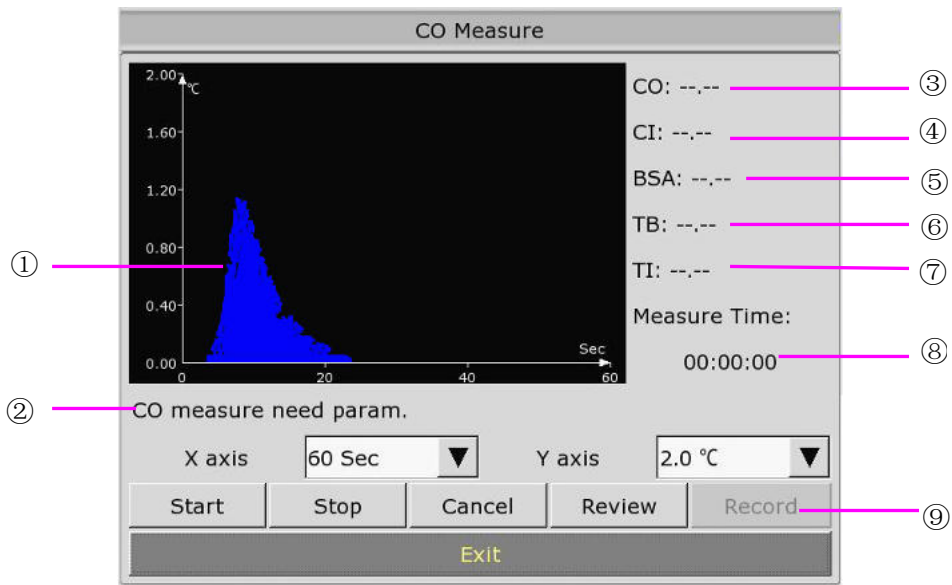
Make sure that the computational constant for the measurement is appropriate to the catheter used.

NOTE:

The blood temperature alarm will not function during C.O. measurement. It will resume automatically when the measurement is over.

17.4 C.O. Measurement Window

Select the **C.O. Option** menu to enter the **C.O. Measure** window and start C.O. measurement. If C.O. transducer is not connected, the monitor will display **No Sensor** on the screen.



C.O. Measure Window

| | |
|---|-------------------------------|
| ① | Measurement curve |
| ② | Prompt message area |
| ③ | Cardiac Output |
| ④ | Cardiac Index |
| ⑤ | Body Surface Area |
| ⑥ | Blood Temperature |
| ⑦ | Injectate Temperature |
| ⑧ | Start time of the measurement |
| ⑨ | Function keys |

The functional keys on the C.O. measure window are explained in the following table:

| | |
|---------------|--|
| Start | Start a measurement |
| Stop | If the blood temperature cannot resume in a considerably long time, the measurement could not stop automatically. Use this button to stop the measurement and display the C.O., CI calculation result. |
| Cancel | Cancel the processing measurement or cancel the result after measurement. |
| Record | Print out the curve. |
| Y axis | Change the scale Y (temperature) value. Three modes are available: 0~0.5°C, 0~1°C, 0~2.0°C. Adjust the scale by the temperature differences. A smaller scale results in a larger curve. |

| | |
|---------------|--|
| X axis | Change the Scale X (time) value. Two modes are available: 0~30s, 0~60s. If you start measurement in the 0~30s mode, it will be switched to 0~60s mode automatically if the measurement can not finish within 30 seconds. After the switch, no further adjustment can be made to the Scale X. |
| Review | Enter the Review window |
| Exit | Press the item to exit C.O. Measure . |

17.5 Measurement Process

Measurement should be taken when the message “**Ready for new measurement**” appears on the screen. Press the **Start** button, and then start injection. The thermodilution curve, current blood temperature and the injectate temperature are displayed during the measurement. Curve drawing will stop automatically when the measurement finishes, and the the C.O. and CI (③ and ④ in the above figure) will be calculated and displayed on the screen. The monitor will display C.O. in the parameter area and the start measurement time (⑧ in the above figure).

To ensure the accuracy of the measurement, it is suggested that a reasonable interval should take place between two consecutive measurements. The length of the interval can be set in the C.O. Setup menu (Time unit: second). The interval time counter is displayed on the screen. The next measurement can not be performed until the time reduces to zero and a message **Ready for new measurement** appears.

NOTE:

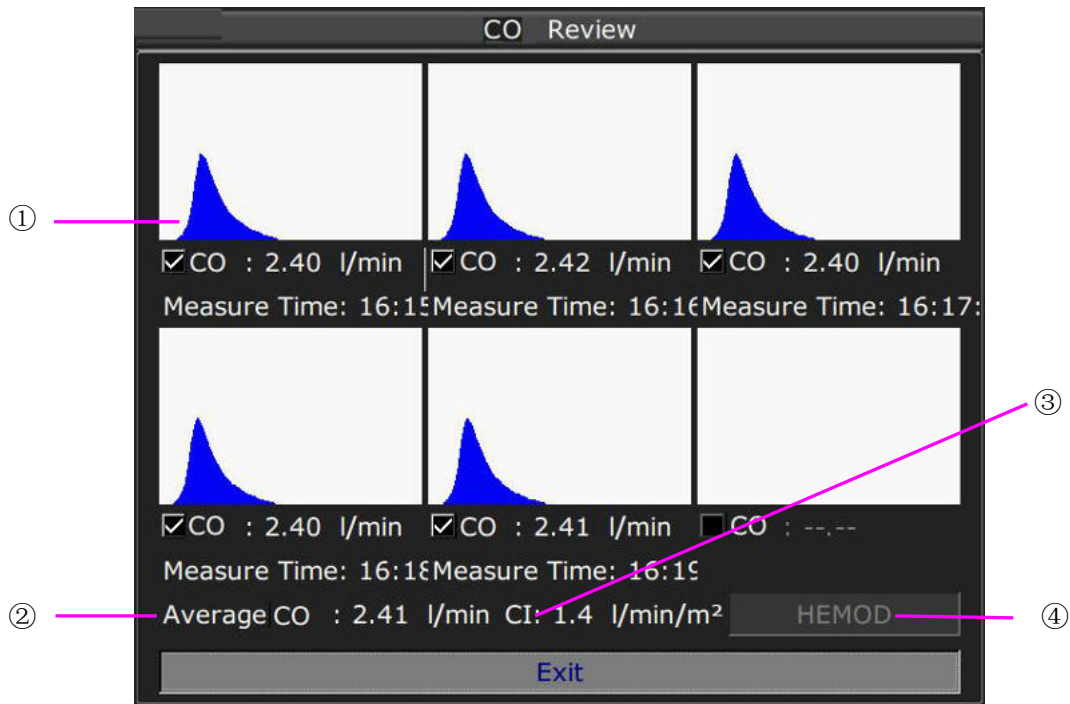
- 1 It is strongly recommended that the user must push the injector within four seconds after pressing the **Start** button.
- 2 It is strongly recommended that you wait at least 1 minute (or longer depending on the patient’s clinical condition) before starting the next measurement.

Repeat this procedure until you have completed the measurements you want.

You can perform a maximum of six measurement editing. If you perform additional measurements the earliest measurement each time will be deleted. If any of the curves in the editing window is not selected for calculation (excluded from the averaging calculations), the place will be taken by the new measurement.

17.6 Editing C.O.

Pick the **Review** button on the **C.O. Measure** menu to access the **Review** as shown below:



Window For C.O. Edit

◆ Contents displayed in the window:

| | |
|---|---|
| ① | Six curves of the six measurements and C.O. value |
| ② | Average value of C.O. |
| ③ | Average value of CI |
| ④ | Function button in the edit window |

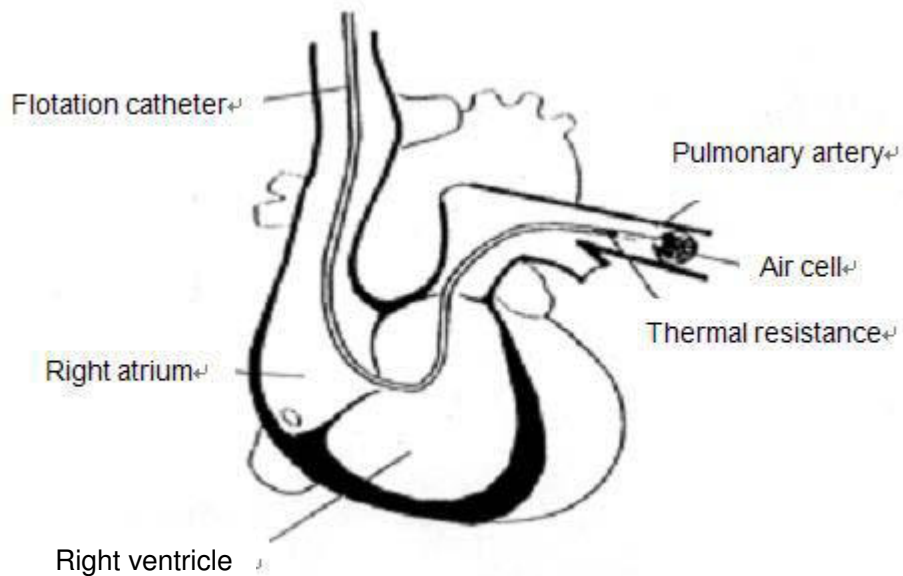
Values of selected measurements can be averaged and stored in the C.O. item in the HEMOD menu as the basis for Hemodynamic calculations.

17.7 Blood Temperature Monitoring

Blood temperature monitoring can function when C.O. measurement is not taken. The blood temperature is measured by the thermistor situated in the distal end of the flotation catheter in the pulmonary artery.

The blood temperature alarm function will not work during the C.O. measurement. When the measurement ends, the function will automatically resume.

The current blood temperature is displayed in the C.O. parameter area.



Thermodilution Catheter Site

17.8 Setting the Computation Constant

The computation constant is associated with catheter and injectate volume. When the catheter is changed, please adjust **Constant** in the **C.O. Settings** menu based on product description provided by the manufacturer.

17.9 Recording C.O. Measurements

C.O. measurement can be recorded by the recorder. To record the C.O. measurement, please select **Record** in the **C.O. Measure** menu.

17.10 Setting INJ. TEMPSource

To change the INJ Temp Source, please:

- 1 Select **Inj Temp Source** in the **C.O. Settings** menu;
 - 2 Select **Auto** or **Manual** from the list;
- **Manual**: directly displaying the injectate temperature from INJ. TEMP.
 - **Auto**: indicating the system obtains the injectate temperature through sampling.

Chapter 18 Monitoring AG

18.1 Overview

The monitor uses ISA sidestream gas analyzer (hereinafter called ISA analyzer) and IRMA mainstream module (hereinafter called IRMA module) to monitor the anesthetic gas which can be used to measure the gases of adult, pediatric and neonatal patients during anesthesia, recovery and respiratory care. And the anesthetic gas includes Halothane (HAL), Isoflurane (ISO), Enflurane (ENF), Sevoflurane (SEV), Desflurane (DES), CO₂ and N₂O.

18.2 Safety Information

18.2.1 Safety Information for ISA Analyzer

WARNING

- 1 The ISA analyzer is intended for use by authorized and trained medical personnel only.
 - 2 Use only Nomoline sampling lines manufactured by PHASEIN.
 - 3 The ISA analyzer must not be used with flammable anesthetic agents.
 - 4 Carefully route the sampling line to reduce the risk of patient entanglement or strangulation.
 - 5 Do not re-use disposable sampling line.
 - 6 Do not lift the monitor by the sampling line as it could disconnect from the monitor, causing the monitor to fall on the patient.
 - 7 Used disposable sampling lines shall be disposed of in accordance with local regulations for medical waste.
 - 8 Do not use adult/pediatric type sampling line configurations with infants, as this may add dead space to the patient circuit.
 - 9 Do not use infant type sampling line configurations with adults, as this may cause excessive flow resistance.
 - 10 Do not use the ISA analyzer with metered-dose inhalers or nebulized medications as this may clog the bacteria filter.
 - 11 Check that the gas sample flow is not too high for the present patient category.
 - 12 Since a successful zeroing requires the presence of ambient air (21% O₂ and 0% CO₂) in the gas analyzer, ensure that the ISA is placed in a well ventilated place. Avoid breathing near the ISA sidestream gas analyzer before or during the zeroing procedure.
 - 13 The Nomoline sampling line and its interfaces are non-sterile devices. To avoid damage, do not autoclave any part of the sampling line.
-

WARNING

- 14 Never sterilize or immerse the ISA analyzer in liquid.
 - 15 Measurements can be affected by mobile and RF communications equipment. Make sure that the ISA analyzer is used in the electromagnetic environment specified in this manual.
 - 16 ISA analyzer is intended only as an adjunct in patient assessment. It must be used in conjunction with other assessments of clinical signs and symptoms.
 - 17 Replace the sampling line if the sampling line input connector starts flashing red, or a Nomoline occlusion message is displayed on the host.
 - 18 No modification of this equipment is allowed without authorization of the manufacturer. If this equipment is modified, appropriate inspection and testing must be conducted to ensure continued safe operation.
 - 19 ISA analyzers are not designed for MRI environments.
 - 20 During MRI scanning, the monitor must be placed outside the MRI suite.
 - 21 Use of high frequency electrosurgical equipment in the vicinity of the monitor may produce interference and cause incorrect measurements.
 - 22 Do not use external ambient cooling of the ISA device.
 - 23 Do not apply negative pressure to the Nomoline to remove condensed water.
 - 24 Too strong positive or negative pressure in the patient circuit might affect the sample flow.
 - 25 Exhaust gases should be returned to the patient circuit or a scavenging system.
 - 26 Always use a bacteria filter on the evac side if sampled gas is intended to be re-breathed.
 - 27 Do not place the ISA analyzer in any position that might cause it to fall on the patient.
-
-

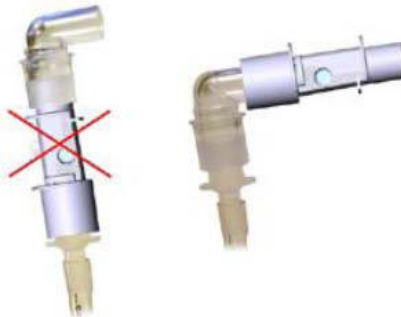
CAUTION

- 1 The ISA analyzers should be securely mounted in order to avoid the risk of damage to the ISA.
 - 2 Do not apply tension to the ISA analyzer cable.
 - 3 Do not operate the ISA analyzer outside the specified operating temperature environment.
-
-

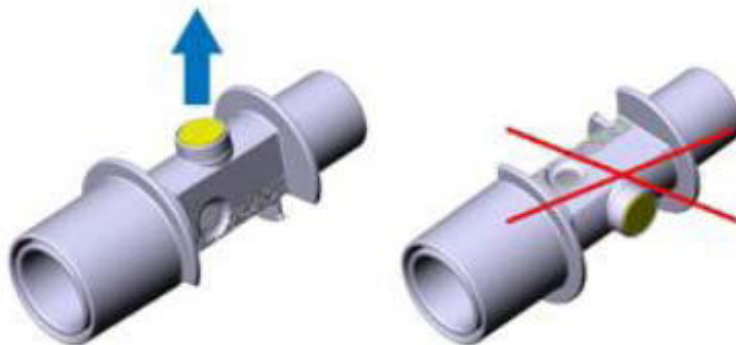
18.2.2 Safety Information for IRMA Module

WARNING

- 1 The IRMA probe is intended for use by authorized and trained medical personnel only.
- 2 The IRMA probe must not be used with flammable anesthetic agents.
- 3 Disposable IRMA airway adapters shall not be reused. Used disposable airway adapters shall be disposed of in accordance with local regulations for medical wastes.
- 4 Use only PHASEIN manufactured oxygen sensor cells. Depleted oxygen sensors shall be disposed of in accordance with local regulations for batteries.
- 5 Do not use the IRMA Adult/Pediatric adapter with infants as the adapter adds 6 ml dead space to the patient circuit.
- 6 Do not use the IRMA airway adapter with adults as this may cause excessive flow resistance.
- 7 Measurements can be affected by mobile and RF communications equipment. It should be assured that the IRMA probe is used in the electromagnetic environment specified in this manual.
- 8 Do not place the IRMA airway adapter between the endotracheal tube and an elbow as this may allow patient secretions to block the adapter windows and result in incorrect operation.



- 9 To keep secretions and moisture from pooling on the windows or oxygen sensor port, always position the IRMA probe in a vertical position with the LED pointing upwards.



WARNING

- 10 Do not use the IRMA airway adapter with metered dose inhalers or nebulized medications as this may affect the light transmission of the airway adapter windows.
 - 11 Do not try to open the oxygen sensor assembly. The oxygen sensor is a disposable product and contains a caustic electrolyte and lead.
 - 12 The IRMA probe is intended only as an adjunct in patient assessment. It must be used in conjunction with other assessment of clinical signs and symptoms.
 - 13 Incorrect probe zeroing will result in false gas readings.
 - 14 Incorrect agent selection by the user for IRMA AX (no automatic agent identification) will result in false agent readings.
 - 15 Using IRMA AX (no automatic identification) with gas mixtures containing more than one agent will result in false agent readings.
 - 16 Replace the adapter if rainout/condensation occurs inside the airway adapter.
 - 17 Use only PHASEIN manufactured IRMA airway adapters.
-
-

CAUTION

- 1 Do not apply tension to the probe cable.
 - 2 Do not operate the IRMA probe outside the specified operating temperature environment.
 - 3 Do not leave depleted oxygen sensors mounted in the IRMA probe, even if the probe is not in use.
-
-

18.3 Monitoring Steps

18.3.1 Monitoring Steps for ISA Analyzer

18.3.1.1 Performing a Pre-use Check

Before connecting the Nomoline sampling line to the breathing circuit, do the following:

1. Connect the sampling line to the ISA gas inlet connector (LEGI).
2. Check that the LEGI shows a steady green light (indicating that the system is OK).
3. For ISA OR+ and ISA AX+ module with O₂ option fitted: Check that the O₂ reading on the monitor is correct (21%).
4. Breathe into the sampling line and check that valid CO₂ waveforms and values are displayed on the monitor.
5. Occlude the sampling line with a fingertip and wait for 10 seconds.
6. Check that an occlusion alarm is displayed and that the LEGI shows a flashing red light.

7. If applicable: Perform a tightness check of the patient circuit with the sampling line attached.

18.3.1.2 System Setup for Analyzer

If your system is using the plug-in and measure ISA analyzer, please follow the setup instructions below:

1. Connect the ISA analyzer interface cable to the monitor.
2. Connect a Nomoline sampling line to the ISA analyzer input connector.
3. Connect the gas sample exhaust port to a scavenging system or return the gas to the patient circuit.
4. Power up the monitor.
5. A green LED indicates that the ISA analyzer is ready for use.
6. Perform a pre-use check as described in section Perform a pre-use Check.

18.3.1.3 Zeroing

The infrared module needs to establish a zero reference level for the CO₂, N₂O and anesthetic agent gas measurement. This zero calibration is here referred to as "zeroing".

ISA analyzer performs zeroing automatically by switching the gas sampling from the respiratory circuit to ambient air. The automatic zeroing is performed every 24 hours, and takes less than 3 seconds for ISA CO₂ module and less than 10 seconds for ISA analyzer.

If the ISA analyzer is fitted with an oxygen sensor, the automatic zeroing will also include room air calibration of the oxygen sensor.

WARNING

1. Since a successful zeroing requires the presence of ambient air (21% O₂ and 0% CO₂) in the ISA analyzer, ensure that the ISA analyzer is placed in a well ventilated place. Avoid breathing near the ISA analyzer before or during the zeroing procedure.
 2. The sampling line should be replaced every two weeks, otherwise it is clogged.
-
-

18.3.1.4 Maintenance

GAS readings should be verified by conducting the recommended maintenance checks. For details, please refer to Chapter *Maintenance*.

WARNING

1. The Nomoline sampling lines are non-sterile devices. To avoid damage, do not autoclave any parts of the sampling line.
 2. Never sterilize or immerse the ISA analyzer in liquid.
-
-

18.3.1.5 MAC Calculation

The MAC value may be calculated and displayed by using end-tidal (Et) gas concentrations according to the following formula:

$$MAC = \frac{\%Et(AA1)}{X(AA1)} + \frac{\%Et(AA2)}{X(AA2)} + \frac{\%Et(N2O)}{100}$$

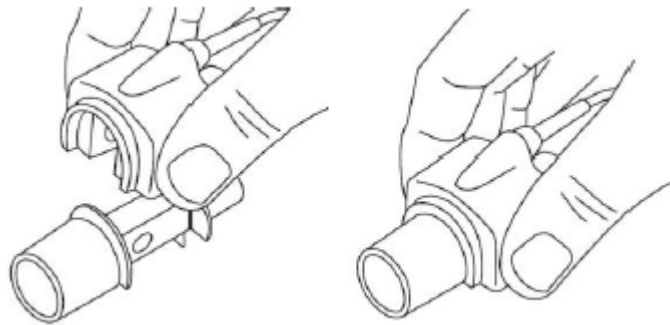
X (AA): HAL=0.75%, ENF=1.7%, ISO=1.15%, SEV=2.05%, DES=6.0%

NOTE:

Altitude, patient age and other individual factors are not considered in the formula above.

18.3.2 Monitoring Steps for IRMA Module

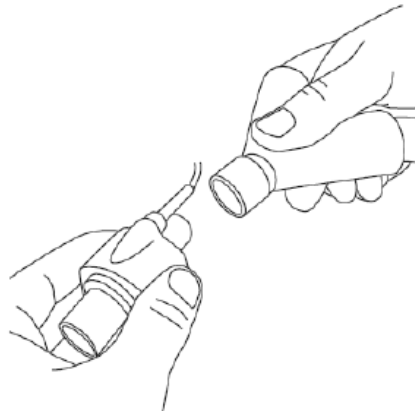
- 1 Plug the IRMA connector into the IRMA input and switch the power on.
- 2 Snap the IRMA sensor head on the top of the IRMA airway adapter. It will click into place when properly seated.



- 3 A green LED indicates that the IRMA probe is ready for use.



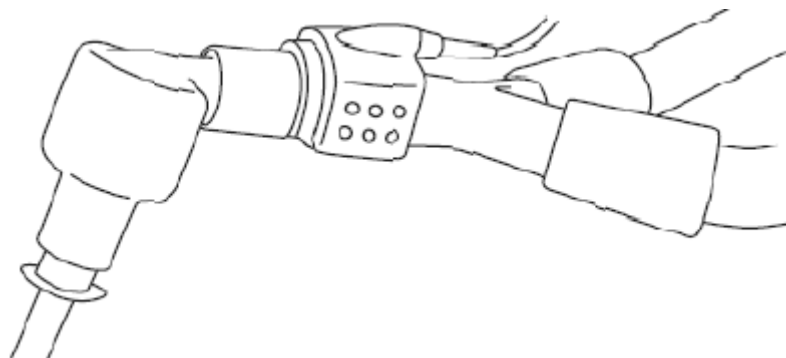
- 4 Connect IRMA/airway adapter 15mm male connector to the breathing circuit Y-piece.



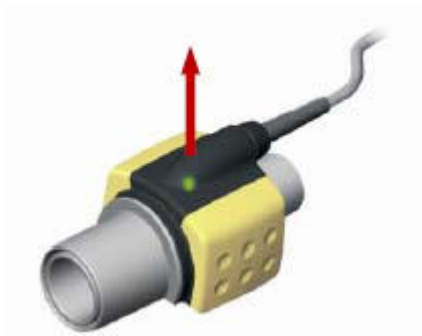
- 5 Connect the IRMA/airway adapter 15mm female connector to the patient's endotracheal tube.



Alternatively, connect an HME (Heat Moisture Exchanger) between the patient's endotracheal tube and the IPMA probe. Placing an HME in front of the IRMA probe protects the airway adapter from secretions and effects of water vapor and eliminates the need of changing the adapter. It allows free positioning of the IRMA probe as well.



- 6 Unless the IRMA probe is protected with an HME always position the IRMA probe with the status LED pointing upwards.



18.3.2.1 Placement of IRMA Probe

When connecting IRMA probe to an infant patient circuit, it is important to avoid a direct contact between the IRMA probe and the infant's body. If, for whatever the reason, the IRMA probe is in direct contact with any parts of the infant's body, an insulation material shall be placed between the IRMA probe and the body.

WARNING

The IRMA probe is not intended to be in long term skin contact.

18.3.2.2 Performing a Pre-use Check

Prior to connecting the IRMA airway adapter to the breathing circuit, verify the O₂ calibration by checking that the O₂ reading on the monitor is correct (21%). See the following section on how to perform air calibration.

Always verify gas readings and waveforms on the monitor before connecting the airway adapter to the patient circuit.

Perform the tightness check of the patient circuit with the IRMA probe snapped on the IRMA airway adapter.

18.3.2.3 Zeroing

WARNING

Incorrect probe zeroing will result in false gas readings.

In order to secure high precision of the IRMA probe measurements the following zeroing recommendations should be followed.

Zeroing is performed by snapping a new IRMA airway adapter onto the IRMA probe, without connecting the airway adapter to the patient circuit, and then using the host instrument to transmit a zero reference command to the IRMA probe.

Special care should be taken to avoid breathing near the airway adapter before or during the zeroing procedure. The presence of ambient air (21% O₂ and 0% CO₂) in the IRMA airway

adapter is of crucial importance for a successful zeroing. If a “ZERO-REQ” alarm should appear directly after a zeroing procedure, the procedure has to be repeated.

Always perform a pre-use check after zeroing the probe.

Zeroing for IRMA CO₂ probes:

Zeroing needs to be performed ONLY when an offset in gas values is observed, or when an unspecified accuracy message is displayed.

Allow 10 seconds for warm up of the IRMA CO₂ probe after power on and after changing the IRMA airway adapter before proceeding with the Zeroing Procedure. The green LED on the probe will be blinking for approximately 5 seconds while zeroing is in progress.

Zeroing for IRMA AX+ probes:

Zeroing should be performed every time the IRMA airway adapter is replaced, or whenever an offset in gas values or an unspecified gas accuracy message is displayed.

Allow 30 seconds for warm up of the IRMA AX+ probes after power on and after changing the IRMA airway adapter before proceeding with the Zeroing Procedure. The green LED on the probe will be blinking for approximately 5 seconds while zeroing is in progress.

18.3.2.4 Cleaning

The IRMA probe can be cleaned using a cloth moistened with maximum 70% ethanol or maximum 70% isopropyl alcohol.

Remove the disposable IRMA airway adapter prior to cleaning the IRMA probe.

CAUTION

- 1 The IRMA oxygen sensor cell and IRMA airway adapters are non-sterile devices. Do not autoclave the devices as this will damage them.
 - 2 Never sterilize or immerse the IRMA probe in liquid.
-
-

18.3.2.5 MAC Calculation

The MAC value may be calculated and displayed by using end-tidal (ET) gas concentrations according to the following formula:

$$\text{MAC} = \% \text{ET}(\text{AA}_1) / \text{X}(\text{AA}_1) + \% \text{ET}(\text{AA}_2) / \text{X}(\text{AA}_2) + \% \text{ET}(\text{N}_2\text{O}) / 100$$

X(AA): HAL=0.75%, ENF=1.7%, ISO=1.15%, SEV=2.05%, DES=6.0%

18.4 Setting Work Mode

There are two woke modes: **Measure** and **Standby**. To change the work mode, please refer to the following steps:

- 1 Select the **AG Setup** menu;
- 2 Select **Work Mode** on the interface and select **Measure** or **Standby** from the popup list.

18.5 Setting Alarms

Here we take CO₂ alarm for example. This refers to CO₂ specific alarms. See the Alarms Chapter for general alarm information. To change the alarm, please refer to the following steps:

- 1 Select the **CO₂ Setup** menu;
- 2 Select **EtCO₂ Alarm High Limit** or **EtCO₂ Alarm Low Limit** to adjust the alarm limit.

18.6 Setting Apnea Alarm Time

This determines the time limit after which the monitor gives an alarm if the patient stops breathing.

- 1 Select the **AG Setup** menu to open it;
- 2 Select **Apnea Alarm** from the menu;
- 3 Choose the apnea alarm time from the pop-up list.

18.7 Working Status of ISA analyzer

Working status of the ISA analyzer can be indicated by the indicator. For the detailed information, please refer to the following table.

| Indication | Status |
|----------------------|--------------------------|
| Steady green light | System OK |
| Blinking green light | Zeroing in progress |
| Steady blue light | Anesthetic agent present |
| Steady red light | Sensor error |
| Blinking red light | Check sampling line |

18.8 Working Status of IRMA Module

The working status of the IRMA module can be transmitted by the IRMA probe. For the detailed information, please refer to the following table.

| Indication | Status |
|----------------------|--------------------------|
| Steady green light | System OK |
| Blinking green light | Zeroing in progress |
| Steady blue light | Anesthetic agent present |
| Steady red light | Sensor error |
| Blinking red light | Check adapter |

18.9 N₂O and O₂ Compensations

The following models need O₂ compensation: IRMA AX+, IRMA CO₂, ISA AX+, ISA CO₂. The following models need N₂O compensation: IRMA CO₂ and ISA CO₂. For the compensation details, please refer to the following table.

| O ₂ Range | SetO ₂ Range |
|----------------------|-------------------------|
| 0~30 vol% | 21 |
| 30~70 vol% | 50 |
| 70~100 vol% | 85 |

| N ₂ O Range | Set N ₂ O Range |
|------------------------|----------------------------|
| 0~30 vol% | 0 |
| 30~70 vol% | 50 |

18.10 Effects of humidity

The partial pressure and the volume percentage of CO₂, N₂O, O₂ and anesthetic agents depend on the amount of water vapor in the measured gas. The O₂ measurement will be calibrated to show 20.8 vol% at actual ambient temperature and humidity level, instead of showing actual partial pressure. 20.8 vol% O₂ corresponds to the actual O₂ concentration in room air with 0.7 vol% H₂O concentration (at 1013 hPa this equals for example 25°C and 23% RH). The measurement of CO₂, N₂O, and anesthetic agents (e.g. all gases measured by the IR-bench) will always show the actual partial pressure at the current humidity level.

In the alveoli of the patient, the breathing gas is saturated with water vapor at body temperature (BTPS).

When the breathing gas flows through the sampling line, the gas temperature will adapt to ambient before reaching the gas analyzer. As the NOMO section removes all condensed water, no water will reach the ISA gas analyzer. The relative humidity of the sampled gas will be about 95%.

If CO₂ values at BTPS are required, the following equation can be used:

$$EtCO_2(BTPS) = EtCO_2 * \left(1 - \left(\frac{3.8}{P_{amb}} \right) \right)$$

where:

$EtCO_2$ = EtCO₂ value sent from ISA [vol %]

P_{amb} = Ambient pressure sent from ISA [kPa]

3.8 = Typical partial pressure of water vapor condensed between patient circuit and ISA [kPa]

EtCO₂(BTPS) = EtCO₂ gas concentration at BTPS [vol%]

O₂ is assumed to be room air calibrated at a humidity level of 0.7 vol% H₂O.

Chapter 19 Freeze


19.1 Overview

When monitoring a patient, the user may freeze the waveforms and examine them. Generally, the user can review a frozen waveform of a maximum of 120 seconds. The freeze function of this monitor has the following features:

- Freeze status can be activated on any operating screen.
- Once entering the Freeze status, the system exits all other operating menus. Besides, the system freezes all waveforms in the Waveform area of the Basic Screen, and also freezes Full Lead ECG waveforms and extra waveforms on the Full Lead ECG interface (if any). Nevertheless the Parameter area refreshes normally.
- The frozen waveforms can be reviewed and recorded.


19.2 Entering/Exiting Freeze Status

19.2.1 Entering Freeze Status


In the Non-Freeze status, press the  button on the control panel of the monitor to exit the current menu. Freeze status is entered and the popup **Freeze** menu is displayed. In Freeze status, all waveforms are frozen and will no longer be refreshed.

19.2.2 Exiting Freeze Status

In the Freeze status, executing any of the following operations will command the system to exit the Freeze status:


- Select the **Exit** option in/from the **Freeze** menu;
- Press the  button on the control panel again;
- Execute any operation that may trigger the adjustment of the screen or the display of a new menu.

After exiting Freeze status, the system will clear screen waveforms and resume displaying real-time waveforms. In the Screen Refresh mode, the system will sweep the waveforms from left to right in the Waveform Area.

Press the  button on the control panel, and the **Freeze** menu will appear on the bottom part of the screen. At the same time, the system freezes the waveforms.

- **REC WAVE:** it can be set to any waveform of 8s, such as **IBP1**, **CO₂**, **PLETH** etc. It can also be set to **OFF**.
- **Review:** Used to review frozen waveforms.
- **Exit:** The system closes the **Freeze** menu and exits the Freeze status.

NOTE:

Pressing the  button repeatedly over a short period of time may result in discontinuous waveforms on the screen.

19.3 Reviewing Frozen Waveform

By moving the waveform, you may review a waveform of 120 seconds before it is frozen. For a waveform of less than 60 seconds, the remaining part is displayed as a straight line. Use the rotary knob on the control panel to move the cursor to the **Review** option in the **Freeze** menu. Press the knob. By turning the knob left or right, frozen waveforms on the screen will move left or right correspondingly. There is an arrow indicating upward on the right side of the last waveform.

Chapter 20 Review

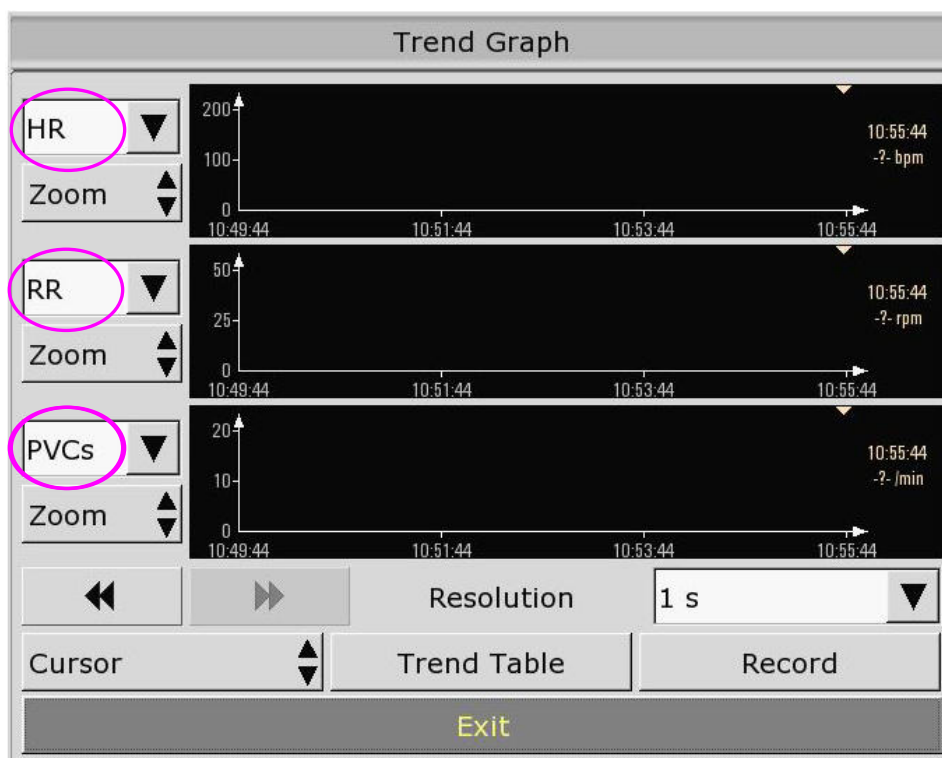
The monitor provides 120-hour trend data of all parameters, storage of 1200 NIBP measurement results and 60 alarm events. This chapter gives detailed instruction for review of all data.

20.1 Trend Graph Review

- The latest 1-hour trend is displayed every 1 or 5 seconds.
- The latest 120-hour trend is displayed every 1, 5 or 10 minutes.

To review Trend Graph, please press the **Trend Graph** key on the screen or select **Menu > Review > Trend Graph**, then the trend graph interface is displayed.

In the trend graph, the y-axis stands for the measurement value and x-axis stands for the time.



20.1.1 Selecting Trend Graph of Specific Parameter



The monitor can review trend graph of different parameters. To change the existing trend graph, please select **Menu > Review > Trend Graph** and select a required parameter name from the popup list (as shown in red text in the above figure).

20.1.2 Setting Resolution

The monitor can support five kinds of resolutions. To set an appropriate resolution, please select **Menu > Review > Trend Graph** and an interface is displayed. Choose **Resolution** on the interface to open the list and select an appropriate resolution among **1 sec**, **5 sec**, **1 min**, **5 min** and **10 min**.

20.1.3 Scrolling Left and Right the Screen

All trend graphs can't be displayed on the current screen due to the screen limitation. The user

can scroll left and right the screen manually to see measurement trends that do not fit in the current view by selecting and pressing the symbol  and  displayed on the trend graph.

20.1.4 Switching to the Trend Table

The user can switch to the trend table interface on the **Trend Graph** interface. To do so, please select **Menu > Review > Trend Graph** and select the **Trend Table** option from the popup interface.

20.1.5 Record

The monitor can make a tabular trend recording of the data in the current trend graph window. The report will use the current trend interval settings. For the detailed information about recording the trend graph, please refer to Chapter *Recording*.

20.2 Trend Table Review

To review the trend table, please press the **Trend Table** key on the screen or select **Menu > Review > Trend Table**, then the trend table is displayed.





NOTE:

The CO₂ module and AG module can not be measured at the same time, so their trend graph can not be displayed at the same time.

20.2.1 Setting Resolution

The monitor can support seven kinds of interval. To set an appropriate resolution, please select **Menu > Review > Trend Table** and an interface is displayed. Choose **Resolution** on the interface to open the list and select an appropriate interval among **1 sec, 5 sec, 1 min, 5 min, 10 min, 30 min** and **60 min**.

20.2.2 Scrolling the Screen

All trend tables can't be displayed on the current screen due to the screen limitation. The user can scroll left, right, up and down the screen manually to see measurement trend tables that do not fit in the current view by selecting and pressing the symbol , ,  and  displayed on the trend graph.

20.2.3 Switching to Trend Graph

The user can switch to the trend graph on the **Trend Table** interface. To do so, please select **Menu > Review > Trend Table** and select the **Trend Graph** option from the popup interface.

20.2.4 Recording



The monitor can make a tabular trend recording of the data in the current trend graph window. The report will use the current trend interval settings. For the detailed information about recording the trend table, please refer to Chapter *Recording*.

20.3 NIBP Review

To review the NIBP measurement data, select the **NIBP Review** key on the screen or select

Menu > Review > NIBP Review, then the **NIBP Review** window is displayed.

20.3.1 Scrolling the Screen

All measurement data can't be displayed on the current screen due to the screen limitation. The user can scroll up and down the screen manually to see measurement data that doesn't fit in the current view by selecting and pressing the symbol  and  displayed on the **NIBP Review** interface.

20.3.2 Recording



The monitor can record the measurement data in the NIBP review window. For the detailed information about recording the NIBP review, please refer to Chapter *Recording*.

20.4 Alarm Review

The monitor can display up to 10 technical alarm events in the current screen.

To review the alarm event, select the **Alarm Review** key on the screen or select **Menu > Review > Alarm Review**, then the **Alarm Review** Window is displayed.

20.4.1 Scrolling the Screen

All alarm events can't be displayed on the current screen due to the screen limitation. The user can scroll up and down the screen manually to see alarm events that don't fit in the current view by selecting and pressing the symbol  and  displayed on the **Alarm Review** interface.

20.4.2 Selecting Alarm Event of Specific Parameter

The monitor can review alarm event of the specific parameters. To view the alarm event of the specific parameter, please select **Menu > Review > Alarm Event** and choose **Event Type** to select the required parameter name from the popup list.

20.4.3 Setting Time Index

The user can set end time of alarm review by selecting the **Time Index** option displayed on the alarm review interface.

If the user selects **Current Time** on the popup interface, the alarm events occurring before the current time are displayed on the alarm event review interface.



If the user selects **User Define**, he can define the review time by setting time box displayed on the interface. The alarm events occurring before the **User Define** option are displayed on the alarm event review interface.

20.5 Arr Review

Select **ECG Setup > Arr Analysis > Arr Review** or **Menu > Review > Arr Review** to open the Arr review interface. The interface displays the latest arrhythmia events.

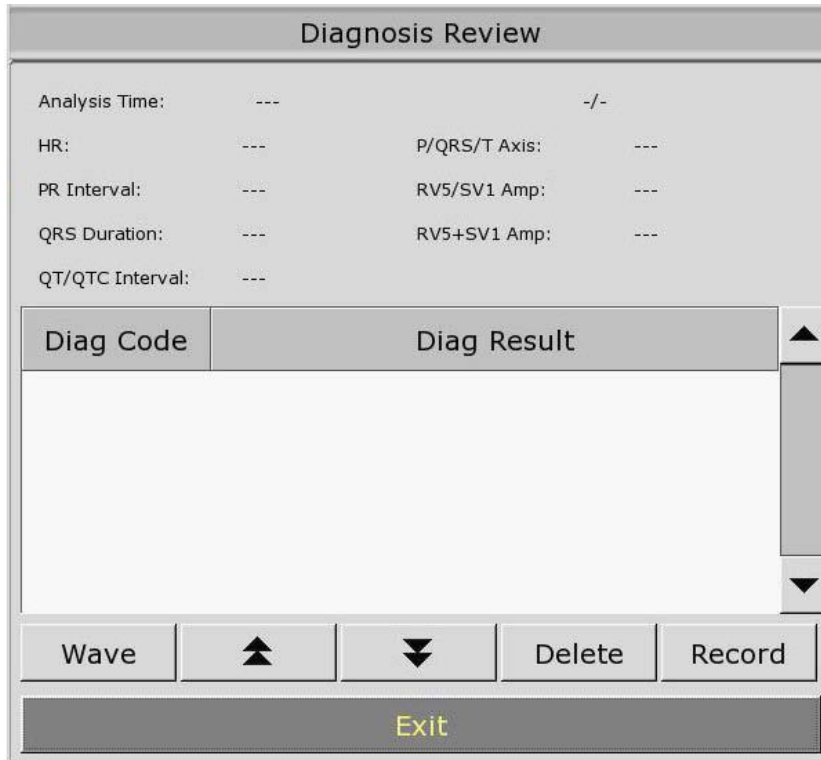
20.5.1 Scrolling the Screen

All arrhythmia events can't be displayed on the current screen due to the screen limitation. The user can scroll up and down the screen manually to see the other arrhythmia events that do not fit



in the current view by selecting and pressing the symbol  and  displayed on the **Arrhythmia Review** interface.

20.6 12-lead Diagnosis Review

Select **Menu > Review > Analysis Review** to open the 12-lead analysis review interface. If there is no analysis result, the prompt of **No Analysis Result** is displayed on the interface.



20.6.1 Scrolling the Screen

All analysis results or waveforms can't be displayed on the current screen due to the screen limitation. The user can scroll up and down the screen manually to see the analysis results or waveforms that do not fit in the current view by selecting and pressing the symbol  and  displayed on the 12-lead analysis review interface.

20.6.2 Deleting Diagnosis Results

The user can delete the analysis results displayed on the current screen by selecting **Delete** on the interface.

20.6.3 Switching Between Waveforms and Results

The user can review the analysis waveforms on the analysis result interface by selecting the **Wave** option and review the analysis results on the analysis waveform interface by selecting the **Results** option.

20.6.4 Recording

The monitor can record the 12-lead diagnosis waveforms or results displayed on the current screen. To do so, press **Record** on the interface. For the detailed information about recording the diagnosis waveforms or results, please refer to Chapter *Recording*.

Chapter 21 Calculation and Titration Table

The monitor provides calculation function and titration table. Calculations are patient data that are not directly measured but calculated by the monitor.

The monitor can perform drug calculation. Hemodynamic calculation is not described in this Instructions for Use. To perform one calculation, select **Menu > Common Function > Drug Dose**.

NOTE:

The drug calculation function acts only as a calculator. The patient weights in Drug Calculation menu and in Patient Information menu are independent of each other. Therefore changing the Weight in Drug Calculation menu will not change the weight in the Patient Information menu.

21.1 Drug Calculation

21.1.1 Calculation Procedures

1. The drug calculation window is displayed by selecting **Menu > Common Function > Drug Dose**.
2. Select the right pull-down box of the **Drug** option and select the required drug name among the 15 drugs which are listed as follows. And the drug name of **Drug A, Drug B, Drug C, Drug D** and **Drug E** can be defined by the user.
 - Drug A, Drug B, Drug C, Drug D and Drug E
 - AMINOPHYLLINE
 - DOBUTAMINE
 - DOPAMINE
 - EPINEPHRINE
 - HEPARIN
 - ISUPREL
 - LIDOCAINE
 - NIPRIDE
 - NITROGLYCERIN
 - PITOCIN
3. The system generates values that can't be treated the calculation results. The user must enter the correct parameter value based on the doctor's instruction.
4. Enter the patient's weight.
5. Enter the correct parameter value.
6. Confirm whether the calculation result is correct.

21.1.2 Calculation Unit

Each drug has the fixed unit or unit series to calculate. Among the same unit series, the unit binary varies with the entered parameter value.

The calculation units of the drugs are listed as follows:

| Drug | Unit |
|--|--------------|
| DRUG A, DRUG B, DRUG C, AMINOPHYLLINE, DOBUTAMINE, DOPAMINE, EPINEPHRINE, ISUPREL, LIDOCAINE, NIPRIDE, NITROGLYCERIN | g, mg, mcg |
| DRUG D, PITOCIN, HEPARIN | Ku, mu, Unit |
| DRUG E | mEq |

When defining a drug, select Drug A, Drug B, Drug C, Drug D, and Drug E based on the unit series.

NOTE:

- 1 The drug calculation is displayed as invalid value before the user edits the drug name and patient weight, and the user can't enter any value.
- 2 Drip Rate and Drop Size are invalid in the neonatal mode.



21.2 Titration Table

After completing the drug calculation, the user can open the **Titration** on the **Drug Dose** interface.

The user can change the following items in the titration table:

- Basic
- Step
- Dose Type

The data in the trend table will vary with the changes above. And the user can perform the following:

- Scroll up and down the screen by selecting and pressing the symbol  and  displayed on the trend graph.
- Record the data displayed in the current window by selecting **Record**.

Chapter 22 Recording

22.1 General Information

A thermal dot matrix recorder is used for the monitor and can support many recording types and output patient information, measurement data, review data waveform and so forth.



| | |
|---|--|
| 1 | Recording indicator |
| 2 | Recording key, press this key to start or stop recording task. |
| 3 | Paper outlet |
| 4 | Recorder Door |

22.2 Performance of the Recorder

- Waveform record is printed at the rate of 25 mm/s or 50 mm/s.
- 48mm wide printout paper.
- It can record up to three waveforms.
- User-selectable real-time recording time and waveform.
- Auto recording interval is set by the user, and the waveform is in accordance with the real time recording.

NOTE:

It is suggested that the user should not use the recorder when the low battery displays, or the monitor may be turned off automatically.

22.3 Recording Type

The monitor provides several types of stripe recording:

- Continuous real-time recording
- Time recording
- Alarm recording
- Trend graph, trend table recording
- Arrhythmia review recording
- Drug calculation titration recording
- NIBP review recording
- Alarm review recording
- Hemodynamic Calculation result recording
- 12-lead analysis recording

22.4 Starting and Stopping Recording


You can start and stop the recording in the following ways:

| | |
|--------------------------------|--|
| Continuous real-time recording | Press the Record button on the front panel to start the recording, and repress it to stop the recording. |
| Auto recording | Record three waveforms selected in Recorder Setup menu according to the setup time interval in Recorder Setup menu. It will automatically stop in 8 seconds. |
| Trend graph recording | Enter the Menu > Review > Trend Graph menu, and press the Record button to start recording. |
| Trend table recording | Enter the Menu > Review > Trend Table menu, and press the Record button to start recording. |
| NIBP review recording | Enter the Menu > Review > NIBP Review menu, then press the Record button to start recording. |
| 12-lead diagnosis recording | Select Record on the diagnosis review interface to start the recording. To stop the recording, press the Record button on the front panel. |

The recorder will stop recording in the following situations:

- The recording task is finished.
- No paper in the recorder.
- Malfunction stops the recorder from running properly.

NOTE:

You can press the  button on the front panel to stop the currently recording process.

22.5 Recorder Operations and Status Messages

22.5.1 Record Paper Requirement

Only standard thermosensitive record paper can be used: otherwise the recorder may not function, the recording quality may be poor, and the thermosensitive printhead may be damaged.

22.5.2 Proper Operation

- When the recorder is working, the record paper goes out steadily. Do not pull the paper outward with force: otherwise the recorder may be damaged.
- Do not operate the recorder without record paper.

22.5.3 Paper Out

When the **Recorder Out OF Paper** alarm is displayed, the recorder cannot start. Please insert record paper properly.

22.5.4 Replacing Paper

1. Pull outwards the upper arc part of the recorder casing to release the casing, shown in the following figure.



2. Insert a new roll of paper into the paper cassette, printing side facing upwards.



3. Ensure proper position and tidy margin.



4. Pull about 2cm of the paper out, and close the recorder casing.

NOTE:

Be careful when inserting papers. Avoid damaging the thermo-sensitive print head. Unless when inserting papers or shooting troubles, do not leave the recorder catch open.

22.5.5 Removing Paper Jam

When the recorder functions or sounds improperly, you should open the recorder casing to check for a paper jam. Remove the paper jam in the following way:

- Cut the record paper from the feeding edge.
- Open the recorder casing.
- Re-insert the paper.

NOTE:

1. If the monitor is not installed with a recorder, it will indicate **RECORDER SETUP NEEDED** after pressing the **Record** button.
2. Do not touch the thermo-sensitive print head when performing continuous recording.

Chapter 23 Other Functions

23.1 Nurse Call

The monitor provides dedicated nurse call port which is connected to nurse call system through the nurse call cable to perform the nurse call function.



Chapter 24 Using Battery


24.1 Battery Power Indicator

The indicator labeled Battery on the front panel of the monitor illuminates in green when the monitor is battery powered and illuminates in yellow when battery is being charged. The indicator is not illuminated when the monitor is not powered or when AC power is applied.

24.2 Battery Status on the Main Screen

Battery status shows the status of each battery detected and the combined battery power remaining, with an estimate of the monitoring time this represents.

There is a sign  in the lower left corner of screen to show the charging status, and the yellow part is the electric energy of battery. When the monitor is not equipped with battery, the battery status will be shown as the sign , which means no battery.

When the monitor is powered by the battery, the monitor will switch off automatically if there is no electric energy in the battery. When there isn't enough electric energy, a sign  is displayed on the screen.

When the monitor is battery powered, the monitor switches off automatically if there is no power.

24.3 Checking Battery Performance

The performance of rechargeable batteries may deteriorate over time. Battery maintenance as recommended here can help to slow down this process.

1. Disconnect the patient from the monitor and stop all monitoring and measurement.
2. Switch the monitor power on and charge the battery for more than 6 hours continuously.
3. Disconnect monitor from mains power and let the monitor run until there is no battery power left and the monitor shuts off.
4. The running time of the battery reflects the battery performance.

If the running time is obviously less than the specified time in the specification, please change the battery or contact the service personnel.

WARNING

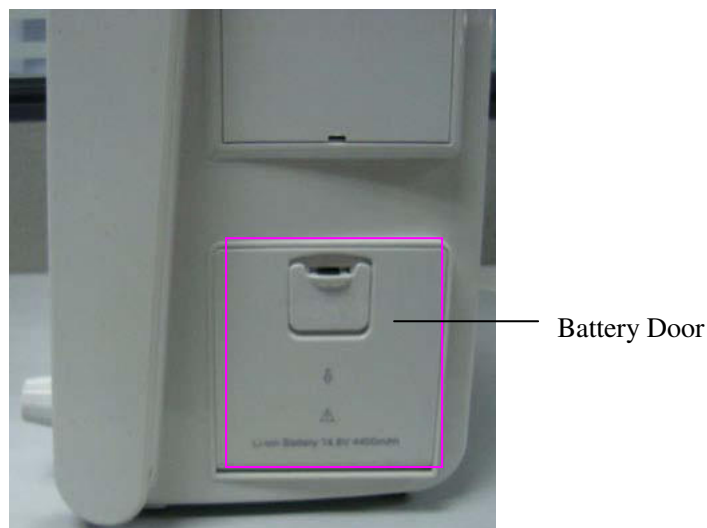
- 1 Before using the rechargeable lithium-ion battery (hereinafter called battery), be sure to read the user manual and safety precautions thoroughly.
 - 2 Do not connect the positive (+) and negative (-) terminals with metal objects, and do not put the battery together with metal object, which can result in short circuit.
 - 3 Do not unplug the battery when monitoring.
 - 4 Do not heat or throw battery into a fire.
-
-

WARNING

- 5 Do not use, leave battery close to fire or other places where temperature may be above 60°C.
 - 6 Do not immerse, throw, or wet battery in water/seawater.
 - 7 Do not destroy the battery: do not pierce battery with a sharp object such as a needle; Do not hit with a hammer, step on or throw or drop to cause strong shock; Do not disassemble or modify the battery.
 - 8 Use the battery only in the monitor. Do not connect battery directly to an electric outlet or cigarette lighter charger.
 - 9 Do not solder the leading wire and the battery terminal directly.
 - 10 If liquid leaking from the battery gets into your eyes, do not rub your eyes. Wash them well with clean water and go to see a doctor immediately. If liquid leaks of the battery splash onto your skin or clothes, wash well with fresh water immediately.
 - 11 Keep away from fire immediately when leakage or foul odor is detected.
 - 12 Stop using the battery if abnormal heat, odor, discoloration, deformation or abnormal condition is detected during use, charge, or storage. Keep it away from the monitor.
 - 13 Do not use a battery with serious scar or deformation.
-
-

24.4 Replacing the Battery

To install or replace the battery, please follow the procedure:



1. Pull the battery door downwards to open it according to indication on it.
2. Pull the metal retainer until the battery can be removed.
3. Insert the new battery into the battery compartment.
4. Pull the metal retainer downward to fix the battery and close the battery door.

24.5 Recycling the Battery

When the battery no longer holds a charge, it should be replaced. Remove the old battery from the monitor and recycle it properly.

NOTE:

To prolong the life of rechargeable battery, it is recommended to charge it at least once every month, and it must be done after the electric energy runs out.

24.6 Maintaining the Battery

Batteries should be conditioned regularly to maintain their useful life.

Discharge the battery completely once every month. And remove the battery from the monitor when the monitor isn't used for a long time.

Chapter 25 Care and Cleaning

Use only the EDAN-approved substances and methods listed in this chapter to clean or disinfect your equipment. Warranty does not cover damage caused by using unapproved substances or methods.

EDAN makes no claims regarding the efficacy of the listed chemicals or methods as a means for controlling infection. Consult your hospital's Infection Control Officer or Epidemiologist for control infection.

25.1 General Points

Keep your monitor, cables and accessories free of dust and dirt. To prevent the device from damage, please follow the procedure:

- Always dilute according to the manufacturer's instructions or use lowest possible concentration.
- Do not immerse any part of the equipment or any accessories in liquid.
- Do not pour liquid onto the system.
- Never use bleach.
- Do not allow liquid to enter the case.
- Never use abrasive material (such as steel wool or silver polish).

CAUTION

If you spill liquid on the equipment, battery, or accessories, or they are accidentally immersed in liquid, contact your service personnel or EDAN service engineer.

25.2 Cleaning

WARNING

Before cleaning the monitor or the sensor, make sure that the equipment is switched off and disconnected from the power line.

25.2.1 Cleaning the Monitor

Regular cleaning of the monitor enclosure and the screen is strongly recommended.

The solutions recommended for monitor cleaning are: soft soap water, Tensides, Ethylate and Acetaldehyde.

WARNING

- 1 Unplug the monitor from the AC power source and detach all accessories before cleaning. Do not immerse the unit in water or allow liquids to enter the case.

- 2 If liquid is splashed on or into the main unit inadvertently, or enters the conduit, stop using the monitor and contact the manufacturer for service immediately.
-
-

Clean the monitor enclosure with soft cloth and diluent non-caustic detergents recommended above.

Clean the screen with a dry soft cloth.

Clean the touch screen with a soft cloth and neutral detergent or isopropyl alcohol. Do not use chemical solvents, acidic or alkali solutions.

CAUTION

- 1 Although the monitor is chemically resistant to most common hospital cleaners and non-caustic detergents, different cleaners are not recommended and may stain the monitor.
 - 2 Many cleansers must be diluted before use. Follow the manufacturer's directions carefully to avoid damaging the monitor.
 - 3 Don't use strong solvent, for example, acetone.
 - 4 Never use an abrasive such as steel wool or metal polish.
 - 5 Do not allow any liquid to enter the product, and do not immerse any part of the monitor into any liquid.
 - 6 Avoid pouring liquids on the monitor while cleaning.
 - 7 Don't remain any cleaning solution on the surface of the monitor.
-
-

NOTE:

- 1 The monitor surface can be cleaned with hospital-grade ethanol and dried in air or with crisp and clean cloth.
- 2 The manufacturer has no responsibility for the effectiveness of controlling infectious disease using these chemical agents. Please contact infectious disease experts in your hospital for details.

25.2.2 Cleaning the Accessories

25.2.2.1 Cleaning the ECG Cables and Lead Wires

NOTE:

1. Use only recommended cleaning substances and disinfectants listed in this document. Others may cause damage (not covered by warranty), reduce product lifetime or cause safety hazards.
2. Keep the cable and lead wires free of dust and dirt.

3. Never immerse or soak the ECG cable.
4. Inspect the cables after cleaning.

CAUTION

Do not allow a cleaning or disinfecting agent to leave residues on any equipment surface. After allowing the appropriate time for the agent to work (as indicated by the manufacturer), wipe off residues with a cloth dampened with water.

Clean with a lint-free cloth, moistened with warm water (40°C/104°F) and substances listed below. Never use strong solvents such as acetone or trichloroethylene.

Approved Cleaning Substances

- Mild Soaps
- Tenside (as active cleaning agent)

Cables and leadwires can be cleaned with a warm, damp cloth and mild soap, or isopropyl alcohol.

25.2.2.2 Cleaning the Blood Pressure Cuff

Wipe cuffs with a solution of mild soap and water. If the cover requires more rigorous cleaning, remove the air bladder first. Allow the cover to thoroughly air dry before use.

Cuffs have been tested to withstand the following recommended disinfectants: cidex, sporicidin, microzid, isopropyl-alcohol 70%, ethanol 70% buraton liquid.

25.2.2.3 Cleaning the SpO₂ Sensor

These reusable sensors should be cleaned and disinfected, but never sterilized. The validated cleaning agents are listed below:

- Mild Detergent
- Salt Solution (1%)

25.2.2.4 Cleaning the IBP/C.O. Cables

NOTE:

1. Keep the cable free of dust and dirt.
2. Never immerse or soak the cable.
3. Inspect the cables after cleaning.

CAUTION

Do not allow a cleaning or disinfecting agent to leave residues on any equipment surface. After allowing the appropriate time for the agent to work (as indicated by the manufacturer), wipe off residues with a cloth dampened with water.

Clean with a lint-free cloth, moistened with warm water (40°C/104°F) and substances listed

below. Never use strong solvents such as acetone or trichloroethylene.

Approved Cleaning Substances

- Mild Soaps
- Tenside (as active cleaning agent)

Cables can be cleaned with a warm, damp cloth and mild soap, or isopropyl alcohol.

25.2.2.5 Cleaning TEMP/Quick TEMP Sensor

CAUTION

Do not allow a cleaning or disinfecting agent to leave residues on any equipment surface. After allowing the appropriate time for the agent to work (as indicated by the manufacturer), wipe off residues with a cloth dampened with water.

Clean with a lint-free cloth, moistened with warm water (40°C/104°F) and substances listed below. Never use strong solvents such as acetone or trichloroethylene.

Approved Cleaning Substances are Hospital-Grade Ethanol.

25.2.2.6 Other Accessories

For cleaning other accessories, please contact the manufactures for details.

25.3 Disinfection

WARNING

Do not mix disinfecting solutions (such as bleach and ammonia), or it may produce hazardous gases.

To avoid extended damage to the equipment, disinfection is only recommended when stipulated as necessary in the Hospital Maintenance Schedule. Disinfection facilities should be cleaned first.

Recommended types of disinfecting agents are:

- Alcohol
- Aldehyde

CAUTION

Do not use EtO gas or formaldehyde to disinfect the monitor.

Chapter 26 Maintenance

WARNING

- 1 Failure on the part of the responsible individual hospital or institution employing the use of this equipment to implement a satisfactory maintenance schedule may cause undue equipment failure and possible health hazards.
- 2 If you discover a problem with any of the equipment, contact your service personnel, or your authorized supplier.

26.1 Inspecting

The overall check of the monitor, including the safety check, should be performed only by qualified personnel every 24 months, and each time after fix up.

The following items should be checked:

- If the environment condition and power supply meet requirement.
- If the power supply cord has damage and insulativity meets requirement.
- If the device and accessories have damage.
- Specified accessories.
- If the alarm system can work properly.
- If the recorder can work properly and the paper meets the requirement.
- Battery performance
- If all monitoring functions are in good conditions.
- If the grounding resistance and leakage current meet requirement.

If any damage or abnormality is found, please don't use the monitor and contact local Customer Service Center.

26.2 Maintenance Task and Test Schedule

The following tasks are for EDAN-qualified service professionals only. Contact an EDAN-qualified service provider if your monitor needs a safety or performance test. Clean and disinfect equipment to decontaminate it before testing or maintaining it.

| Maintenance and Test Schedule | Frequency |
|--|---|
| Safety checks. Selected tests on the basis of IEC60601-1 | At least once every two years, or as needed, after any repairs where the power supply is removed or replaced, or if the monitor has been dropped. |
| Check ECG synchronization of the monitor and defibrillator | At least once every two years, or as needed. |

| | |
|--|---|
| NIBP Leakage Inspection | At least once every two years, or as specified by local laws. |
| NIBP Pressure Calibration | At least once every two years, or as specified by local laws. |
| NIBP Calibration | At least once every two years, or as specified by local laws. |
| CO ₂ Calibration and Performance Test | At least once every two years, or if you suspect the measurement values are incorrect. |
| AG Calibration | If you suspect the measurement values are incorrect and need to calibrate, please contact the manufacturer. |

Chapter 27 Warranty and Service

27.1 Warranty

EDAN warrants that EDAN's products meet the labeled specifications of the products and will be free from defects in materials and workmanship that occur within warranty period.

The warranty is void in cases of:

- a) damage caused by mishandling during shipping.
- b) subsequent damage caused by improper use or maintenance.
- c) damage caused by alteration or repair by anyone not authorized by EDAN.
- d) damage caused by accidents.
- e) replacement or removal of serial number label and manufacture label.

If a product covered by this warranty is determined to be defective because of defective materials, components, or workmanship, and the warranty claim is made within the warranty period, EDAN will, at its discretion, repair or replace the defective part(s) free of charge. EDAN will not provide a substitute product for use when the defective product is being repaired.

27.2 Contact information

If you have any question about maintenance, technical specifications or malfunctions of devices, contact your local distributor.

Alternatively, you can send an email to EDAN service department at: support@edan.com.cn.

Chapter 28 Accessories

You can order accessories from EDAN supplies at www.edan.com.cn or consult your local Edan representative for details.

WARNING

- 1 Never reuse disposable transducers, sensors, accessories and so forth that are intended for single use, or single patient use only. Reuse may compromise device functionality and system performance and cause a potential hazard.
- 2 Use only Edan-approved accessories. Using non-edan-approved accessories may compromise device functionality and system performance and cause a potential hazard.
- 3 Do not use a sterilized accessory if its packaging is damaged.

NOTE:

Transducers and sensors have a limited shelf life. Refer to the package labeling.

The following cables may not all be available in all countries. Please check availability with your local Edan supplier.

28.1 ECG Accessories

The following table lists the optional configuration for the monitor:

| Part Number | Accessories |
|-----------------|--|
| 01.57.471002-11 | ECG cable, IEC, 3-lead, clip, Defib, PVC |
| 01.57.101027-11 | ECG cable, AHA, 5-lead, snap, Defib, PVC |
| 01.57.109100-11 | ECG trunk cable, AHA, 10-lead, Defib, TPU |
| 01.57.109101 | ECG limb wires,10-lead,snap,AHA,0.9m,reusable(only applicable to M80) |
| 01.57.040206 | ECG trunk cable, IEC, 5-lead, Defib, TPU |
| 01.57.040207 | ECG limb wires,5-lead,snap,IEC,0.9m,reusable |
| 01.57.040208 | ECG limb wires,5-lead,clip,IEC,0.9m,reusable |
| 01.57.471022 | ECG trunk cable, AHA, 5-lead, Defib, TPU |
| 01.57.471023 | ECG limb wires,5-lead,snap,AHA,0.9m,reusable |
| 01.57.471024 | ECG trunk cable, IEC, 3-lead, Defib, TPU |
| 01.57.471025 | ECG limb cable,3-lead ,clip,IEC,0.9m, reusable |
| 01.57.040202-11 | ECG trunk cable, IEC, 10-lead, Defib, TPU |
| 01.57.040203 | ECG limb wires,10-lead,snap,IEC,0.9m,reusable |

| | |
|-----------------|--|
| 01.57.471095-10 | ECG cable, AHA, 3-lead, snap, Defib, TPU |
| 01.57.471087-10 | ECG cable, AHA, 3-lead, clip, Defib, TPU |
| 01.57.471096-10 | ECG cable, AHA, 5-lead, snap, Defib, TPU |
| 01.57.471097-10 | ECG cable, AHA, 5-lead, clip, Defib, TPU |
| 01.57.471098-10 | ECG cable, IEC, 3-lead, snap, Defib, TPU |
| 01.57.471099-10 | ECG cable, IEC, 3-lead, clip, Defib, TPU |
| 01.57.471089-10 | ECG cable, IEC, 5-lead, snap, Defib, TPU |
| 01.57.471088-10 | ECG cable, IEC, 5-lead, clip, Defib, TPU |
| 11.57.471056 | ECG Electrodes, adult, disposable, 30 pieces |
| 11.57.471057 | ECG Electrodes, child, neo disposable, 50 pieces |
| 11.57.471060 | ECG Electrodes, adult, disposable, 100 pieces |

28.2 SpO₂ Accessories

| Part Number | Accessories |
|----------------|---|
| EDAN | |
| 02.01.210119 | LEMO SH1 adult reusable SpO ₂ sensor (Only compatible with EDAN SpO ₂ module), (finger type, patient weight more than 40kg) |
| 02.01.210120 | EDAN SH1 adult reusable SpO ₂ finger sensor (DB9), (finger type, patient weight more than 40kg) |
| 01.13.210001 | EDAN SpO ₂ extension cable, DB9 to LEMO, TPU, 2M |
| 02.01.210122 | EDAN SH4 adult soft-tip SpO ₂ sensor, TPU, (finger type, patient weight more than 50kg) |
| 02.01.210123 | EDAN SH4 adult silicone soft-tip SpO ₂ sensor, (finger type, patient weight more than 50kg) |
| 02.01.210121 | EDAN SH5 child silicone soft-tip SpO ₂ sensor, (finger type, patient weight from 10kg to 50kg) |
| 01.57.040196 | Adult disposable SpO ₂ sensor (patient weight more than 30kg) |
| 01.57.040197 | Pediatric Disposable SpO ₂ sensor (patient weight from 10 to 50kg) |
| 01.57.040198 | Infant Disposable SpO ₂ sensor (patient weight from 3 to 20kg) |
| 01.57.040199 | Neonatal Disposable SpO ₂ Sensor (patient weight less than 3kg) |
| NELLCOR | |
| 11.15.30043 | Nellcor Reusable Adult SpO ₂ Sensor (DS-100A OxiMax) |
| 11.15.40096 | Nellcor Reusable Adult/Neonate SpO ₂ Sensor (OXI-A/N OxiMax) |
| 11.13.30131-11 | Nellcor SpO ₂ Extension Cable (Compatible with Nellcor OXI-Max SpO ₂ Module and Nellcor Sensor) |

28.3 NIBP Accessories

| Part Number | Accessories |
|-----------------|---|
| EDAN | |
| 01.59.036118 | NIBP Tube, 3m, Grey |
| 01.59.36036 | NIBP Tube, 3m, Black |
| 01.57.471021 | NIBP Tube for neonatal cuff, 3m |
| 01.57.040210-11 | NIBP Cuff, Larger Adult, 33cm-47cm, reusable |
| 01.57.040205-11 | NIBP Cuff, Adult, 25cm-35cm, reusable |
| 01.57.040211-11 | NIBP Cuff, Child, 18cm-26cm, reusable |
| 01.57.040212-11 | NIBP Cuff, Infant, 10cm-19cm, reusable |
| 11.57.40074 | NIBP Cuff, Larger Adult, 33cm-47cm, reusable |
| 11.57.40029 | NIBP Cuff, Adult, 25cm-35cm, reusable |
| 11.57.40020 | NIBP Cuff, Infant, 10cm-19cm, reusable |
| 11.57.40018 | NIBP Cuff, Child, 18cm-26cm, reusable |
| 01.57.471157 | NIBP Cuff Neonatal, 3cm-6cm, disposable, Suntech |
| 01.57.471158 | NIBP Cuff Neonatal, 4cm-8cm, disposable, Suntech |
| 01.57.471159 | NIBP Cuff Neonatal, 6cm-11cm, disposable, Suntech |
| 01.57.471160 | NIBP Cuff Neonatal, 7cm-13cm, disposable, Suntech |
| 01.57.471161 | NIBP Cuff Neonatal, 8cm-15cm, disposable, Suntech |
| M3600 | |
| 01.59.102099 | OMRON NIBP Tube (3.5m) /CUFF HOSE (NO.1) length3.5m, CE |
| 01.57.471078-10 | OMRON CUFF/CUFF (NO.1) arm12—18cm, width7cm, LATEX, CE |
| 01.57.471079-10 | OMRON CUFF/CUFF (NO.2) arm17—23cm, width9cm, LATEX, CE |
| 01.57.102100 | OMRON CUFF/CUFF (NO.3) arm23—33cm, width12cm, LATEX, CE |
| 01.57.471080-10 | OMRON CUFF/CUFF (NO.4) arm30—40cm, width14cm, LATEX, CE |
| 01.57.471081-10 | OMRON Neonatal disposable cuff/CUFF (NO.10) arm3.5—6cm, width2.5cm, CE |
| 01.57.471082-10 | OMRON Neonatal disposable cuff/CUFF (NO.11) arm5—7.5cm, width3cm, CE |
| 01.57.471083-10 | OMRON Neonatal disposable cuff/CUFF (NO.12) arm7.5—10.5cm, width4cm, CE |

| | |
|-----------------|--|
| 01.57.471084-10 | OMRON Neonatal disposable cuff/CUFF (NO.13) arm8.5—13cm, width5cm, CE |
| 01.59.473003-10 | Connecting Tube for Neonatal Cuff (Only compatible with Neonatal Disposable and NIBP Tube)/CUFF HOSE (NO.3) length3.5m, CE |

28.4 Temp Accessories

| Part Number | Accessories |
|--------------|--|
| 01.57.040185 | Skin Temperature Probe (2.252K Ω) |
| 01.57.040187 | Skin Temperature Probe (10K Ω) |
| 01.57.040184 | Rectal / Oral Temperature Probe (2.252K Ω) |
| 01.57.040186 | Rectal / Oral Temperature Probe (10K Ω) |

28.5 Quick Temp Accessories

| Part Number | Accessories |
|--------------|----------------------|
| 02.04.110140 | Oral/Auxiliary Probe |
| 02.04.110139 | Rectal Probe |
| 11.57.110159 | Probe Covers |

28.6 IBP Accessories

| Part Number | Accessories |
|-----------------|--|
| 01.57.471014 | IBP cable/ BD-IBP cable, used with 682000 sensor |
| 01.57.471013 | Aidehua IBP cable, used with PX260 sensor |
| 11.57.40121 | Disposable pressure transducer kit (BD DT-4812) |
| 01.57.471027-10 | IBP cable, used with 42584 sensor |
| 01.57.471028-10 | IBP cable, used with DPT-248 sensor |

28.7 CO₂ Accessories

| Part Number | Accessories |
|--------------|---|
| 12.08.078137 | Respironics EtCO ₂ Module/ (Side-stream) 1022054 |
| 12.08.078166 | LoFlo™ Module Mounting Bracket (Respironics 1027730) |
| 11.57.078139 | Disposable CO ₂ Nasal Cannula – Adult (Respironics 3468ADU-00) |
| 11.57.078151 | Adult/Pediatric Airway adapter kit with dehumidification tubing |
| 11.57.078154 | Disposable Sampling Line Kit with Dehumidification Tubing (Respironics 3475-00) |
| 11.15.040143 | Respironics CAPNOSTAT 5 EtCO ₂ (Main-stream) Module 1015928 |

| | |
|--------------|--|
| 11.57.471019 | Airway Adapter, reusable, adult/children |
| 11.57.471020 | Airway Adapter, reusable, neonate/premature baby |
| 11.59.078155 | Disposable Adult Airway Adapter (6063-00) |
| 11.59.078156 | Disposable Neonatal (Infant/Pediatric) Airway Adapter (6312-00) |
| 11.57.078142 | Adult Nasal CO ₂ with O ₂ Delivery Sampling Cannula |
| 11.57.078143 | Pediatric Nasal CO ₂ with O ₂ Delivery Sampling Cannula |
| 11.57.078144 | Infant Nasal CO ₂ with O ₂ Delivery Sampling Cannula |
| 11.57.101019 | Adult Nasal/Oral CO ₂ Sampling Cannula |
| 11.57.101020 | Pediatric Nasal/Oral CO ₂ Sampling Cannula |
| 11.57.101021 | Adult Nasal/Oral CO ₂ with O ₂ Delivery Sampling Cannula |
| 01.12.031598 | Adult/Pediatric Airway Adapter Kit |
| 11.57.078140 | Children Nasal Sampling Cannula, sidestream, 2.5m |
| 11.57.078141 | Infant Nasal Sampling Cannula, sidestream, 2.5m |
| 11.57.078152 | Pediatric/Infant Airway Adapter Kit with Dehumidification Tubing |
| 11.57.078158 | Pediatric Mask /Mainstream 9960PED-00 |
| 11.57.078159 | Adult Standard Mask /Mainstream 9960STD-00 |
| 11.57.078160 | Adult Large Mask /Mainstream 9960STD-00 |
| 11.57.078161 | Band /Mainstream 8751-00 |
| 11.12.078162 | Card Slot /Mainstream 6934-00 |

28.8 C.O. Accessories

| Part Number | Accessories | Remark |
|--------------|---|--------------------|
| 11.15.40119 | In-line Injection temperature probe | BD 684056-SP4042 |
| 11.15.40120 | In-line Injection temperature probe housing | BD 680006-SP5045 |
| 11.57.100175 | Control Syringe | Medex MA387 |
| 01.57.471012 | Cardiac output main cable | Yuanhe 98ME07GB106 |

28.9 AG Accessories

| Part Number | Accessories | Remark |
|-----------------|--|------------|
| 11.57.471042-10 | IRMA Airway Adapter, Disposable, adult/children, 25 pcs/package, PHASEIN | Mainstream |
| 11.57.471043-10 | Nomoline with Luer Lock connector, L=2m, 50ml/min, 25 pcs/package, PHASEIN | Sidestream |
| 12.08.208006 | Multi-gas analyzer, IRMA AX+ | |

| | | |
|--------------|-----------------------------|--|
| 12.08.208005 | Multi-gas analyzer, ISA AX+ | |
| 12.08.208007 | Multi-gas analyzer, ISA OR+ | |

28.10 Other Accessories

| Part Number | Accessories | Remark |
|--------------|--------------------------------------|--|
| 11.21.064142 | Rechargeable Lithium-Ion Battery | 14.8V, 2.1Ah |
| 11.21.064143 | Rechargeable Lithium-Ion Battery | 14.8V, 4.2Ah |
| 01.57.78035 | Recorder paper | |
| 12.01.19084 | Thermal printer | |
| 11.21.64056 | Vehicle-carried Inverter | |
| 12.01.30493 | Wall Mount | Simple |
| 02.01.30164 | Wall Mount | |
| 02.01.101043 | Basket | Only compatible with Wall Mount MS3R-30164 |
| 03.28.101952 | Rolling Stand (MT-207) | |
| 02.04.101976 | Rolling Stand Basket (in the bottom) | |
| 11.13.114214 | Ground Cable | |
| 02.01.109592 | Pole clamp | 1 set /package |
| 02.01.109636 | Pole clamp | 4 sets/package |
| 11.18.078205 | Security lock | |
| 11.23.068003 | USB barcode scanner | |

A Product Specification

A.1 Classification

| | |
|-----------------------------------|--|
| Anti-electroshock type | Class I equipment and internal powered equipment |
| EMC type | Group 1, Class A |
| Anti-electroshock degree | ECG (RESP), TEMP, IBP, C.O., Quick Temp CF SpO ₂ , NIBP, CO ₂ , AG BF |
| Ingress Protection | IPX1 (No protection against ingress of water if configured with Quick TEMP module) |
| Disinfection/sterilization method | Refer to Chapter Care and Cleaning for details. |
| Working system | Continuous operation equipment |
| Compliant with Standards | IEC 60601-1:1988+A1+A2, EN 60601-1:1990+A1+A2, IEC/EN 60601-1-2:2001+A1, ISO 9919, ISO 21647, IEC/EN 60601-2-27, IEC/EN 60601-2-30, IEC/EN 60601-2-34, IEC/EN 60601-2-49, ANSI/AAMI SP10, IEC/EN 60601-2-25, AAMI/ANSI EC13, EN12470-4 EN1060-1 EN1060-3, EN1060-4 |

A.2 Physical Specifications

A.2.1 Size and Weight

| Product | Size | Weight |
|---------|--------------------------------------|--------|
| iM80 | 370 mm (L) × 175 mm (W) × 320 mm (H) | 7 kg |
| iM50 | 260 mm (L) × 140 mm (W) × 205 mm (H) | 3.6kg |

A.3 Environment Specification

The monitor may not meet the performance specifications given here if stored or used outside the specified temperature and humidity ranges.

When the monitor and related products have differing environmental specifications, the effective range for the combined products is that range which is common to the specifications for all products.

| | |
|-----------------------|----------------------------|
| Temperature | |
| Working | +5°C ~ +40°C |
| Transport and Storage | -20°C ~ +55°C |
| Humidity | |
| Working | 25% ~ 80% (non-condensing) |
| Transport and Storage | 25% ~ 93% (non-condensing) |

| | |
|-----------------------|--|
| Altitude | |
| Working | 860hPa ~ 1060hPa |
| Transport and Storage | 700hPa ~ 1060hPa |
| Power Supply | 100V-240V~, 50Hz/60Hz Current=1.0A-0.5A(iM50), Pmax=1.4A-0.7A(iM80) FUSE T 1.6AL, 250V |

A.4 Display

| Product | Display | Messages |
|---------|---|---|
| iM50 | Display screen: 8.4 inch colour TFT Resolution: 800×600 | A maximum of 11 waveforms One power LED Two alarm LED One charge LED |
| iM80 | Display screen: 15 inch color TFT Resolution: 1024 × 768 | A maximum of 13 waveforms One power LED Two alarm LED One charge LED |

A.5 Battery Specification

| | | | |
|------|----------------------------|----------------|---|
| iM50 | 2.1Ah | Operating Time | 180 min (At 25°C, continuous SpO ₂ measuring and NIBP automatic measuring mode at interval of 15 minutes, recording at interval of 10 minutes) |
| | | Charge Time | 200 min (Monitor is on or in standby mode.) |
| | 4.2Ah | Operating Time | 420 min (At 25°C, continuous SpO ₂ measuring and NIBP automatic measuring mode at interval of 15 minutes, recording at interval of 10 minutes) |
| | | Charge Time | 380 min (Monitor is on or in standby mode.) |
| iM80 | one battery (4.2Ah) | Operating Time | 120 min (At 25°C, continuous SpO ₂ measuring and NIBP automatic measuring mode at interval of 15 minutes, recording at interval of 10 minutes) |
| | | Charge Time | 320 min (Monitor is on or in standby mode) |
| | Two batteries (2*4.2Ah) | Operating Time | 240 min (At 25°C, continuous SpO ₂ measuring and NIBP automatic measuring mode at interval of 15 minutes, recording at interval of 10 minutes) |
| | | Charge Time | 560 min (Monitor is on or in standby mode) |

A.6 Recorder

| | |
|-----------------|--|
| Record Width | 48 mm |
| Paper Speed | 25 mm/s, 50 mm/s |
| Trace | Up to 3 |
| Recording types | 8 second real-time recording Auto 8 second recording Parameter alarm recording Trend recording Titration table recording |

A.7 Review

| | |
|--------------|---------------------------------------|
| Trend Review | |
| Short | 1 hrs, 1 Second Resolution |
| Long | 120 hrs, 1 min. Resolution |
| Review | 1200 sets NIBP measurement data |
| | 50 sets 12-lead ECG diagnosis results |

A.8 ECG

A.8.1 3, 5-lead Monitoring

| | |
|------------------------------------|--|
| Lead Mode | 3-Lead: I, II, III 5-Lead: I, II, III, aVR, aVL, aVF, V |
| Waveform | 3-Lead: 1-channel waveform 5-Lead: 2-channel waveform, max. seven waveforms; |
| Lead naming style | AHA, IEC |
| Display Sensitivity | 1.25mm/mV (×0.125), 2.5mm/mV (×0.25), 5mm/mV (×0.5), 10mm/mV (×1), 20mm/mV (×2), AUTO gain |
| Bandwidth (-3dB) | Diagnosis: 0.05Hz to 150Hz Monitor: 0.5Hz to 40Hz Surgery: 1Hz to 20Hz |
| CMRR (Common Mode Rejection Ratio) | Diagnosis: >95dB (the Notch filter is off) Monitor: >105dB (the Notch filter is on) Surgery: >105dB (the Notch filter is on) |

| | |
|---|--|
| Notch | 50Hz/60Hz (Notch filter can be turned on or off manually) |
| Differential Input Impedance | >5M Ω |
| Input Signal Range | \pm 8mV PP |
| Accuracy of Input Signal Reconstruction | The total error and frequency response comply with ANSI/AAMI EC13:2002, Sect. 4.2.9.8. |
| Electrode Offset Potential Tolerance | \pm 500mV |
| Auxiliary Current (Leads off detection) | Active electrode: <100nA Reference electrode: <900nA |
| Input Offset Current | \leq 0.1 μ A |
| Recovery time after Defibrillation | <5s |
| Leakage current of patient | <10 μ A |
| Scale signal | 1mVPP, accuracy is \pm 5% |
| System noise | <30 μ VPP |
| ESU Protection | Incision mode: 300W Congelation mode: 100W Restore time: \leq 10s Meets the requirements of ANSI/AAMI EC13-2002: Sect. 4.1.2.1 a) |
| Noise Suppression of Electrotome | Tested according to the test method in EC13: 2002 Sect.5.2.9.14, it accords with the standard. |
| Pace Pulse | |
| Pulse indicator | Pulse is marked if the requirements of ANSI/AAMI EC13:2002, Sect. 4.1.4.1 are met: Amplitude: \pm 2 mV ~ \pm 700 mV Width: 0.1 ms ~2 ms Ascending time: 10 μ s ~ 100 μ s |
| Pulse Rejection | Pulse is rejected if the requirements of ANSI/AAMI EC13-2002: Sect. 4.1.4.1 are met: Amplitude: \pm 2 mV ~ \pm 700 mV Width: 0.1 ms ~2 ms Ascending time: 10 μ s ~100 μ s |

| | |
|------------------------------|---|
| Minimum input slew rate | >2.5V/S |
| Heart rate | |
| Range | ADU: 15 bpm ~ 300 bpm PED/NEO: 15 bpm ~ 350 bpm |
| Accuracy | ±1% or 1 bpm, whichever is greater |
| Resolution | 1 bpm |
| Sensibility | ≥300 μVPP |
| PVC | |
| Range | ADU: 0~300 PVCs/ min PED/NEO: 0~350 PVCs/ min |
| Resolution | 1 PVCs/min |
| ST value | |
| Range | -2.0 mV ~ +2.0 mV |
| Accuracy | The max. of ±0.02 mV or 10% (-0.8 mV ~ +0.8 mV), whichever is greater. |
| Resolution | 0.01 mV |
| HR averaging method | |
| Method 1 | Normally, heart rate is computed by excluding the minimum and maximum values from the 12 most recent RR intervals and averaging the residual 10 RR intervals. |
| Method 2 | If each of three consecutive RR intervals is greater than 1200ms, then the four most recent RR intervals are averaged to compute the HR. |
| Range of Sinus and SV Rhythm | |
| Tachy | ADU: 120 bpm ~ 300 bpm PED/NEO: 160 bpm ~ 350 bpm |
| Normal | ADU: 41 bpm ~ 119 bpm PED/NEO: 61 bpm ~159 bpm |
| Brady | ADU: 15 bpm ~ 40 bpm PED/NEO: 15 bpm ~ 60 bpm |
| Range of Ventricular Rhythm | |
| Ventricular Tachycardia | The interval of 5 consecutive ventricular wave is less than 600 ms |

| | | | |
|---|--|--------------|---------------|
| Ventricular Rhythm | The interval of 5 consecutive ventricular wave ranges from 600 ms to 1000 ms | | |
| Ventricular Bradycardia | The interval of 5 consecutive ventricular wave is more than 1000 ms | | |
| Startup time for Tachycardia | | | |
| Ventricular Tachycardia 1 mV 206bpm | Gain 1.0: 10 s Gain 0.5: 10 s Gain 2.0: 10 s | | |
| Ventricular Tachycardia 2 mV 195bpm | Gain 1.0: 10 s Gain 0.5: 10 s Gain 2.0: 10 s | | |
| Response time of Heart Rate Meter to Change in HR | HR range: 80 bpm ~ 120 bpm Range : 7s ~ 8s, average is 7.5s HR range: 80bpm ~ 40bpm Range : 7s ~ 8s, average is 7.5s | | |
| Tall T-wave Rejection | Exceeds ANSI/AAMI EC13-2002 Sect. 4.1.2.1 C) minimum recommended 1.2mV T-Wave amplitude | | |
| Accuracy of Heart Rate Meter and Response to Irregular Rhythm | Complies with ANSI/AAMI EC13-2002 Sect.4.1.2.1 e) The HR value displays after a stable period of 20s: Ventricular bigeminy: 80bpm±1bpm Slow alternating ventricular bigeminy: 60bpm±1bpm Rapid alternating ventricular bigeminy: 120bpm±1bpm Bidirectional systoles: 91bpm±1bpm | | |
| 16 different arrhythmia analyses | Non-Paced Patient | | Paced Patient |
| | ASYSTOLE | R on T | ASYSTOLE |
| | VFIB/VTAC | PVC | TACHY |
| | COUPLET | TACHY | BRADY |
| | VT>2 | BRADY | PNC |
| | BIGEMINY | MISSED BEATS | PNP |
| | TRIGEMINY | IRR | |
| | VENT | VBRADY | |

| ECG Analog Output | |
|--|--|
| Bandwidth (-3dB) | Diagnosis: 0.05Hz ~ 100Hz Monitor: 0.5Hz ~ 40Hz Surgery: 1Hz ~ 20Hz |
| Maximum transmission delay | 500ms (in diagnostic mode, and with notch off) |
| Sensitivity | 1V/mV $\pm 10\%$ |
| PACE rejection/enhancement | Without Pace enhancement or pace rejection |
| Defib Sync Pulse | |
| Output wave | Square pulse |
| Output impedance | 50 Ω |
| Delay from R-wave peak to start of pulse | 35ms |
| Amplitude | High level: 3.5 to 5 V, providing a maximum of 1 mA output current; Low level: < 0.5V, receiving a maximum of 5 mA input current. |
| Minimum required R wave amplitude | 0.3mV |
| Pulse width | 100ms $\pm 10\%$ |
| Limited current | 15 mA rating |
| Rising and falling time | < 1 ms |

A.8.2 12-lead Monitoring

| | |
|---------------------|--|
| Lead Mode | 3-Lead: I, II, III 5-Leads: I, II, III, aVR, aVL, aVF, V 12-Lead: I, II, III, aVR, aVL, aVF, V1, V2, V3, V4, V5, V6 |
| Waveform | 3-Lead: 1-channel waveform 5-Lead: 2-channel waveform, a maximum of seven waveforms; 12-Lead: 2-channel waveform, a maximum of 13 waveforms. |
| Lead naming style | AHA, IEC |
| Display Sensitivity | 1.25mm/mV ($\times 0.125$), 2.5mm/mV ($\times 0.25$), 5mm/mV ($\times 0.5$), 10mm/mV ($\times 1$), 20mm/mV ($\times 2$), AUTO gain |
| Bandwidth (-3dB) | Diagnosis: 0.05Hz to 150Hz |

| | |
|---|--|
| | Monitor: 0.5Hz to 40Hz Surgery: 1Hz to 20Hz |
| CMRR (Common Mode Rejection Ratio) | Diagnosis: >95dB (the Notch filter is off) Monitor: >105dB (the Notch filter is on) Surgery: >105dB (the Notch filter is on) |
| Notch | 50Hz/60Hz (Notch filter can be turned on or off manually) |
| Differential Input Impedance | >5M Ω |
| Input Signal Range | \pm 8mV PP |
| Accuracy of Input Signal Reconstruction | The total error and frequency response comply with ANSI/AAMI EC13:2002, Sect. 4.2.9.8. |
| Electrode Offset Potential Tolerance | \pm 500mV |
| Auxiliary Current (Leads off detection) | Active electrode: <100nA Reference electrode: <900nA |
| Input Offset Current | \leq 0.1 μ A |
| Recovery time after Defibrillation | <5s |
| Leakage current of patient | <10 μ A |
| Scale signal | 1mVPP, accuracy is \pm 5% |
| System noise | <30 μ VPP |
| ESU Protection | Incision mode: 300W Congelation mode: 100W Restore time: \leq 10s Meets the requirements of ANSI/AAMI EC13-2002: Sect. 4.1.2.1 a) |
| Noise Suppression of Electrotome | Tested according to the test method in EC13: 2002 Sect.5.2.9.14, it accords with the standard. |
| Pace Pulse | |
| Pulse indicator | PACE pulses are marked by the PACE indicator if the following conditions are met: Amplitude: \pm 2 mV ~ \pm 700 mV (12-lead) |

| | |
|------------------------------|--|
| | Width: 0.1 ms ~2 ms Ascending time: 10 μ s ~ 100 μ s |
| Pulse Rejection | Pulse is rejected if the requirements of ANSI/AAMI EC13-2002: Sect. 4.1.4.1 are met: Amplitude: ± 2 mV ~ ± 700 mV Width: 0.1 ms ~2 ms Ascending time: 10 μ s ~100 μ s |
| Minimum input slew rate | >2.5V/S |
| Heart rate | |
| Alarm Range | ADU: 15 bpm ~ 300 bpm PED/NEO: 15 bpm ~ 350 bpm |
| Accuracy | $\pm 1\%$ or 1 bpm, whichever is greater |
| Resolution | 1 bpm |
| Sensibility | ≥ 300 μ VPP |
| PVC | |
| Range | ADU: 0~300 PVCs/ min PED/NEO: 0~350 PVCs/ min |
| Resolution | 1 PVCs/min |
| ST value | |
| Range | -2.0 mV ~ +2.0 mV |
| Accuracy | The max. of ± 0.02 mV or 10% (-0.8 mV ~ +0.8 mV), whichever is greater. |
| Resolution | 0.01 mV |
| HR averaging method | |
| Method 1 | Normally, heart rate is computed by excluding the minimum and maximum values from the 12 most recent RR intervals and averaging the residual 10 RR intervals. |
| Method 2 | If each of three consecutive RR intervals is greater than 1200ms, then the four most recent RR intervals are averaged to compute the HR. |
| Range of Sinus and SV Rhythm | |
| Tachy | ADU: 120 bpm ~ 300 bpm PED/NEO: 160 bpm ~ 350 bpm |

| | | | |
|---|--|--------|---------------|
| Normal | ADU: 41 bpm ~ 119 bpm PED/NEO: 61 bpm ~159 bpm | | |
| Brady | ADU: 15 bpm ~ 40 bpm PED/NEO: 15 bpm ~ 60 bpm | | |
| Range of Ventricular Rhythm | | | |
| Ventricular Tachycardia | The interval of 5 consecutive ventricular wave is less than 600 ms | | |
| Ventricular Rhythm | The interval of 5 consecutive ventricular wave ranges from 600 ms to 1000 ms | | |
| Ventricular Bradycardia | The interval of 5 consecutive ventricular wave is more than 1000 ms | | |
| Startup time for Tachycardia | | | |
| Ventricular Tachycardia 1 mV 206bpm | Gain 1.0: 10 s Gain 0.5: 10 s Gain 2.0: 10 s | | |
| Ventricular Tachycardia 2 mV 195bpm | Gain 1.0: 10 s Gain 0.5: 10 s Gain 2.0: 10 s | | |
| Response time of Heart Rate Meter to Change in HR | HR range: 80 bpm ~ 120 bpm Range : 7s ~ 8s, average is 7.5s HR range: 80bpm ~ 40bpm Range : 7s ~ 8s, average is 7.5s | | |
| Tall T-wave Rejection | Exceeds ANSI/AAMI EC13-2002 Sect. 4.1.2.1 C) minimum recommended 1.2mV T-Wave amplitude | | |
| Accuracy of Heart Rate Meter and Response to Irregular Rhythm | Complies with ANSI/AAMI EC13-2002 Sect.4.1.2.1 e) The HR value displays after a stable period of 20s: According with ANSI/AAMI EC13-2002 Sect.4.1.2.1 e) Ventricular bigeminy: 80bpm±1bpm Slow alternating ventricular bigeminy: 60bpm±1bpm Rapid alternating ventricular bigeminy: 120bpm±1bpm Bidirectional systoles: 91bpm±1bpm | | |
| 16 different arrhythmia analyses | Non-Paced Patient | | Paced Patient |
| | ASYSTOLE | R on T | ASYSTOLE |
| | VFIB/VTAC | PVC | TACHY |

| | | | |
|--|---|--------------|-------|
| | COUPLET | TACHY | BRADY |
| | VT>2 | BRADY | PNC |
| | BIGEMINY | MISSED BEATS | PNP |
| | TRIGEMINY | IRR | |
| | VENT | VBRADY | |
| 12-lead ECG Synchronization Analysis | Average parameters of heart beat | | |
| | Heart rate (bpm) | | |
| | Time limit of P wave (ms) | | |
| | PR interval (ms) | | |
| | QRS interval (ms) | | |
| | QT/QTc (ms) | | |
| | P-QRS-T AXIS | | |
| ECG Analog Output | | | |
| Bandwidth (-3dB) | Diagnosis: 0.05Hz ~ 100Hz | | |
| | Monitor: 0.5Hz ~ 40Hz | | |
| | Surgery: 1Hz ~ 20Hz | | |
| Max transmission delay | 500ms (in diagnostic mode, and with notch off) | | |
| Sensitivity | 1V/mV ±10% | | |
| PACE rejection/enhancement | Without Pace enhancement or pace rejection | | |
| Defib Sync Pulse | | | |
| Output wave | Square pulse | | |
| Output impedance | 50Ω | | |
| Delay from R-wave peak to start of pulse | 35ms | | |
| Amplitude | High level: 3.5 to 5 V, providing a maximum of 1 mA output current; | | |
| | Low level: < 0.5V, receiving a maximum of 5 mA input current. | | |
| Minimum required R wave amplitude | 0.3mV | | |
| Pulse width | 100ms ± 10% | | |
| Limited current | 15 mA rating | | |
| Rising and falling time | < 1 ms | | |

A.9 RESP

| | |
|---------------------------------|---|
| Method | Trans-thoracic impedance: R-F(RA-LL), R-L (RA-LA) |
| Respiration excitation waveform | < 300 μ A, sinusoid, 62.8 kHz (\pm 10%) |
| Measuring sensitivity | 0.3 Ω (base impedance 200 to 4500 Ω) |
| Base impedance range | 200 to 2500 Ω (cable resistance = 0 K) |
| | 2200 to 4500 Ω (leads cables 1K Ω resistance) |
| Differential input impedance | > 2.5M Ω |
| Waveform bandwidth | 0.2 to 2.5 Hz (-3 dB) |
| RR measuring range: | |
| Adult | 0 to 120 rpm |
| Neo/Ped | 0 to 150 rpm |
| Resolution | 1 rpm |
| Accuracy | \pm 2 rpm |
| Apnea Alarm Time Setup | 10s, 15s, 20s, 25s, 30s, 35s, 40s |

A.10 NIBP

A.10.1 NIBP from EDAN Module

| | |
|---------------------------------|---|
| Method | Oscillometric |
| Mode | Manual, Auto, Continuous |
| Measuring interval in AUTO Mode | 1/2/3/4/5/10/15/30/60/90/120/240/480 min |
| Continuous | 5min, interval is 5s |
| Measuring type | SYS, DIA, MAP |
| Measuring Range | |
| Adult mode | SYS: 40 mmHg ~ 270 mmHg DIA: 10 mmHg ~ 215 mmHg MAP: 20 mmHg ~ 235 mmHg |
| Pediatric mode | SYS: 40 mmHg ~ 200 mmHg DIA: 10 mmHg ~ 150 mmHg MAP: 20 mmHg ~ 165 mmHg |
| Neonatal mode | SYS: 40 mmHg ~ 135 mmHg DIA: 10 mmHg ~ 100 mmHg |

| | |
|--|---|
| | MAP: 20 mmHg ~ 110 mmHg |
| Cuff pressure measuring range | 0 mmHg ~ 300 mmHg |
| Pressure resolution | 1mmHg |
| Maximum mean error | ±5mmHg |
| Maximum standard deviation | 8mmHg |
| Maximum measuring period | |
| Adult/Pediatric | 120s |
| Neonate | 90s |
| Typical measuring period | 30s ~ 45s (depend on HR/motion disturbance) |
| Overpressure protection (Dual overpressure protection) | |
| Adult | 297±3mmHg |
| Pediatric | 240±3mmHg |
| Neonatal | 147±3mmHg |
| PR | |
| Measuring range | 40 bpm ~240bpm |
| Accuracy | ±3bpm or 3.5%, whichever is greater |

A.10.2 NIBP from M3600 Module

| | |
|---------------------------------|---------------------------------------|
| Method | Oscillometric |
| Mode | Manual, Auto, Continuous |
| Measuring Interval in AUTO Mode | 1/2/3/4/5/10/15/30/60/90 min, 2/4/8h |
| PR Range | Adult/ Pediatric mode: 40bpm ~ 200bpm |
| | Neonatal mode: 40 bpm ~ 240bpm |
| PR Accuracy | ± 2 bpm or 2% of the readings |
| Measuring Type | SYS, DIA, MAP |
| Measuring Range | |
| Adult/ Pediatric Mode | SYS: 60 mmHg ~ 250 mmHg |
| | DIA: 40 mmHg ~ 200 mmHg |
| | MAP: 45 mmHg ~ 235 mmHg |
| Neonatal Mode | SYS: 40 mmHg ~ 120 mmHg |
| | DIA: 20 mmHg ~ 90 mmHg |
| | MAP: 30 mmHg ~ 100 mmHg |

| | | | | | |
|-------------------------------|------------------|------------------------|------------------|------------------------|--|
| Alarm Type | SYS, DIA, MAP | | | | |
| Pressure Resolution | 1mmHg | | | | |
| Measuring Accuracy | | | | | |
| Maximum Mean Error | ±5mmHg | | | | |
| Maximum Standard Deviation | 8mmHg | | | | |
| | Adult/Pediatric | | | Neonate | |
| | Normal Condition | Single Fault Condition | Normal Condition | Single fault Condition | |
| Cuff Pressure Measuring Range | 300 mmHg | 330 mmHg | 150 mmHg | 165 mmHg | |
| Maximum Measuring Period | Less than 160s | Less than 180s | Less than 80s | Less than 90s | |

A.11 SpO₂

A.11.1 SpO₂ from EDAN Module

| | |
|-----------------------------|-------------------------------------|
| Measuring Range | 0 ~ 100 % |
| Resolution | 1 % |
| Accuracy | |
| Adult (including Pediatric) | ±2 % (70%~100% SpO ₂) |
| | Undefined (0~69% SpO ₂) |
| Neonate | ±3 % (70%~100% SpO ₂) |
| | Undefined (0~69% SpO ₂) |
| Pulse Rate | |
| Measuring Range | 25bpm ~ 300bpm |
| Resolution | 1bpm |
| Accuracy | ±2bpm |
| Data update period | 1s |
| Wave length | |
| Red light | 660±3 nm |
| Infrared light | 905±5 nm |
| Emitted light energy | ≤15 mW |

A.11.2 SpO₂ from Nellcor Module

| | | |
|---|--|--|
| Measuring Range | | 1% ~ 100% |
| Resolution | | 1% |
| Accuracy | Sensor Type | Accuracy |
| | MAX-A, MAX-AL, MAX-N, MAX-P, MAX-I, MAX-FAST | ± 2 (70% ~ 100% SpO ₂) |
| | OxiCliq A, OxiCliq P, OxiCliq N (Adult), OxiCliq N (Neonate), OxiCliq I | ± 2.5 (70% ~ 100% SpO ₂) |
| | D-YS (Infant to Adult), DS-100A, OXI-A/N, OXI-P/I | ± 3(70% ~ 100% SpO ₂) |
| | D-YS (including D-YSE ear clip), D-YS (including D-YSPD spotclip) | ± 3.5(70% ~ 100% SpO ₂) |
| * When the sensor is used to neotate as recommendation, the specified accuracy range of the neotate is always higher ±1 than adult. | | |
| Pulse Rate | | |
| Measuring Range | | 20bpm ~ 300bpm |
| Resolution | | 1bpm |
| Accuracy | | ± 3bpm (20bpm ~ 250bpm) |
| Sensor | | Wave length: approximately 660 and 900nm |
| | | Emitted light energy: ≤15mW |

A.12 TEMP

| | |
|---------------------------|--------------------------------|
| Channel | 2 |
| Sensor type | YSI-10K and YSI-2.252K |
| Measuring Range | 0 °C ~ 50 °C(32 ° F ~ 122 ° F) |
| Resolution | 0.1°C (0.1 ° F) |
| Accuracy (Without sensor) | ±0.1°C or ±0.2 ° F |
| Refresh Time | Every 1s ~ 2s |

A.13 Quick TEMP

| | |
|-----------------|--|
| Measuring Range | 25°C ~ 45°C (77 ° F ~113 ° F) |
| Operating Temp | 10°C ~ 40°C (50 ° F ~104 ° F) |
| Sensor Type | Oral/Axillary sensor, Rectal sensor |
| Resolution | 0.1°C (0.1 ° F) |
| Accuracy | ±0.1°C (25°C ~ 45°C) or ± 0.2°F (77 ° F ~113 ° F) |
| Response time | < 60s |
| Update time | 1s ~ 2s |

A.14 IBP

| | |
|----------------------------|--|
| Pressure measuring range | -50 to +300 mmHg |
| Resolution | 1 mmHg |
| Accuracy (without sensor) | ± 2 % or ±1 mmHg, whichever is greater |
| Pressure sensor | |
| Sensitivity | 5 (μV/V/mmHg) |
| Impedance | 300 to 3000 Ω |
| Frequency response | d.c. to 12.5 or 40 Hz |
| Zero | Range: ±200 mmHg |
| | Accuracy: 1 mmHg |
| Measuring range | |
| Art | 0 mmHg to 300 mmHg |
| PA | -6 to +120mmHg |
| CVP/RAP/LAP/ICP | -10 to +40 mmHg |
| P1/P2 | -50 to +300 mmHg |
| Volume displacement of MSI | $4.5 \times 10^{-4} \text{ in}^3 / 100 \text{ mmHg}$ |

A.15 CO₂

| | |
|-------------------|--------------------------------|
| Method | Infra-red Absorption Technique |
| Unit | mmHg, %, Kpa |
| Measuring Range | |
| EtCO ₂ | 0 mmHg ~ 150 mmHg |
| FiCO ₂ | 3 mmHg ~50 mmHg |

| | |
|----------------------------------|---|
| AwRR | 0 rpm ~ 150 rpm |
| Resolution | |
| EtCO ₂ | 1mmHg |
| FiCO ₂ | 1mmHg |
| AwRR | 1 rpm |
| EtCO ₂ Accuracy | ± 2 mmHg, 0 to 40 mmHg |
| | ± 5 % of reading, 41 to 70 mmHg |
| | ± 8 % of reading, 71 to 100 mmHg |
| | ± 10 % of reading, 101 to 150 mmHg |
| AwRR Accuracy | ± 1 rpm |
| Suffocation Alarm Delay | 10s, 15s, 20s, 25s, 30s, 35s, 40s, default value is 20 second. |
| Sample Gas Flowrate | 50ml/min |
| Stability | |
| Short Term Drift | Drift over 4 hours < 0.8 mmHg |
| Long Term Drift | 120 hour period |
| O ₂ Compensation | |
| Range | 0 ~ 100% |
| Resolution | 1% |
| Default | 16% |
| Initialization time | It displays the value within 15s and meets the requirement for measurement accuracy within 2min. (Mainstream) |
| | It displays the value within 20s and meets the requirement for measurement accuracy within 2min. (Sidestream) |
| Response time | 60ms (Mainstream) |
| Calibration | Not required. |
| Barometric pressure compensation | User setup |
| Apnea Alarm Delay | 10s, 15s, 20s, 25s, 30s, 35s, 40s, 45s; default value is 20s. |

Interfering Gas and Vapor Effect on EtCO₂ Measurement Values:

| Gas or vapor | Gas level (%) | Quantitative effect/Comments |
|---------------|---------------|--|
| Nitrous oxide | 60 | Dry and Saturated Gas |
| Halothane | 4 | 0 – 40 mmHg: ± 1 mmHg additional error |

| | | |
|-------------|----|---|
| Enflurane | 5 | 41 – 70 mmHg: $\pm 2.5\%$ additional error |
| Isoflurane | 5 | 71 – 100 mmHg: $\pm 4\%$ additional error |
| Sevoflurane | 5 | 101 – 150 mmHg: $\pm 5\%$ additional error |
| Xenon | 80 | <p>*Additional worst case error when compensation for P_B, O_2, N_2O, anesthetic agents, or helium is correctly selected for the actual fractional gas constituents present.</p> <p>Desflurane:</p> <p>The presence of desflurane in the exhaled breath at concentrations greater than 5% will positively bias Carbon Dioxide values by up to an additional 3 mmHg at 38mmHg.</p> <p>Xenon:</p> <p>The presence of Xenon in the exhaled breath will negatively bias Carbon Dioxide values by up to an additional 5 mmHg at 38mmHg.</p> |
| Helium | 50 | |
| Desflurane | 15 | |

Barometric Pressure on EtCO₂ Measurement Values:

| |
|--|
| Quantitative effect |
| Ambient Barometric, Operational |
| 0 – 40 mmHg: ± 1 mmHg additional error |
| 41 – 70 mmHg: $\pm 2.5\%$ additional error |
| 71 – 100 mmHg: $\pm 4\%$ additional error |
| 101 – 150 mmHg: $\pm 5\%$ additional error |
| *Additional worst case error when compensation for P_B , O_2 , N_2O , anesthetic agents, or helium is correctly selected for the actual fractional gas constituents present. |

A.16 C.O.

| Method | Thermodilution Technique |
|-----------------|--------------------------------------|
| Measuring range | |
| C.O. | 0.1 L/min ~ 20L/min |
| TB | 23°C ~ 43°C(73.4 ° F ~109.4 ° F) |
| TI | -1°C ~ 27°C(30.2 ° F ~80.6 ° F) |
| Resolution | |
| C.O. | 0.1L/min |
| TB, TI | +0.1°C (+0.1 ° F) |
| Accuracy | |
| C.O. | For C.O. ≥ 4.0 L/min: $\pm 5\%$ |

| | |
|----|---------------------------------|
| | For C.O. < 4.0 L/min: 0.2 L/min |
| TB | $\pm 0.1^{\circ}\text{C}$ |
| TI | $\pm 0.1^{\circ}\text{C}$ |

NOTE:

At least 90% of the C.O. data should reside inside the bounded region, and the lower 95% confidence interval should not exceed 85%.

A.17 AG**A.17.1 Phasein Sidestream**

| | | |
|-------------------------------------|---|--|
| Module Type | ISA AX+ Analyzer | Displaying the concentration of CO ₂ , N ₂ O, and two anaesthesia agent and identifying the anaesthesia agent automatically (portable module) |
| | ISA OR+ Analyzer | Displaying the concentration of CO ₂ , O ₂ , N ₂ O, and two anaesthesia agent and identifying the anaesthesia agent automatically (portable module) |
| Measurement Parameters | CO ₂ , N ₂ O , O ₂ , Halothane (HAL), Isoflurane(ISO), Enflurane(ENF), Sevoflurane(SEV) , Desflurane(DES), awRR, MAC | |
| Measurement Principle | CO ₂ , N ₂ O, Anaesthesia Agent: infra-red absorption characteristic; O ₂ : Paramagnetic method | |
| Sampling Flow Rate | 50 ml/min | |
| Work Mode | Measurement, Standby | |
| Warm-up Time | < 20s | |
| Typical Rise Time | CO ₂ ≤ 200ms O ₂ ≤ 350ms N ₂ O ≤ 350ms O ₂ ≤ 450ms | |
| Primary Anaesthesia Agent Threshold | ≤ 0.15 vol% | |
| Second Anaesthesia Agent Threshold | 0.2 vol% + 10% | |
| Agent Identificaiton Time | < 20 seconds (typically < 10 seconds) | |

| | | | | | |
|-----------------------------------|----------------------------|------------------------------|--------------------------|-----------------|------------------|
| Response Time | < 3 seconds | | | | |
| Accuracy(Standard Conditions) | | | | | |
| GAS | Measurement Range | Accuracy | | | |
| CO ₂ | 0 to 15 vol% | ±(0.2 vol% + 2% of reading) | | | |
| | 15 to 25 vol% | Unspecified | | | |
| N ₂ O | 0 to 100 vol% | ±(2 vol% + 2% of reading) | | | |
| HAL, ENF, ISO | 0 to 8 vol % | ±(0.15 vol% + 5% of reading) | | | |
| | 8 to 25 vol % | Unspecified | | | |
| SEV | 0 to 10 vol % | ±(0.15 vol% + 5% of reading) | | | |
| | 10 to 25 vol % | Unspecified | | | |
| DES | 0 to 22 vol % | ±(0.15 vol% + 5% of reading) | | | |
| | 22 to 25 vol % | Unspecified | | | |
| O ₂ | 0 to 100 vol % | ±(1 vol% + 2% of reading) | | | |
| Accuracy (All Conditions) | | | | | |
| Gas | Accuracy | | | | |
| CO ₂ | ±(0.3kPa + 4% of reading) | | | | |
| N ₂ O | ±(2kPa + 5% of reading) | | | | |
| Agents | ±(0.2kPa + 10% of reading) | | | | |
| O ₂ | ±(2kPa + 2 of reading) | | | | |
| Apnea Alarm Delay | 20s~60s | | | | |
| Item | Operating conditions | | Storage conditions | | |
| Temperature(°C) | 5 to 50 | | -40 ~ 70 | | |
| Relative humidity | <95% RH(non-condensing) | | 5 ~ 100% RH(Condensing) | | |
| Atmospheric pressure | 525 ~ 1200hPa | | 200 ~ 1200hPa | | |
| Interfering gas and vapor effects | | | | | |
| Gas or vapour | Gas level | CO ₂ | | Agents | N ₂ O |
| | | ISA CO ₂ | ISA AX+ | | |
| N ₂ O ⁴⁾ | 60 vol% | - ²⁾ | - ¹⁾ | - ¹⁾ | - ¹⁾ |
| HAL ⁴⁾ | 4 vol% | - ¹⁾ | - ¹⁾ | - ¹⁾ | - ¹⁾ |
| ENF, ISO, SEV ⁴⁾ | 5 vol% | +8% of | - ¹⁾ | - ¹⁾ | - ¹⁾ |

| | | | | | |
|--|---|-------------------------------|-----------------|-----------------|-----------------|
| | | reading ³⁾ | | | |
| DES ⁴⁾ | 15 vol% | +12% of reading ³⁾ | - ¹⁾ | - ¹⁾ | - ¹⁾ |
| Xe(Xenon) ⁴⁾ | 80 vol% | -10% of reading ³⁾ | | - ¹⁾ | - ¹⁾ |
| He(Helium) ⁴⁾ | 50 vol% | -6% of reading ³⁾ | | - ¹⁾ | - ¹⁾ |
| Metered dose inhaler propellants ⁴⁾ | Not for use with metered dose inhaler propellants | | | | |
| C ₂ H ₅ OH(Ethanol) ⁴⁾ | 0.3 vol% | - ¹⁾ | - ¹⁾ | - ¹⁾ | - ¹⁾ |
| C ₃ H ₇ OH (Isopropanol) ⁴⁾ | 0.5 vol% | - ¹⁾ | - ¹⁾ | - ¹⁾ | - ¹⁾ |
| CH ₃ COCH ₃ (Acetone) ⁴⁾ | 1 vol% | - ¹⁾ | - ¹⁾ | - ¹⁾ | - ¹⁾ |
| CH ₄ (Methane) ⁴⁾ | 3 vol% | - ¹⁾ | - ¹⁾ | - ¹⁾ | - ¹⁾ |
| CO(Carbon monoxide) ⁵⁾ | 1 vol% | - ¹⁾ | - ¹⁾ | - ¹⁾ | - ¹⁾ |
| NO(Nitrogen monoxide) | 0.02 vol% | - ¹⁾ | - ¹⁾ | - ¹⁾ | - ¹⁾ |
| O ₂ ⁵⁾ | 100 vol% | - ²⁾ | - ²⁾ | - ¹⁾ | - ¹⁾ |

Note 1: Negligible interference, effect included in the specification “Accuracy, all conditions” above.

Note 2: Negligible interference with N₂O / O₂ concentrations correctly set, effect included in the specification “Accuracy, all conditions” above.

Note 3: Interference at indicated gas level. For example, 50 vol% Helium typically decreases the CO₂ readings by 6%. This means that if measuring on a mixture containing 5.0 vol% CO₂ and 50 vol% Helium, the actual measured CO₂ concentration will typically be (1-0.06)*5.0 vol% =4.7 vol% CO₂.

Note 4: According to the EN ISO 21647 standard.

Note 5: In addition to the EN ISO 21647 standard.

A.17.2 Phasein Mainstream

| | | |
|-------------|--|---|
| Module Type | IRMA AX+ | Displaying the concentration of CO ₂ , N ₂ O and two anaesthesia agent and indentifying two anaesthesia agent |
| Measurement | CO ₂ , N ₂ O, O ₂ , HAL, Isoflurane(ISO), Enflurane(ENF), | |

| | | |
|-------------------------------|--|------------------------------|
| Parameters | Sevoflurane(SEV), Desflurane(DES), awRR, MAC | |
| Measurement Principle | CO ₂ , N ₂ O, anaesthesia agent: infra-red absorption characteristic | |
| Warm-up Time | Concentrations are reported and the automatic agent identification is running within 10 seconds. 20 seconds for IRMA AX+. | |
| Rise Time | CO ₂ ≤ 90ms O ₂ ≤ 300ms N ₂ O ≤ 300ms HAL, ISO, ENF, SEV, DES ≤ 300ms | |
| Primary Agent Threshold | 0.15 vol% | |
| Secondary Agent Threshold | 0.2 vol% + 10% of total agent concentration | |
| Agent Identificaiton Time | < 20 seconds (typically less than 10 seconds) | |
| Response Time | < 1 second | |
| Accuracy(Standard Conditions) | | |
| Gas | Range | Accuracy |
| CO ₂ | 0 ~ 10 vol% | ±(0.2 vol% + 2% of reading) |
| | 10 ~ 15vol% | ±(0.3 vol% + 2% of reading) |
| | 15 ~ 25 vol% | Unspecified |
| N ₂ O | 0 to 100% | ±(2 vol% + 2% of reading) |
| HAL | 0 to 8 | ±(0.15 vol% + 5% of reading) |
| ISO | 8 to 25 | Unspecified |
| ENF | | |
| SEV | 0 to 10 | ±(0.15 vol% + 5% of reading) |
| | 10 to 25 | Unspecified |
| DES | 0 to 22 | ±(0.15 vol% + 5% of reading) |
| | 22 to 25 | Unspecified |
| Accuracy (All Conditions) | | |
| GAS | Accuracy | |

| CO ₂ | ±(0.3 vol% + 4 % of reading) | | | | |
|--|---|-------------------------------|-----------------------|-----------------|------------------|
| N ₂ O | ±(2 vol% + 4 % of reading) | | | | |
| Agents | ±(0.2 vol% + 10 % of reading) | | | | |
| Apnea Alarm Delay | 20s ~ 60s | | | | |
| Temperature(°C) | 10 to 40 | -20 ~ 75 | | | |
| Relative humidity | 10 ~ 95% RH(non-condensing) | 5 ~ 100% RH(condensing) | | | |
| Atmospheric pressure | 525 ~ 1200hPa | 500 ~ 1200hPa | | | |
| Interfering gas and vapour effects | | | | | |
| Gas or vapour | Gas level | CO ₂ | | Agents | N ₂ O |
| | | IRMA CO ₂ | IRMA AX+ | | |
| N ₂ O ⁴⁾ | 60 vol% | - ^{1&2)} | - ^{1&2)} | - ¹⁾ | - ¹⁾ |
| HAL ⁴⁾ | 4 vol% | - ¹⁾ | - ¹⁾ | - ¹⁾ | - ¹⁾ |
| ENF, ISO, SEV ⁴⁾ | 5 vol% | +8% of reading ³⁾ | - ¹⁾ | - ¹⁾ | - ¹⁾ |
| DES ⁴⁾ | 15 vol% | +12% of reading ³⁾ | - ¹⁾ | - ¹⁾ | - ¹⁾ |
| Xe(Xenon) ⁴⁾ | 80 vol% | -10% of reading ³⁾ | | - ¹⁾ | - ¹⁾ |
| He(Helium) ⁴⁾ | 50 vol% | -6% of reading ³⁾ | | - ¹⁾ | - ¹⁾ |
| Metered dose inhaler propellants ⁴⁾ | Not for use with metered dose inhaler propellants | | | | |
| C ₂ H ₅ OH(Ethanol) ⁴⁾ | 0.3 vol% | - ¹⁾ | - ¹⁾ | - ¹⁾ | - ¹⁾ |
| C ₃ H ₇ OH (Isopropanol) ⁴⁾ | 0.5 vol% | - ¹⁾ | - ¹⁾ | - ¹⁾ | - ¹⁾ |
| CH ₃ COCH ₃ (Acetone) ⁴⁾ | 1 vol% | - ¹⁾ | - ¹⁾ | - ¹⁾ | - ¹⁾ |
| CH ₄ (Methane) ⁴⁾ | 3 vol% | - ¹⁾ | - ¹⁾ | - ¹⁾ | - ¹⁾ |
| CO(Carbon monoxide) ⁵⁾ | 1 vol% | - ¹⁾ | - ¹⁾ | - ¹⁾ | - ¹⁾ |

| | | | | | |
|------------------------------|----------|-------|-------|-----|-----|
| O ₂ ⁵⁾ | 100 vol% | -1&2) | -1&2) | -1) | -1) |
|------------------------------|----------|-------|-------|-----|-----|

Note 1: Negligible interference, effect included in the specification “Accuracy, all conditions” above.

Note 2: For probes not measuring N₂O and/or O₂ the concentrations shall be set from monitor. (IRMA CO₂ measures neither N₂O, nor O₂. IRMA AX+ does not measure O₂.)

Note 3: Interference at indicated gas level. For example, 50 vol% Helium typically decreases the CO₂ readings by 6%. This means that if measuring on a mixture containing 5.0 vol% CO₂ and 50 vol% Helium, the measured CO₂ concentration will typically be $(1-0.06)*5.0 \text{ vol\%} = 4.7 \text{ vol\% CO}_2$.

Note 4: According to the EN ISO 21647 standard.

Note 5: In addition to the EN ISO 21647 standard.

A.18 Wireless Network

| | |
|---------------------------------------|---|
| Compliant with Standard and Directive | IEEE802.11b/g, R&TTE Directive (99/5/EEC) |
| Frequency Range | 2.412 GHz ~2.462 GHz (America) 2.412 GHz ~2.484 GHz (Japan) 2.412 GHz ~2.472 GHz (ETSI) |
| Working frequency segment | Ch1 ~ 11 (America) Ch1 ~ 14 (Japan) Ch1 ~ 13 (ETSI) |

B EMC Information

- Guidance and Manufacture's Declaration

B.1 Electromagnetic Emissions - for all EQUIPMENT and SYSTEMS

| Guidance and manufacture's declaration – electromagnetic emission | | |
|---|-------------------|---|
| The monitor is intended for use in the electromagnetic environment specified below. The customer or the user of the monitor should assure that it is used in such an environment. | | |
| Emission test | Compliance | Electromagnetic environment – guidance |
| RF emissions CISPR 11 | Group 1 | The monitor uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment. |
| RF emission CISPR 11 | Class A | The monitor 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. |
| Harmonic emissions IEC/EN 61000-3-2 | Class A | |
| Voltage fluctuations/ flicker emissions IEC/EN 61000-3-3 | Complies | |

B.2 Electromagnetic Immunity - for all EQUIPMENT and SYSTEMS

| Guidance and manufacture's declaration – electromagnetic immunity | | | |
|---|--------------------------------|-----------------------------|--|
| The monitor is intended for use in the electromagnetic environment specified below. The customer or the user of the monitor should assure that it is used in such an environment. | | | |
| Immunity test | IEC/EN 60601 test level | Compliance level | Electromagnetic environment - guidance |
| Electrostatic discharge (ESD) IEC/EN 61000-4-2 | ±6 kV contact ±8 kV air | ±6 kV contact ±8 kV air | Floors should be wood, concrete or ceramic tile. If floor are covered with synthetic material, the relative humidity should be at least 30%. |
| Electrical fast transient/burst | ±2 kV for power supply lines | ±2kV for power supply lines | Mains power quality should be that of a |


| | | | |
|---|---|---|--|
| IEC/EN 61000-4-4 | ± 1 kV for input /output signal | ± 1 kV for input /output signal | typical commercial or hospital environment. |
| Surge IEC/EN 61000-4-5 | ± 1 kV for line to line ± 2 kV for line to ground | ± 1 kV for line to line ± 2 kV for line to ground | Mains power quality should be that of a typical commercial or hospital environment. |
| Power frequency (50/60Hz) magnetic field IEC/EN 61000-4-8 | 3A/m | 3A/m | Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment. |
| Voltage dips, short interruptions and voltage variations on power supply input lines IEC/EN 61000-4-11 | $<5\%$ U_T ($>95\%$ dip in U_T) for 0.5 cycle 40% U_T (60% dip in U_T) for 5 cycles 70% U_T (30% dip in U_T) for 25 cycles $<5\%$ U_T ($>95\%$ dip in U_T) for 5 sec | $<5\%$ U_T ($>95\%$ dip in U_T) for 0.5 cycle 40% U_T (60% dip in U_T) for 5 cycles 70% U_T (30% dip in U_T) for 25 cycles $<5\%$ U_T ($>95\%$ dip in U_T) for 5 sec | Mains power quality should be that of a typical commercial or hospital environment. If the user of the Patient Monitor requires continued operation during power mains interruptions, it is recommended that the Patient Monitor be powered from an uninterruptible power supply or a battery. |

NOTE U_T is the a.c. mains voltage prior to application of the test level.

B.3 Electromagnetic Immunity - for EQUIPMENT and SYSTEMS that are not LIFE-SUPPORTING

Guidance and manufacture's declaration – electromagnetic immunity

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

| environment. | | | |
|--|---|---------------------------------------|---|
| Immunity test | IEC/EN 60601 test level | Compliance level | Electromagnetic environment - guidance |
| <p>Conducted RF IEC/EN 61000-4-6</p> <p>Radiated RF IEC/EN 61000-4-3</p> | <p>3 V_{rms} 150 kHz to 80 MHz</p> <p>3 V/m 80 MHz to 2.5 GHz</p> | <p>3 V_{rms}</p> <p>3 V/m</p> | <p>Portable and mobile RF communications equipment should be used no closer to any part of the Patient Monitor, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter.</p> <p>Recommended separation distance</p> $d = \left[\frac{3.5}{V_1} \right] \sqrt{P}$ $d = \left[\frac{3.5}{E_1} \right] \sqrt{P} \quad 80 \text{ MHz to } 800 \text{ MHz}$ $d = \left[\frac{7}{E_1} \right] \sqrt{P} \quad 800 \text{ MHz to } 2.5 \text{ GHz}$ <p>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 metres (m).</p> <p>Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey,^a should be less than the compliance level in each frequency range.^b</p> <p>Interference may occur in the vicinity of equipment marked with the following symbol:</p>  |

NOTE 1 At 80 MHz and 800 MHz, the higher frequency range applies.

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

^a Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless)

telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the Patient Monitor is used exceeds the applicable RF compliance level above, the Patient Monitor should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as reorienting or relocating the Patient Monitor.

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

B.4 Recommended Separation Distances

| Recommended separation distances between portable and mobile RF communications equipment and the monitor | | | |
|---|---|---|---|
| The monitor is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the monitor can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the monitor as recommended below, according to the maximum output power of the communications equipment. | | | |
| Rated maximum output power of transmitter (W) | Separation distance according to frequency of transmitter(m) | | |
| | 150 kHz to 80 MHz | 80 MHz to 800 MHz | 800 MHz to 2.5 GHz |
| | $d = \left[\frac{3.5}{V_1} \right] \sqrt{P}$ | $d = \left[\frac{3.5}{E_1} \right] \sqrt{P}$ | $d = \left[\frac{7}{E_1} \right] \sqrt{P}$ |
| 0.01 | 0.12 | 0.12 | 0.23 |
| 0.1 | 0.36 | 0.37 | 0.74 |
| 1 | 1.16 | 1.17 | 2.33 |
| 10 | 3.69 | 3.69 | 7.38 |
| 100 | 11.67 | 11.67 | 23.33 |
| For transmitters rated at a maximum output power not listed above, the recommended separation distance d in metres (m) can be estimated using the equation applicable to the frequency of the transmitter, where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer. | | | |
| NOTE 1 At 80 MHz and 800 MHz, the separation distance for the higher frequency range applies. | | | |
| NOTE 2 These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people. | | | |

C Default Settings

This appendix documents the most important default settings of your monitor as it is delivered from the factory.

Note: If your monitor has been ordered preconfigured to your requirements, the settings at delivery will be different from those listed here.

C.1 Patient Information Default Settings

| Patient Information Settings | |
|------------------------------|-------|
| Patient Type | Adult |
| Pace | Off |

C.2 Alarm Default Settings

| | |
|------------------|---------|
| Alarm Settings | |
| Pause Time | 120s |
| Mute | On |
| Sensor Off Alarm | On |
| Alarm Latch | Unlatch |

C.3 ECG Default Settings

| ECG Settings | ADU | PED | NEO |
|------------------|---------|-----|-----|
| Alarm Switch | On | | |
| Alarm Record | Off | | |
| Alarm Level | Medium | | |
| Alarm High Limit | 120 | 160 | 200 |
| Alarm Low Limit | 50 | 75 | 100 |
| Pace | Off | | |
| Lead Type | 5 Leads | | |
| Display | Normal | | |
| Filter | Monitor | | |
| Smart Lead Off | Off | | |
| Heart Volume | 2 | | |
| ST Analysis | ADU | PED | NEO |
| ST Analysis | Off | | |
| Alarm Switch | Off | | |

| | | | |
|--|--------------|-------------|--------------|
| Alarm Level | Medium | | |
| Alarm Record | Off | | |
| Alarm High Limit (ST-X) | 0.2 | | |
| Alarm Low Limit (ST-X) | -0.2 | | |
| X stands for I, II, III, aVR, aVL, aVF, V, V1, V2, V3, V4, V5 or V6. | | | |
| ARR Analysis | | | |
| ARR Analysis | On | | |
| PVCs Alarm Level | Medium | | |
| PVCs Alarm Switch | Off | | |
| PVCs Alarm Record | Off | | |
| ARR Alarm Settings | Alarm Switch | Alarm Level | Alarm Record |
| ASYSTOLE | On | High | Off |
| VFIB/VTAC | On | High | Off |
| R ON T | On | Medium | Off |
| VT > 2 | On | Medium | Off |
| COUPLET | On | Medium | Off |
| PVC | On | Medium | Off |
| BIGEMINY | On | Medium | Off |
| TRIGEMINY | On | Medium | Off |
| TACHY | On | Medium | Off |
| BRADY | On | Medium | Off |
| MISSEDBEATS | On | Medium | Off |
| IRR | On | Medium | Off |
| PNC | On | Medium | Off |
| PNP | On | Medium | Off |
| VBRADY | On | Medium | Off |
| VENT | On | Medium | Off |

C.4 RESP

| | | | |
|---------------|-----|-----|-----|
| RESP Settings | ADU | PED | NEO |
| Alarm Switch | On | | |

| | | | |
|------------------|----------|----|-----|
| Alarm Record | Off | | |
| Alarm Level | Medium | | |
| Alarm High Limit | 30 | 30 | 100 |
| Alarm Low Limit | 8 | 8 | 30 |
| Apnea Time | 20s | | |
| Calculation Type | Auto | | |
| Resp Type | II | | |
| Sweep | 12.5mm/s | | |
| Amplitude | 1 | | |

C.5 SpO₂

| | | | |
|---------------------------|----------|-----|-----|
| SpO ₂ Settings | ADU | PED | NEO |
| Alarm Switch | On | | |
| Alarm Record | Off | | |
| Alarm Level | Medium | | |
| Alarm High Limit | 100 | 100 | 95 |
| Alarm Low Limit | 90 | 90 | 88 |
| Pitch Tone | On | | |
| Sweep | 12.5mm/s | | |

C.6 PR

| | | | |
|------------------|------------------|-----|-----|
| PR Settings | ADU | PED | NEO |
| PR Source | SpO ₂ | | |
| Alarm Switch | On | | |
| Alarm Record | Off | | |
| Alarm Level | Medium | | |
| Alarm High Limit | 120 | 160 | 200 |
| Alarm Low Limit | 50 | 75 | 100 |
| Pulse Volume | 3 | | |
| Alarm Source | Auto | | |

C.7 NIBP

| | | | |
|---------------|-----|-----|-----|
| NIBP Settings | ADU | PED | NEO |
| Alarm Switch | On | | |

| | | | |
|------------------------|--------|-----|-----|
| Alarm Record | Off | | |
| Alarm Level | Medium | | |
| Alarm High Limit (SYS) | 160 | 120 | 90 |
| Alarm Low Limit (SYS) | 90 | 70 | 40 |
| Alarm High Limit (Map) | 110 | 90 | 70 |
| Alarm Low Limit (Map) | 60 | 50 | 30 |
| Alarm High Limit (Dia) | 90 | 70 | 60 |
| Alarm Low Limit (Dia) | 50 | 40 | 20 |
| EDAN Module | | | |
| Inflation value | 160 | 140 | 100 |
| | | | |
| M3600 Module | | | |
| Inflation value | 180 | 180 | 120 |
| Unit | mmHg | | |
| Interval | Manual | | |

C.8 TEMP

| TEMP Settings | ADU | PED | NEO |
|-----------------------|--------|------|------|
| Alarm Switch | On | | |
| Alarm Record | Off | | |
| Alarm Level | Medium | | |
| Alarm High Limit (T1) | 39.0 | 39.0 | 39.0 |
| Alarm Low Limit (T1) | 36.0 | 36.0 | 36.0 |
| Alarm High Limit (T2) | 39.0 | 39.0 | 39.0 |
| Alarm Low Limit (T2) | 36.0 | 36.0 | 36.0 |
| Alarm High Limit (TD) | 2.0 | 2.0 | 2.0 |
| Unit | °C | | |

C.9 Quick TEMP

| Quick TEMP Settings | ADU | PED | NEO |
|---------------------|-----|-----|-----|
| Alarm Switch | On | | |
| Alarm Record | Off | | |

| | | | |
|-----------------------|--------|------|---|
| Alarm Level | Medium | | |
| Alarm High Limit (T1) | 39.0 | 39.0 | / |
| Alarm Low Limit (T1) | 36.0 | 36.0 | / |
| Unit | °C | | |

C.10 IBP

| IBP Settings | ADU | PED | NEO |
|---------------------------------------|---------------|---------------|---------------|
| Alarm Switch | On | | |
| Alarm Record | Off | | |
| Alarm Level | Medium | | |
| Unit | mmHg | | |
| Filter | 12.5Hz | | |
| | SYS, DIA, MAP | SYS, DIA, MAP | SYS, DIA, MAP |
| Alarm High Limit (ART, P1, P2) | 160, 90, 110 | 160, 90, 110 | 160, 90, 110 |
| Alarm Low Limit (ART, P1, P2) | 90, 50, 70 | 90, 50, 70 | 90, 50, 70 |
| Alarm High Limit (PA) | 35, 16, 20 | 35, 16, 20 | 35, 16, 20 |
| Alarm Low Limit (PA) | 10, 0, 0 | 10, 0, 0 | 10, 0, 0 |
| | MAP | MAP | MAP |
| Alarm High Limit (CVP, RAP, LAP, ICP) | 10 | 10 | 10 |
| Alarm Low Limit (CVP, RAP, LAP, ICP) | 0 | 0 | 0 |

C.11 CO₂

| CO2 Settings | ADU | PED | NEO |
|---------------------------|---------|-----|-----|
| Alarm Switch | On | | |
| Alarm Record | Off | | |
| Alarm Level | Medium | | |
| Work Mode | Standby | | |
| Unit | mmHg | | |
| Apnea Time | 20s | | |
| O ₂ Compensate | 16% | | |

| | | | |
|---------------------------------------|----------|----|-----|
| Anes Agent | 0% | | |
| Alarm High Limit (EtCO ₂) | 50 | 50 | 45 |
| Alarm Low Limit (EtCO ₂) | 15 | 20 | 30 |
| Alarm High Limit (FiCO ₂) | 4 | 4 | 4 |
| Alarm High Limit (AWRR) | 30 | 30 | 100 |
| Alarm Low Limit (AWRR) | 8 | 8 | 30 |
| Sweep | 12.5mm/s | | |
| Amplitude | Low | | |

C.12 C.O.

| C.O. Settings | ADU | PED | NEO |
|------------------------------|--------|------|------|
| Alarm Switch | On | | |
| Alarm Record | Off | | |
| Alarm Level | Medium | | |
| Alarm High Limit (TB) | 43.0 | 43.0 | 43.0 |
| Alarm Low Limit (TB) | 23.0 | 23.0 | 23.0 |
| Injectate Temperature Source | Auto | | |
| Temperature Unit | °C | | |
| Interval | 30 | | |
| Constant | 0.542 | | |

C.13 AG

| AG Settings | ADU | PED | NEO |
|---------------------------|---------|-----|-----|
| Alarm Switch | On | | |
| Alarm Record | Off | | |
| Alarm Level | Medium | | |
| Work Mode | Measure | | |
| Apnea Time | 20s | | |
| Unit | % | | |
| O ₂ Compensate | OFF | | |
| Anes Agent | HAL | | |
| Alarm High Limit (EtAA) | 8.0 | 8.0 | 8.0 |

| | | | |
|---------------------------------------|----------|------|------|
| Alarm Low Limit (EtAA) | 0.0 | 0.0 | 0.0 |
| Alarm High Limit (FiAA) | 6.0 | 6.0 | 6.0 |
| Alarm Low Limit (FiAA) | 0.0 | 0.0 | 0.0 |
| Alarm High Limit (EtN ₂ O) | 55 | 55 | 55 |
| Alarm Low Limit (EtN ₂ O) | 0 | 0 | 0 |
| Alarm High Limit (FiN ₂ O) | 53 | 53 | 53 |
| Alarm Low Limit (FiN ₂ O) | 0 | 0 | 0 |
| Alarm High Limit (EtO ₂) | 90.0 | 90.0 | 90.0 |
| Alarm Low Limit (EtO ₂) | 18.0 | 18.0 | 18.0 |
| Alarm High Limit (FiO ₂) | 88.0 | 88.0 | 88.0 |
| Alarm Low Limit (FiO ₂) | 18.0 | 18.0 | 18.0 |
| Sweep | 12.5mm/s | | |
| Amplitude | 2 | | |

D Abbreviations

| Abbr | English Full Name/Description |
|--------------------|---|
| AC | Alternating current |
| Adu | Adult |
| AG | Anaesthesia gas |
| Art | Arterial |
| aVF | Left foot augmented lead |
| aVL | Left arm augmented lead |
| aVR | Right arm augmented lead |
| awRR | Airway respiration rate |
| BP | Blood pressure |
| BTPS | Body temperature and pressure, saturated |
| CI | Cardiac index |
| C.O. | Cardiac output |
| CISPR | International Special Committee on Radio Interference |
| CMS | Central monitoring system |
| CO ₂ | Carbon dioxide |
| COHb | Carboxyhemoglobin |
| CVP | Central venous pressure |
| DC | Direct current |
| Des | Desflurane |
| Dia | Diastolic |
| ECG | Electrocardiogram |
| EEC | European Economic Community |
| EMC | Electromagnetic compatibility |
| EMI | Electromagnetic interference |
| Enf | Enflurane |
| ESU | Electrosurgical unit |
| Et | End-tidal |
| EtCO ₂ | End-tidal carbon dioxide |
| EtN ₂ O | End-tidal nitrous oxide |

| | |
|--------------------|--|
| Eto | Ethylene oxide |
| EtO ₂ | End-tidal oxygen |
| FCC | Federal Communication Commission |
| FDA | Food and Drug Administration |
| Fi | Fraction of inspired |
| FiCO ₂ | Fraction of inspired carbon dioxide |
| FiN ₂ O | Fraction of inspired nitrous oxide |
| FiO ₂ | Fraction of inspired oxygen |
| Hal | Halothane |
| Hb | Hemoglobin |
| Hb-CO | Carbon mono-xide hemoglobin |
| HR | Heart rate |
| IBP | Invasive blood pressure |
| ICP | Intracranial pressure |
| ICU | Intensive care unit |
| ID | Identification |
| IEC | International Electrotechnical Commission |
| IEEE | Institute of Electrical and Electronic Engineers |
| Iso | Isoflurane |
| LA | Left arm |
| LAP | Left atrial pressure |
| LCD | Liquid crystal display |
| LED | Light emitting diode |
| LL | Left leg |
| MAP | Mean arterial pressure |
| MDD | Medical Device Directive |
| MetHb | Methemoglobin |
| MRI | Magnetic resonance imaging |
| N/A | Not applied |
| N ₂ | Nitrogen |
| N ₂ O | Nitrous oxide |
| Neo | Neonate |

| | |
|----------------|---------------------------------|
| NIBP | Non-invasive blood pressure |
| O ₂ | Oxygen |
| oxyCRG | Oxygen cardio-respirogram |
| PA | Pulmonary artery |
| PAWP | Pulmonary artery wedge pressure |
| Ped | Pediatric |
| Pleth | Plethysmogram |
| PR | Pulse rate |
| PVC | Premature ventricular complex |
| R | Right |
| RA | Right arm |
| RAP | Right atrial pressure |
| Resp | Respiration |
| RHb | Reduced hemoglobin |
| RL | Right leg |
| RR | Respiration Rate |
| Sev | Sevoflurane |
| SYS | Systolic pressure |
| TB | Blood Temperature |
| TD | Temperature difference |
| TEMP | Temperature |
| USB | Universal serial bus |

P/N: 01.54.455473-10

EDANUSA
www.edanusa.com

4901 Morena Blvd., Suite 505
San Deigo, CA 92117
Tel: 888.850.4597