

Operator's Manual

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

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1.0 **Notices**

1.1 **CONTACT INFORMATION**

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1.1.2 **Notified Body**

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1.2 PATENT NOTICE

MOVES[®] SLC[™] is covered by one or more US and international patents and patents pending.

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1.4 TRADEMARK NOTICES

MOVES® is a registered trademark of Thornhill Research Inc.

SLC™ is a trademark of Thornhill Research Inc.



All brand and product names mentioned herein are used for identification purposes only and are trademarks or registered trademarks of their respective holders.

1.5 EMC (ELECTROMAGNETIC COMPATIBILITY) NOTICE

MOVES[®] SLC[™] generates, uses, and can radiate RF (radio frequency) energy. If it is not installed and used in accordance with the instructions in this manual, electromagnetic interference may result. MOVES[®] SLC[™] has been tested and found to comply with the limits set forth in IEC 60601-1-2 (identical to EN 60601-1-2) for medical products. These limits provide reasonable protection against electromagnetic interference in the intended use environments described in this manual.

1.6 MRI (MAGNETIC RESONANCE IMAGING) NOTICE

MOVES[®] SLC[™] contains electromagnetic components whose operation can be affected by intense electromagnetic fields. Do not operate MOVES[®] SLC[™] in an MRI environment or in the vicinity of high-frequency surgical diathermy equipment. Electromagnetic interference could disrupt the operation of the ventilator.

1.7 PHTHALATES NOTICE

MOVES[®] SLC™ and its accessories may contain phthalates. Phthalates are classified as carcinogenic, mutagenic or toxic to reproduction.



WARNING! IN ORDER TO REDUCE THE POTENTIAL RISK FROM PHTHALATES, LONG TERM EXPOSURE SHOULD BE AVOIDED IN THE TREATMENT OF CHILDREN AND PREGNANT OR NURSING WOMEN.

1.8 REGULATORY NOTICE

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

1.9 CLASSIFICATION

Type of Equipment: Medical Equipment

Electrical Classification: Class I, Type BF (unit)

Registration Classification: Class IIb

Type CF Defibrillation Proof (applied parts)

Pediatric and Adult Lung Ventilator

1.10 DECLARATION OF CONFORMITY NOTICE

MOVES[®] SLC™ is declared to conform to the Medical Device Directive of the European community (re: Medical Device Directive 93/42/EEC). This is indicated by the CE Mark shown below.





MOVES[®] SLC[™] also conforms to the following Technical Standards:

IEC 60601-1	IEC 80601-2-30	ISO 8359
IEC 60601-1-2	IEC 60601-2-34	ASTM E1112-00
ISO 80601-2-12	IEC 60601-2-49	BS EN 794-3 (2009)
ISO 80601-2-13	ISO 80601-2-55	MIL-STD-810G
IEC 60601-2-27	ISO 80601-2-61	JECETS

For more information on these standards, see Regulatory Standards Compliance on page 22.

1.10.1 Trade Name

 $\mathsf{MOVES}^{\circledR}\,\mathsf{SLC^{\intercal\!\mathsf{M}}}\;\mathsf{portable}\;\mathsf{life}\;\mathsf{support}\;\mathsf{system}$



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2.0 MOVES® SLC™ Quick Start Guide for Ventilated Patients

Equipment Setup Procedures

- 1. Lift the release/lock latch for the ventilator cartridge door at the top front of the MOVES[®] SLC[™]. Insert the ventilator breathing cartridge, close the door and lock it by depressing the latch.
- 2. Install the hydrocarbon filter in the REAR panel.
- 3. Attach the ventilator breathing circuit to the cartridge door. Connect the Nafion tubing to the sampling line.
- 4. If ventilating patients under 30 kg or using tidal volumes under 150 mL, replace the breathing filter with the pediatric breathing system filter.
- 5. Attach the suction canister and tubing to MOVES[®] SLC™.
- 6. Connect any required patient monitoring accessories to MOVES[®] SLC[™]. **DO NOT CONNECT TO THE PATIENT AT THIS TIME.**
- 7. Verify the battery charge levels and insert both batteries.
- Attach the MOVES[®] SLC[™] AC power unit (includes power supply, battery charger and cables) to the MOVES[®] SLC[™] if required to recharge.
- 9. Ensure that a backup means of ventilation with a high level of oxygen is immediately available.

Pre-use Test Instructions

- Turn the MOVES[®] SLC[™] unit on by pressing the Power Control Button (on the REAR panel).
- 2. The System Test screen is displayed.

NOTE: The <u>Jog Wheel</u> control is used to select options. It moves from selection to selection in a circular clockwise or counterclockwise direction around the screen depending on in which direction it is turned. Pressing the Jog Wheel activates the selected option.



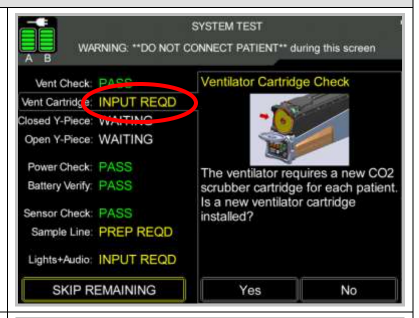


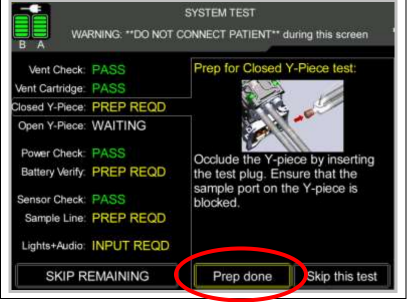


Some tests, including RAM and firmware integrity checking, are AUTOMATIC (note tests initially labeled PASS).
 Others require preparation of the device into a certain configuration (i.e., PREP REQD) or require the user to give feedback (i.e., INPUT REQD). The first test selected, the Ventilator Cartridge Check test, is an INPUT REQD test.

NOTE: The system should not be used until all tests pass. If any test reports a failure, wait until all tests have been run. Then use the Jog Wheel to select the failed test. Troubleshooting instructions will then be presented on the screen.

 Perform the preparation steps indicated if the test is PREP REQD (for other tests that are INPUT REQD, respond with the input requested), then choose Prep done to run the test.







NOTE: In order to save time, certain tests can be run concurrently. For example, while the Closed Y-Piece test is still running, the screen may advance to the Sample Line or Lights + Audio tests.



5. The closed Y-Piece test is completed and passed in the screen shot at the right. To initiate the Alarm Lights + Audio tests, user input is required. Start must be confirmed, and then the operator must continue to observe the device and answer the questions shown on the device.

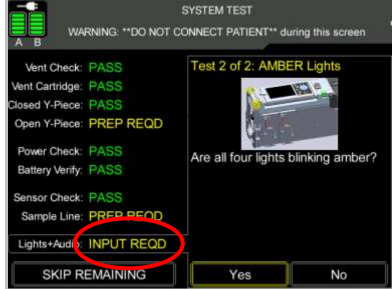




 The <u>first</u> Alarm Lights + Audio test requires input from the user (INPUT REQD test). This test ensures that the high-priority alarm visual and audio indicators are working properly. Respond with Yes or No.



 The <u>second</u> Alarm Lights & Audio test also requires input from the user (INPUT REQD test). This test ensures that the medium-priority alarm visual indicator is working properly. Respond with Yes or No.

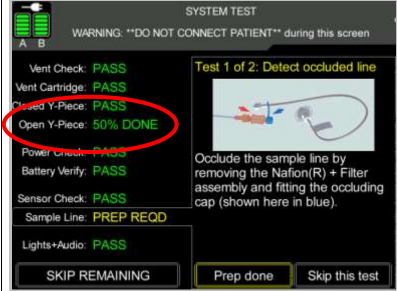




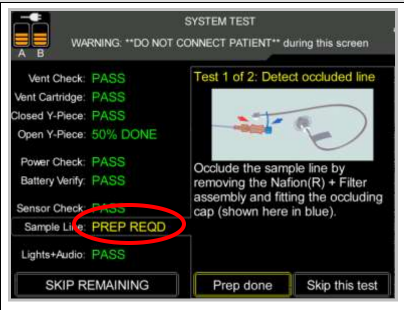
8. The next test, the Open Y-Piece test, is a PREP REQD test. Perform the requested preparations then select Prep done.



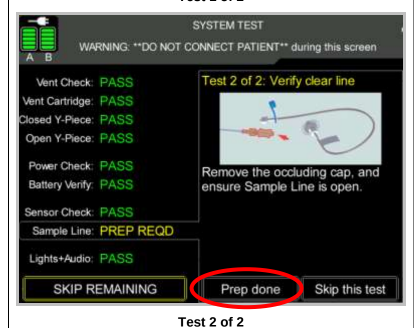
 As noted previously, in order to save time, certain tests can be run concurrently. In this case, while the Open Y-Piece test is still running, the Sample Line tests can be run.



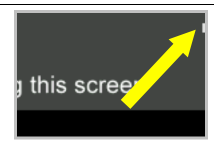
10. The Sample Line test is actually comprised of two tests. Both require user preparation (PREP REQD test). Follow the instructions on screen, perform the preparation required, and then choose Prep done to start the test.



Test 1 of 2



11. When testing is in progress, and the MOVES[®] SLC™ is running, a small white dot can be seen moving up and down at the top right of the screen to indicate that the device is not "frozen". The white "dot" is a way of knowing the device is not "frozen" at any time, not just during startup tests.



Pre-use Test Instructions 12. If all tests are successful, the System SYSTEM TEST Test screen displays a Test Summary WARNING. **DO NOT CONNECT PATIENT** during this screen indicating all tests have passed. Choose CONTINUE. TEST SUMMARY: PASSED Vent Check: PASS Vent Cartridge: PASS Closed Y-Piece: PASS Open Y-Piece: PASS Power Check: PASS The system, as configured at time Battery Verify: PASS of testing, passed all tests. Press CONTINUE to acknowledge Sensor Check: PASS and proceed. Sample Line: PASS Lights+Audio: PASS RETRY ALL TESTS CONTINUE 13. You are taken to the Setup Screen and SETUP placed in Monitor Only mode. No Alarms Monitor Only SYSTEM MODE: Ventilate O2 Supplement Monitor Only **VENTILATOR** Vent Mode: IMV A/C SIMV APRV CPAP/PS Sensitivity: Normal Low CONCENTRATOR Control: Volume Pressure Vent O2: 40% Tidal Volume: 500 mL Control Pressure: 20 cmH2O OTHER PEEP: 0-3 cmH2O ECG HR Mode: Adult Pediatric Frequency: 10 B/M ECG Rate: 10mm/s 25mm/s ECG Range: 2.2 mV 1:E Ratio: 1:2 NIBP Mode: MANUAL Inspire/High Time: 1.0 s Pressure Support: Off Suction: 325 mmHg Apnea Backup: Off ADVANCED... WIFI... INFO..

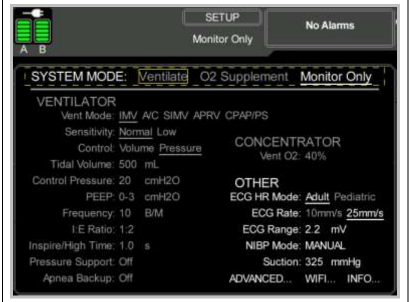


14. Make any changes necessary to the Ventilate mode settings and configure any other settings as necessary. Also, connect any desired patient monitoring cables to the patient.

NOTE: Ventilation and other settings can be changed even though the system is Monitor Only mode.



15. Change the system to Ventilate mode. Ventilation begins using the indicated settings.



16. Attach the patient to the breathing circuit.

BATTERY VERIFY TEST SCREEN

If the MOVES[®] SLC™ is connected to wall power, or has two full batteries, the Battery Verify test will be passed automatically and its test screen will not appear in the startup sequence. However, if neither of these two conditions is satisfied, the Battery Verify test screen will appear.

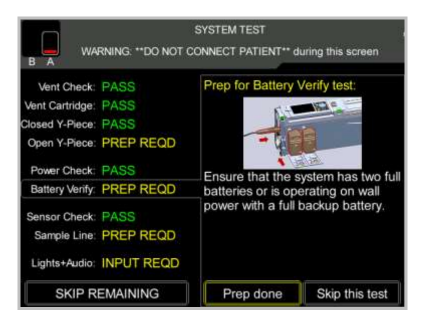


Figure 2-1: Battery Verify Test Screen

Keypad Buttons		
Image	Name	Use
	Screen Brightness Button	Use to adjust screen brightness.
	Flip Screen Button	Use to flip the information displayed on the screen 180°.
	Alarm Audio Pause Button	Use either to silence ALL alarms permanently (if this feature is permitted) or for a temporary period (120 seconds).
	NIBP Control Button	Use to start or abort a Non-Invasive Blood Pressure (NIBP) reading.
0	Suction Control Button	Use to start or abort suction. All suction accessories must be connected and ready to use before activating.



2.1 FURTHER INFORMATION ON PROCEDURES

For further information on procedures, see the sections indicated in the following table.

Table 1: Quick Reference to Information on Procedures

PROCEDURE	Page
Attaching the shoulder strap.	80
2. Installing the ventilator breathing cartridge.	95
Installing the hydrocarbon filter.	98
Installing the ventilator breathing circuit.	99
5. Inspecting the batteries.	115
6. Installing the batteries.	117
7. Preparing the power supply / battery charger	121
8. Connecting MOVES [®] SLC [™] to the power supply / battery charger.	122
9. Connecting AC power.	123
10. Adjusting the screen display orientation.	125
 Changing the brightness of the display (see third item in Table 20: User Interface Controls and Functions). 	128
12. Performing System Tests.	143
13. Attaching the suction tube and suction canister to the MOVES [®] SLC™.	109
14. Using the suction feature.	220
15. Connecting to the patient.	208
16. Making patient monitoring connections between the MOVES [®] SLC™ unit and the patient.	92
17. Connecting ABP, CVP or ICP equipment to the MOVES [®] SLC™	208
18. Selecting the operating mode and associated settings.	144
19. Changing alarm settings.	172
20. Viewing the Main screen.	158
21. Viewing graphs and trends of patient data.	223

PROCEDURE	Page
22. Taking NIBP readings manually.	129
23. Using the remote screen.	177
24. Shutting down MOVES [®] SLC™.	



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3.0 Introduction

This Operator's Manual is a reference guide for the MOVES[®] SLC™ mobile life support system (P/N 122752). This manual includes illustrations, annotated photographs and procedures designed to assist in the operation of various systems, subsystems and components that comprise the unit.

An operator of the MOVES[®] SLC™ system must read this manual and any accompanying accessory manuals and instructions in their entirety prior to use to safely and effectively operate the system. The operator, or a designated healthcare professional, must disclose the risks and associated mitigation steps to the subject on which the system will be used.

Keep this manual in a dry, convenient location for easy access. All information, illustrations, photographed procedures and specifications in this manual represent the most current product information available at the time of publication.

3.1 GENERAL INFORMATION

3.1.1 Manufacturer's Information

For warranty, parts, repair or customer service, please contact Thornhill Research Inc. See *Contact Information on page 1* for full contact details.

3.1.2 Glossary of Terms and Abbreviations

Table 2: Glossary of Terms and Abbreviations

TERM / ABBREVIATION	DEFINITION
AC	Alternating Current. A type of electrical current in which the direction of the electrical flow switches back and forth regularly.
A/C	Assist/Control Ventilation. In A/C mode, the system delivers a specific tidal volume or pressure at specific intervals based on a patient inspiratory trigger or time trigger.
ABP	Arterial Blood Pressure. ABP is measured in millimeters of mercury (mmHg).
APRV	Airway Pressure Release Ventilation. In APRV, the system delivers a high pressure with short, timer-controlled, lower pressure periods.
ВРМ	Beats per Minute
B/M	Breaths per Minute
CFM	Cubic Feet per Minute
Control Pressure	Breath target pressure over PEEP (cmH ₂ O)
CPAP	Continuous Positive Airway Pressure
CVP	Central Venous Pressure. CVP is measured in millimeters of mercury (mmHg).
DC	Direct Current. A type of electrical current in which the electrons always flow in the same direction.
ECG	Electrocardiogram



TERM / ABBREVIATION	DEFINITION
EtCO ₂	End Tidal Carbon Dioxide. EtCO2 is measured in millimeters of mercury (mmHg).
FiO ₂	Fraction of Inspired Oxygen by Volume (%)
FRC	Functional Residual Capacity – the volume of air (about 3 liters in an adult) that is present in the lungs at the end of a normal expiration.
Fresh Gas	Gas which has a negligible concentration of CO ₂ .
Frequency	Machine controlled breaths in a minute
HC Filter	Hydrocarbon Filter
HR	Heart Rate
I/E ratio	Ratio of Inspiratory Time / Expiratory Time
IP	Invasive Pressure. Can refer collectively or individually to ABP, CVP or ICP (which is not a "blood" pressure).
ICP	Intracranial Pressure. ICP is measured in centimeters of water (cmH ₂ O).
IMV	Intermittent Mandatory Ventilation. In IMV mode, the system delivers a specific tidal volume or pressure at specific intervals based on specific time triggers.
LED	Light Emitting Diode
LPM	Liters Per Minute
NATO	North Atlantic Treaty Organization
NIBP	Non-Invasive Blood Pressure. Measured via a blood pressure cuff. NIBP is measured in millimeters of mercury (mmHg).
OEM	Original Equipment Manufacturer
Oxygen Supplementation	The provision of gas containing a higher concentration of oxygen than ambient air.
Paw	Airway Pressure
PC	Pressure Control ventilation
pCO ₂	The partial pressure of CO₂ measured in millimeters of mercury (mmHg).
PEEP	Positive End-Expiratory Pressure in centimeters of water pressure (cmH ₂ O).
PI	Perfusion Index
	(Perfusion Index, or PI, is a relative assessment of the pulse strength at the monitoring site.)
PIP	Peak Inspiratory Pressure. PIP is measured in centimeters of water (cmH ₂ O).
PS	Pressure Support. PS assists a patient's inspiratory effort through the application of an additional set level of pressure above PEEP. PS is measured in centimeters of water (cmH2O).



TERM / ABBREVIATION	DEFINITION
PVI	Pleth Variability Index
	PVI may help clinicians non-invasively assess the fluid status of a patient.
RR	Respiratory Rate. Breaths per minute (B/M).
SGM	Safe Gas Mode
SpCO	Saturation percentage of carbon monoxide attached to hemoglobin.
	CO (carbon monoxide) competes with oxygen for the oxygen-binding sites on hemoglobin. The binding of CO to hemoglobin results in the formation of a compound called Carboxyhemoglobin (COHb). This compound is unable to transport or transfer oxygen.
SpHb	Total hemoglobin concentration in arterial blood.
	Hemoglobin is the part of a red blood cell that carries oxygen to the body. SpHb measures total hemoglobin and indicates the oxygen carrying capacity of the blood.
SpMet	Saturation percentage of methemoglobin.
	(Methemoglobin [MetHb] is an oxidized form of hemoglobin that is unable to carry oxygen.)
SpOC	Total oxygen content.
	(SpHb and SpO ₂ are used together to calculate the actual amount of oxygen in the blood.)
SpO ₂	Oxygen saturation of hemoglobin. Arterial oxygen saturation of hemoglobin as read from a pulse oximeter. It is measured as a percentage (%) of oxyhemoglobin present in arterial blood in relation to total hemoglobin.
SIMV	Synchronized Intermittent Mandatory Ventilation. In SIMV mode, the system delivers breaths synchronized with the patient's and ensures that a minimum number of breaths of a specified tidal volume or PIP are delivered. Additionally, any breaths beyond the minimum set number can be supported with a specific level of pressure.
UI	User Interface
VC	Volume Control ventilation.
Vt	Breath (tidal) volume. Vt is typically measured in milliliters (mL) or liters (L). MOVES [®] SLC™ measures only in milliliters (mL).

3.2 MOVES® SLC™ INTENDED USE

The MOVES[®] SLCTM is a portable ventilator which is computer-controlled and electrically powered. It is intended to provide continuous or intermittent ventilatory support for the care of individuals who require mechanical ventilation for periods up to 24 hours. The MOVES[®] SLCTM is intended to be used as both a transport and short term critical-care ventilator.

 $\mathsf{MOVES}^{\circledR}$ $\mathsf{SLC^{\intercal M}}$ provides the following supplemental functions for patients that it is ventilating or supplying with supplemental oxygen:



a. Suction

The MOVES[®] SLC[™] suction pump is intended for aspiration and removal of fluids, tissue (including bone), gases, bodily fluids or infectious materials from wounds or from a patient's airway or respiratory support system.

b. Supplementary Oxygen

The MOVES[®] SLC[™] is intended to provide supplemental oxygen-enriched air to patients that require supplemental oxygen.

c. Patient Monitoring

MOVES[®] SLC™ is intended to monitor physiological parameters of patients and provide these parameters to a health care provider for interpretation in the form of physiological data and system alarms. Physiological data and system alarms will be available to the care provider from the monitor.

3.2.1 Operating Environment

MOVES[®] SLC™ is intended to be operated in a transport, emergency, hospital, or field hospital setting.

3.2.2 Target Population

The intended patient population is pediatric and adult patients who weigh between 10 kg and 120 kg.



NOTE: The automated sphygmomanometer is not intended for use with pregnant patients.

3.3 INTENDED OPERATOR

MOVES[®] SLC[™] is intended to be used by, or under the supervision of, medically qualified and trained personnel.



4.0 Regulatory Compliance

4.1 REGULATORY SYMBOLS

Regulatory symbols have been added to the labeling on the MOVES[®] SLC™ unit, power supply and battery charger, and accompanying accessories to indicate regulatory compliance. These symbols, along with a brief description, are shown in the following table.

Table 3: Regulatory Symbols Used and Description

SYMBOL	DESCRIPTION	
C US 244588	CSA (Canadian Standards Association) Mark – This symbol appears on the MOVES [®] SLC™ Power Supply / Battery Charger. It indicates that approval by the Canadian Standards Association has been granted for use in Canada (C) and the United States (US) under the Certificate Number 244588.	
C US 2676502	CSA (Canadian Standards Association) Mark – This symbol appears on the MOVES [®] SLC [™] unit. It indicates that approval by the Canadian Standards Association has been granted for use in Canada (C) and the United States (US) under the Certificate Number 2676502.	
	"Conformité Européen" Mark – The CE mark is a self-declaratory mark that indicates the manufacturer or the importer of record has ensured that all of the applicable European safety and conformity directives and standards have been applied to the product.	
	The presence of the following symbols indicates compliance with International Electrotechnical Commission (IEC) standard IEC 60601-1 for Medical Electrical Equipment.	
<u></u>	Consult Accompanying Documents – This symbol appears on the Product Information labels for the MOVES [®] SLC™ unit and directs the operator to consult the accompanying documents. It also appears frequently in this operator's manual as a general warning and caution symbol.	
*	Type BF Applied Part – Indicates the device provides an intermediate degree of protection should the patient come in contact with an unintended source of voltage from an external source, but it is not approved for direct cardiac application.	
H	Type CF Applied Part Defibrillator-Proof - Indicates the applied part provides a high degree of protection should the patient come in contact with an unintended source of voltage from an external source, and that it is approved for direct cardiac application.	



4.2 REGULATORY STANDARDS COMPLIANCE

Table 4: Regulatory Standards Compliance

STANDARD#	DESCRIPTION
IEC 60601-1	Medical Electrical Equipment (Ed 3.0, 2005), General Requirements for basic safety and essential performance.
IEC 60601-1-2	Medical Electrical Equipment – Part 1-2: General requirements for safety – Collateral standard: Electromagnetic compatibility – Requirements and tests (Ed 3.0 2007)
IEC 60601-1-8	Medical Electrical Equipment – Part 1-8: General requirements for safety – Collateral standard: General requirements, tests and guidance for alarm systems in medical electrical equipment and medical electrical systems (Ed 2.1 2012)
ISO 80601-2-12	Medical Electrical Equipment - Part 2: Particular Requirements for basic safety and essential performance of critical care ventilators (Ed. 1.0 2011)
IEC 60601-2-27	Medical Electrical Equipment - Part 2: Particular Requirements for the Safety, including Essential Performance, of Electrocardiographic Monitoring Equipment (Ed. 3.0, 2011)
IEC 80601-2-30	Particular requirements for the safety, including essential performance, of automatic cycling non-invasive blood pressure monitoring equipment (2009)
IEC 60601-2-34	Medical Electrical Equipment - Part 2: Particular Requirements for The Safety, Including Essential Performance, of Invasive Blood Pressure Monitoring Equipment (Ed. 3.0, 2011)
IEC 60601-2-49	Medical Electrical Equipment - Part 2-49: Particular Requirements For The Safety Of Multifunction Patient Monitoring Equipment (Ed. 2.0, 2011)
ISO 80601-2-61	Medical Electrical Equipment - Particular requirements for the basic safety and essential performance of pulse oximeter equipment for medical use (2011)
ISO 80601-2-55	Medical Electrical Equipment - Particular requirements for the basic safety and essential performance of respiratory gas monitors (2011)
ISO 8359	Oxygen concentrators for medical use – Safety requirements (1996 + A1:2012)
ASTM E1112-00	Standard Specification for Electronic Thermometer for Intermittent Determination of Patient Temperature (2006)
BS EN 794-3	Lung ventilators. Particular requirements for emergency and transport ventilators (1998+A2:2009)
MIL-STD-810G	Department of Defense Test Method Standard for Environmental Engineering Considerations and Laboratory Tests
JECETS	Joint Enroute Care Equipment Test Standard

4.3 WiFi COMPLIANCE

WORLDWIDE REGULATORY COMPLIANCE

- FCC (IDs are XF6-RS9113SB, XF6-RS9113DB)
- IC (IDs are 8407A-RS9113SB, 8407A-RS9113DB)



- CE/ETSI
- TELEC
- SRRC



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5.0 MOVES[®] SLC[™] System Overview

5.1 GENERAL OVERVIEW

The MOVES[®] SLC™ system is comprised of six main modules:

- 1. Oxygen Concentrator
- 2. Ventilator
- 3. Suction System
- 4. Patient Monitoring System
- 5. Disposable breathing cartridges and breathing circuits for both intubated and non-intubated patients
- 6. Power system, comprised of removable hot swappable batteries and an AC power supply and battery charger



NOTE: Color shown in pictures may differ from the actual system or accessories.

5.2 SYSTEM ORIENTATION AND EXTERNAL COMPONENTS

Throughout this manual there are references to aid in orienting the caregiver with the positioning of the unit, especially when the orientation must change in order to conduct a given operating procedure. References to key external components may be given to further assist in the operator's orientation.

The following orientations should be remembered:

- FRONT: End of MOVES[®] SLC™ that contains the patient connection panel.
- BACK or REAR: End of MOVES[®] SLC™ that contains the power switch.
- RIGHT and LEFT Sides: As seen looking from FRONT to BACK.
- MOVES[®] SLC[™] should be operated in the upright position.



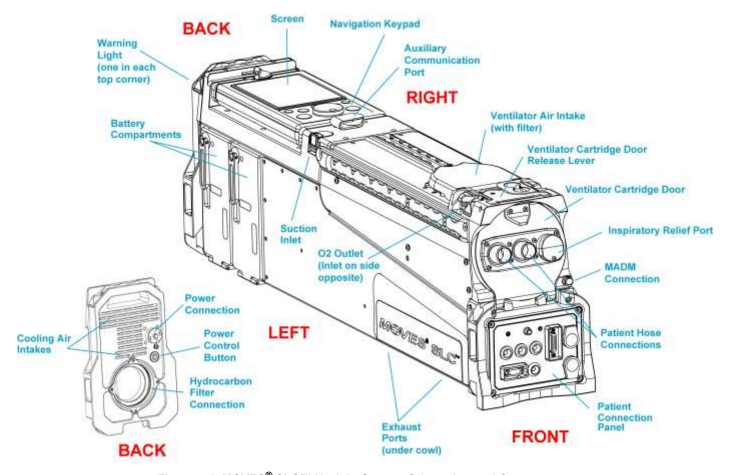


Figure 5-1: MOVES[®] SLC™ Upright System Orientation and Components

5.3 THEORY OF OPERATION

5.3.1 Breathing Circuit and Oxygen Supplement

Intubated patients are ventilated with a circle circuit (called a ventilator circuit) used with a ventilator cartridge. Oxygen and air enter the circuit from either the oxygen concentrator, the air pump or an external O₂ supply. The oxygen concentrator can be set to provide an intubated patient with an inspired O₂ concentration (FiO₂) between 30% and 85%. In Ventilator mode, the air pump serves as a back-up source of air flow if the system concentration of oxygen is too low or carbon dioxide is too high or either is unknown.



NOTE: MOVES[®] SLC™ is not intended to be used in an environment with ambient O₂ lower than 19.5% or with significant CO₂ buildup.

The ventilator cartridge is designed to remove CO₂ from the circle circuit for an approximately 2-hour period when the system operates at room temperature. The system provides a warning when the CO₂ level in inspired gas exceeds 6 mmHg, which is indicative of cartridge exhaustion, so the cartridge can be changed. If the cartridge is not changed, the system will enter Safe Gas Mode and increase oxygen production. It will also increase air supply if FiO₂ is set to ≤ 40%.



5.3.2 Oxygen Concentrator

The concentrator provides up to or greater than 87% O₂ into the ventilator breathing circuit during Ventilate mode or directly to the O₂ outlet during O2 Supplement mode.

5.3.3 Ventilator

The ventilator is comprised of a blower, a sealed ventilator chamber that houses the ventilator bag (air / oxygen reservoir), and a valve block that interfaces with the breathing cartridge and ventilator breathing circuit. The valve block includes the following components:

- Inspiratory and expiratory flow sensors
- One-way valves which are used to direct gas to and from the patient during inhalation and exhalation
- An independent high pressure relief valve ensures that the pressure in the breathing circuit will never
 exceed 100 cmH₂O at 60 LPM of flow. (NOTE: PEEP and PIP values are displayed from 0–70 cmH₂O.
 The pressure relief valve starts to trigger when the pressure exceeds 70 cmH₂O.)
- A port for measuring airway pressure
- A latch that secures the breathing cartridge to the valve block
- A ring mount for the ventilator bag

During patient inhalation, the blower pressurizes the ventilator chamber which displaces the gas from the ventilator bag into the patient's breathing circuit. The blower draws external air through an inlet filter. During exhalation, the patient's expired gas is directed to the ventilator bag. The ventilator bag also receives oxygen from the concentrator or external O₂ source and air from the air pump.



NOTE: For additional information on ventilation, please see <u>Appendix B – Pneumatic Diagram on page</u> 318.

5.3.4 Respiratory Gas Monitoring



NOTE: The system reports O_2/CO_2 gas measurements as ATPD (Ambient temperature [variable] and pressure [variable], dry [no humidity]). O_2/CO_2 readings are corrected based on water vapor pressure measurements.

The Ventilator breathing circuit is equipped with a filter on the Y-piece plus a sampling line and filter which is attached to the GAS SAMPLE port of the patient-connection panel on the MOVES[®] SLC[™]. To prevent equipment failure, only the sampling line and filter supplied with MOVES[®] SLC[™] should be used. The filter should be attached to the GAS SAMPLE port at all times unless the GAS SAMPLE port is capped. In addition, only the filter supplied with the Y-piece should be used with the Y-piece.

The oxygen sensor detects the concentration of oxygen – between 5 and 100% – that is being supplied to and exhaled from the patient and reports it in a plot as a percentage. The CO₂ sensor detects the concentration of CO₂ – between 0 and 10% – that is being supplied to and exhaled from the patient and then, taking into account ambient barometric pressure, reports it in a plot as a partial pressure in mmHg.

Under normal operating conditions (0°C to 40°C), the O₂ and CO₂ sensor should reach specified operating performance in less than two (2) minutes. Until the O₂ and CO₂ sensor reaches specified operating performance, the screen will display "CAL" for O₂ and CO₂ values, and a message "CO₂/O₂ sensors warming up" will be displayed in the alarm queue. When values begin to be displayed, the O₂ and CO₂ sensor has reached specified operating performance.





WARNING! DO NOT REUSE SAMPLING LINES OR FILTERS. THIS COULD PRESENT A DANGER OF INFECTION.



NOTE: Compensation for barometric pressure and temperature is performed by MOVES[®] SLC[™] internal sensing equipment to maintain accuracy of gas calibration over the MOVES[®] SLC[™] environmental operating range. For information on the MOVES[®] SLC[™] environmental operating range see Environmental Specifications on page 299.

ABOUT 02 AND CO2 SENSORS AND CALIBRATION

 O_2 and CO_2 calibration are performed at device startup after the O_2 and CO_2 sensors have initialized and warmed up. "CAL" is displayed in each O_2 and CO_2 parameter (i.e., FiO₂, PetCO₂ and PiCO₂) during warm up and calibration. The calibration is based on measuring the air surrounding the MOVES[®] SLCTM. If the initial attempt at O_2 calibration fails, on each subsequent O_2 calibration check, if the check is in bounds, a calibration is attempted until O_2 calibration succeeds.

A CO_2 calibration is performed every 30 minutes. O_2 calibration is only performed once, and then an O_2 calibration check is performed every 30 minutes. However, if SpO_2 is at or below 85%, the O_2 calibration check is performed every five (5) minutes until the condition clears.

The O_2 calibration check fails if the surrounding air's average O_2 reading \pm one (1) standard deviation (data point taken every 90 ms over a five [5] second period) is not contained within the bounds of 19.4–22.4%. The O_2 calibration check fails *high* if the air's average O_2 reading plus one (1) standard deviation is above the upper bound; otherwise, the O_2 calibration check fails *low* if the air's average O_2 reading minus one (1) standard deviation is below the lower bound

If the O_2 calibration check fails high, then the $MOVES^{@}$ SLCTM enters Safe Gas Mode (SGM) until an O_2 calibration check does not fail high (when the $MOVES^{@}$ SLCTM O_2 calibration is next checked). While the $MOVES^{@}$ SLCTM is in SGM due to an O_2 calibration check failure high, the O_2 calibration recheck occurs every five (5) minutes. If the O_2 calibration check fails low, subsequent O_2 calibration checks are scheduled at ten (10) minute intervals while each subsequent O_2 calibration check continues to fail low.

A high priority alarm stating " O_2 reading may be biased high" will appear in the alarm queue when the O_2 calibration check fails high. A low priority alarm stating " O_2 reading may be biased low" alarm will appear in the alarm queue when the O_2 calibration check fails low.

Whenever a CO_2 calibration is scheduled, an O_2 calibration check is performed at the same time. This minimizes the time that patient gas monitoring is suspended. Whenever an O_2 calibration check is performed, and the CO_2 calibration is scheduled to occur within the next 11 minutes, then a CO_2 calibration is performed at the same time as the O_2 calibration check.

ASTERISK BESIDE SENSOR VALUES

An O₂ calibration check and/or CO₂ calibration requires 15 seconds to complete. O₂ (FiO₂), CO₂ (PetCO₂ and PiCO₂) and RR values are displayed but are "frozen" while the O₂ calibration check and/or CO₂ calibration is performed. During this time, an asterisk is displayed beside each numeric value. If a parameter has no value (i.e., dashes, "---", are displayed), then no asterisk is shown beside that parameter.

ADVERSE EFFECTS

There are no known adverse effects associated with MOVES[®] SLC[™] respiratory gas monitoring in itself. However, sampling lines and filters are not reusable and could present a danger of infection were they to be reused. They should be disposed of in accordance with local biohazard regulations.



5.3.5 Suction

The system provides variable suction between −100 and −325 mmHg with flow rates of 20 L/min. A suction kit, consisting of a wand, two hoses, and an 800 mL suction filtration canister with a canister holder, is attached to the suction port of the MOVES[®] SLC[™]. The aspirated air is vented through the exhaust of the MOVES[®] SLC[™] system. Overfill protection for the suction container is provided by a filter in the lid which blocks flow when canister capacity is reached. The MOVES[®] SLC[™] system also contains a mechanical suction safety relief valve in the suction path which opens at approximately −415 mmHg.

When using a closed suction catheter, do not use patient triggered ventilator modes. Only pressure and volume controlled IMV should be used.



CAUTION! OPERATING SUCTION WITHOUT THE SUPPLIED FILTER, CANISTER, AND HOSES WILL RESULT IN PERMANENT FAILURE OF THE SUCTION SYSTEM AND THE OXYGEN CONCENTRATOR.

5.3.6 Patient Monitoring

NON-INVASIVE BLOOD PRESSURE (NIBP)

Using the appropriate cuff, NIBP measurements can be completed on either an arm or thigh. NIBP measurements made with the MOVES[®] SLC[™] for pediatric and adult patient populations are equivalent to those obtained by trained observers using the cuff/stethoscope auscultatory method within the limits prescribed by IEC 80601-2-30:2009 (difference of ± 3 mmHg or 2% of reading, whichever is greater).

In addition, blood pressure measurements determined with the MOVES[®] SLC™ are equivalent to those obtained by a trained observer using the cuff / stethoscope auscultation method, within the limits prescribed by the American National Standard, electronic or automated sphygmomanometers.

Readings can be affected by the measurement site, the position of the patient, exercise, or the patient's physiologic condition.

To obtain accurate resting blood pressure measurements for conscious patients with hypertension, ensure that the patient is comfortably seated with legs uncrossed, feet flat on the floor, back and arm supported and the middle of the cuff at the level of the right atrium of the heart. Recommend that the patient relax as much as possible and not talk during the measurement procedure. It is recommended that five (5) minutes should elapse before the first reading is taken.

If unexpected readings are obtained, verify the position and integrity of the cuff, make certain that there is no compression or restriction of the connecting tubing, and ensure that the patient is lying down or sitting still during measurement.

PULSE OXIMETRY

The Masimo Rainbow SET® Pulse Co-Oximeter measures the functional oxygen saturation of arterial hemoglobin (% SpO₂). Significant levels of dysfunctional hemoglobin, such as carboxyhemoglobin or methemoglobin, may affect the accuracy of the measurement. Factors that may degrade pulse performance or affect the accuracy of the measurement include the following: excessive ambient light, excessive motion, electrosurgical interference, blood flow restrictors (arterial catheters, blood pressure cuffs, infusing lines, etc.), moisture in the sensor, improperly applied sensor, incorrect sensor type, poor pulse quality, venous pulsations, anemia or low hemoglobin concentrations, cardiogreen or other intravascular dyes, carboxyhemoglobin, methemoglobin, dysfunctional hemoglobin, artificial nails or fingernail polish, or a sensor not at heart level. The pulse oximeter is calibrated by the original manufacturer to display functional oxygen concentration.





NOTE: For more information, see The Masimo Rainbow SET® Pulse CO-Oximeter beginning on page 53.

INVASIVE PRESSURE (IP)

Data is available from three types of Invasive Pressure (IP) sensors: Arterial, Central Venous, and Intracranial. Arterial Blood Pressure (ABP) is shown with numeric systolic and diastolic readings. The display is capable of displaying a pressure range of –10 to 300 mmHg for systolic and diastolic pressure readings. Intracranial Pressure (ICP) displays pressures from –14 to 408 cmH₂O. Central Venous Pressure (CVP) display pressures from –10 to 300 mmHg. For all invasive pressures, if the reading is below the sensor range, '<–10' is shown for ABP and CVP, and '<–14' is shown for ICP. If the reading is above the sensor range, '>300' is shown for ABP and CVP, and '>408' is shown for ICP.

To minimize the performance results due to ageing, and the effects of environmental conditions on the tubing, transducer or cable, always ensure that the IP transducer and tubing are stored appropriately and have not expired.

TEMPERATURE

When the temperature probe is attached to the patient and connected to the MOVES[®] SLC[™], the patient's body temperature is displayed on the Monitor Screen. The temperature can be displayed in degrees Fahrenheit from 82.4°F to 108.0°F or degrees Celsius from 28°C to 42°C. Changing the temperature display from Celsius to Fahrenheit (or vice versa) may only be done by a qualified MOVES[®] SLC[™] service technician.

ELECTROCARDIOGRAM (ECG)

MOVES[®] SLC™ uses a standard 12-lead ECG system for monitoring the heart and produces 12 ECG channels (I, II, III, aVL, aVR, aVF, V1, V2, V3, V4, V5 and V6). ECG data can be displayed on the Graphs / Trends section of the Monitoring Screen with a vertical scale (for data) and a horizontal scale (for time). The HR (Heart Rate) area of the Monitoring Screen displays heart rate in BPM (Beats per Minute) and data can be drawn from any of the ECG channel waveforms shown on the ECG monitor. If the ECG monitor is being used, and the readings are erratic, the accuracy of the heart rate cannot be guaranteed. The ECG heart rate meter's response to irregular rhythms has not been assessed.

5.3.7 Power System

The MOVES[®] SLC[™] operates on either rechargeable batteries (DC) or AC power. The power range of the MOVES[®] SLC[™] power supply is as follows:

- Input: 100–240 VAC, 50–60 Hz, 5.5 A max.
- Output: 28 VDC, 14.3 A max.

The MOVES[®] SLC[™] houses up to two lithium polymer batteries. MOVES[®] SLC[™] will operate on a set of 2 batteries for approximately 2.5 hours minimum. Under typical clinical use (ventilator and monitors running, concentrator on for 30 seconds / off for 90 seconds, assuming no leaks), MOVES[®] SLC[™] should operate at least 4 hours on a set of 2 batteries. Battery run time is highly dependent on the use of the oxygen concentrator or suction.

Power is supplied to the MOVES[®] SLC[™] from an AC source via the MOVES[®] SLC[™] power supply and battery charger, which provides 28 VDC at up to 14.3 A. Batteries installed in the system are charged automatically when external AC power is connected. The system's batteries should take no more than 2.5 hours to fully charge when the system is idle. Batteries can be charged while the system is running although charge times may be longer.



NOTE: The battery charge level may not appear to increase for approximately 3 hours. This is normal for the battery's initial charge and after extended periods without use.



5.3.8 System Auto Resume on Power Loss

If the system has been shut down for a period of time less than or equal to three (3) minutes, the system will auto resume using the system settings configuration prior to shutdown. A temporary loss of power is assumed here.

If the system has been shut down for a period of time greater than three (3) minutes but less than 30 minutes, the system will query the user as to whether the patient is new or continuing. Selecting YES will return the system settings to default values (see *System Default Settings on page 283* for details). Selecting NO will keep the last system settings used.

NO should only be selected if the operator is aware of the last system settings configuration, or the operator should review the settings on the Setup, Alarm Limits, Alarm ON/OFF and Advanced screens.



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6.0 Safety Information

6.1 MANUAL SYMBOLS AND MESSAGES

6.1.1 Notes, Cautions and Warnings

This manual contains important messages with symbols labeled **NOTE**, **CAUTION** and **WARNING**. These messages have the following format and meaning:

Table 5: Symbols and Messages Used in Manual



Supplies additional information that will help complete, offer an alternative to, or explain a portion of a given procedure.

NOTE:



PROVIDES INFORMATION TO PREVENT ERRORS OR INDISCRETIONS THAT COULD RESULT IN EQUIPMENT, SYSTEM, OR COMPONENT DAMAGE.

CAUTION!



INDICATES AREAS WHERE INSUFFICIENT KNOWLEDGE OF A GIVEN PROCEDURE, IMPROPER HANDLING, OR LACK OF ATTENTION COULD RESULT IN PERSONAL INJURY OR LOSS OF LIFE!

WARNING!

Read each labeled message carefully, and follow its instructions during operation to reduce the risk of system or component damage and/or personal injury.



WARNING! IMPROPER OPERATION OF THE MOVES[®] SLC™ COULD ENDANGER A PATIENT!

Since it is virtually impossible to foresee all of the possible consequences resulting from the failure to follow instructions and adhere to safety procedures, the NOTES, CAUTIONS and WARNINGS contained in this manual are not exhaustive. It is the responsibility of the individual operating MOVES[®] SLC™ to make safety the number one priority during operating procedures.



6.1.2 Operational Symbols

Table 6: Operational Symbols and Descriptions

SYMBOL	DESCRIPTION
~	Single Phase Alternating Current
===	Direct Current

6.1.3 Label Warning Symbols

Table 7: Label Warning Symbols and Descriptions

SYMBOL	DESCRIPTION
3	Fire Hazard: Do not smoke near unit.
	Fire Hazard: Do not operate device near open flame.

6.1.4 Product Labels Symbols

Table 8: Product Label Symbols and Descriptions

SYMBOL	DESCRIPTION
\triangle	Caution. Read accompanying documentation.
(3)	Follow instructions for use.
	Fragile item, handle carefully. Sensitive to mechanical shock.
2	Do not reuse (single use only)



SYMBOL	DESCRIPTION
LATEX	No latex used in the manufacture of this product
	Power indicator
	Battery indicator
	Class II equipment
EC REP	Authorized Representative in the European Community
•••	Manufacturer
	Date of manufacture
2011 -11	Date of expiration (use by)
LOT 000001	Lot number
REF	Reference or Model Number
SN	Serial Number
	Phone
	Fax



SYMBOL	DESCRIPTION
\bowtie	E-mail
A	The equipment shall not be disposed of as unsorted municipal waste and shall be collected as electrical and electronic equipment, as applicable, separately as specified by Waste Electrical and Electronic Equipment (WEEE).
	Battery condition indicator
NON	Non sterile; material cannot be guaranteed to be free of contamination.
1	Temperature limitation range for usage. Both upper and lower limits are indicated adjacent to horizontal lines.
Æ	Humidity limitation range for usage. Both upper and lower limits are indicated adjacent to horizontal lines.
600	Pressure limitation range for usage. Both upper and lower limits are indicated adjacent to horizontal lines.
<u><</u>	Gas sample line port
PHT DEHP	Indicates labeled item contains phthalates.
KONLY	For use by or on the order of a physician
	Locked
	Unlocked
Ť	Protect from rain



SYMBOL	DESCRIPTION
类	Do not expose to sunlight
[]i	Read usage instructions

6.2 GENERAL WARNINGS

Table 9: General Warnings

SYMBOL	GENERAL WARNING
	WARNING! THE MOVES [®] SLC™ SHOULD NOT BE USED IN AN EXPLOSIVE GAS ENVIRONMENT.
	WARNING! THE POWER SUPPLY / CHARGER CORD IS SUPPLIED WITH A GROUNDING PRONG ON THE MALE CONNECTOR. TO REDUCE THE RISK OF ELECTRICAL SHOCK, THIS PRONG SHOULD NEVER BE REMOVED OR COMPROMISED.
	WARNING! TO AVOID BREATH STACKING WHEN RUNNING VOLUME CONTROLLED IMV, THE RELEASE PRESSURE MUST BE APPROPRIATELY SET.
	WARNING! THE OPERATOR SHOULD ALWAYS HAVE AVAILABLE AN ALTERNATE MEANS OF SUPPLYING A HIGH CONCENTRATION OF O2 IN THE EVENT OF POWER FAILURE, MECHANICAL FAILURE, OR SERIOUS OCCLUSION IN THE CONCENTRATOR SYSTEM.
	WARNING! THE OPERATOR SHOULD ALWAYS HAVE AVAILABLE AN ALTERNATIVE MEANS OF VENTILATION CAPABLE OF SUPPLEMENTING A HIGH CONCENTRATION OF O2 IN THE EVENT OF POWER FAILURE, MECHANICAL FAILURE, OR SERIOUS OCCLUSION IN THE VENTILATOR SYSTEM. LACK OF IMMEDIATE ACCESS TO ALTERNATE VENTILATION CAN RESULT IN PATIENT DEATH.
	WARNING! THE MOVES [®] SLC™ SHOULD NOT BE COVERED OR POSITIONED IN SUCH A WAY THAT THE OPERATION OR PERFORMANCE OF THE VENTILATOR IS ADVERSELY AFFECTED. (EG. DO NOT COVER WITH A BLANKET TO REDUCE SOUND OR LIGHT)
	WARNING! THE OPERATOR SHOULD USE AN ALTERNATIVE MEANS OF VENTILATION UPON EXPERIENCING A PROLONGED APNEA ALARM.
	WARNING! THE VENTILATOR SHOULD NOT BE USED WITH HELIUM OR NITROUS OXIDE.
	WARNING! THE MOVES [®] SLC™ SHOULD NOT BE USED IN A HYPERBARIC CHAMBER.
	WARNING! THE OPERATOR SHOULD ALWAYS HAVE AVAILABLE AN ALTERNATE MEANS OF SUCTION IN THE EVENT OF POWER FAILURE, MECHANICAL FAILURE OR SERIOUS OCCLUSION IN THE SUCTION CIRCUIT.



SYMBOL	GENERAL WARNING
	WARNING! IT MAY BE NECESSARY TO USE AN ALTERNATIVE MEANS OF OXYGEN SUPPLEMENTATION SHOULD THE HYDROCARBON FILTER REQUIRE REPLACEMENT WHILE TREATING A PATIENT.
	WARNING! THE MOVES [®] SLC [™] SHOULD NOT BE RUN CONTINUOUSLY IN SAFE GAS MODE. SAFE GAS MODE IS INTENDED FOR <u>SHORT TERM USE ONLY</u> TO COMPLETE TRANSPORTS.
	WARNING! IF LEFT IN A HOT ENVIRONMENT OR DIRECT SUNLIGHT FOR A CONSIDERABLE LENGTH OF TIME, THE MOVES [®] SLC™ ACCESSORIES CASE AND THE ACCESSORIES IN IT CAN BECOME QUITE HOT. MONITORING OF CASE TEMPERATURE IS RECOMMENDED. ALWAYS MAKE SURE THAT ACCESSORIES THAT WILL BE APPLIED DIRECTLY TO THE PATIENT ARE SUITABLE FOR SKIN CONTACT AND WILL NOT CAUSE BURNS.
	WARNING! BATTERY EXPOSURE TO TEMPERATURES IN EXCESS OF 130°C (266°F) WILL RESULT IN THE BATTERY VENTING FLAMMABLE LIQUID AND GASES.
	WARNING! THE MOVES [®] SLC [™] OXYGEN CONCENTRATOR DOES NOT FUNCTION WHILE SUCTION IS ON. AN ALTERNATIVE MEANS OF SUPPLYING O ₂ WILL BE NECESSARY IF A HIGH PERCENTAGE OF O ₂ IS CRITICAL WHILE SUCTIONING.
	WARNING! ONE CHARGED BATTERY MUST BE PRESENT IN THE MOVES [®] SLC™ UNIT WHENEVER IT IS IN OPERATION, EVEN WHEN IT IS RUNNING ON EXTERNAL POWER. THIS REDUCES THE RISK TO THE PATIENT IN THE EVENT OF A POWER FAILURE.
	WARNING! CHARGING AND DISCHARGING THE BATTERIES WITH OTHER THAN MOVES [®] SLC™ SYSTEM EQUIPMENT, AND/OR IMPROPER HANDLING, CAN RESULT IN FIRE, EXPLOSION, TOXIC GASES AND SMOKE.
	WARNING! BATTERY TIME SHOWN REMAINING IS APPROXIMATE AND HIGHLY DEPENDENT ON OPERATING CONDITIONS! PUT SAFETY FIRST – ALWAYS CARRY SPARE BATTERIES!
	WARNING! DO NOT OPERATE THE MOVES [®] SLC™ SYSTEM WITH A DEFECTIVE BATTERY.
	WARNING! DO NOT USE OR CHARGE A DAMAGED BATTERY!
	WARNING! DO NOT OPERATE THE MOVES [®] SLC™ SYSTEM UNTIL ALL SYSTEM TEST FAILURES HAVE BEEN RESOLVED, AND ALL TESTS HAVE BEEN REPEATED AND PASSED.
	WARNING! ONLY AUTHORIZED SERVICE AND MAINTENANCE PERSONNEL SHOULD REMOVE ANY COVERS FROM MOVES [®] SLC™. UNAUTHORIZED REMOVAL OF COVERS FROM MOVES [®] SLC™ MAY RESULT IN ELECTRIC SHOCK AND POSSIBLY DEATH, AND MAY DAMAGE THE SYSTEM COMPONENTS.
	WARNING! BECAUSE THE MOVES [®] SLC [™] CONTAINS AN OXYGEN CONCENTRATOR, IT SHOULD ONLY BE USED IN A WELL-VENTILATED ENVIRONMENT AWAY FROM POLLUTANTS, FLAMES, SPARKS, OR FUMES.
	WARNING! LEAKS IN THE SAMPLING LINE CAN CAUSE LOW PCO ₂ AND/OR LOW O ₂ LEVELS.



SYMBOL	GENERAL WARNING
	WARNING! WHEN MOVES [®] SLC™ IS NOT IN OPERATION, BATTERIES SHOULD BE REMOVED FROM THE UNIT AND STORED IN A DRY AREA AT ROOM TEMPERATURE. LEAVING BATTERIES INSTALLED IN A NON-OPERATIONAL UNIT MAY CAUSE THEM TO DRAIN TO AN UNRECHARGEABLE LEVEL.
	WARNING! THE CLAMPS HAVE NUMEROUS MOVING PARTS THAT MAY PRESENT A PINCHING OR CRUSHING HAZARD. ALWAYS USE CAUTION WHEN HANDLING BOTH THE FRONT AND BACK CLAMPS.
	WARNING! WHEN SETTING ALARM LIMITS, DO NOT SET TO EXTREME VALUES THAT CAN RENDER THE ALARM SYSTEM USELESS.
	WARNING! DO NOT MODIFY THIS EQUIPMENT IN ANY WAY.
	WARNING! ONLY APPROVED NETWORK/DATA COUPLINGS ARE TO BE CONNECTED TO THE MOVES [®] SLC™ SYSTEM OR COMPONENTS.

6.3 ELECTRICAL WARNINGS

Table 10: Electrical Warnings

SYMBOL	ELECTRICAL WARNING
	WARNING! WHEN USING EXTERNAL POWER WITH THE MOVES [®] SLC™, THE POWER CORD MUST ALWAYS BE READILY ACCESSIBLE.
	TO AVOID THE RISK OF ELECTRIC SHOCK, THIS EQUIPMENT MUST ONLY BE CONNECTED TO A SUPPLY MAINS WITH PROTECTIVE EARTH.
	WARNING! THE MOVES [®] SLC™ SYSTEM IS NOT ELECTROSURGERY COMPATIBLE.
	WARNING! THE USE OF MULTIPLE (NON-) MEDICAL ELECTRICAL EQUIPMENT CONNECTED TO THE SAME PATIENT MAY POSE A SAFETY HAZARD DUE TO THE SUMMATION OF LEAKAGE CURRENTS FROM EACH INSTRUMENT. ANY COMBINATION OF (NON-) MEDICAL ELECTRICAL EQUIPMENT SHOULD BE EVALUATED BY LOCAL SAFETY PERSONNEL BEFORE BEING PUT INTO SERVICE.
	WARNING! CONDUCTIVE PARTS OF ELECTRODES AND ASSOCIATED CONNECTORS FOR APPLIED PART, INCLUDING THE NEUTRAL ELECTRODE, SHOULD NOT CONTACT OTHER CONDUCTIVE PARTS AND EARTH.



6.4 PATIENT-SPECIFIC WARNINGS

Table 11: Patient-Specific Warnings

SYMBOL	PATIENT-SPECIFIC WARNINGS
	WARNING! IN THE EVENT OF DISAGREEMENT BETWEEN THE DEVICE AND THE REMOTE SCREEN THE DEVICE SHALL BE CONSIDERED CORRECT.
	WARNING! IF THE REMOTE SCREEN BECOMES INOPERABLE, LOCKS UP OR BEHAVES ERRATICALLY, IT SHOULD NO LONGER BE USED AND BE DISCONNECTED FROM THE SYSTEM.
	WARNING! DO NOT CONNECT MOVES [®] SLC™ DIAGNOSTICS SOFTWARE TO THE DEVICE WHEN THE DEVICE IS CONNECTED TO A PATIENT.
	WARNING! WHEN USING O2 SUPPLEMENTATION, AN O2 SAT MONITOR MUST BE USED.
	WARNING! THE AUTOMATED SPHYGMOMANOMETER IS NOT INTENDED FOR USE WITH PREGNANT PATIENTS.
	WARNING! WHEN USING O2 SUPPLEMENT MODE, THE MOVES [®] SLC™ GAS SAMPLING PORT MUST BE CONNECTED TO THE OXYGEN DELIVERY CIRCUIT (E.G., O2 MASK SAMPLE PORT) AND AN OXYGEN SHUT-OFF DEVICE, SUCH AS BPR'S FIRESAFE™ CANNULA VALVE, SHOULD BE USED IN THE OXYGEN SUPPLY LINE AS CLOSE TO THE PATIENT AS POSSIBLE.
	WARNING! DO NOT CONNECT A PATIENT TO MOVES [®] SLC™ UNTIL THE MOVES [®] SLC™ SYSTEM IS PROPERLY WARMED UP AND O ₂ VALUES ARE DISPLAYED.
	WARNING! ACCURACY OF ANY BLOOD PRESSURE MEASUREMENT MAY BE AFFECTED BY THE POSITION OF THE SUBJECT, HIS OR HER PHYSICAL CONDITION AND USE OUTSIDE OF THE OPERATING INSTRUCTIONS DETAILED IN THIS MANUAL. INTERPRETATION OF BLOOD PRESSURE MEASUREMENTS SHOULD BE MADE ONLY BY A PHYSICIAN OR TRAINED MEDICAL STAFF.
	WARNING! DO NOT ATTACH THE BLOOD PRESSURE CUFF TO A LIMB WHERE INTRAVASCULAR ACCESS OR THERAPY OR AN ARTERIO-VENOUS SHUNT IS PRESENT. CUFF INFLATION CAN INTERFERE WITH THE BLOOD FLOW AND COULD RESULT IN INJURY TO THE PATIENT.
	WARNING! IF THE BLOOD PRESSURE CUFF IS ON THE SAME LIMB AS A PULSE OXIMETER PROBE, THE OXYGEN SATURATION RESULTS WILL BE ALTERED WHEN THE CUFF OCCLUDES THE BRACHIAL ARTERY.
	WARNING! IF THE BLOOD PRESSURE CUFF IS ON THE SAME LIMB AS ANOTHER PATIENT MONITOR, A TEMPORARY LOSS OF FUNCTION OF THE OTHER MONITOR MAY OCCUR WHEN THE CUFF IS PRESSURIZED.
	WARNING! DO NOT USE BLOOD PRESSURE CUFF ON THE ARM ON THE SIDE OF A MASTECTOMY.
	WARNING! TO OBTAIN ACCURATE BLOOD PRESSURE READINGS, THE BLOOD PRESSURE CUFF MUST BE THE CORRECT SIZE, AND ALSO BE CORRECTLY FITTED TO THE PATIENT. INCORRECT SIZE OR INCORRECT FITTING MAY RESULT IN INCORRECT READINGS.



SYMBOL	PATIENT-SPECIFIC WARNINGS
	WARNING! WHEN A BLOOD PRESSURE CUFF IS TO BE POSITIONED ON A PATIENT FOR AN EXTENDED LENGTH OF TIME, BE SURE TO OCCASIONALLY CHECK THE LIMB FOR PROPER CIRCULATION.
	WARNING! USING A BLOOD PRESSURE CUFF OVER A WOUND MAY CAUSE FURTHER INJURY.
	WARNING! USING A BLOOD PRESSURE CUFF TOO FREQUENTLY MAY CAUSE INJURY TO THE PATIENT DUE TO BLOOD FLOW INTERFERENCE.
	WARNING! IRREGULAR HEART RHYTHMS SUCH AS ATRIAL OR VENTRICULAR PREMATURE BEATS, ATRIAL FIBRILLATION, ARTERIALSCLEROSIS, POOR PERFUSION OR DIABETES MAY AFFECT BLOOD PRESSURE PERFORMANCE AND READING.
	WARNING! BEFORE VENTILATING A PATIENT, ENSURE THAT A SPARE VENTILATOR CARTRIDGE AND BREATHING CIRCUIT ARE READILY AVAILABLE.
	WARNING! CARE SHOULD BE TAKEN WHEN MONITORING PATIENTS WITH PACEMAKERS SINCE HEART RATE METERS MAY FALSELY COUNT PACEMAKER PULSES.
	WARNING! IRREGULAR HEART RHYTHMS SUCH AS PREMATURE ATRIAL OR VENTRICULAR BEATS MAY CAUSE THE HEART RATE TO BE UNDERESTIMATED.
	WARNING! DO NOT REUSE SAMPLING LINES OR FILTERS. THIS COULD PRESENT A DANGER OF INFECTION.
	WARNING! DO NOT REUSE PARTS MARKED FOR SINGLE USE ONLY. THIS COULD PRESENT A DANGER OF INFECTION.
	WARNING! WHEN MONITORING PACEMAKER PATIENTS, HEART RATE METERS MAY CONTINUE TO COUNT THE PACEMAKER RATE DURING OCCURRENCES OF CARDIAC ARREST OR SOME ARRHYTHMIAS. DO NOT RELY ENTIRELY UPON HEART RATE METER ALARMS. KEEP PACEMAKER PATIENTS UNDER CLOSE SURVEILLANCE. SEE 16.7.1 HEART RATE MONITORING SPECIFICATIONS ON PAGE 301 FOR DISCLOSURE OF THE PACEMAKER PULSE REJECTION CAPABILITY OF MOVES® SLCTM.
	WARNING! IMPROPER OPERATION OF THE MOVES [®] SLC™ SYSTEM COULD ENDANGER A PATIENT!
	WARNING! MOVES [®] SLC™ IS INTENDED FOR USE ON ONE PATIENT AT A TIME. FOR EXAMPLE, IT SHOULD NOT BE USED TO VENTILATE ONE PATIENT WHILE MONITORING ANOTHER.
	WARNING! DO NOT CONNECT ANY SENSORS, MONITORS, OR THE BREATHING CIRCUIT TO THE PATIENT WHILE PERFORMING SYSTEM TESTS! DOING SO COULD ENDANGER THE PATIENT!
	WARNING! FAILURE TO CHANGE THE VENTILATOR CARTRIDGE WHEN INDICATED MAY LEAD TO THE PATIENT'S SUFFERING FROM AN INCREASE IN INSPIRED CO ₂ .



SYMBOL	PATIENT-SPECIFIC WARNINGS
	WARNING! ALWAYS CARRY ALTERNATE MEANS OF VENTILATING, SUCTIONING, AND OXYGENATING THE PATIENT.
	WARNING! ALWAYS CARRY BACKUPS OF CONSUMABLES SUCH AS CARTRIDGES AND FILTERS.
	WARNING! DEFIBRILLATOR PROTECTION REQUIRES USE OF SPECIFIED ACCESSORIES, INCLUDING PATIENT CABLES AND TRANSDUCERS.
	WARNING! WHEN VENTILATING PATIENTS UNDER 30 KG OR WITH TIDAL VOLUMES UNDER 150 ML, REPLACE THE AIRWAY FILTER WITH PEDIATRIC BREATHING SYSTEM FILTER (P/N 125245) TO REDUCE DEAD SPACE VENTILATION.
	WARNING! NEVER LEAVE A PATIENT UNATTENDED WHEN RUNNING MOVES [®] SLC™.
	WARNING! MOVES [®] SLC™ PATIENT TEMPERATURE MEASUREMENT PERFORMANCE MAY DEGRADE IF THE PATIENT TEMPERATURE IS BELOW THE AMBIENT TEMPERATURE.
	EXTREME WARNING! WHEN USING FLUID FILLED PRESSURE TRANSDUCERS TO MONITOR INTRACRANIAL PRESSURE (ICP), MAKE SURE THAT THE TRANSDUCER AND THE LINE CONNECTING TO THE PATIENT'S DRAIN ARE FREE OF ANY AIR BUBBLES!
	EXTREME WARNING! AFTER COMPLETING FILLING THE ICP TRANSDUCER AND THE LINE, DISCONNECT THE FLUID BAG FROM THE TRANSDUCER, AND CAP THE END WITH THE STERILE CAP PRIOR TO CONNECTING THE TRANSDUCER TO THE PATIENT'S BRAIN!
	EXTREME WARNING! NEVER FLUSH THE ICP TRANSDUCER WHILE CONNECTED TO THE PATIENT!
	EXTREME WARNING! FAILURE TO OBSERVE THE PREVIOUS THREE PRECAUTIONS MAY RESULT IN SERIOUS INJURY OR DEATH!

6.5 MASIMO RAINBOW SET® PULSE CO-OXIMETER WARNINGS

Table 12: Masimo Rainbow SET® Pulse CO-Oximeter Warnings

SYMBOL	MASIMO RAINBOW SET® PULSE CO-OXIMETER WARNINGS
	WARNING! THE MASIMO RAINBOW SET [®] PULSE CO-OXIMETER PARAMETERS ARE FOR REFERENCE ONLY, AND THERAPEUTIC DECISIONS NEED TO BE MADE IN THE CONTEXT OF CLINICAL ASSESSMENT.
	WARNING! PULSE RATE MEASUREMENT IS BASED ON THE OPTICAL DETECTION OF A PERIPHERAL FLOW PULSE AND THEREFORE MAY NOT DETECT CERTAIN ARRHYTHMIAS. THE PULSE OXIMETER SHOULD NOT BE USED AS A REPLACEMENT OR SUBSTITUTE FOR ECGBASED ARRHYTHMIA ANALYSIS.
	WARNING! A PULSE CO-OXIMETER SHOULD BE CONSIDERED AN EARLY WARNING DEVICE. AS A TREND TOWARDS PATIENT HYPOXEMIA IS INDICATED, BLOOD SAMPLES SHOULD BE ANALYZED BY LABORATORY INSTRUMENTS TO COMPLETELY UNDERSTAND THE PATIENT'S CONDITION.



SYMBOL | MASIMO RAINBOW SET® PULSE CO-OXIMETER WARNINGS



WARNING! FOR MEASUREMENTS OF HIGH OR LOW SPHB READINGS, BLOOD SAMPLES SHOULD BE ANALYZED BY LABORATORY INSTRUMENTS TO COMPLETELY UNDERSTAND THE PATIENT'S CONDITION.



WARNING! SPO2 IS EMPIRICALLY CALIBRATED TO FUNCTIONAL ARTERIAL OXYGEN SATURATION IN HEALTHY ADULT VOLUNTEERS WITH NORMAL LEVELS OF CARBOXYHEMOGLOBIN (COHB) AND METHEMOGLOBIN (METHB). A PULSE OXIMETER CAN NOT MEASURE ELEVATED LEVELS OF COHB OR METHB. INCREASES IN EITHER COHB OR METHB WILL AFFECT THE ACCURACY OF THE SPO2 MEASUREMENT.

FOR INCREASED COHB: COHB LEVELS ABOVE NORMAL TEND TO INCREASE THE LEVEL OF SPO2. THE LEVEL OF INCREASE IS APPROXIMATELY EQUAL TO THE AMOUNT OF COHB THAT IS PRESENT.

NOTE: HIGH LEVELS OF COHB MAY OCCUR WITH A SEEMINGLY NORMAL SPO2. WHEN ELEVATED LEVELS OF COHB ARE SUSPECTED, LABORATORY ANALYSIS (CO-OXIMETRY) OF A BLOOD SAMPLE SHOULD BE PERFORMED.

FOR INCREASED METHB: THE SPO₂ MAY BE DECREASED BY LEVELS OF METHB OF UP TO APPROXIMATELY 10% TO 15%. AT HIGHER LEVELS OF METHB, THE SPO₂ MAY TEND TO READ IN THE LOW TO MID 80S. WHEN ELEVATED LEVELS OF METHB ARE SUSPECTED, LABORATORY ANALYSIS (CO-OXIMETRY) OF A BLOOD SAMPLE SHOULD BE PERFORMED.



WARNING! INTERFERING SUBSTANCES: DYES, OR ANY SUBSTANCE CONTAINING DYES, THAT CHANGE USUAL BLOOD PIGMENTATION MAY CAUSE ERRONEOUS READINGS.



WARNING! HEMOGLOBIN SYNTHESIS DISORDERS MAY CAUSE ERRONEOUS SPHB READINGS.



WARNING! ELEVATED LEVELS OF TOTAL BILIRUBIN MAY LEAD TO INACCURATE SPO₂, SPMET, SPCO, SPHB, AND SPOC MEASUREMENTS.



WARNING! MOTION ARTIFACT MAY LEAD TO INACCURATE SPMET, SPCO, SPHB, AND SPOC MEASUREMENTS.



WARNING! SEVERE ANEMIA MAY CAUSE ERRONEOUS SPO2 READINGS.



WARNING! VERY LOW ARTERIAL OXYGEN SATURATION (SPO2) LEVELS MAY CAUSE INACCURATE SPCO AND SPMET MEASUREMENTS.



WARNING! WITH VERY LOW PERFUSION AT THE MONITORED SITE, THE READINGS MAY READ LOWER THAN CORE ARTERIAL OXYGEN SATURATION.



WARNING! DO NOT USE TAPE TO SECURE THE SENSOR TO THE SITE; THIS CAN RESTRICT BLOOD FLOW AND CAUSE INACCURATE READINGS. USE OF ADDITIONAL TAPE CAN CAUSE SKIN DAMAGE OR DAMAGE THE SENSOR.



WARNING! IF THE SENSOR IS WRAPPED TOO TIGHTLY, OR SUPPLEMENTAL TAPE IS USED, VENOUS CONGESTION / PULSATIONS MAY OCCUR, CAUSING ERRONEOUS READINGS.



SYMBOL	MASIMO RAINBOW SET® PULSE CO-OXIMETER WARNINGS
	WARNING! VENOUS CONGESTION MAY CAUSE UNDER READING OF ACTUAL ARTERIAL OXYGEN SATURATION. THEREFORE, ASSURE PROPER VENOUS OUTFLOW FROM MONITORED SITE. SENSOR SHOULD NOT BE BELOW HEART LEVEL (E.G., SENSOR ON HAND OF A PATIENT IN A BED WITH ARM DANGLING TO THE FLOOR).
	WARNING! VENOUS PULSATIONS MAY CAUSE ERRONEOUS LOW READINGS (E.G.,TRICUSPID VALVE REGURGITATION).
	WARNING! LOSS OF PULSE SIGNAL CAN OCCUR WHEN:
	THE SENSOR IS TOO TIGHT. THE PATIENT HAS HYPOTENSION, SEVERE VASOCONSTRICTION, SEVERE ANEMIA, OR HYPOTHERMIA. THERE IS ARTERIAL OCCLUSION PROXIMAL TO THE SENSOR. THE PATIENT IS IN CARDIAC ARREST OR IS IN SHOCK.
	WARNING! THE PULSATIONS FROM INTRA-AORTIC BALLOON SUPPORT CAN BE ADDITIVE TO THE PULSE RATE ON THE OXIMETER PULSE RATE DISPLAY. VERIFY PATIENT'S PULSE RATE AGAINST THE ECG HEART RATE.
	WARNING! MISAPPLIED SENSORS OR SENSORS THAT BECOME PARTIALLY DISLODGED MAY CAUSE EITHER OVER OR UNDER READING OF ACTUAL ARTERIAL OXYGEN SATURATION.
	WARNING! AVOID PLACING THE SENSOR ON ANY EXTREMITY WITH AN ARTERIAL CATHETER OR BLOOD PRESSURE CUFF.
	WARNING! HIGH INTENSITY EXTREME LIGHTS (INCLUDING PULSATING STROBE LIGHTS) DIRECTED ON THE SENSOR MAY NOT ALLOW THE PULSE CO-OXIMETER TO OBTAIN READINGS.
	WARNING! THE PULSE CO-OXIMETER CAN BE USED DURING DEFIBRILLATION, BUT THE READINGS MAY BE INACCURATE FOR UP TO 20 SECONDS.
	WARNING! BEFORE USE, CAREFULLY READ THE MASIMO SENSOR <i>DIRECTIONS FOR USE</i> .
	WARNING! TISSUE DAMAGE CAN BE CAUSED BY INCORRECT APPLICATION OR USE OF A SENSOR, FOR EXAMPLE BY WRAPPING THE SENSOR TOO TIGHTLY. INSPECT THE SENSOR SITE AS DIRECTED IN THE SENSOR'S <i>DIRECTIONS FOR USE</i> TO ENSURE SKIN INTEGRITY AND CORRECT POSITIONING AND ADHESION OF THE SENSOR.
	WARNING! THE PULSE CO-OXIMETER IS NOT INTENDED FOR USE AS AN APNEA MONITOR.
	WARNING! TO AVOID CROSS CONTAMINATION USE ONLY MASIMO SINGLE USE SENSORS ON THE SAME PATIENT.
	WARNING! UNLESS OTHERWISE SPECIFIED, DO NOT STERILIZE SENSORS OR PATIENT CABLES BY IRRADIATION, STEAM, AUTOCLAVE OR ETHYLENE OXIDE. SEE THE CLEANING INSTRUCTIONS IN THE <i>DIRECTIONS FOR USE</i> FOR THE MASIMO RE-USEABLE SENSORS.



CVMDOL	MASIMO RAINBOW SET® PULSE CO-OXIMETER WARNINGS
SYMBOL	
	WARNING! DO NOT USE THE PULSE CO-OXIMETER OR OXIMETRY SENSORS DURING MAGNETIC RESONANCE IMAGING (MRI) SCANNING. INDUCED CURRENT COULD POTENTIALLY CAUSE BURNS. THE PULSE CO-OXIMETER MAY AFFECT THE MRI IMAGE, AND THE MRI UNIT MAY AFFECT THE ACCURACY OF THE OXIMETRY MEASUREMENTS.
	WARNING! IF USING PULSE CO-OXIMETRY DURING FULL BODY IRRADIATION, KEEP THE SENSOR OUT OF THE IRRADIATION FIELD. IF THE SENSOR IS EXPOSED TO THE IRRADIATION, THE READING MIGHT BE INACCURATE, OR THE UNIT MIGHT READ ZERO FOR THE DURATION OF THE ACTIVE IRRADIATION PERIOD.
	WARNING! EXERCISE CAUTION WHEN APPLYING A SENSOR TO A SITE WITH COMPROMISED SKIN INTEGRITY. APPLYING TAPE OR PRESSURE TO SUCH A SITE MAY REDUCE CIRCULATION AND/OR CAUSE FURTHER SKIN DETERIORATION.
	WARNING! CIRCULATION DISTAL TO THE SENSOR SITE SHOULD BE CHECKED ROUTINELY.
	WARNING! A FUNCTIONAL TESTER CANNOT BE UTILIZED TO ASSESS THE ACCURACY OF THE PULSE CO-OXIMETER OR ANY SENSORS.
	WARNING! DO NOT MODIFY OR ALTER A PULSE CO-OXIMETER SENSOR IN ANY WAY. ALTERATIONS OR MODIFICATION MAY AFFECT PERFORMANCE AND/OR ACCURACY.
	WARNING! DO NOT USE DAMAGED SENSORS OR PATIENT CABLES. DO NOT USE A SENSOR OR PATIENT CABLE WITH EXPOSED OPTICAL OR ELECTRICAL COMPONENTS.
	WARNING! DO NOT ATTEMPT TO REPROCESS, RECONDITION OR RECYCLE ANY MASIMO SENSORS OR PATIENT CABLES AS THESE PROCESSES MAY DAMAGE THE ELECTRICAL COMPONENTS, POTENTIALLY LEADING TO HARM.
	WARNING! EXPLOSION HAZARD. DO NOT USE THE PULSE CO-OXIMETER IN THE PRESENCE OF FLAMMABLE ANESTHETICS OR OTHER FLAMMABLE SUBSTANCE IN COMBINATION WITH AIR, OXYGEN-ENRICHED ENVIRONMENTS, OR NITROUS OXIDE.
	WARNING! AS WITH ALL MEDICAL EQUIPMENT, CAREFULLY ROUTE PATIENT CABLING TO REDUCE THE POSSIBILITY OF PATIENT ENTANGLEMENT OR STRANGULATION.
	WARNING! ALWAYS REMOVE THE SENSOR FROM THE PATIENT AND COMPLETELY DISCONNECT THE PATIENT FROM THE PULSE CO-OXIMETER BEFORE BATHING THE PATIENT.
	WARNING! DO NOT IMMERSE THE SENSOR OR PATIENT CABLE IN WATER, SOLVENTS, OR CLEANING SOLUTIONS (THE SENSORS AND CONNECTORS ARE NOT WATERPROOF).
	WARNING! INTRAVASCULAR DYES OR EXTERNALLY APPLIED COLORING (SUCH AS NAIL POLISH) MAY LEAD TO INACCURATE SPO ₂ MEASUREMENTS.
	WARNING! PATIENT SAFETY – IF A SENSOR IS DAMAGED IN ANY WAY, DISCONTINUE USE IMMEDIATELY.
	WARNING! FAILURE TO APPLY THE SENSOR PROPERLY MAY CAUSE INCORRECT MEASUREMENTS.



SYMBOL	MASIMO RAINBOW SET® PULSE CO-OXIMETER WARNINGS
	WARNING! USE ONLY MASIMO SENSORS FOR PULSE OXIMETRY OR PULSE CO-OXIMETRY MEASUREMENTS.
	WARNING! SPO ₂ SENSOR APPLICATION SITES SHOULD BE INSPECTED AT LEAST EVERY FOUR (4) HOURS, OR AS DIRECTED IN THE SENSOR'S <i>DIRECTIONS FOR USE</i> , TO ENSURE CORRECT SENSOR ALIGNMENT AND SKIN INTEGRITY. IF CIRCULATION OR SKIN INTEGRITY IS COMPROMISED, THE SENSOR SHOULD BE APPLIED TO A DIFFERENT SITE. PATIENT SENSITIVITY MAY VARY DUE TO MEDICAL STATUS OR SKIN CONDITION. DISCONTINUE THE USE OF ADHESIVE TAPE STRIPS IF THE PATIENT EXHIBITS AN ALLERGIC REACTION TO THE ADHESIVE MATERIAL.
	WARNING! WHEN USING SPO ₂ ABOVE 41°C, TAKE EXTRA CAUTION TO ENSURE SENSOR IS PLACED WITHOUT EXCESSIVE PRESSURE AND CHANGE THE APPLICATION SITE MORE

6.6 GENERAL CAUTIONS

FREQUENTLY.

Table 13: General Cautions

SYMBOL	GENERAL CAUTION
<u>^i</u>	CAUTION! ALWAYS MAKE SURE THAT MOVES [®] SLC™ IS CLAMPED AND FULLY SECURED WHEN IN USE.
<u> </u>	CAUTION! OPERATION OF MOVES [®] SLC™ OUTSIDE OF SPECIFIED LIMITS MAY CAUSE INACCURATE RESULTS.
<u> </u>	CAUTION! OPERATING SUCTION WITHOUT THE SUPPLIED FILTER, CANISTER, AND HOSES WILL RESULT IN PERMANENT FAILURE OF THE SUCTION SYSTEM AND THE OXYGEN CONCENTRATOR.
	CAUTION! MOVES [®] SLC™ SHOULD NOT BE USED ADJACENT TO OR STACKED WITH OTHER EQUIPMENT. IF ADJACENT OR STACKED USE IS NECESSARY, MOVES [®] SLC™ SHOULD BE OBSERVED TO VERIFY NORMAL OPERATION IN THE CONFIGURATION IN WHICH IT WILL BE USED.
	CAUTION! THE USE OF ACCESSORIES AND CABLES OTHER THAN THOSE SPECIFIED, WITH THE EXCEPTION OF ACCESSORIES AND CABLES QUALIFIED AND SOLD BY THORNHILL RESEARCH INC., MAY RESULT IN INCREASED EMISSIONS OR DECREASED IMMUNITY OF MOVES [®] SLC™ AND MAY CAUSE THE SYSTEM TO BE NON-COMPLIANT WITH THE REQUIREMENTS OF IEC 60601-1-2:2007 (ED 3.0).
<u>^i</u>	CAUTION! SOME PATIENT MONITORING ACCESSORIES MAY NOT FUNCTION PROPERLY OR MAY RELAY INACCURATE READINGS IF OPERATED OUTSIDE OF THEIR NORMAL OPERATING CONDITIONS.
	CAUTION! USE ONLY THE ACCESSORIES THAT ARE PROVIDED WITH THE MOVES [®] SLC™ SYSTEM! OTHER ACCESSORIES MAY ADVERSELY AFFECT THE VENTILATOR PERFORMANCE.
	CAUTION! THE MOVES® SLC™ ACCESSORIES PROVIDED WITH THE MOVES® SLC™ SYSTEM ARE FOR USE ONLY WITH THE MOVES® SLC™ SYSTEM.



SYMBOL	GENERAL CAUTION
	CAUTION! BEFORE INSTALLING A HYDROCARBON FILTER, CHECK THE FOUR-DIGIT DATE CODE PRINTED ON THE CARTRIDGE. THE CARTRIDGE LABEL IS STAMPED WITH FOUR CHARACTERS "XXYY", WHERE "XX" IS THE WEEK OF THE YEAR AND "YY" IS THE YEAR. A CARTRIDGE MORE THAN THREE YEARS OLD SHOULD BE DISCARDED SINCE IT MAY DEGRADE THE PERFORMANCE OF OR CAUSE DAMAGE TO THE MOVES [®] SLC™ OXYGEN CONCENTRATOR.
	CAUTION! OPERATING MOVES [®] SLC™ WITHOUT A HYDROCARBON FILTER WILL DAMAGE THE UNIT. DO NOT OPERATE WITHOUT A HYDROCARBON FILTER!
	CAUTION! A MOVES [®] SLC™ UNIT SHOULD NEVER BE PUT INTO TRANSPORT SERVICE WITH LESS THAN A 95% CHARGE IN BOTH BATTERIES.
<u> </u>	CAUTION! NEVER CHARGE BATTERIES IN AMBIENT TEMPERATURES BELOW 32°F (0°C) OR ABOVE 104°F (40°C).
	CAUTION! IF INTENDING TO RUN ON BATTERIES, ENSURE THAT THERE IS SUFFICIENT POWER FOR THE LENGTH OF TIME REQUIRED, OR REPLACE THE BATTERIES.
<u> </u>	CAUTION! ONLY SELECT "SKIP TESTS" IF A SYSTEM TEST HAS PREVIOUSLY BEEN PERFORMED AND PASSED, AND THE SYSTEM HAS JUST BEEN RESTARTED DUE TO POWER FAILURE (E.G., LOSS OF BATTERY OR EXTERNAL POWER).
	CAUTION! THE SUCTION WAND SUPPLIED WITH THE MOVES [®] SLC™ IS DESIGNED TO MINIMIZE TOTAL OCCLUSION. IT IS STRONGLY ADVISED THAT ONLY THE WAND SUPPLIED BE USED WITH THE MOVES [®] SLC™ SYSTEM.
<u> </u>	CAUTION! ACCURACY OF ANY BLOOD PRESSURE MEASUREMENT MAY BE AFFECTED IF MOVES [®] SLC™ IS USED OR STORED OUTSIDE THE RELEVANT TEMPERATURE OR HUMIDITY RANGES DESCRIBED IN THE ENVIRONMENTAL SPECIFICATIONS (SEE SECTION 16.6.7 ON PAGE 298).
	CAUTION! SUBSTITUTION OF COMPONENTS DIFFERENT FROM THOSE SUPPLIED WITH THE MOVES [®] SLC™ MAY RESULT IN MEASUREMENT ERROR.
	CAUTION! THE SUCTION CANISTER AND SUCTION FILTER ARE INTENDED FOR SINGLE USE ONLY AND SHOULD BE DISPOSED OF IN ACCORDANCE WITH LOCAL BIOHAZARD REGULATIONS.
	CAUTION! BREATHING CIRCUITS, SAMPLE LINES (<u>BUT NOT THE NAFION TUBE</u>) AND FILTERS, BREATHING CARTRIDGES, ECG ADHESIVE SENSOR PADS, ABP/CVP/ICP TRANSDUCER AND SUCTION WAND AND TUBING ARE DISPOSABLE AND SHOULD BE DISPOSED OF IN ACCORDANCE WITH LOCAL BIOHAZARD REGULATIONS.
	CAUTION! ALL MOVES [®] SLC™ ACCESSORIES' PACKAGING AND DISPOSABLE ACCESSORIES SHOULD BE DISPOSED OF RESPONSIBLY IN ACCORDANCE WITH LOCAL WASTE DISPOSAL STANDARDS AND / OR LOCAL BIOHAZARD REGULATIONS.
	CAUTION! DO NOT SUBMERGE THE MOVES [®] SLC™ OR POUR CLEANING LIQUIDS OVER OR INTO THE MOVES [®] SLC™.



SYMBOL	GENERAL CAUTION
	CAUTION! THE LABEL ON THE PACKAGE OF THE VENTILATOR CARTRIDGE CONTAINS AN EXPIRY DATE. ALWAYS CHECK THE EXPIRY DATE ON THE VENTILATOR CARTRIDGE BEFORE USING IT TO MAKE SURE THAT THE VENTILATOR CARTRIDGE HAS NOT EXPIRED. AS WELL, MONITOR SPARE CARTRIDGES WITH REGARD TO THEIR REMAINING "SHELF LIFE".
<u> </u>	CAUTION! THE SURFACE OF THE MOVES [®] SLC™ SYSTEM CAN BECOME HOT, ESPECIALLY IF IT IS BEING OPERATED IN DIRECT SUNLIGHT. CARE SHOULD BE TAKEN WHEN TOUCHING OR CONTACTING THE SURFACE OF THE MOVES [®] SLC™ SYSTEM.
<u> </u>	CAUTION! ECG CABLES SHOULD BE DISCARDED AND REPLACED AFTER TWO (2) YEARS OF CONTINUOUS USE. CHECK CABLE USE BY RECORDING THE DATE THE CABLE WAS FIRST USED.
	CAUTION! CHECK THE EXPIRY DATE ON THE ECG ELECTRODES PACKAGE BEFORE USING. ELECTRODES ARE GOOD FOR 45 DAYS ONCE PACKAGE IS OPENED.
<u>^</u>	CAUTION! PAY SPECIAL ATTENTION TO THE TYPE OF ECG ELECTRODES USED. SOME ELECTRODES MAY BE SUBJECT TO LARGE OFFSET POTENTIALS. RECOVERY TIME AFTER DEFIBRILLATION MAY BE ESPECIALLY COMPROMISED.
	CAUTION! WHEN YOU DETACH THE MOVES [®] SLC™ CLAMPS, AND RETURN THEM TO THE ACCESSORIES CASE, BE CAREFUL NOT TO PINCH THE WIRES ATTACHED TO THE PINS IN THE CLAMP APPARATUS AS THIS CAN CAUSE WEAR, ABRADING, AND EVENTUAL BREAKAGE OF THE WIRES.
	CAUTION! WHEN BATTERIES ARE DISCHARGED AND LEFT IN MOVES [®] SLC™ FOR A PROLONGED PERIOD, A <u>COMPLETELY</u> DISCHARGED BATTERY (NO LED LIGHTS) CAN RESULT. THE BATTERY CAN STILL BE RECHARGED, BUT IT MAY TAKE MORE THAN THE NORMAL 2.5 HOURS. IT HAS BEEN OBSERVED TO TAKE ANYWHERE FROM 6 TO 48 HOURS TO FULLY CHARGE.
	CAUTION! IF MOVES [®] SLC [™] IS EXPOSED TO SIGNIFICANT AMOUNTS OF SAND OR DUST, IT SHOULD BE CLEANED BY AN AUTHORIZED TECHNICIAN IN ACCORDANCE WITH THE MAINTENANCE MANUAL.
	CAUTION! THE VENTILATOR BREATHING CIRCUIT AND THE SAMPLING FILTER CONNECTED TO THE NAFION TUBING SHOULD BE INSPECTED EVERY FOUR (4) HOURS FOR CONDENSATION AND DRAINED AS REQUIRED. THE CIRCUIT AND FILTER SHOULD BE CHANGED AFTER 24 HOURS OF CONTINUOUS USE.
	CAUTION! NO LUBRICANTS OTHER THAN THOSE RECOMMENDED BY THE MANUFACTURER SHALL BE USED ON THE MOVES [®] SLC™.

6.7 ELECTRICAL CAUTIONS

Table 14: Electrical Cautions

SYMBOL	ELECTRICAL CAUTION
<u></u>	CAUTION! MOVES [®] SLC™ IS INTENDED FOR USE BY HEALTHCARE PROFESSIONALS ONLY. MOVES [®] SLC™ MAY CAUSE RADIO INTERFERENCE OR MAY DISRUPT THE OPERATION OF NEARBY EQUIPMENT. IT MAY BE NECESSARY TO TAKE MITIGATION MEASURES, SUCH AS REORIENTING OR RELOCATING MOVES [®] SLC™ OR SHIELDING THE LOCATION.



SYMBOL	ELECTRICAL CAUTION
	CAUTION! MEDICAL ELECTRICAL EQUIPMENT NEEDS SPECIAL PRECAUTIONS REGARDING EMC (ELECTROMAGNETIC COMPATIBILITY) AND NEEDS TO BE INSTALLED AND PUT INTO SERVICE ACCORDING TO THE EMC INFORMATION PROVIDED IN THE MOVES [®] SLC™ OPERATOR'S MANUAL.
<u></u>	CAUTION! THE POWER SUPPLY/CHARGER CORD IS A SPECIAL MEDICAL-GRADE POWER CORD AND SHOULD NOT BE REPLACED WITH A NON-MOVES [®] SLC™ SUPPLIED PART.
	CAUTION! PORTABLE AND MOBILE RF (RADIO FREQUENCY) COMMUNICATIONS EQUIPMENT CAN AFFECT MEDICAL ELECTRICAL EQUIPMENT.
	CAUTION! INTERFERENCE MAY OCCUR IN THE VICINITY OF KNOWN RADIO FREQUENCY (RF) TRANSMITTING DEVICES AND EQUIPMENT MARKED WITH THE FOLLOWING SYMBOL:
	CAUTION! OXIMETER READINGS MAY BE AFFECTED BY THE USE OF AN ELECTROSURGICAL UNIT (ESU).

6.8 PATIENT-SPECIFIC CAUTIONS

Table 15: Patient-Specific Cautions

SYMBOL	PATIENT-SPECIFIC CAUTION
	CAUTION! ALL COMPONENTS IN MOVES [®] SLC [™] THAT COME IN CONTACT WITH RESPIRATORY GASES ARE NON-STERILE.

6.9 GENERAL SAFETY

- 1. This equipment should be operated only by a trained medical practitioner.
- 2. The operator must inspect the MOVES[®] SLC[™] unit and all accessories for visible physical damage (cracks, holes, leaks; missing components, structural hardware and protective covers; signs of tampering, etc.) prior to each use. All system tests must be completed prior to connecting the MOVES[®] SLC[™] unit to a patient.
- 3. The operator must be fully familiar with the contents of this manual before operating the MOVES[®] SLC™ system.
- 4. The MOVES[®] SLC[™] unit must be serviced only by qualified personnel. There are no user serviceable parts inside the MOVES[®] SLC[™] unit.
- 5. The MOVES® SLCTM system must not be used for any purpose other than as stated in section 3.2 MOVES® SLC^{TM} Intended Use on page 19.
- 6. The MOVES[®] SLC[™] unit must always be used and stored in accordance with the environmental specifications listed in the subsection *Environmental Specifications on page 299*.
- 7. Any storage or use outside of these conditions may cause system degradation and harm to patients. After any such storage or use, the MOVES[®] SLC™ unit must be serviced by qualified personnel. Do not use the MOVES[®] SLC™ unit in contaminated environments.



- 8. The operator must ensure that the MOVES[®] SLC[™] Unit and Accessories Cases contain all the necessary components for successful use. See Section 9.1: MOVES® SLC[™] System Contents on page 61.
- 9. Always ensure that system batteries are fully charged, and always maintain alternative methods of ventilation, O₂ supplementation and suction in case of failure of part or all of the MOVES[®] SLC™ system.
- 10. If the MOVES[®] SLC™ unit and accessories are placed in storage, they must be inspected and serviced every 12 months at a minimum.
- 11. The MOVES[®] SLC[™] system must be operated using only accessories specified in this manual, and supplied by the manufacturer, or their equivalent.
- 12. The accessories provided with the MOVES[®] SLC™ system are for use only with the MOVES[®] SLC™ system. Use of MOVES[®] SLC™ accessories with another system may result in failure of the accessories and present a risk to and/or harm a patient, operator or bystander.
- 13. The MOVES[®] SLC[™] system must not be connected to a patient unless the operator has thoroughly read and understood this manual, the operator is qualified or is under qualified supervision, all system tests have been performed, and all of the Safety Procedures contained within this manual have been read and adhered to.
- 14. The MOVES[®] SLC[™] unit should be placed in *Monitor Only* mode when it is operating in 'Ventilator' or 'O2 Supplement' mode and filters or cartridges need to be changed or if the breathing circuit is disconnected for any reason.
- 15. The MOVES[®] SLC™ unit must not be operated without filters and cartridges present and properly installed.
- 16. Failure to change the ventilator cartridge when indicated will lead to hypercarbia. Alternative ventilation and/or external O₂ supplementation may be required while changing the cartridge.
- 17. Cartridges are for single patient use only. A Ventilator Cartridge must be replaced after two (2) hours of continuous use under standard temperature and pressure conditions. However, challenging environmental and patient conditions may reduce the life of the Ventilator Cartridge to less than two (2) hours. Always have a spare cartridge and an alternative ventilation method available.
- 18. The Ventilator Cartridge has an expiry date listed on the package label. Before using the Ventilator Cartridge, always check the expiry date to make sure that the cartridge has not expired.
- 19. Always have available an alternative means of suction in the event of a power failure, mechanical failure or serious occlusion in the suction circuit.
- 20. The MOVES[®] SLC[™] system is not recommended for use in treating patients with pacemakers or other implanted medical devices.
- 21. If the MOVES[®] SLC[™] unit should experience ingress of any particulate matter (dirt, dust, sand, etc.) through suction, O₂ inlet, ventilator driving gas or valve block ports, the unit should be cleaned and serviced by authorized personnel.
- 22. Should the MOVES[®] SLC[™] unit experience ingress of any liquids or become contaminated by bodily fluids through suction, O₂ inlet, air intake, ventilator driving gas or valve block ports, the unit should be transferred to authorized personnel for cleaning, sterilization and/or component replacement.
- 23. Should the MOVES[®] SLC™ unit experience a sudden shock (by being dropped or roughly handled), or exposed externally to high voltage, the unit should be inspected and serviced by authorized personnel.
- 24. The MOVES[®] SLC[™] system outputs concentrated oxygen. No smoking or open flame is permitted near the unit.
- 25. Do not use oil or grease on or near the MOVES[®] SLC™ unit or its components.
- 26. The surface of the MOVES[®] SLC™ system can become hot, especially if it is being operated in direct sunlight. Care should be taken when touching or contacting the surface of the MOVES[®] SLC™ system.
- 27. There are no known toxic effects of any materials used in the system that can come in contact with the patient, the operator or the gas delivered to the patient.



6.10 ELECTRICAL SAFETY

- Connect the MOVES[®] SLC[™] power supply / charger only to an AC source from 100–240V, 50–60 Hz,
 A max. Fluctuations in voltage and current can have adverse effects on the performance of the MOVES[®] SLC[™] system.
- 2. Do not modify the power supply provided with the MOVES[®] SLC[™] system with additional voltage regulators or similar equipment. The MOVES[®] SLC[™] power supply/charger detects the supply automatically.
- 3. Do not route power cables immediately adjacent to patient connection cables. Power cables can produce voltage transients ("crosstalk") that seriously affect data collection cables, especially ECG sensor connections. Voltage transients can resemble ECG readings.
- 4. Do not operate the MOVES[®] SLC™ unit if the main AC cord or the power-supply cord to MOVES[®] SLC™ unit shows any sign of damage such as frayed insulation, or if there are cracked or damaged plugs or receptacles, or a missing grounding plug, or if the 'PWR" light on the power supply/charger does not illuminate.
- 5. Do not operate the MOVES[®] SLC[™] system on battery power if:
 - a. A battery shows any sign of damage such as cracks, holes, or leakage.
 - b. Any battery is known to be defective.
 - c. The battery discharge indicator on either battery fails to illuminate.
 - d. The power source indicator on the startup screen of the MOVES[®] SLC™ unit fails to detect a battery when a battery is installed, or it indicates a charge that is not in agreement with the indicators on the batteries.
 - e. The power source indicator on the startup screen of the MOVES[®] SLC™ fails to show a decrease in charge while running.
 - f. The 'PWR" light on the power supply/charger does not illuminate when attempting to charge the batteries.
- 6. The main external AC cord is supplied with a grounding prong on the male connector to reduce the risk of electrical shock. Never remove or compromise this prong.
- 7. Use only the MOVES[®] SLC[™] power supply P/N 111422, batteries P/N 126071, and supplied external power cables to operate the MOVES[®] SLC[™] system.

6.11 PREPARING FOR EMERGENCY OPERATION

The operator must always be prepared for emergency situations such as power failure, mechanical failure or serious occlusions in the MOVES[®] SLC™ or its accessories that may require alternative means of treating a patient.

The following guidelines should always be adhered to:

- 1. The operator must always have prepared two completely charged battery packs.
- 2. The operator must always have prepared an alternative means of ventilating a patient.
- 3. The operator must always have prepared an alternative means of providing a high concentration of oxygen to a patient.
- 4. The operator must always have prepared an alternative means of monitoring a patient's vital signs (e.g., heart rate, blood pressure, respiratory rate, etc.).
- 5. The operator must always have prepared an alternative means of applying suction to a patient.
- 6. When treating a patient and operating the MOVES[®] SLC[™] on external power, the operator must always have at least one charged battery installed in the MOVES[®] SLC[™] unit. In the event of an AC power failure, the MOVES[®] SLC[™] unit will immediately switch to battery power.



6.12 RADIO INTERFERENCE



CAUTION! INTERFERENCE MAY OCCUR IN THE VICINITY OF KNOWN RADIO FREQUENCY (RF) TRANSMITTING DEVICES AND EQUIPMENT MARKED WITH THE FOLLOWING SYMBOL:



6.13 BATTERY HANDLING

Please note the following important cautions regarding battery handling:

- The batteries should not be opened, destroyed nor incinerated since they may leak or rupture and release into the environment the ingredients they contain.
- The batteries are designed to be recharged. However, improperly charging a battery may cause the battery to flame.
- Use only the MOVES[®] SLC[™] battery charger (P/N 111422) and MOVES[®] SLC[™] battery charging procedures.
- NEVER disassemble a battery or bypass any safety device.
- Do not crush, pierce, or short battery terminals with conductive (i.e., metal) goods.
- · Do not directly heat or solder.
- Do not throw into fire.

6.14 BATTERY DISPOSAL

MOVES[®] SLC[™] batteries are based on Lithium Polymer (LiPo) chemistry. Always dispose of batteries in accordance with local municipal, state, and federal regulations.



WARNING! CHARGING AND DISCHARGING THE LITHIUM POLYMER BATTERIES WITH OTHER THAN MOVES[®] SLC™ SYSTEM EQUIPMENT, AND/OR IMPROPER HANDLING, CAN RESULT IN FIRE, EXPLOSION, TOXIC GASES AND SMOKE.



WARNING! DO NOT OPERATE THE MOVES® SLC™ SYSTEM WITH A DEFECTIVE BATTERY.



7.0 The Masimo Rainbow SET® Pulse CO-Oximeter

7.1 NO IMPLIED LICENSE

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

7.2 OVERVIEW

The MOVES[®] SLC[™] uses the Masimo Rainbow SET[®] Pulse CO-Oximeter to provide non-invasive monitoring that measures arterial oxygen saturation (SpO₂₎, pulse rate (PR), and perfusion index (PI), along with optional measurements of hemoglobin (SpHb), total oxygen content (SpOC), carboxyhemoglobin (SpCO), methemoglobin (SpMet), and pleth variability index (PVI).

The Masimo Rainbow SET[®] Pulse CO-Oximeter and accessories have been validated and are indicated for use with pediatric and adult patients during both no motion and motion conditions, and for patients who are well or poorly perfused in hospitals, hospital-type facilities, and transport.

7.3 KEY FEATURES

- Masimo SET[®] is clinically proven to satisfy all sensitivity and specificity requirements for pulse oximeter technology.
- Rainbow technology uses 7+ wavelengths of light to continuously and noninvasively measure carboxyhemoglobin (SpCO), methemoglobin (SpMet), and total hemoglobin (SpHb), as well as providing a more reliable probe-off detection.
- Total oxygen content (SpOC) provides a calculated measurement of the amount of oxygen in arterial blood, which may provide useful information about oxygen both dissolved in plasma and combined with hemoglobin.
- Pleth Variability Index (PVI) may show changes that reflect physiologic factors such as vascular tone, circulating blood volume, and intrathoracic pressure excursions. [The utility of PVI is unknown at this time and requires further clinical studies. Technical factors that may affect PVI include probe malposition and patient motion.]
- Perfusion Index (PI) with trending capability indicates arterial pulse signal strength and may be used as a diagnostic tool during low perfusion.

7.4 INDICATIONS FOR USE

Masimo Rainbow SET[®] Pulse CO-Oximeter and accessories are indicated for the continuous noninvasive monitoring of functional oxygen saturation of arterial hemoglobin (SpO₂), pulse rate (PR), carboxyhemoglobin saturation (SpCO), methemoglobin saturation (SpMet) and total hemoglobin concentration (SpHb).

Masimo Rainbow SET[®] Pulse CO-Oximeter and accessories have been validated and are indicated for use with pediatric and adult patients during both no motion and motion conditions, and for patients who are well or poorly perfused in hospitals, hospital-type facilities and transport.

In addition, the Masimo Rainbow SET^{\circledR} Pulse CO-Oximeter and accessories are indicated to provide continuous noninvasive monitoring data of functional oxygen saturation of arterial hemoglobin (SpO₂) and pulse rate (PR) to multi-parameter devices for the display of those devices.





WARNING! THE MASIMO RAINBOW SET® PULSE CO-OXIMETER IS NOT INTENDED FOR USE AS AN APNEA MONITOR.

7.5 PULSE OXIMETER TECHNOLOGY OVERVIEW

7.5.1 Signal Extraction Technology (SET®)

Masimo Signal Extraction Technology's signal processing differs from that of conventional pulse oximeters. Conventional pulse oximeters assume that arterial blood is the only blood moving (pulsating) in the measurement site. During patient motion, however, the venous blood also moves, causing conventional pulse oximeters to read low values, because they cannot distinguish between the arterial and venous blood movement (sometimes referred to as noise).

Masimo SET[®] pulse oximetry utilizes parallel engines and adaptive digital filtering. Adaptive filters are powerful because they are able to adapt to the varying physiologic signals and/or noise and separate them by looking at the whole signal and breaking it down to its fundamental components. The Masimo SET signal processing algorithm, Discrete Saturation Transform[®] (DST[®]), in parallel with Fast Saturation Transform (FST[®]), reliably identifies the noise, isolates it and, using adaptive filters, cancels it. It then reports the true arterial oxygen saturation for display on the monitor.

7.5.2 General Description for Oxygen Saturation (SpO₂)

Pulse oximetry is governed by the following principles:

- Oxyhemoglobin (oxygenated blood) and deoxyhemoglobin (non-oxygenated blood) differ in their absorption of red and infrared light (spectrophotometry).
- 2. The amount of arterial blood in tissue changes with your pulse (photoplethysmography). Therefore, the amount of light absorbed by the varying quantities of arterial blood changes as well.

7.5.3 Successful Monitoring for SpO₂, PR, and PI

Stability of the SpO₂ readings may be a good indicator of signal validity. Although stability is a relative term, experience will provide a good feeling for changes that are artifactual or physiological and the speed, timing, and behavior of each.

The stability of the readings over time is affected by the averaging mode being used. The longer the averaging time, the more stable the readings tend to become. This is due to a dampened response as the signal is averaged over a longer period of time than during shorter averaging times. However, longer averaging times delay the response of the oximeter and reduce the measured variations of SpO₂ and pulse rate.

7.5.4 Functional Oxygen Saturation

The Masimo Rainbow SET® Pulse CO-Oximeter is calibrated to measure and display functional oxygen saturation (SpO2): the amount of oxyhemoglobin expressed as a percentage of the hemoglobin that is available to transport oxygen.

Note that carboxyhemoglobin is not capable of transporting oxygen, but is recognized as oxygenated hemoglobin by conventional pulse oximetry.

7.5.5 General Description for Pulse Rate (PR)

Pulse rate (PR), measured in beats per minute (BPM), is based on the optical detection of peripheral flow pulse.



7.5.6 General Description for Perfusion Index (PI)

The Perfusion Index (PI) is the ratio of the pulsatile blood flow to the non-pulsatile or static blood in peripheral tissue. PI thus represents a noninvasive measure of peripheral perfusion that can be continuously and noninvasively obtained from a pulse oximeter.

7.5.7 General Description for Pleth Variability Index (PVI)

The pleth variability index (PVI) is a measure of the dynamic changes in the perfusion index (PI) that occur during the respiratory cycle. The calculation is accomplished by measuring changes in the PI over a time interval where one or more complete respiratory cycles have occurred. PVI is displayed as a percentage (0–100%).

The utility of PVI is unknown at this time and requires further clinical studies. Technical factors that may affect PVI include probe malposition and patient motion.

7.5.8 Rainbow Pulse CO-Oximetry Technology

Rainbow Pulse CO-Oximetry technology is governed by the following principles:

- 1. Oxyhemoglobin (oxygenated blood), deoxyhemoglobin (non-oxygenated blood), carboxyhemoglobin (blood with carbon monoxide content), methemoglobin (blood with oxidized hemoglobin) and blood plasma constituents differ in their absorption of visible and infrared light (using spectrophotometry).
- 2. The amount of arterial blood in tissue changes with pulse (photoplethysmography). Therefore, the amount of light absorbed by the varying quantities of arterial blood changes as well.

The Masimo Rainbow SET[®] Pulse CO-Oximeter uses a multi-wavelength sensor to distinguish between oxygenated blood, deoxygenated blood, blood with carbon monoxide, oxidized blood and blood plasma.

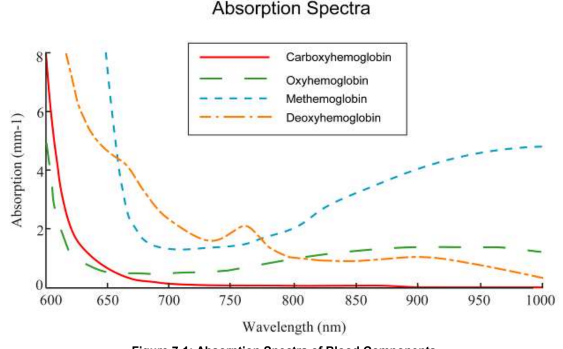


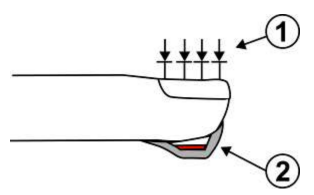
Figure 7-1: Absorption Spectra of Blood Components



NOTE: The wavelength of blood plasma, which begins near the 1000 nm range and peaks in the 1400 nm range, is omitted from the above graph since most of its wavelength falls outside of the graph parameters.



The Masimo Rainbow SET[®] Pulse CO-Oximeter utilizes a sensor with various light-emitting diodes (LEDs) that pass light through the site to a diode (detector). Signal data is obtained by passing various visible and infrared lights (LEDs, 500 to 1400 nm) through a capillary bed (for example, a fingertip, a hand, a foot) and measuring changes in light absorption during the blood pulsatile cycle. This information may be useful to clinicians. The maximum radiant power of the strongest light is rated at ≤ 25 mW. The detector receives the light, converts it into an electronic signal and sends it to the Masimo Rainbow SET[®] Pulse CO-Oximeter for calculation.



- 1. Light Emitting Diodes (LEDs) (7 + wavelengths)
- 2. Detector

Figure 7-2: LEDs and Detector

Once the Masimo Rainbow SET® Pulse CO-Oximeter receives the signal from the sensor, it utilizes proprietary algorithms to calculate the patient's functional oxygen saturation (SpO₂ [%]), blood levels of carboxyhemoglobin (SpCO [%]), methemoglobin (SpMet [%]), total hemoglobin concentration (SpHb [g/dL]) and pulse rate (PR). The SpCO, SpMet and SpHb measurements rely on a multi-wavelength calibration equation to quantify the percentage of carbon monoxide and methemoglobin and the concentration of total hemoglobin in arterial blood. In an ambient temperature of 35° C the maximum skin surface temperature has been measured at less than 106° F (41° C), verified by Masimo sensor skin temperature test procedure.

7.5.9 Pulse CO-Oximetry vs. Drawn Whole Blood Measurements

When SpO₂, SpCO, SpMet, and SpHb measurements obtained from the Masimo Rainbow SET[®] Pulse CO-Oximeter (noninvasive) are compared to drawn whole blood (invasive) measurements by blood gas and/or laboratory CO-Oximetry methods, caution should be taken when evaluating and interpreting the results.

The blood gas and/or laboratory CO-Oximetry measurements may differ from the SpO₂, SpCO, SpMet, SpHb, and SpOC measurements of the Masimo Rainbow SET[®] Pulse CO-Oximeter. Any comparisons should be simultaneous, meaning the measurement on the device should be noted at the exact time that blood is drawn.

In the case of SpO₂, different results are usually obtained from the arterial blood gas sample if the calculated measurement is not appropriately corrected for the effects of variables that shift the relationship between the partial pressure of oxygen (PO₂) and saturation, such as: pH, temperature, the partial pressure of carbon dioxide (PCO₂), 2,3-DPG, and fetal hemoglobin. In the case of SpCO, different results are also expected if concentration of methemoglobin in the blood gas sample is abnormal (greater than 2% for methemoglobin concentration).

High levels of bilirubin may cause erroneous SpO₂, SpMet, SpCO, and SpHb readings. As blood samples are usually taken over a period of 20 seconds (the time it takes to draw the blood) a meaningful comparison can only be achieved if the oxygen saturation, carboxyhemoglobin, and methemoglobin concentration of the patient are stable and not changing over the period of time that the blood gas sample is taken. Subsequently, blood gas and laboratory CO-Oximetry measurements of SpO₂, SpCO, SpMet, SpHb, and SpOC may vary with the rapid administration of fluids and in procedures such as dialysis. Additionally, drawn whole blood testing can be affected by sample handling methods and time elapsed between blood draw and sample testing.

Measurements with Low Signal IQ should not be compared to laboratory measurements.



7.5.10 General Description for Total Hemoglobin (SpHb)

Pulse CO-Oximetry is a continuous and noninvasive method of measuring the levels of total hemoglobin (SpHb) in arterial blood. It relies on the same principles of pulse oximetry to make its SpHb measurement. The measurement is taken by a sensor capable of measuring SpHb, usually on the fingertip for pediatric and adult patients.

The sensor connects directly to the Pulse CO-Oximeter or with a patient cable. The sensor collects signal data from the patient and sends it to the instrument. The instrument displays the calculated data as measurement of total hemoglobin concentration.

7.5.11 Successful Monitoring for SpHb

A stable SpHb reading is associated with correct sensor placement, small physiological changes during the measurement and acceptable levels of arterial perfusion at the measurement site. Physiological changes at the measurement site are mainly caused by fluctuations in the oxygen saturation, blood concentration and perfusion.

7.5.12 General Description for SpOC

The following is the equation for oxygen content via the Pulse CO-Oximeter:

SpOC
$$(ml/dL^*) = 1.31 (ml O_2/g Hb) x SpHb (g/dL) x SpO_2 + 0.3 ml/dL$$

7.5.13 General Description for Carboxyhemoglobin (SpCO)

Pulse CO-Oximetry is a continuous and noninvasive method of measuring the levels of carboxyhemoglobin concentration (SpCO) in arterial blood. It relies on the same basic principles of pulse oximetry (spectrophotometry) to make its SpCO measurement.

The measurement is obtained by placing a sensor on a patient, usually on the fingertip for pediatric and adult patients. The sensor connects either directly to the Pulse CO-Oximetry instrument or through an instrument patient cable.

The sensor collects signal data from the patient and sends it to the instrument. The instrument displays the calculated data as percentage value for the SpCO, which reflect blood levels of carbon monoxide bound to hemoglobin.

7.5.14 Successful Monitoring for SpCO

A stable SpCO reading is associated with correct sensor placement, small physiological changes during the measurement and acceptable levels of arterial perfusion in the patient's fingertip (measurement site). Physiological changes at the measurement site are mainly caused by fluctuations in the oxygen saturation, blood concentration and perfusion.

7.5.15 General Description for Methemoglobin (SpMet)

Pulse CO-Oximetry is a continuous and noninvasive method of measuring the levels of methemoglobin concentration (SpMet) in arterial blood. It relies on the same basic principles of pulse oximetry (spectrophotometry) to make its SpMet measurement.

The measurement is obtained by placing a sensor on a patient, usually on the fingertip for pediatric and adult patients. The sensor connects either directly to the Pulse CO-Oximetry instrument or through a patient cable. The sensor collects signal data from the patient and sends it to the instrument. The instrument displays the calculated data as percentage value for the SpMet.



^{*}When mI O2/g Hb is multiplied by g/dL of SpHb, the gram unit in the denominator of mI/g cancels the gram unit in the numerator of g/dL resulting in mI/dL (mI of oxygen in one dL of blood) as the unit of measure for SpOC.

7.5.16 Successful Monitoring for SpMet

A stable SpMet reading is associated with correct sensor placement, small physiological changes during the measurement and acceptable levels of arterial perfusion in the patient's fingertip (measurement site).

Physiological changes at the measurement site are mainly caused by fluctuations in the oxygen saturation, blood concentration and perfusion.

7.5.17 SpCO, SpMet, and SpHb Measurements During Patient Motion

The MOVES[®] SLC™ displays measurements of SpCO, SpMet, and SpHb during patient motion. However, because of the changes in the physiological parameters such as blood volume, arterial-venous coupling, etc., that occur during patient motion, the accuracy of such measurements may not be reliable during excessive motion. In this case, the measurement value for SpCO, SpMet, or SpHb displays as dashes (---) and a message displays to alert the clinician that the instrument does not have confidence in the value due to poor signal quality caused by excessive motion or other signal interference.



8.0 Troubleshooting the Masimo Pulse Oximeter

The following chapter contains information about troubleshooting the pulse oximeter.

8.1 TROUBLESHOOTING MEASUREMENTS

The operation of the Pulse Oximeter can be verified by using a function simulator such as the Fluke ProSim 8 Vital Signs Simulator with the ProSim RAINBOW test cable (#4034609).

For more information, see Masimo Rainbow SET® Pulse CO-Oximeter Warnings beginning on page 42.

8.2 LOW MEASUREMENT CONFIDENCE

The Pulse CO-Oximeter maintains an internal measure of the signal quality of each of the displayed parameters (i.e., SpO₂, HR, PI, SpCO, SpMet, etc). When the signal quality is low enough, the accuracy of the displayed measurement may be compromised. A low confidence alarm is generated for the parameter with low signal quality to indicate that the measurement accuracy may be compromised (e.g., for SpO₂ the alarm "PulseOx: SpO₂ reading confidence poor" is generated). This will also cause the parameter to be highlighted.

When parameters have associated low confidence alarms, proceed with caution and do the following:

- Assess the patient.
- Check the sensor and ensure proper sensor application. The sensor must be well secured to the site for the Pulse CO-Oximeter to maintain accurate readings. Misalignment of the sensor's emitter and detector can result in smaller signals and cause erroneous readings.
- Determine if an extreme change in the patient's physiology and blood flow at the monitoring site occurred, (e.g., an inflated blood-pressure cuff, a squeezing motion, sampling of an arterial blood specimen from the hand containing the pulse oximetry sensor, severe hypotension, peripheral vasoconstriction in response to hypothermia, medications, or an episode of Raynaud's syndrome.)
- After performing the above, if the low perfusion and/or low confidence alarm(s) occur frequently or continuously, obtaining an arterial blood specimen for CO-Oximetry analysis may be considered to verify the oxygen saturation value.

8.3 LOW PERFUSION

It has been suggested that at extremely low perfusion levels, pulse oximeters can measure peripheral saturation, which may differ from central arterial saturation. This "localized hypoxemia" may result from the metabolic demands of other tissues extracting oxygen proximal to the monitoring site under conditions of sustained peripheral hypoperfusion. This may occur even with a pulse rate that correlates with the ECG heart rate.

8.4 LOW SIGNAL QUALITY

Improper sensor type or application.

Next steps: Excessive motion relative to perfusion. Sensor is damaged or not functioning. Check and see if blood flow to the site is restricted. Check the placement of the sensor. Reapply sensor or move to a different site.



8.5 SPO2 VALUES DO NOT CORRELATE WITH CLINICAL ASSESSMENT OR ARTERIAL BLOOD GAS MEASUREMENTS

Low perfusion or sensor displacement.

Next steps: Check for pulse oximeter alarm messages. See *Alarm Conditions and Causes beginning on page 238*. Check placement of sensor or if it is too tight. Reapply sensor or select a new site. Set to MAX sensitivity and confirm that the sensor is securely on the patient. See the directions for use provided with your sensor.

8.6 UNEXPECTED SPO2, SPCO, SPMET, OR SPHB READING

Low SIQ or PI values.

Next steps: Reposition sensor to site with strong SIQ and PI. Average readings taken from three different sites to improve accuracy. Submit blood sample for laboratory CO-Oximetry test for comparison.

Inappropriate sensor size or sensor measurement location.

Next steps: Verify proper sensor for patient size. Verify proper sensor site.

8.7 UNEXPECTEDLY HIGH SPCO READING

Possible elevated methemoglobin level.

Next steps: Submit blood sample for laboratory CO-Oximetry test.

8.8 DIFFICULTY OBTAINING A READING

Interference from line frequency induced noise.

Next steps: Verify/set 50/60 Hz Line Filter setting. See the section Advanced Screen on page 151.

Inappropriate sensor or sensor size.

Next steps: Verify proper sensor and sensor size for the patient.

Excessive ambient or strobing light.

Next steps: Shield the sensor from excessive or strobing light. Minimize or eliminate motion at the monitoring site.

8.9 SPCO READING DISPLAYS AS DASHES

• SpO2 value below 90%

Next steps: Assess/address patient condition.

SpMet value greater than 2%

Next steps: Laboratory analysis of a blood sample should be performed.



9.0 Getting Started

The following section provides information and instructions on installing and connecting various parts and accessories and preparing the MOVES[®] SLC™ for activation.



CAUTION! THESE ACTIVITIES SHOULD BE CARRIED OUT ONLY BY AUTHORIZED / TRAINED PERSONNEL.

9.1 MOVES® SLC™ SYSTEM CONTENTS

9.1.1 MOVES® SLC™ Accessories & Remote Screen Warnings and Cautions

Table 16: MOVES[®] SLC[™] Accessories Warnings and Cautions



WARNING! ALWAYS CARRY BACKUPS OF ACCESSORIES SUCH AS CARTRIDGES AND FILTERS.



WARNING! IF LEFT IN A HOT ENVIRONMENT OR DIRECT SUNLIGHT FOR A CONSIDERABLE LENGTH OF TIME, THE MOVES[®] SLC™ ACCESSORIES CAN BECOME QUITE HOT. MONITORING OF TEMPERATURE IS RECOMMENDED. ALWAYS MAKE SURE THAT ACCESSORIES THAT WILL BE APPLIED DIRECTLY TO THE PATIENT ARE SUITABLE FOR SKIN CONTACT AND WILL NOT CAUSE BURNS.



WARNING! IN THE EVENT OF DISAGREEMENT BETWEEN THE DEVICE AND THE REMOTE SCREEN THE DEVICE SHALL BE CONSIDERED CORRECT.



WARNING! IF THE REMOTE SCREEN BECOMES INOPERABLE, LOCKS UP OR BEHAVES ERRATICALLY, IT SHOULD NO LONGER BE USED AND BE DISCONNECTED FROM THE SYSTEM.



CAUTION! SOME PATIENT MONITORING ACCESSORIES MAY NOT FUNCTION PROPERLY, OR MAY RELAY INACCURATE READINGS, IF OPERATED OUTSIDE OF THEIR NORMAL OPERATING CONDITIONS.



CAUTION! USE ONLY ACCESSORIES THAT ARE PROVIDED OR AUTHORIZED BY THORNHILL RESEARCH INC!



CAUTION! THE MOVES[®] SLC™ ACCESSORIES PROVIDED BY THORNHILL RESEARCH INC. ARE FOR USE ONLY WITH THE MOVES[®] SLC™ SYSTEM.



CAUTION! ALL MOVES[®] SLC™ ACCESSORIES' PACKAGING SHOULD BE DISPOSED OF RESPONSIBLY IN ACCORDANCE WITH LOCAL WASTE DISPOSAL STANDARDS AND / OR LOCAL BIOHAZARD REGULATIONS.



NOTE: Where applicable, refer to accessories documentation for specific instructions for use, warnings, storage and operating guidelines.



9.1.2 System Contents



NOTE: All items listed in the following table are **multiple use items**. They are intended to be **reused**.

Table 17: MOVES[®] SLC™ System Contents

PICTURE	TRI P/N	DESCRIPTION
	122752	MOVES [®] SLC™ System Manufacturer: Thornhill Research Inc.
	111422	Power Supply / Battery Charger (both together in single unit) and AC cable. Manufacturer: Thornhill Research Inc.

PICTURE	TRI P/N	DESCRIPTION
A WARNING WAND WARNING WARNING WARNING WARNING WARNING WARNING WARNING WARNING WAND WAND WAND WAND WAND WAND WAND WAND	126071	Manufacturer: Thornhill Research Inc.
	101238	Manufacturer: Thornhill Research Inc.



PICTURE	TRI P/N	DESCRIPTION
	111462	Manufacturer: Thornhill Research Inc.
Front Clamp – Rear View	126440	Manufacturer: Thornhill Research Inc. NOTE: The front clamp is labeled FRONT.

PICTURE	TRI P/N	DESCRIPTION
Front Clamp – Side View		NOTE: The front clamp can be distinguished by its <u>flat</u> bottom portion near the locking mechanism.
Tront Gamp – Side View	126441	Back Clamp
		Manufacturer: Thornhill Research Inc. NOTE: The back clamp is labeled BACK.
Back Clamp – Rear View		



PICTURE	TRI P/N	DESCRIPTION
Back Clamp – Side View		NOTE: The back clamp can be distinguished by its <u>curved</u> bottom portion near the locking mechanism.
Sun Tech CHILD 12-19 cm	126024	Sensor, NIBP Cuff Child: 12-19 cm Manufacturer: SunTech Medical, (K051904) P/N # 98-0080-02
Auntral	100832	Sensor, NIBP Cuff Small Adult 17–25 cm Manufacturer: SunTech Medical, (K051904) P/N # 98-0080-04
Suited South and South and Suited South and South and Suited South and South a	100834	Sensor, NIBP Cuff Adult 23–33 cm Manufacturer: SunTech Medical, (K051904) P/N # 98-0080-06
	100836	Sensor, NIBP Cuff Large Adult 31–40 cm Manufacturer: SunTech Medical, (K051904) P/N # 98-0080-08

PICTURE	TRI P/N	DESCRIPTION
	100838	Sensor, NIBP Cuff Thigh 38-50 cm Manufacturer: SunTech Medical, (K051904)
	124848	P/N # 98-0080-10 NIBP Hose (3 m with Lemo connector) Manufacturer: Thornhill Research Inc.
	101048	ABP/CVP/ICP Short Cable (4 ft.) Manufacturer: Thornhill Research Inc.



PICTURE	TRI P/N	DESCRIPTION
	111442	ABP/CVP/ICP Long Cable (13 ft.) Manufacturer: Thornhill Research Inc.
	125350	12 Lead ECG Cable Manufacturer: Thornhill Research Inc.
	126760	Adult SpO ₂ (only) Finger Clip Manufacturer: Masimo Corporation, K120657 Prod. Ref: M-LNCS DCI, 3 ft. P/N # 2501 NOTE: Red RC-4 adapter cable required.



126761 | Pediatric SpO₂ (only) Finger Clip

Manufacturer: Masimo Corporation, K120657

Prod. Ref: M-LNCS DCIP, 3 ft.

P/N # 2502

NOTE: Red RC-4 adapter cable required.



126762 SpO₂ Ear Clip

Manufacturer: Masimo Corporation, K120657

Prod. Ref: M-LNCS TC-I, 3 ft.

P/N # 2503

NOTE: Red RC-4 adapter cable required.

NOTE: Sensor has not been validated under

motion conditions.

NOTE: Sensor is contraindicated for patients with

pierced ears at the measuring site.



125738 Rainbow® Patient Cable

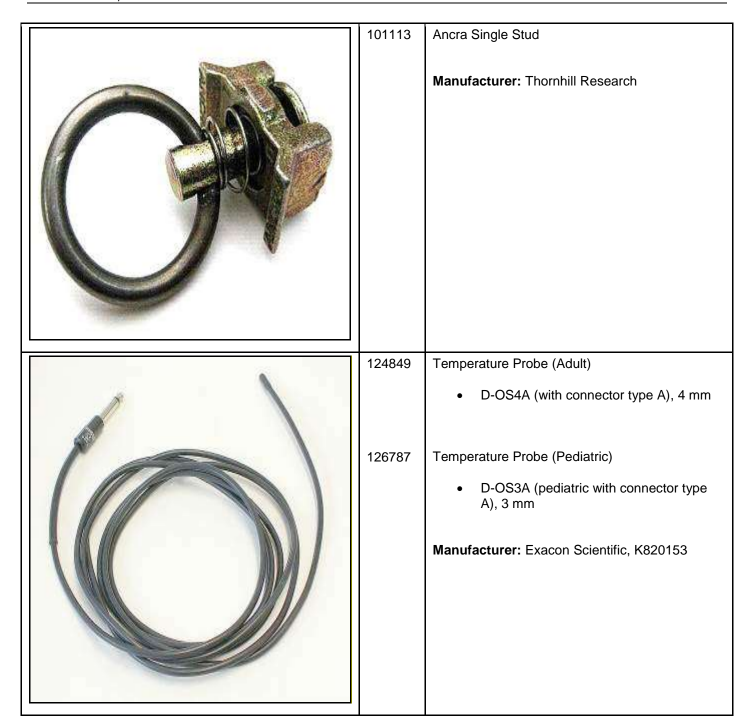
Rainbow® RC-4, 4 ft.

Manufacturer: Masimo Corporation, K110028

Prod. Ref: Rainbow® RC-4

P/N # 2406





PICTURE	TRI P/N	DESCRIPTION
P CO NOT DISCARDI	124241	Manufacturer: Thornhill Research Inc.
	125780	O2 Outlet Sampling Adaptor Manufacturer: Thornhill Research Inc.

PICTURE	TRI P/N	DESCRIPTION
Operator's Manual Content Patricular restricts the device to save by or on the order of a physician. Cylin Special Research to of Agric Signment The Bester (1833) See C. 1846-200	124826	MOVES [®] SLC™ Operator's Manual Manufacturer: Thornhill Research Inc.
Physical and the second	126581	MOVES [®] SLC™ Remote Screen (Optional) Manufacturer: Thornhill Research Inc.

PICTURE	TRI P/N	DESCRIPTION
	126779	Manufacturer: Thornhill Research Inc. The trap is composed of the following two parts which are periodically separated to drain the contents: • Top • Collection Cylinder



9.2 MOVES® SLC™ CONSUMABLES



NOTE: All consumables listed in the table below are **single patient use items**. They are **not intended** to be **reused**.

Table 18: MOVES[®] SLC™ Consumables

PICTURE	TRI P/N	DESCRIPTION
	126645	Ventilator Cartridge, SLC™ Manufacturer: Thornhill Research Inc.
	126647	Ventilator Cartridge, SLC™, 3 pack
	101114	Suction Canister Manufacturer: Bemis Health Care, K771737, P/N # 424410
AND STATE OF THE PARTY OF THE P	124359	Suction Canister, 10 pack
All Districts of the second of		Manufacturer: Bemis Health Care, K771737, P/N # 424410

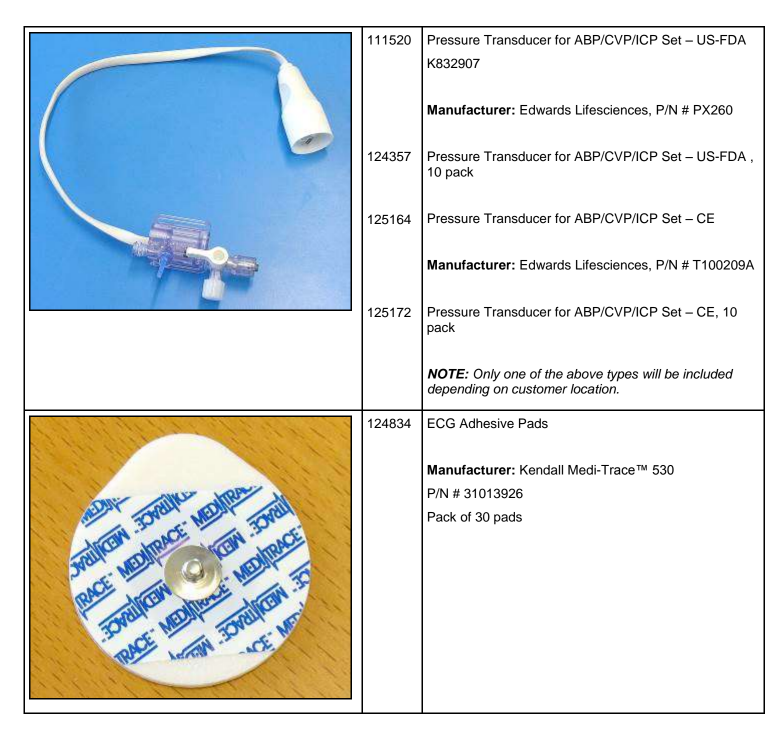
	111458	Suction Wand, Yankauer – FDA
		Manufacturer: Cardinal Healthcare, P/N # K82
	124358	Suction Wand, Yankauer, 10 pack
	125168	Suction Wand, Yankauer – CE
		Manufacturer: Covidien PLC, 8888505024
	125171	Suction Wand, Yankauer – CE, 10 pack
		NOTE: Only one of the above types will be included depending on customer location.
	100915	Hydrocarbon Filter
		Manufacturer: Airgas Inc. P/N # 815182
	124354	Hydrocarbon Filter, 6 pack
ur lictroctions ur long des respirator.		



	400504	Ventileten Eilen
	126504	Manufacturer: Pall Medical, P/N BB25
	126245	Pediatric Breathing System Filter
		Manufacturer: PALL Medical, P/N # BB25
		NOTE: Use as a replacement filter in MOVES® SLC™ <u>Ventilator Breathing Circuit (P/N 101210)</u> when ventilating patients under 30 kg or tidal volumes under 150 mL.
	101243	Tube, Suction (short, canister to system) – FDA
		Manufacturer: Cardinal Health Canada, P/N # N52A
	124355	Tube, Suction (short, canister to system) – FDA, 10 pack
	125166	Tube, Suction (short, canister to system) – CE
		Manufacturer: Covidien PLC, 8888301507
	125169	Tube, Suction (short, canister to system) – CE, 10 pack
		NOTE: Only one of the above types will be included depending on customer location.

101244	Tube, Suction (long, canister to patient) – FDA Manufacturer: Cardinal Health Canada, P/N # 66A
124356	Tube, Suction (long, canister to patient) – FDA, 10 pack
125167	Tube, Suction (long canister to patient) – CE
	Manufacturer: Covidien PLC, 8888301606
125170	Tube, Suction (long canister to patient) – CE, 10 pack
	NOTE: Only one of the above types will be included depending on customer location.
101210	Ventilator Breathing Circuit
	Manufacturer: Thornhill Research Inc.
124352	Ventilator Breathing Circuit , 12 pack
	NOTE: Nafion tube is shown attached – this tube is NOT included in the breathing circuit packaging but should be attached prior to use.





9.3 LIFTING THE MOVES® SLC™



NOTE: It is recommended that two people lift MOVES[®] SLC™.

The MOVES[®] SLC[™] has a handle at the front and a recessed slot at the back which should be used when lifting the unit. The position of these features is shown in the illustration below, and the features themselves are shown in the photos following the illustration.



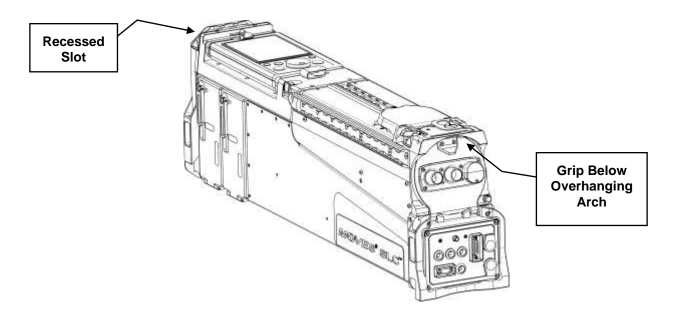
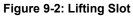


Figure 9-1: Lifting Points on MOVES[®] SLC™





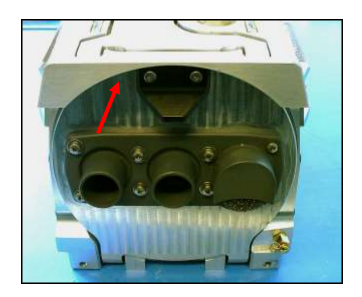


Figure 9-3: Lifting Arch



CAUTION! THE MOVES $^{\otimes}$ SLC $^{\text{TM}}$ WEIGHS APPROXIMATELY 37.5 LBS. BE SURE TO FOLLOW PROPER LIFTING PROCEDURES WHEN LIFTING THE MOVES.

9.4 ATTACHING THE SHOULDER STRAP TO MOVES® SLC™

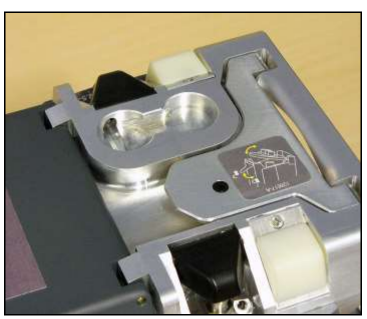


NOTE: The shoulder strap supplied with the MOVES[®] SLC[™] is attached using two Ancra Single Stud anchors. To attach the shoulder strap, the two stud anchors supplied with the MOVES[®] SLC[™] must be attached first and then the shoulder strap is clipped to them.

 Locate the two short rails at either end of MOVES[®] SLC™



Short Rail at Back



Short Rail at Front

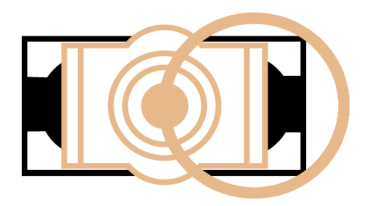
2. Depress the pin in one of the Ancra Single Stud anchors.



3. Slide the anchor along one of the short rails at either end of MOVES[®] SLC[™] until it is in the center position between two circular or semicircular openings.



4. The anchor will lock into place in the space between the two circular openings in the row (as shown in the illustration at the right).



- 5. Repeat the above actions for a second stud at the other end of $MOVES^{@}SLC^{TM}$.
- Attach the clips on the two ends of the shoulder strap to the two anchors now fastened to MOVES[®] SLC[™].



7. Strap attached to MOVES[®] SLC™.



9.5 ATTACHING CLAMPS TO THE MOVES® SLC™

Clamps can be attached to the MOVES[®] SLC™ which will then enable it to be clamped to a gurney, bed frame or stretcher to enable patient transport. Depending on where the clamps are attached, MOVES[®] SLC™ will fasten either to the top or the side of the receiving structure.

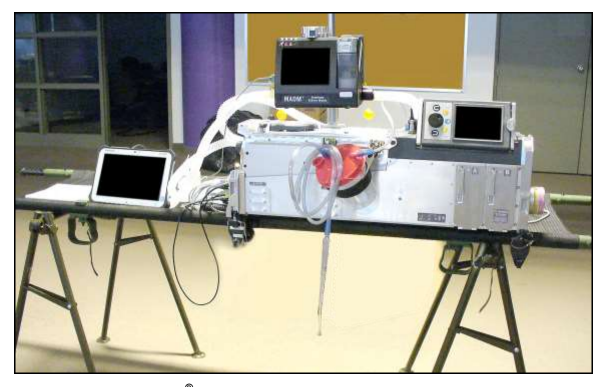


Figure 9-4 : MOVES[®] SLC™ with Accessories and Clamped to Top of a Stretcher



CAUTION! ALWAYS MAKE SURE THAT MOVES® SLC™ IS CLAMPED AND FULLY SECURED WHEN IN USE.



WARNING! THE CLAMPS HAVE NUMEROUS MOVING PARTS THAT MAY PRESENT A PINCHING OR CRUSHING HAZARD. ALWAYS USE CAUTION WHEN HANDLING BOTH THE FRONT AND BACK CLAMPS.

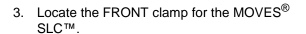
9.5.1 Attaching Clamps to Top Mount MOVES® SLC™

1. Orient the MOVES[®] SLC[™] with its FRONT to the operator's left.

NOTE: The FRONT is where the ventilator cartridge is inserted and the patient connections panel is located.



2. Next, lay the MOVES[®] SLC™ on its side.

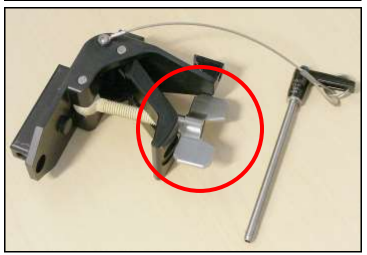


NOTE: The word FRONT is embossed on it.

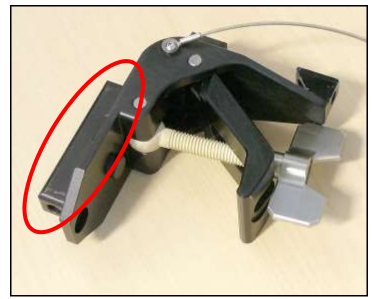




4. Undo the large wing bolt to its furthest extent (by turning it counterclockwise) to open the jaw of the clamp as wide as possible.



5. Identify the fitting projection circled in the photo at the right. It should fit into the receiving slot on the side of MOVES[®] SLC™ shown in the photo in the following step.



6. The receiving slot is shown circled in the photo at the right.

NOTE: The circular holes also shown are for the clamp's anchoring pin.

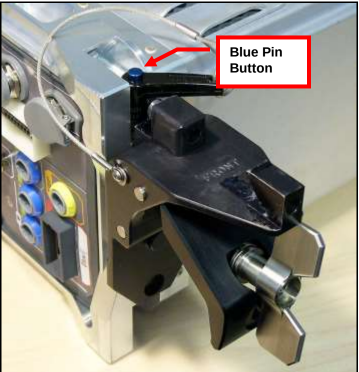




7. Orient the clamp as shown in the photo at right with the word FRONT embossed on the clamp facing up.

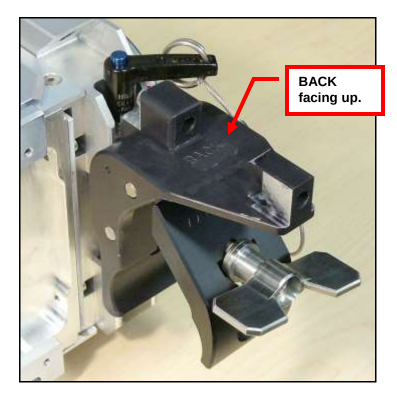


 Insert the anchoring pin attached to the left side of the clamp as shown in the photo at right.
 Press the blue pin button while inserting the pin.
 Make sure the pin inserts fully into its intended slot.



 Repeat the above procedure to mount the BACK clamp at the other end of MOVES[®] SLC™.

NOTE: Do not worry if the clamps feel slightly loose. When they are clamped to a gurney, bed rail or stretcher, the lateral object to which they are clamped adds the required reinforcement.



9.5.2 Attaching Clamps to Side Mount MOVES[®] SLC™

 Lay the MOVES[®] SLC[™] on either one of its sides. Attaching the clamps for side mounting can be done on either the RIGHT or LEFT side.

NOTE: In the following instruction, the clamps will be mounted on the RIGHT side.





2. Locate the FRONT clamp for the MOVES $^{\otimes}$ SLC $^{\text{TM}}$.

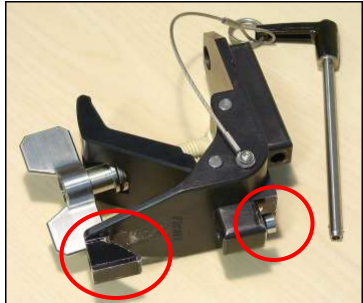
NOTE: The word FRONT is embossed on it.



3. Undo the large wing bolt to its furthest extent (by turning it counterclockwise) to open the jaw of the clamp as wide as possible.



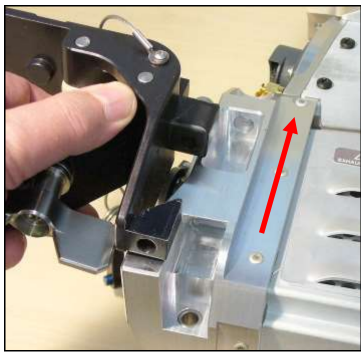
 Note the two connectors circled in the photo at the right. These will mate with receiving slots on the face of MOVES[®] SLC[™].



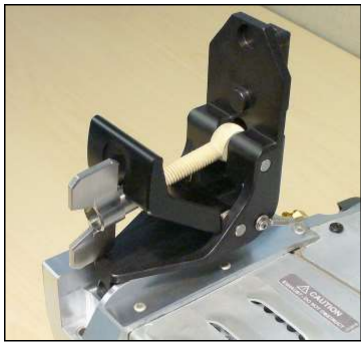
5. The receiving slots are shown in the photo at the right.



6. Align the clamp as shown in the photo at the right. Insert it into the slots and push the clamp upward to engage the connectors.



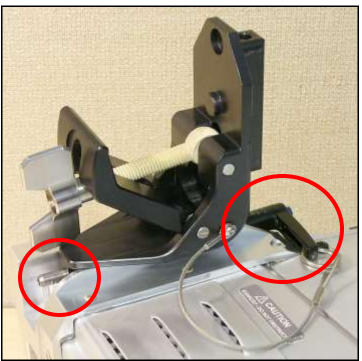
7. The clamp is shown properly seated in the photo at the right.



8. Insert the anchoring pin attached to the right of the clamp as shown in the photo at right. Press the blue pin button while inserting the pin. Make sure the pin inserts fully into its intended slot.



9. The anchoring pin is shown fully inserted in the photo at the right.



10. Repeat the above procedure to mount the BACK clamp at the other end of the MOVES[®] SLC™.

NOTE: Do not worry if the clamps feel slightly loose. When they are clamped to a gurney, bed rail or stretcher, the lateral object to which they are clamped adds the required reinforcement.



11. Both clamps for side mounting of MOVES[®] SLC[™] are shown attached in the photo at the right.



9.6 PATIENT MONITORING ACCESSORIES

This section briefly describes how the patient monitoring accessories are to be connected to the MOVES[®] SLC™ unit. The operator must thoroughly read all of the procedures contained within this manual and must fully understand the MOVES[®] SLC™ system operation before connecting the system to a patient. Prior to connecting the system to a patient, the operator must have alternative methods of treating the patient available should a power or mechanical failure occur.



WARNING! DO NOT CONNECT ANY MONITORS TO PATIENT WHILE PERFORMING SYSTEM TESTS! DOING SO COULD ENDANGER THE PATIENT!



CAUTION! USE ONLY ACCESSORIES PROVIDED WITH THE MOVES[®] SLC™ SYSTEM!

All MOVES[®] SLC™ patient monitors are connected to the patient connection panel (pictured below) as indicated by the labels.





Figure 9-5 : Patient Connection Panel



NOTE: The unlabeled port above IP 1 is the Calibration port.



NOTE: The unlabeled port above IP 3 is the <u>Barometric Sensing port</u>.

Table 19: Patient Connector Labels and Accessories

CONNECTOR LABEL	ACCESSORY
GAS SAMPLE	Sampling line Luer connection
ECG	Electrocardiogram – 12 Lead – Thornhill Research Inc.
NIBP	Non-invasive blood pressure – SunTech
Temp 1 & Temp 2	Temperature – Exacon Scientific
IP 1–3	Ports can be interchangeably used for any of the following: ABP — Arterial Blood Pressure transducer (Edwards TruWave PX Series) CVP — Central Venous Pressure transducer (Edwards TruWave PX Series) ICP — Intracranial Pressure transducer (Edwards TruWave PX Series)



CONNECTOR LABEL	ACCESSORY
SpO ₂	Pulse CO-Oximeter – Masimo Corporation
Barometric Sensing Port	No connectors are fitted to this port. Do Not Block.
Calibration Port	No connectors are fitted to this port. Do Not Block.

9.7 INSTALLING THE VENTILATOR CARTRIDGE AND BREATHING CIRCUIT



WARNING! LEAKS IN THE SAMPLING LINE CAN CAUSE LOW pCO2 AND/OR LOW O2 LEVELS.

9.7.1 About the Ventilator Cartridge

The ventilator cartridge is intended to be used with the MOVES[®] SLC™ system to provide positive pressure ventilation for patients who are *intubated*. MOVES[®] SLC™ recycles exhaled oxygen. This cartridge is made of CO₂ absorbent material to remove CO₂ from any re-breathed gas. Because this material has a *shelf life*, there is an expiry date printed on the cartridge. Always check the expiry date on the ventilator cartridge before using to make sure the cartridge has not expired. As well, monitor your supply of spare cartridges with regard to their remaining "shelf life".



NOTE: The ventilator cartridge should be kept in its packaging until use.

The ventilator cartridge is:

- For use with the ventilator breathing circuit, which includes the ventilator hoses, patient filter, endotracheal tube connector, and sampling line.
- For single patient use ONLY. The cartridge should be discarded and replaced between patients and when MOVES[®] SLC™ triggers an audible or visual alarm indicating that the level of CO₂ in the system is above 6 mmHg on inspiration.



WARNING! FAILURE TO CHANGE THE VENTILATOR CARTRIDGE WHEN INDICATED MAY LEAD TO THE PATIENT'S SUFFERING FROM AN INCREASE IN INSPIRED CO₂.



NOTE: Used breathing cartridges and breathing circuits should be disposed of in accordance with local biohazard regulations.





Figure 9-6: Ventilator Cartridge

9.7.2 Installing the Ventilator Cartridge

1. Lift the cartridge door lock/release latch to the vertical position.



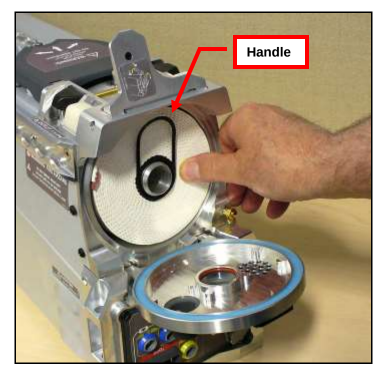
2. Open the cartridge door by pulling it forward using its black plastic handle.





3. Insert the cartridge into the cavity as shown. Press hard to ensure it is fully and securely seated.

NOTE: If you are replacing a used cartridge, use the black plastic handle indicated in the photo at right to pull it out.

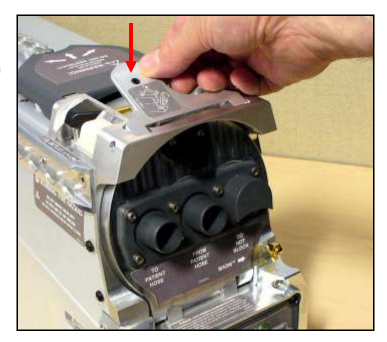


4. Close the door and push it in firmly.

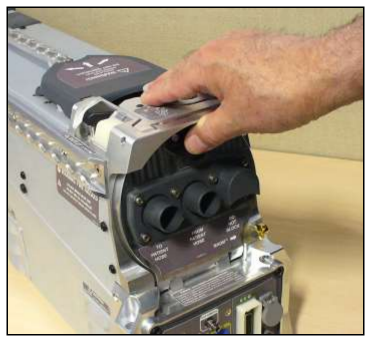




 Depress the cartridge door lock/release latch. It should close smoothly. If you feel any resistance, the door is not completely closed because the cartridge is not fully inserted. Open the door and push the cartridge in further and try again.



6. In the picture at the right, the latch is fully closed securing the cartridge door and the ventilator cartridge.



9.7.3 Installing the Hydrocarbon Filter



CAUTION! BEFORE INSTALLING A HYDROCARBON FILTER, CHECK THE FOUR-DIGIT DATE CODE PRINTED ON THE CARTRIDGE. THE CARTRIDGE LABEL IS STAMPED WITH FOUR CHARACTERS "XXYY", WHERE "XX" IS THE WEEK OF THE YEAR AND "YY" IS THE YEAR OF MANUFACTURE. A CARTRIDGE MORE THAN THREE YEARS OLD SHOULD BE DISCARDED SINCE IT MAY DEGRADE THE PERFORMANCE OF OR CAUSE DAMAGE TO THE MOVES[®] SLC™ OXYGEN CONCENTRATOR.



- Always install a hydrocarbon filter before powering up MOVES[®] SLC[™]. MOVES[®] SLC[™] will alert you with an alarm if it becomes clogged.
- 2. Insert the hydrocarbon filter as shown at right into the REAR panel of the MOVES[®] SLC™ unit.



NOTE: If the hydrocarbon filter needs to be changed when MOVES[®] SLC[™] is in operation, a replacement filter will need to be readily available for quick insertion after the old one is removed.



CAUTION! OPERATING MOVES[®] SLC™ WITHOUT A FILTER WILL DAMAGE THE UNIT. DO NOT OPERATE MOVES[®] SLC™ WITHOUT A FILTER!

3. Rotate the hydrocarbon filter clockwise to install, counterclockwise to remove.





WARNING! IT MAY BE NECESSARY TO USE AN ALTERNATIVE MEANS OF OXYGEN SUPPLEMENTATION SHOULD THE HYDROCARBON FILTER REQUIRE REPLACEMENT WHILE A PATIENT IS BEING TREATED.



NOTE: Use only the filter cartridges supplied with MOVES[®] SLC™. These NIOSH-certified filters are specifically selected for use with MOVES[®] SLC™. These filter cartridges are not for use in atmospheres posing immediate danger to life or health or atmospheres containing less than 19.5% oxygen by volume.

9.7.4 Installing the Ventilator Breathing Circuit

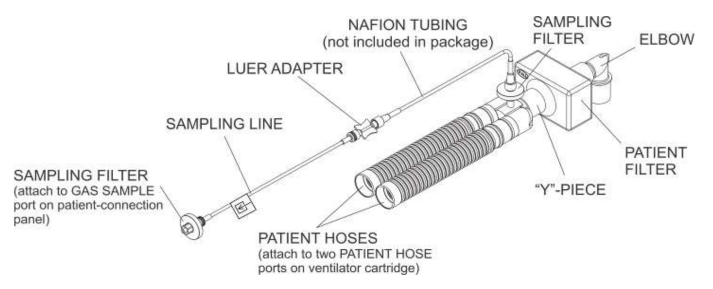


Figure 9-7: Ventilator Breathing Circuit

Refer to the figure above when assembling the Ventilator Breathing circuit. Remove the **single use** circuit from its sealed package.





WARNING! THE CIRCUIT FILTER SUPPLIED WITH THE VENTILATOR BREATHING CIRCUIT IS INTENDED FOR USE WHEN DELIVERING TIDAL VOLUMES OF 150 ML AND OVER. IF DELIVERING TIDAL VOLUMES UNDER 150 ML OR VENTILATING PATIENTS UNDER 30 KG, REPLACE THE BREATHING FILTER WITH THE PEDIATRIC BREATHING SYSTEM FILTER.



CAUTION! THE VENTILATOR BREATHING CIRCUIT AND THE SAMPLING FILTER CONNECTED TO THE NAFION TUBING SHOULD BE INSPECTED EVERY FOUR (4) HOURS FOR CONDENSATION AND DRAINED AS REQUIRED. THE CIRCUIT AND FILTER SHOULD BE CHANGED AFTER 24 HOURS OF CONTINUOUS USE.



NOTE: The Nafion tube is NOT a single use item and DOES NOT come with the circuit. It should be housed in the MOVES[®] SLC[™] accessory case. DO NOT DISCARD THE NAFION TUBE AFTER USE. It has a sample filter to prevent contamination.



NOTE: Replace the Nafion tube at six (6) month intervals or as needed. The Nafion tube should be inspected prior to use for signs of physical damage including cracking and kinking. Premature failure of the Nafion tube can present as either an occluded sample line or, more commonly, as a leaky sample line that will produce a dampened PCO₂ trace and lower PetCO₂ readings.

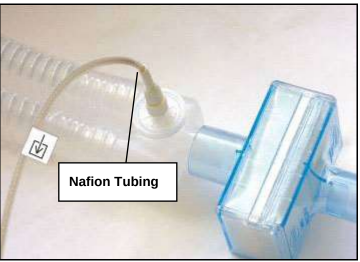
1. Make sure there is a sample filter connected to the "Y" piece. If there is not, connect one.



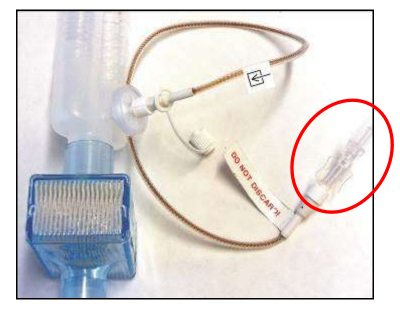
NOTE: This sample filter, which is used to keep the Nafion tubing clean between uses, should be discarded after each use.



Connect one end of the Nafion tubing to the sample filter.

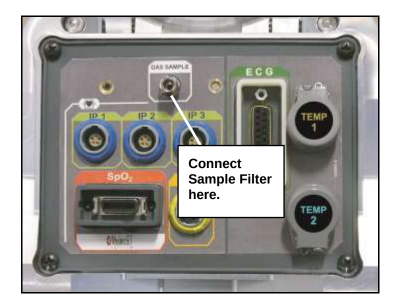


3. Connect the other end of the Nafion tubing to the Luer adapter (circled at right), and then the Luer adapter to one end of the sampling line.



4. Connect the other end of the sampling line (with another sample filter on its end) to the 'GAS SAMPLE' port on the patient connections panel. The sample filter should be connected directly to the patient connection panel.







5. Attach the water trap to the "From Patient Hose" port on the ventilator cartridge door.



6. Attach one patient breathing circuit hose to the "To Patient Hose" port on the ventilator cartridge door and the other to the water trap.



9.8 DELIVERING SUPPLEMENTARY OXYGEN (O2)



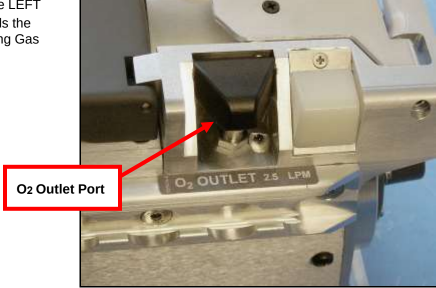
WARNING! WHEN USING O2 SUPPLEMENTATION, AN O2 SATURATION MONITOR MUST BE USED.



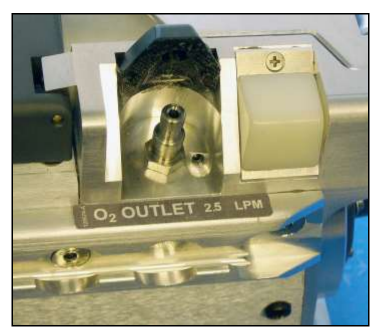
WARNING! WHEN USING OXYGEN SUPPLEMENTATION MODE, THE MOVES[®] SLC™ GAS SAMPLING PORT MUST BE CONNECTED TO THE OXYGEN DELIVERY CIRCUIT (E.G., O₂ MASK SAMPLE PORT) AND AN OXYGEN SHUT-OFF DEVICE, SUCH AS BPR'S FIRESAFE™ CANNULA VALVE, SHOULD BE USED IN THE OXYGEN SUPPLY LINE AS CLOSE TO THE PATIENT AS POSSIBLE.

The concentrator delivers up to a nominal 90% O₂ at a nominal flow rate of 2.5 LPM directly to the patient during O₂ Supplement mode. Follow these instructions to supply supplementary oxygen to the patient.

 Locate the O₂ Outlet port located on the LEFT side of the MOVES[®] SLC[™] unit towards the FRONT and beside the Ventilator Driving Gas Inlet.



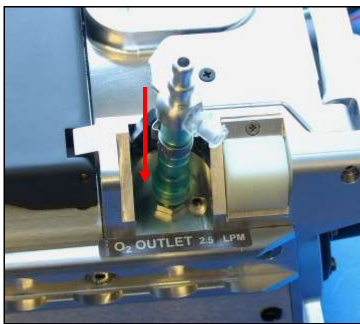
2. Open the protective cover of the O₂ Outlet port (by pulling it up) to access the O₂ Outlet.



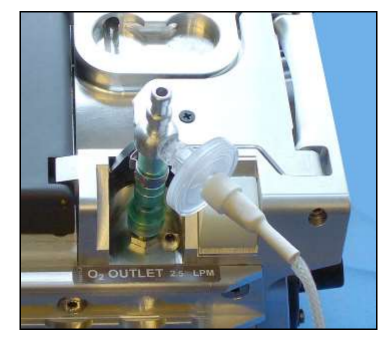
3. Locate the O₂ Outlet Sampling Adaptor.



4. Attach the green flexible tubing end of the O₂
Outlet Sampling Adaptor to the O₂ Outlet by placing it over the outlet and applying downward pressure until the adaptor fits snugly.

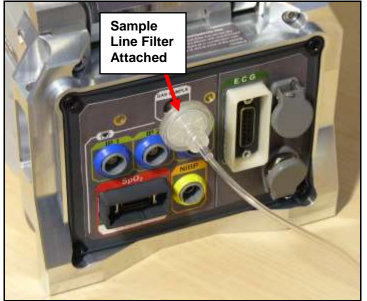


5. Attach the Sample Line Filter that is connected to the Nafion tube end of the Sampling Line to the O2 Outlet Sampling Adaptor by fitting it onto the clear, hard-plastic port on the O2 Outlet Sampling Adaptor and rotating it clockwise until it is snug.



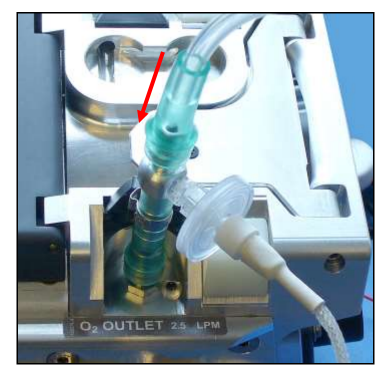
 Attach the Sample Line Filter at the other end of the Sampling Line to the Gas Sample port in the Patient Connections Panel by placing it over the port and rotating the filter clockwise until it is snug.

NOTE: In both O2 Supplement mode and Ventilate mode, the sample system should be connected from the sample port in the following sequence: filter, sample line, adapter, Nafion tube, filter.





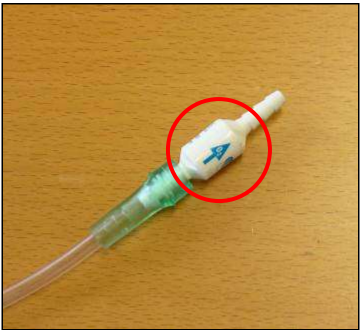
 Attach one end of the green O₂ tube to the metallic outlet on the O₂ Outlet Sampling Adaptor by placing it over the metallic outlet and applying downward pressure until the tube end fits snugly.



8. Insert a fire suppression device (such as BPR's Firesafe™ Cannula Valve shown in the photo at the right) onto the other end of the green O₂ tube.

NOTE: Place the fire suppression device as close to the patient as possible.

NOTE: Make sure that any O_2 directional indicator on the fire suppression device is pointing in the direction of the patient and not back toward the O_2 outlet.



9. On the Setup screen, set the system to run in O2 Supplement mode.

NOTE: In O2 Supplement mode, the Vent O2 setting (under CONCENTRATOR) is grayed out, indicating that it (and its value) does not apply in O2 Supplement mode. As mentioned above, the concentrator provides 2.5 LPM of >87% O₂ directly to the patient.

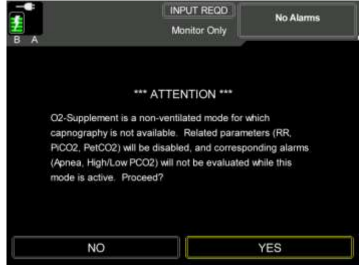
 When you switch to O2 Supplement mode, a confirmation message is displayed to ensure that this switch is intentional. User input is required (choose NO or YES) in order to proceed.

The confirmation message states:

"O2 Supplement is a non-ventilated mode for which capnography is not available. Related parameters (RR, PiCO2, PetCo2) will be disabled, and corresponding alarms (Apnea, High/Low PCO2) will not be evaluated while this mode is active. Proceed?"

11. Attach the O₂ Mask line or the Nasal Cannula line to remaining available end of the fire suppression device.



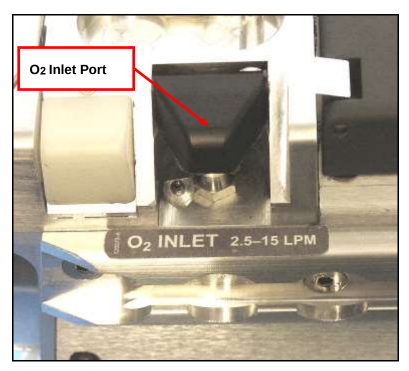


9.9 USING AN EXTERNAL GAS SUPPLY

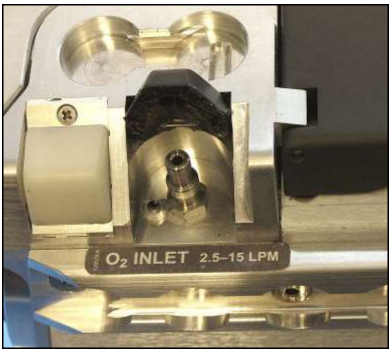


WARNING! EXTERNAL OXYGEN SUPPLY IS ONLY USED FOR VENTILATE MODE. FOR O_2 SUPPLEMENTATION, USE THE OXYGEN SOURCE DIRECTLY.

 Locate the O₂ Inlet port located on the RIGHT side of the MOVES[®] SLC[™] unit near the Ventilator Driving Gas Inlet.



2. Open the protective cover of the O₂ Inlet port (by raising it) to access the O₂ Inlet.



Attach the External Gas Supply tube to the O2 Inlet nipple

Note the following:

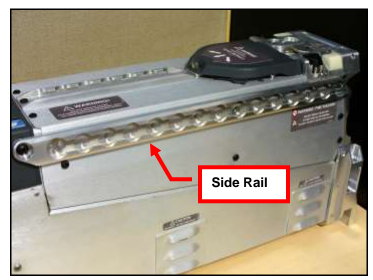
• When using the O₂ Inlet, concentrations other that 100% are controlled with an external air/oxygen mixer. No other gasses can be used.



- MOVES[®] SLC[™] uses an oxygen conserving ventilation system that normally requires less than 1LPM of oxygen to maintain an FiO₂ of 100% to the patient. Therefore, when using the O₂ Inlet, gas flows of 1 to 2 LPM can be used to conserve external oxygen tanks, with higher flows used briefly to pre-charge or flush the circuit, or if the desired FiO₂ is not being achieved.
- When using the O₂ Inlet, the system should be set to a Vent FiO₂ slightly under the external gas mix and NEVER set to Air or Maximum. For example, setting the system to a Vent FiO₂ of 85% while applying 100% O₂ into the external gas inlet will only run the concentrator if the FiO₂ drops to 82%, working as a backup for the external O₂ supply.

9.10 INSTALLING SUCTION ACCESSORIES

 The clips of suction canister holder attach to the SIDE rail on either side of MOVES[®] SLC™.



Attach the suction canister holder to MOVES[®]
 SLC™ by depressing the clips and sliding them
 along the SIDE rail into the desired position.
 Release the clips and ensure that they lock into
 place.



- 3. The anchor will lock into place in the space **between** any two of the holes in the row (as shown in the illustration at the right).
- 4. The suction canister holder is shown correctly attached in the photo at the right.





5. Insert the suction canister into the holder as shown. Make sure that the large red plug indicated at right is <u>firmly</u> in place.



6. Connect the short suction hose to the 'SUCTION INLET' port of the suction canister.

NOTE: The <u>short</u> suction hose connects the canister to the $MOVES^{\circledR}$ SLCTM unit.



7. Open the protective cover of the 'SUCTION' port located on the LEFT side of the MOVES[®] SLC™



8. 'SUCTION' port shown at right.



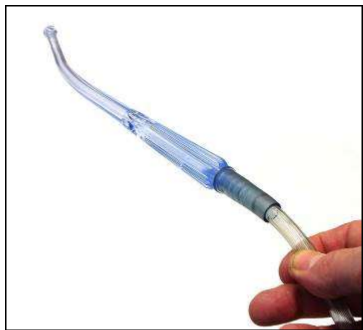
9. Connect the other end of the short suction hose to the 'SUCTION' port.



10. Connect the patient suction hose (long hose) to 'PATIENT PORT' on the suction canister.



11. Connect the patient suction wand to the other end of the patient suction hose.





CAUTION! THE SUCTION WAND AVAILABLE FROM THORNHILL RESEARCH IS DESIGNED TO MINIMIZE TOTAL OCCLUSION. ONLY THIS WAND OR EQUIVALENT SHOULD BE USED WITH THE MOVES SLC TM SYSTEM.



WARNING! THE OPERATOR SHOULD ALWAYS HAVE AVAILABLE AN ALTERNATE MEANS OF SUCTION IN THE EVENT OF POWER FAILURE, MECHANICAL FAILURE OR SERIOUS OCCLUSION IN THE SUCTION SYSTEM.



WARNING! THE MOVES[®] SLC™ OXYGEN CONCENTRATOR DOES NOT FUNCTION WHILE SUCTION IS ON. AN ALTERNATIVE MEANS OF SUPPLYING O₂ WILL BE NECESSARY IF A HIGH PERCENTAGE OF O₂ IS CRITICAL.

9.11 USING AN OPTIONAL HUMIDIFIER

The MOVES[®] SLC[™] is intended for use in short term critical care, transport and emergency situations and utilizes a circle breathing circuit. Therefore, the use of an optional humidifier is not required nor recommended.

9.12 PREPARING MOVES® SLC™ FOR USE

9.12.1 Checking the Charge

There is a Battery Condition Indicator on the front of each battery.



Figure 9-8: Battery with Condition Indicator Shown

Pressing and holding the button on the Battery Condition Indicator (white button in circled area above) shows the battery condition. As battery power decreases, the illuminated LEDs extinguish from the top (green – meaning the battery power is at a high level), through middle indicators (orange – meaning that the battery power is at an intermediate stage), the bottom (red – meaning that the battery power is very low or exhausted).

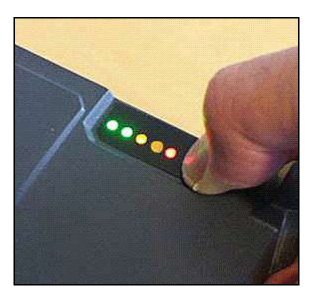


Figure 9-9: Condition Indicator Showing High Level Charge

When the power supply / battery charger is connected to a live AC power supply and to MOVES[®] SLC™, any installed batteries will charge.



NOTE: The battery charge level may not appear to increase for approximately three (3) hours. This is normal for the battery's initial charge and after extended periods without use.



NOTE: It is possible for batteries to be exhausted beyond normal recovery. If pushing a battery's status button does not cause any LEDs to be illuminated, place the battery in a MOVES[®] SLC[™] to charge for up to 48 hours. If this does not recover the battery's charge, the battery is non-functional and should be returned to the manufacturer.



WARNING! TO REDUCE RISK TO THE PATIENT IN THE EVENT OF A POWER FAILURE, WHEN RUNNING ON EXTERNAL POWER, ONE CHARGED BATTERY SHOULD BE PRESENT IN THE MOVES[®] SLC™ UNIT. IN ADDITION, ALWAYS CARRY ALTERNATE MEANS OF VENTILATING, SUCTIONING, AND OXYGENATING THE PATIENT.



CAUTION! A MOVES[®] SLC™ UNIT SHOULD NEVER BE PUT INTO TRANSPORT SERVICE WITH LESS THAN A 95% CHARGE IN BOTH BATTERIES.



WARNING! BATTERY TIME SHOWN REMAINING IS APPROXIMATE AND HIGHLY DEPENDENT ON OPERATING CONDITIONS! PUT SAFETY FIRST – ALWAYS CARRY SPARE BATTERIES!



CAUTION! NEVER CHARGE BATTERIES IN AMBIENT TEMPERATURES BELOW 32°F (0°C) OR ABOVE 104°F (40°C).

9.12.2 Inspecting the Batteries

Inspect the battery for physical damage such as cracks, holes, and leaks.





WARNING! DO NOT USE OR CHARGE A DAMAGED BATTERY!

9.12.3 Checking for Battery Installation

Whether or not a battery (or batteries) is installed in MOVES[®] SLC[™] can be determined by looking through the horizontal window on the battery bay door. If a yellow handle is visible, then a battery is installed.



Battery Charge Level Visible Here

Figure 9-10: Visible Yellow Handle Indicates Battery Present



NOTE: When the MOVES[®] SLC[™] system has a battery installed and is plugged in, the battery charge level can also be determined by looking through the <u>vertical</u> window on the side of the battery bay door (even if the system is turned off).

9.12.4 Installing the Batteries

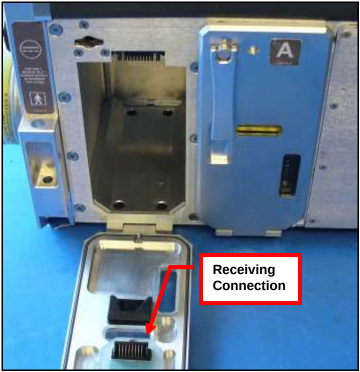
1. Lift the battery compartment latch and turn it fully counter-clockwise to open the battery bay door.



2. Open the battery bay door.



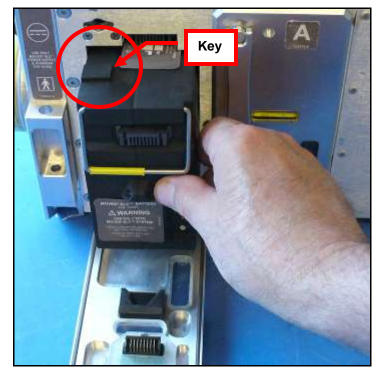
CAUTION! THERE ARE BATTERY COMPARTMENT DOORS ON BOTH SIDES OF THE MOVES® SLC™. THEREFORE, A BATTERY CAN BE INSERTED FROM EITHER SIDE. HOWEVER, THE BATTERY CONNECTIONS MUST BE PROPERLY ORIENTED WITH THE RECEIVING CONNECTIONS INSIDE THE BATTERY COMPARTMENT (ON THE DOORS). CONNECTIONS ARE NOT IDENTICAL ON EITHER END OF THE BATTERY.





3. Push the battery into the battery compartment until it engages. You should not have to force it. If you feel resistance, the battery is not properly oriented. Pull the battery out, then check the location of the connections and reorient the battery.

NOTE: The battery is keyed to be inserted in only one orientation. If you are experiencing trouble inserting the battery, check that "key" on the top of the battery matches the slot at the top of the battery cavity.



4. The battery fully inserted and properly seated is shown in the photo at the right.

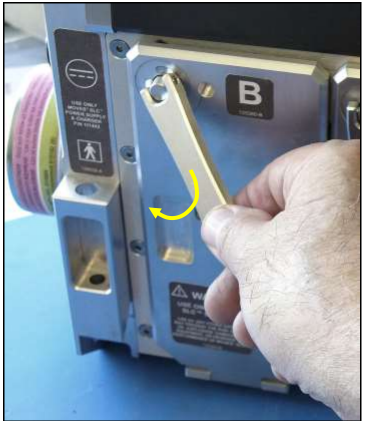


 To close the battery bay door, lift the latch and turn it as far as it goes counterclockwise to the horizontal position. This will align the key on the back of the door to the receiving slot on the MOVES[®] SLC™.



6. Close the door and then turn the lever fully clockwise as far as it goes.

NOTE: Also make sure that the battery latch on the opposite side is closed completely.



7. Lastly, push the lever in to lock it



8. Repeat procedures 1–7 for the second battery.



NOTE: It is recommended that both batteries be installed during all operational times. If one battery requires a charge, it can be recharged simultaneously during normal operation when the MOVES[®] SLC™ is connected to the power supply / charger.



NOTE: The MOVES[®] SLCTM can operate from one battery to allow uninterrupted operation during battery replacement (since the system uses only one battery at a time). When a battery reaches an exhausted state, remove the battery and replace with a charged battery.



WARNING! WHEN MOVES[®] SLC™ IS NOT IN OPERATION, BATTERIES SHOULD BE REMOVED FROM THE UNIT AND STORED IN A DRY AREA AT ROOM TEMPERATURE. LEAVING BATTERIES INSTALLED IN A NON-OPERATIONAL UNIT MAY CAUSE THEM TO DRAIN TO AN <u>UN-RECHARGEABLE</u> LEVEL.



9.12.5 Preparing the Power Supply / Battery Charger

- 1. The power supply / battery charger is shown to the right. Note, there are two cords:
- A light grey cord this cord delivers power from a wall socket, or line supply (like a generator), to the power supply / battery charger.



CAUTION! THIS CORD COMES WITH A GROUNDING PRONG ON THE MALE CONNECTOR. TO REDUCE THE RISK OF ELECTRICAL SHOCK, THIS PRONG SHOULD NEVER BE REMOVED OR COMPROMISED.



CAUTION! THIS CORD IS A SPECIAL MEDICAL-GRADE POWER CORD AND SHOULD NOT BE REPLACED WITH A NON-MOVES[®] SLC™ SUPPLIED PART.

- A black cord permanently affixed at one end to the power supply/charger, and with a special nine-pin female connector on the other end. This cord delivers power from the power supply / battery charger to the MOVES[®] SLC™ unit. The female connector that attaches to the MOVES[®] SLC™ unit is keyed to ensure proper connection.
- If it is not already inserted, insert the light grey power cord into the front of the power supply / battery charger.







9.12.6 Connecting MOVES[®] SLC[™] to the Power Supply / Battery Charger

 Remove the protective cap from the receptacle of the REAR panel of the MOVES[®] SLC™ by turning it counterclockwise .



 Insert the nine-pin connector into the receptacle. Note the chevrons which indicate the top of the connector. Make sure the chevrons face straight up when inserting the connector, which is keyed to match the connector on MOVES[®] SLC™.



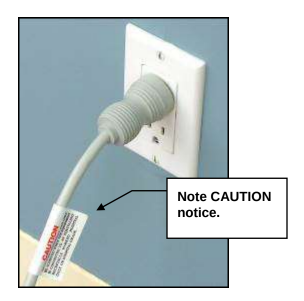
3. Rotate the locking collar clockwise to secure the connector.



9.12.7 Connecting AC Power

- 1. Insert the power supply / battery charger AC connector into a wall socket or line supply.
- Verify the connection by checking for an illuminated PWR LED on the power supply / battery charger.
- Note the CAUTION notice attached to the cord:
- 4. GROUNDING RELIABILITY CAN ONLY BE ACHIEVED WHEN THE EQUIPMENT IS CONNECTED TO AN EQUIVALENT RECEPTACLE MARKED HOSPITAL ONLY OR HOSPITAL GRADE.

NOTE: The photo at right shows a North American power connection. However, interchangeable power cords suitable for other regions are also available.



9.12.8 Battery Storage



CAUTION! WHEN BATTERIES ARE DISCHARGED AND LEFT IN MOVES[®] SLC™ FOR A PROLONGED PERIOD, A <u>COMPLETELY</u> DISCHARGED BATTERY CAN RESULT. THE BATTERY CAN STILL BE RECHARGED, BUT IT MAY TAKE MORE THAN THE NORMAL 2.5 HOURS – AND UP TO 48 HOURS – TO FULLY CHARGE.

When MOVES[®] SLC[™] is not in operation, its batteries should be removed from the unit and stored in a dry area at room temperature.







Figure 9-11: Battery Disconnected (compartment open)

Figure 9-12: Battery Removed

Conversely, to engage the battery close the door fully then turn the exterior handle clockwise until it locks.

Also, please note the following important cautions (and final Warning) about storing MOVES[®] SLC™ batteries:

- Do not store batteries above 60°C (140°F) or below –20°C (–4°F).
- Store batteries in a cool (below 30°C [86°F]), dry area that is subject to little temperature change. Elevated temperatures can result in reduced battery service life.
- Do not place batteries near heating equipment, nor expose to direct sunlight for long periods.



WARNING! BATTERY EXPOSURE TO TEMPERATURES IN EXCESS OF 130°C (266°F) WILL RESULT IN THE BATTERY VENTING FLAMMABLE LIQUID AND GASES.



10.0 Startup



CAUTION! THE SURFACE OF THE MOVES[®] SLC™ SYSTEM CAN BECOME HOT, ESPECIALLY IF IT IS BEING OPERATED IN DIRECT SUNLIGHT. CARE SHOULD BE TAKEN WHEN TOUCHING OR CONTACTING THE SURFACE OF THE MOVES[®] SLC™ SYSTEM.

10.1 ADJUSTING THE SCREEN DISPLAY ORIENTATION

The display screen can be locked or adjusted to face either the RIGHT or LEFT side of the MOVES[®] SLC™ to facilitate operation from either side of the unit.

 In the picture at the right, the screen is locked. Turning the catch indicated 90 degrees to either the left or the right will unlock the screen and allow it to be raised into a working position.



2. In the picture at the right, the screen is unlocked. The arrow on the lock points to the direction of the screen that can be raised. This is shown in the following photograph.







3. In the picture at the right, the screen is unlocked. The arrow on the lock points to the direction of the screen that can be raised. This is shown in the following photograph.







NOTE: The <u>on-screen</u> display orientation can be flipped by pressing the <u>Flip Button</u> on the Keypad.



Figure 10-1: Flip Button

10.2 USER INTERFACE (UI) CONTROLS AND FUNCTIONS

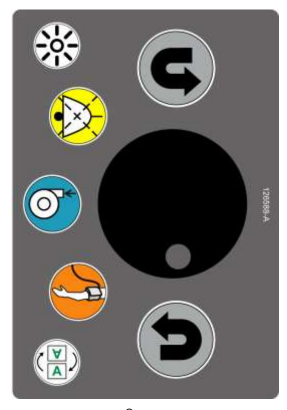


Figure 10-2: MOVES[®] SLC™ UI (User Interface)

The following table describes the physical appearance, function and the effect or use of each of the components used to operate the $MOVES^{@}$ SLC^{TM} unit.

Table 20: User Interface Controls and Functions

Button / Icon	Function
Not Applicable	Main Liquid Crystal Display (LCD) Screen
	This screen is the main component of the user interface. The operator interacts with the display to operate the MOVES [®] SLC [™] unit, getting information from the screen while selecting modes, options and numerical values for settings using the UI buttons and the jog wheel.
	Power Control Button
	This push button (which is next to the power connection on the BACK end of the MOVES [®] SLC™ unit) is used to activate or deactivate the unit. When the button is initially pressed, the system activates, the UI screen illuminates and an audible indicator sounds. When pressed and held for five (5) seconds when the unit is running, the button turns the unit OFF. As well, a message appears on the UI screen indicating this functionality.

Button / Icon	Function
	Brightness Button This keypad button is used to cycle the screen display and visual alarm indicator lights through six (6) diminishing levels of brightness.
	Alarm Audio Pause Button The keypad's Alarm Audio Pause button can be used either to silence ALL alarms permanently or for a temporary period (120 seconds). Pressing the Alarm Audio Pause button for one (1) second and then releasing it will silence ALL alarms temporarily. Pressing the Alarm Audio Pause button for at least three (3) seconds and then releasing it will silence ALL alarms permanently (if allowed, see note below). Pressing the Alarm Audio Pause button for one (1) second again then releasing it will re-enable ALL alarm audio. When alarm audio is paused temporarily, a dashed "X" is shown through the bell icon shown on the Status Bar. When alarm audio is permanently disabled, a solid "X" is shown through the bell icon on the Status Bar. If alarm audio is enabled no icon is shown. If an alarm is turned off on the Alarm On/Off screen, its alarm will not sound. NOTE: The option to silence all alarms permanently must be enabled by System Administrator.
	NIBP Control Button The NIBP Control button on the keypad is used to start or abort a Non-Invasive Blood Pressure (NIBP) reading. Prior to taking an NIBP reading, the NIBP cuff is fitted to the patient with the free end of the sensor cord attached to the patient-connector panel. The NIBP Control button is pressed to begin taking a reading. The data taken from the sensor appears in numerical form on the Monitor screen (if currently shown) once the reading is complete. Pressing the NIBP Control button while a reading is in progress aborts the reading. The button icon will be shown on the Status Bar if an NIBP reading is in progress (i.e., cuff is pressurized). Otherwise, the icon will not be shown. **NOTE: Using the Setup screen, you can set the NIBP control to manual, or to automatically update at set intervals: every 1−5 minutes, 10 minutes, 15 minutes or STAT. STAT, an abbreviation for the Latin word Statim which means "immediately", will set MOVES® SLC™ to take readings as often as is considered safe (i.e., a minimum time of 5 seconds is enforced between the end of one NIBP reading and the beginning of the next reading) for a maximum of 15 minutes. After 15 minutes in STAT mode the MOVES® SLC™ will automatically transition the update interval to 5 minutes. Pressing the NIBP Control Button manually to take a reading, or to abort a reading, temporarily overrides any automatic setting; however, the timer is



Button / Icon	Function	
	Suction Control Button	
	The keypad's Suction Control button is used to activate and deactivate the suction pump. All suction accessories must be connected and ready to use before activating. When the button is pressed, the suction pump activates. When the button is pressed again, the suction pump deactivates.	
	While suction is ON, the Suction icon and the current suction pressure (in mmHg) are displayed in the Status Bar.	
	NOTE: When in Suction mode, the concentrator/suction pump operates at a default (maximum) intake pressure level of 325 mmHg. This value can be changed (lowered) on the Setup screen.	
	WARNING! THE O ₂ CONCENTRATOR DOES NOT PRODUCE OXYGEN WHEN THE SUCTION FEATURE IS IN USE. IF SUCTION USE IS PROLONGED, THE PATIENT WILL BECOME HYPOXIC UNLESS EXTERNAL O ₂ IS SUPPLIED.	
	Flip Screen Button	
	Press the Flip Screen button to flip the on-screen display orientation by 180°.	
	Jog Wheel	
	The Jog Wheel is used to make selections on the various screens. Turning it clockwise or counter clockwise selects items in a circular manner going around the screen like the sweep of a clock's hands. The Jog Wheel does not select in a left to right, line by line manner as if scrolling through a list. To activate a selection, press the Jog Wheel.	
G	Back Button	
	The Back Button takes you "up" one level and cancels the current entry. It ultimately is trying to get back to the screen selection button (MAIN / SETUP / etc.).	
Buttons on the Remote Screen		
	Screen Button Press the Screen button to move to the next screen in sequence. If there are outstanding queries or parameters that need to be satisfied, the functionality of the Screen button will be inhibited until these queries or parameters are satisfied.	
	NOTE: This button can also function as an Accelerator. Pressing and holding the button will take you from any screen to the Monitoring screen. If you are already on the Monitoring screen, pressing and holding the button will take you to the Setup screen.	
8	Cancel Button Press this button to reject the current value that is being changed and return to the previous value.	



Button / Icon	Function
	Check Button Press the Check button to start or end editing a currently selected item, or change the currently selected item. The Check button is also used to acknowledge messages.
*+	Next Button Selects the next item in a group or increases a numeric value.
	Previous Button Selects the previous item in a group or decreases a numeric value.

10.3 SYSTEM VISUAL INDICATORS (ALARMS / SYSTEM STATUS)

There are four (4) System Visual Indicators, one at each top corner of the MOVES[®] SLCTM. These indicators display three (3) different colors: green, red and yellow. The severity of the highest active alarm is show. If an alarm is turned off, the display defaults to the severity of the next active alarm. These System Visual Indicators provide the user with the ability to see alarms, or system alarm status, from a greater distance and angle than would be possible with only screen display. The brightness of the System Visual Indicators is controlled by the front-panel brightness control. System Visual Indicator states are explained in the following table.

Table 21: System Visual Indicator States and Explanations

System Visual Indicator State	Indicates
Off	System off or in startup testing
Solid Green	No alarms active
Solid Yellow	Low Priority Alarm active
Flashing Yellow	Medium Priority Alarm active
Flashing Red	High Priority Alarm active



NOTE: Even when the audio of active alarms is temporarily silenced using the Alarm Audio Pause button, the System Visual Indicators continue to display the corresponding visual alarm signal. If all active alarms are turned OFF (using the Alarm ON/OFF screen), or if all alarm conditions become satisfied, the indicator returns to a steady green illumination.

10.4 CHANGING SETTINGS AND DATA VIEWS

MOVES[®] SLC[™] has settings that the user can modify, for example, Vent O₂, Maximum Airway Pressure, etc. As well, there are customizable interactions to review data. All settings/data that can be manipulated are identified the same way, selected the same way, and are changed/viewed the same way. All modifiable settings/data views are shown on the screen as either a *screen button* or a *selectable area* with multiple elements displayed. These modifiable settings/data views can be selected and made active using the Jog Wheel.



10.4.1 Display of Settings and Views

Screen buttons and areas are displayed in one of four states Disabled, Selectable, Selected, and Active.

Table 22: Screen Buttons and Descriptions

Screen Button / Area State	Description	Example
Disabled	Shown with a double-line dark grey border and dark grey text. Identifies the button or area as something that is a button but is currently disabled. A disabled button cannot be selected.	NO
Selectable	Shown with a double-line white border. Identifies the button or area as something that can be selected for modification.	ECG lead:II
Selected	Shown the same as Selectable but with yellow lines forming the double-line border. Identifies the currently selected button or area.	MAIN
Active	Shown with a single, dashed line. Identifies the currently selected button or area as active and that changing of a setting is in progress.	ALARM 2 of 3 dismiss Ventilator failure

10.4.2 Modifying a Setting

To modify a setting:

- 1. Select the desired button or area by using the Jog Wheel until the desired selection is reached.
- 2. Press the Jog Wheel to initiate modification of the setting.
- 3. Use the Jog Wheel to select/modify the setting.
- 4. Press the Jog Wheel again once to accept the modified setting.



NOTE: In the case of the Alarm On/Off settings:

- Pressing the Jog Wheel toggles between the two states (i.e., on and off).
- To finalize and exit to the top level navigation, press the Back Button.

10.5 CONFIRMING DISPLAY VALIDITY

Note the following in regard to the display of valid information on the MOVES $^{\circledR}$ SLC $^{\intercal M}$ UI Screen:

The display should NOT show any elements that are partially written.

The display should ALWAYS show information while MOVES[®] SLC™ is on.





CAUTION! IF ANY OF THE ABOVE CONDITIONS ARE NOT SATISFIED, THE MOVES[®] SLC™ UNIT SHOULD BE DELIVERED TO AUTHORIZED PERSONNEL FOR SERVICING.

10.6 STARTUP SEQUENCE



NOTE: There should be **NO external oxygen** connected to the MOVES[®] SLCTM system during startup tests. Having O_2 connected creates a flow in the inspiratory limb which causes the open-circuit test to fail.

1. Locate the Power Control button on the rear of the MOVES[®] SLC™ unit (the end of the unit with the hydrocarbon filter and power-cord connection).



Figure 10-3: Power Control Button

2. Press the Power Control button to activate the unit – an audible alert is sounded and the user interface displays either the New Patient screen or the System Test screen

New Patient Screen

If the system has been run within the last 30 minutes, a prompt screen (New Patient Screen) will appear asking the user if the system should be configured for a new patient (i.e., begin with default values or restore previously used ones). The user should only select "NO" if the user is aware of what the previously used settings are, and the machine is still being used for the same patient.

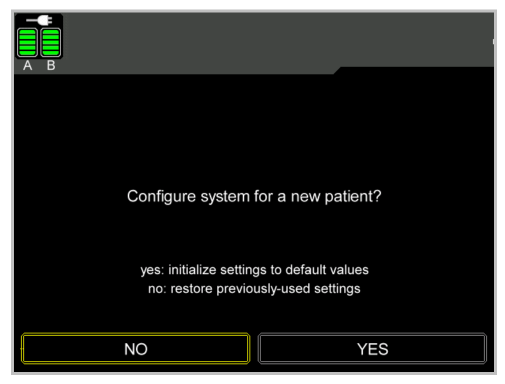


Figure 10-4: New Patient Screen



If there has been data loss and/or corruption of previous data, the user will NOT be allowed to restore previously used values. The "NO" option will be unavailable. The user will have to reconfigure the system manually to return to the previously used settings.

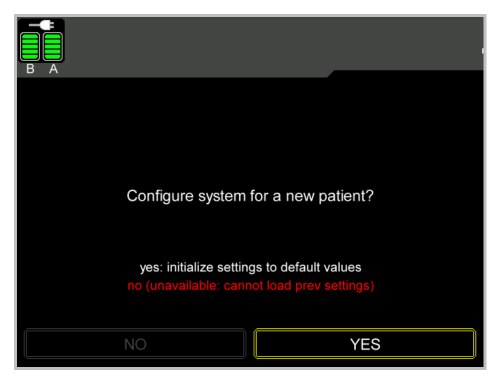


Figure 10-5: New Patient Screen – Restore Settings Unavailable

System Test Screen



NOTE: There should be **NO external oxygen** connected to the MOVES[®] SLCTM system during startup tests. Having O_2 connected creates a flow in the inspiratory limb which causes the open-circuit test to fail.

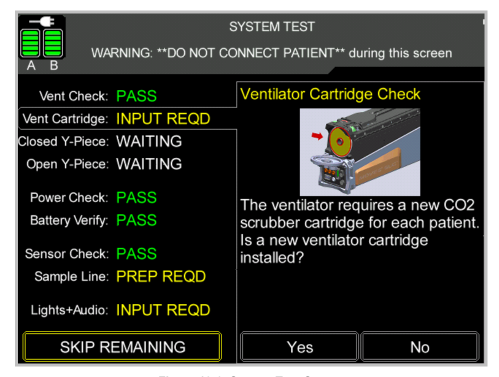


Figure 10-6: System Test Screen

- 3. If not all initial tests are passed, you will be offered certain choices such as SKIP REMAINING, PREP DONE, and SKIP THIS TEST along with Yes and No options.
- 4. Press the Jog Wheel to confirm a selection. Turn the Jog Wheel to move forward to the next or previous item in the list.



CAUTION! ONLY SKIP A TEST OR TESTS IF A SYSTEM TEST HAS PREVIOUSLY BEEN PERFORMED AND PASSED, AND THE SYSTEM HAS JUST BEEN RESTARTED DUE TO POWER FAILURE (E.G., LOSS OF BATTERY OR EXTERNAL POWER).



WARNING! DO NOT CONNECT ANY SENSORS, MONITORS, OR THE BREATHING CIRCUIT TO THE PATIENT WHILE PERFORMING SYSTEM TESTS! DOING SO COULD ENDANGER THE PATIENT!



WARNING! DO NOT OPERATE THE MOVES SLC $^{\mathrm{IM}}$ SYSTEM UNTIL ALL SYSTEM TEST FAILURES HAVE BEEN RESOLVED, AND ALL TESTS HAVE BEEN REPEATED AND PASSED.

Setup Screen

5. Once all prerequisites have been satisfied, the user must select Continue to move to the Setup screen.



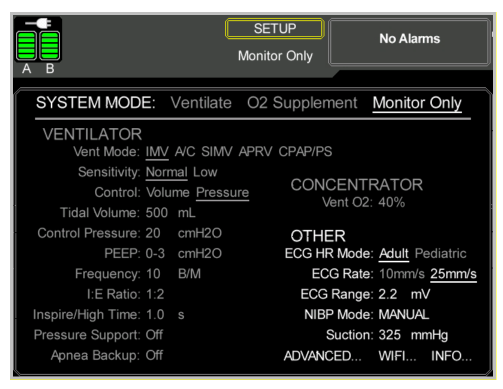


Figure 10-7: Setup Screen

6. The initial system mode will be Monitor Only. If a different system mode is desired, first configure the appropriate settings and then choose the new system mode (i.e., Ventilate or O2 Supplement).



NOTE: If an ABP/CVP/ICP transducer is to be connected to the system, it will be necessary to reset or 'zero' the transducer. For more information, see <u>Zeroing the Pressure in the IP Transducer on page 219</u>.



NOTE: All active alarms in the alarm queue should be reviewed for patient monitoring fault alarms immediately after completing the startup test procedure and before connecting to the patient.



NOTE: If a battery appears to be installed but is shown as missing on the display, inspect the battery and replace or reinstall. Also ensure that the battery doors are closed and latched correctly on both sides of the machine.



CAUTION! IF INTENDING TO RUN ON BATTERIES, ENSURE THAT THERE IS SUFFICIENT POWER FOR THE LENGTH OF TIME REQUIRED, OR REPLACE THE BATTERIES.



WARNING! THE OPERATOR SHOULD ALWAYS HAVE AVAILABLE AN ALTERNATIVE MEANS OF VENTILATION CAPABLE OF SUPPLEMENTING A HIGH CONCENTRATION OF O₂ IN THE EVENT OF POWER FAILURE, MECHANICAL FAILURE, OR SERIOUS OCCLUSION IN THE VENTILATOR SYSTEM.



WARNING! THE OPERATOR SHOULD ALWAYS HAVE AVAILABLE AN ALTERNATE MEANS OF SUPPLYING A HIGH CONCENTRATION OF O₂ IN THE EVENT OF POWER FAILURE, MECHANICAL FAILURE, OR SERIOUS OCCLUSION IN THE CONCENTRATOR CIRCUIT.



WARNING! THE OPERATOR SHOULD USE AN ALTERNATIVE MEANS OF VENTILATION UPON EXPERIENCING A PROLONGED APNEA ALARM.





WARNING! ONE CHARGED BATTERY MUST BE PRESENT IN THE MOVES SLC $^{\rm IM}$ Unit at all times, even when running on external power. This reduces the risk to the patient in the event of a power failure.

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11.0 MOVES® SLCTM User Screens

MOVES[®] SLC[™] has multiple user screens.

11.1 STATUS BAR

At the top of all screens is the Status Bar. The Status Bar is used to display system status and alarms and the name of the current screen. The Status Bar contains two buttons: the Screen Select button and the Alarm View button. The Status Bar shows the system power status (including battery charge and external power status), the number of alarms turned off, the number of alarm limits changed from their default value, alarm audio status, suction status, NIBP status, alarm status, WiFi status, and the system state.



Figure 11-1: Status Bar

In addition, a small white dot can be seen moving up and down at the top right of the Status Bar to indicate that the device is not "frozen" and that displayed information can be trusted to be current. If the white bar stops moving then the system display can no longer be trusted. If this occurs on the remote screen then discontinue use of the remote screen and only use the information displayed on the system screen. If the system screen is frozen, restart the MOVES[®] SLCTM. If freezing persists, discontinue use of the MOVES[®] SLCTM.

The following table explains the Status Bar items.



WARNING! WHEN SETTING ALARM LIMITS, DO NOT SET TO EXTREME VALUES THAT CAN RENDER THE ALARM SYSTEM USELESS.

Table 23: MOVES[®] SLC[™] Status Bar Items and Descriptions

Label/Name	Items on the Status Bar	Description
Plug icon with battery icons	Battery charge for each battery, and charge status.	 Graphic showing charge state of each battery. Shown with red color when battery life is low and with highlight background when one or more alarms is associated with system power. (NOTE: See the Battery Status Icon Table that follows for more information.)
7 7		 If the state of a battery cannot be determined, its outline is shown with the fault icon. If the entire power state cannot be determined, neither battery is shown, and
		a fault icon is displayed only.
Alarms OFF icon	Icon shows that some (or all) alarms are off. Number below indicates	The Alarm OFF icon is shown if any alarms have been turned off by the user on the Alarm ON/OFF screen.
	how many alarms are off.	The number of alarms 'XX OFF' is shown in text below the icon.
X	,,,	No icon or text is shown if all alarms are ON.



Label/Name	Items on the Status Bar	Description
Number of limits changed icon	Icon shows that some (or all) alarm limits have changed from their default values. Number below indicates how many alarm limits have been changed.	 The Alarm Limits Changed icon is shown if any alarm limits have been changed from their default values by the user on the Alarm Limits screen. The text below the icon indicates the number of alarm limits changed: '1 limit chgd' shown if only 1 limit is not set to its default value. 'X lims chgd' shown if 2-9 limits are not set to their default values. 'XX lim chgd' shown if 10 or more limits are not set to their default values. No icon or text is shown if all alarm limits are set to their default values.
Audio PAUSE icon	Icon shows that alarm audio is PAUSED. Numbers display how many minutes and/or seconds remain for audio pause. OR Icon shows that alarm	 The Audio PAUSE icon is shown if audio pause is active. This function is controlled by quickly pressing and releasing the Audio PAUSE button on the front panel. PAUSE time is 120 seconds max. The time remaining for audio pause is shown in text below the icon as 'm:ss', for example, 1:23 (1 minute 23 seconds); or, 0:03 (3 seconds). The Audio OFF icon is shown if audio OFF is active. This function is controlled by pressing and holding the Audio PAUSE button on the front panel for three (3) seconds. Audio PAUSE can only be active if Audio OFF is not active.
OFF icon	audio is OFF. Text describes that audio is OFF.	No icon or text is shown if Audio PAUSE / OFF is not active.
Suction Icon	Icon shows that Suction is active. Text below icon shows the suction pressure.	 Numerical values below the Suction icon show 1 to 999 mmHg in steps of 1 and for an unknown value or value less than or equal to 0. If a value is greater than 999, the value 999 is still shown. The Suction fault icon is shown if the status of suction cannot be determined.
NIBP icon	Icon shows that NIBP is active.	 The NIBP icon is shown if NIBP is obtaining a reading, which is controlled by the NIBP button on the front panel and/or by the auto NIBP setting on the Setup Screen. No text is displayed below the icon. Note that the cuff press is displayed with the NIBP reading on the main screen. No icon or text is shown if NIBP is not active. The NIBP fault icon is shown if the status of NIBP cannot be determined or is known to be in a fault condition.
Monitor Mode WiFi Status Icon	Icon shows that WiFi connection is active in monitor mode.	 The WiFi icon is shown if a WiFi connection is active (connecting or connected). A green icon indicates that the remote screen connection is only monitoring (i.e., no remote control). A blue icon indicates that the remote screen connection has remote control capabilities. The System Administrator configures whether the MOVES[®] SLC™ allows a wireless remote screen to have remote control capabilities.



Label/Name	Items on the Status Bar	Description
Remote Control Mode WiFi Status Icon	Icon shows that WiFi connection is active remote control mode.	When a wireless remote screen connection is in progress, all bars will be grey with the colored circle blinking. Once a connection is established the colored circle will no longer blink, and the number of colored bars indicates the signal strength of the WiFi connection.
Screen Select Button	Button indicates which screen is active. MAIN	When the Screen Select button is selected (double yellow lines), pressing the Jog Wheel makes it active (dashed single yellow line). Then turning the Jog Wheel cycles through the list of screens. Pressing the Jog Wheel a second time confirms the screen choice and causes the chosen screen to be displayed.
System State	Text indicates which mode is active. Monitor Only	 Text describes the system mode: Monitor Only O2 Supplement When the system mode is Ventilate mode, the system state text will describe the specific ventilation configuration (e.g., VC-SIMV+PS, which indicates the system is in Volume Controlled Synchronized Intermittent Mandatory Ventilation with spontaneous breaths receiving Pressure Support).
Alarm View Button	Alarm Queue	Three (3) line alarm message queue at the top right of screen. The alarm queue shows the highest priority alarm. LARM 1 of 4 Low expired GO2 To view additional alarms (if existing), use the Jog Wheel to select the Alarm Queue (double yellow lines). Then, press the Jog Wheel to make it active (dashed yellow line). Then, turning the Jog Wheel cycles through the alarms. Pressing the Back Button will exit the Alarm Queue and return to top level navigation.

11.1.1 Battery Status Icon

The Battery status icon shows the charge of the two system batteries individually; it also indicates if external power is connected. The batteries are drawn beside each other and labeled A and B. Charge is shown by filling in each chamber accordingly, indicated by 5 charge segments (the same number of segments as the physical battery LED status on the battery). When an alarm associated with system power is active, the area behind the battery status icons is highlighted with a red background if highest associated alarm is High priority. Otherwise, a yellow highlight is used for Medium/Low alarm priority. When the system is connected to external power, a plug icon is drawn above the battery icons.

Note the following:

- Each battery is individually hot swappable.
- If a battery has no charge, no segments will be shown.
- If a battery is not present, no segments and no battery outline will be shown.
- If MOVES[®] SLC[™] is evaluating a battery's status, a question mark will be drawn over that battery's status area.



- If a battery's status cannot be determined, a fault icon will be drawn over the battery's status area.
- If a battery is charging, a 'lightning bolt' will be drawn over the battery icon and its segments.
- If the power status is not known, a large fault icon will be displayed.

The color of the segments for a battery depends on the number of segments shown. If one segment is shown, the color is red; if two or three, the color is orange; if more than three segments are shown, the color is green. This matches the color of the top most LED on the battery charge indicator on the battery itself.

Table 24: Battery Status Icon Table

Battery Status Icon Table Battery A Full Charge Battery A Full Charge Battery A Unknown Battery B Half Charge Battery B Full Charge Battery B Near Empty Running on Battery Running on Battery **Running on Battery** High priority alarm No associated alarm(s) No associated alarm(s) **Battery A Missing Battery A Missing Battery A Unknown Battery B Half Charge Battery B Missing** Battery B Half Charge, Charging **Running on External Power Running on External Power Running on Battery** High priority alarm High priority alarm High priority alarm **Power Status Unknown Battery A Full Charge Battery A Near Empty Battery B Full Charge Battery B Near Full Running on External Power Power Status Unknown** Running on Battery No associated alarm(s) Alarm status ignored Medium/Low priority alarm



11.2 SYSTEM TEST SCREEN



NOTE: There should be **NO external oxygen** connected to the MOVES[®] SLCTM system during startup tests. Having O_2 connected creates a flow in the inspiratory limb which causes the open-circuit test to fail.



NOTE: The system will only operate if the power-on system memory and system firmware check was successful. The startup tests perform functional testing on components of the system that require operator intervention to validate – other parts of the system are continually tested and will show failure status in the alarm queue.



NOTE: The operator should wait until the O_2/CO_2 sensor has warmed up and O_2/CO_2 values are displayed before connecting a patient.



NOTE: All active alarms in the alarm queue should be reviewed for patient monitoring fault alarms immediately after completing the startup test procedure and before connecting to the patient.

Upon starting MOVES[®] SLC[™] the user, in most cases, will be presented with the System Test screen.

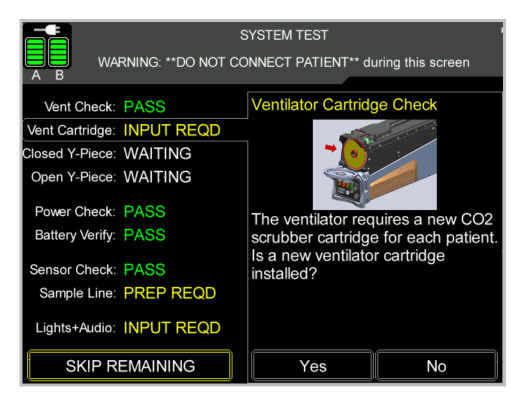


Figure 11-2: System Test Screen

Some tests are of the type INPUT REQD and the user is presented with Yes/No buttons. Use the Jog Wheel to select either option and then press the Jog Wheel to confirm it. Other tests require preparation, and the user is presented with the *Prep done* button. The System Test screen will indicate to the user which tests have passed and which need preparation before they can be conducted. A description of the preparation required is given at the right of the screen. Once preparation has been done, use the Jog Wheel to select *Prep done* and then press the Jog Wheel to initiate the test.



The user is also presented with the options of skipping a particular test or all remaining tests.



NOTE: In order to save time, certain tests can be run concurrently. For example, while the Closed Y-Piece test is still running, the screen will advance to the Lights + Audio test, which can be run concurrently.

The System Test screen will not immediately appear upon startup if:

- 1. The system has been shut down for a period of time less than or equal to 3 minutes. In this case, the system will auto resume using the system settings configuration prior to shutdown.(NOTE: A temporary loss of power is assumed here.)
- 2. The system has been shut down for a period of time greater than 3 minutes but less than 30 minutes. In this case, the system will query the user as to whether the patient is new or continuing (and the following screen will be displayed).

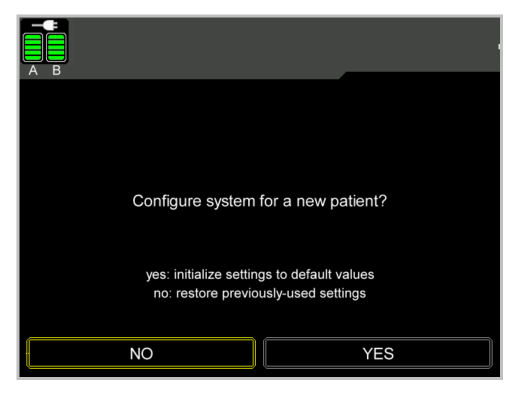


Figure 11-3: Configure for a New Patient Screen

Selecting YES will return the system settings to default values (see *Section 16.1 System Default Settings on page 283* for details). Selecting NO will keep the last system settings used. **NOTE:** NO should only be selected if the operator is aware of the last system settings configuration or the operator should review the settings on the Setup, Alarm Limits, Alarm ON/OFF and Advanced screens.

11.3 SETUP SCREEN

11.3.1 Overview

The Setup Screen is used to view and modify the primary operational settings of MOVES[®] SLC™. The color of a field is dependent on the mode selected. If a field is white, it is currently in use; otherwise, the field is gray. This lets the user see all the primary settings, including those that are not used by the current system mode.



11.3.2 Changing Settings



NOTE: Settings can be changed even in a <u>mode where the setting is not active.</u>

To change system settings:

- Use the Jog Wheel to navigate to the System Mode area (it becomes surrounded by a double yellow line).
- 2. Press the Jog Wheel (the System Mode area becomes surrounded by a dashed yellow line).
- 3. Use the Jog Wheel to navigate to the setting you want to change (it becomes surrounded by a single yellow line).
- 4. Press the Jog Wheel (the setting becomes surrounded by a dashed yellow line).
- 5. Use the Jog Wheel to select a new setting.
- 6. Press the Jog Wheel to confirm the new setting (the setting becomes surrounded by a single yellow line).
- 7. Press the Back Button to exit the System Mode area.

11.3.3 Setup Screen – Ventilate Mode



NOTE: Because there are five (5) Vent Modes (IMV, A/C, SIMV, APRV, and CPAP/PS), certain settings will be grayed out under each. However, these settings can still be changed even though the mode they apply to is not currently selected.

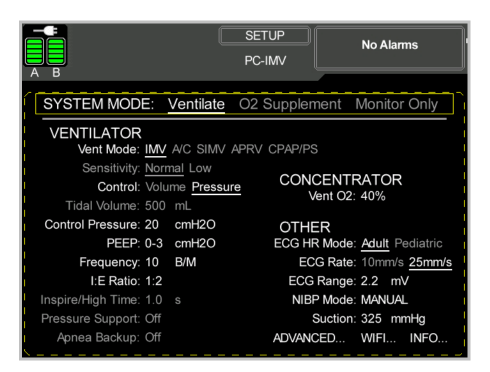


Figure 11-4: Setup Screen - Ventilate Mode

11.3.4 Setup Screen – O2 Supplement

When switching to O2 Supplement mode, you will first be presented with the following warning.



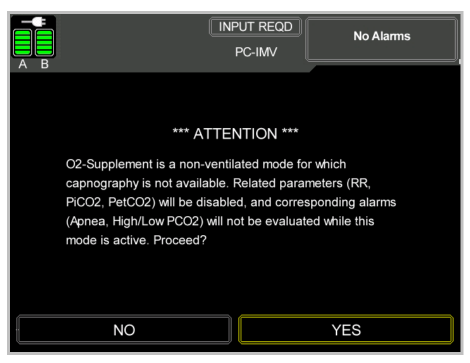


Figure 11-5: Setup Screen - No Capnography Warning

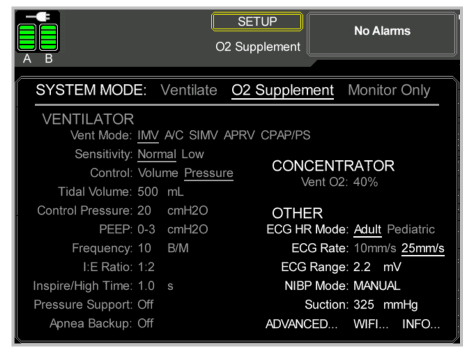


Figure 11-6: Setup Screen - O2 Supplement

11.3.5 Setup Screen – Monitor Only

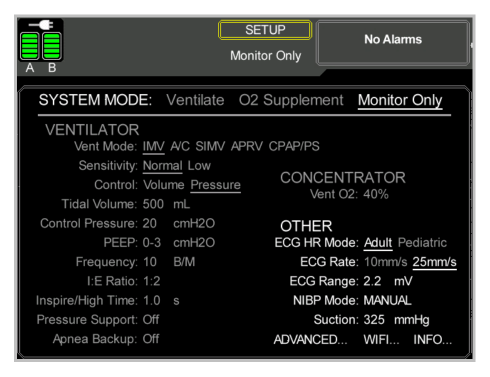


Figure 11-7: Setup Screen - Monitor Only

11.3.6 Setup Screen Options

Table 25: Setup Screen Options and Descriptions

Setup Screen Options		
Fields on the Setup Screen	Used by System Mode	Description
System Mode	All	 Ventilate Mode, O2 Supplement, Monitor Only. Default: Monitor Only Change immediately after change confirmed
Vent Mode	Ventilate Mode	 IMV, A/C, SIMV, APRV, CPAP/PS Default: IMV Change at start of next inhalation after change confirmed.
(Breath) Sensitivity	Ventilate Modes: A/C, SIMV, CPAP+PS	 Normal, Low. Default: Normal Change immediately after change confirmed
Control	IMV, A/C, SIMV	 Volume, Pressure Default: Pressure Change at start of next inhalation after change confirmed.



Setup Screen Options			
Fields on the Setup Screen	Used by System Mode	Description	
Tidal Volume	Ventilate Modes: IMV, A/C, SIMV when Control is set to Volume, and in CPAP+PS with Pressure Support on and Apnea Backup on and Control is set to Volume	 50 to 250 ml in 10 ml intervals 250 to 750 ml in 25 ml intervals Default: 500 ml After a change is accepted, the option selected will be active after the patient's current breath. 	
Control Pressure	Ventilate Modes: IMV, A/C, SIMV when Control is set to Pressure, and in CPAP+PS with Pressure Support on and Apnea Backup on and Control is set to Pressure	 Pressure Control (over PEEP) 10 to 55 cmH₂O in intervals of 1 cmH₂O Default: 20 cmH₂O After a change is accepted, the option selected will be active after the patient's current breath. NOTE: Control Pressure = PIP (Peak Inspiratory Pressure) – PEEP (Positive End Expiratory Pressure) 	
PEEP	Ventilate Mode	 Positive End-Expiratory Pressure 0-3, 4 to 20 cmH₂O in intervals of 1 cmH₂O Default: 0-3 cmH₂O NOTE: 0-3 cmH₂O represents the minimum PEEP setting for the system. When PEEP is set at: 0-3 cmH₂O, the system will maintain a slight bias PEEP pressure of 3 cmH₂O for improved trigger accuracy and system responsiveness. After a change is accepted, the option selected will be active after the patient's current breath. NOTE: When the PEEP value is increased by 10 cmH₂O or more some additional breath triggers may be observed. This includes starting the ventilator at a PEEP setting of 10 cmH₂O or more. 	
Frequency	All Ventilate modes except CPAP/PS	 Respiratory Rate 6 to 40 B/M in steps of 1 B/M Default: 10 B/M Change at start of next inhalation after change confirmed. 	
I/E ratio	Ventilate Modes: IMV, A/C, and in CPAP+PS with Pressure Support on and Apnea Backup on	 Inspiratory/Expiratory Ratio 1:1, 1:1.5, 1:2, 1:2.5, 1:3. Default: 1:2. Change at start of next inhalation after change confirmed. 	



Setup Screen Options		
Fields on the Setup Screen	Used by System Mode	Description
Inspire / High Time	Ventilate Mode: SIMV, APRV	 Inspiratory Time in SIMV High Time in APRV 0.3 to 5.0 seconds in steps of 0.1 seconds Default: 1.0 second Change at start of next inhalation after change confirmed.
Pressure Support	Ventilate Modes: SIMV, CPAP/PS	 Off, 5 to 40 cmH₂O in intervals of 1 cmH₂O Default: Off Change at start of next inhalation after change confirmed.
Apnea Backup	CPAP+PS	Off, OnDefault: Off
Vent O2	Ventilator Mode Only	 Ventilate Mode: Air, 30%, 40%, 50%, 60%, 70%, 85%, Maximum (always on) Default: 40% Change immediate after change confirmed.
ECG Rate	All	 Sweep speed of ECG. ECG is displayed on graphs on Monitor screen or dedicated ECG screen. 10 mm/s or 25mm/s* Default: 25 mm/s 10 mm/s is 60 Hz data point display 25mm/s is 150 Hz Change immediate after change confirmed. * NOTE: ECG Rate listed is for MOVES® SLC™, the actual rate on the MOVES® SLC™ remote screen will be different. Refer to Section 14.2.1 for more details.
ECG Range	All	 Maximum amplitude range of ECG. ECG is displayed on graphs. 2.2 mV, 3 mV, 6 mV or 12mV range Default: 2.2 mV Change immediately after change confirmed. NOTE: Range is centered about zero (e.g., 3 mV is +/-1.5 mV). NOTE: A dashed reference line indicating 1 mV is always displayed on graphs



Setup Screen Options		
Fields on the Setup Screen	Used by System Mode	Description
NIBP	All	 NIBP Measurement Period Manual, Stat, 1, 2, 3, 4, 5, 10, 15 min Default: Manual In Manual mode no automatic NIBP measurements are taken. Press the NIBP button to start a measurement. Stat mode acquires NIBP measurements as fast as possible while guaranteeing 5 seconds between the end of a measurement and the start of the next measurement. Change immediately after change confirmed. NOTE: After 15 minutes of Stat mode, the mode will automatically switch to 5 minute mode. IEC 80601-2-30 201.105.2 states that Stat mode can only last for 15 minutes, and then must go to manual or long term automatic. NOTE: If NIBP measurement period is set to anything other than Manual, and three automatic NIBP measurements fail to obtain a reading, then the system will automatically switch the mode to Manual and notify the operator of this change.
Suction	All	 100 to 325 mmHg in steps of 25 mmHg Default: 325 mmHg Change immediately after change confirmed.
ADVANCED	N/A	 Conveys user to the Advanced screen where advance system settings can be viewed and modified (e.g., SpO₂ Average Time, SpO₂ Sensitivity Mode, ECG EMG Filter, etc.). Additionally, PulseOx feature availability can be verified.
WIFI	N/A	Conveys user to the WIFI setup screen where the wireless connection can be configured and established.
INFO	N/A	 Conveys user to the Info screen which displays: System serial number Total concentrator and ventilator runtimes System firmware versions Masimo information Masimo feature availability



11.4 ADVANCED SCREEN

11.4.1 Accessing the Advanced Screen

The Advanced Screen is accessed from the Setup Screen. The user navigates to the System Mode area using the Jog Wheel and presses the Wheel to make it active. The user then navigates to the ADVANCED... option and presses the Jog Wheel again to proceed to the Advanced Screen.



Figure 11-8: Accessing the Advanced Screen



11.4.2 The Advanced Screen

The Advanced screen opens with the first option selected.

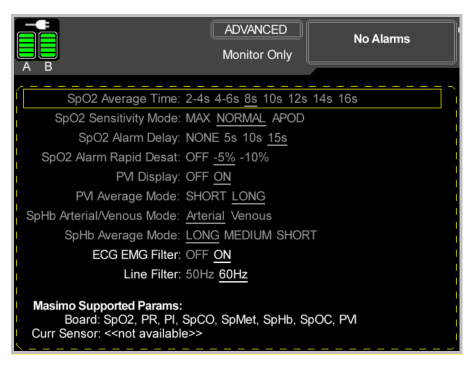


Figure 11-9: The Advanced Screen

Options are selected as on other screens, by navigating to an option with the Jog Wheel and the pressing the Wheel to make it active. The following table lists the options and values available on the Advanced Screen.

Table 26: Advanced Screen Options

Advanced Screen Options		
Option	Values Available	Default Value
SpO2 Average Time	2-4s, 4-6s, 8s, 10s, 12s, 14s, 16s	8s
SpO2 Sensitivity Mode	MAX, NORMAL, APOD	NORMAL
SpO2 Alarm Delay	NONE, 5s, 10s, 15s	15s
SpO2 Alarm Rapid Desat	OFF, -5%, -10%	-5%
PVI Display	OFF, ON	ON
PVI Average Mode	SHORT, LONG	LONG
SpHb Arterial/Venous Mode	Arterial, Venous	Arterial
SpHb Average Mode	LONG, MEDIUM, SHORT	LONG
ECG EMG Filter	OFF, ON	ON
Line Filter:	50 Hz, 60 Hz	Initial setting is 60 Hz; however, if changed, last setting is remembered. Does not default to 60 Hz with a new patient.



11.4.3 SpO2 Average Time

The user-selectable SpO2 averaging feature allows the operator to select the desired level of visibility applied to subtle variations in the measured value. Eight (8) second averaging is generally considered the most common averaging interval, and it is recommended for most patients since it is short enough to provide visibility to subtle desaturations while also being long enough to minimize major changes in SpO2 due to quick, transitory desaturations.

11.4.4 SpO2 Sensitivity Modes

The three sensitivity settings allow the operator to adapt the SpO₂ measurement sensitivity to the patient's level of SpO₂ signal strength and quality at the measurement site.

MAX

MAX mode is used to obtain and display data even when the signal is very weak due to impaired perfusion (can be used, for example, during treatment or examination, i.e., when someone is with the patient). If the sensor becomes detached from the patient, this mode makes virtually no provision for measurements that are displayed erroneously.

NORMAL

NORMAL is the recommended mode for typical monitoring, such as intensive care units.

APOD (Adaptive Probe Off Detection)

APOD is the recommended mode when it is not possible to detect immediately if a sensor has become detached. This mode offers better protection against erroneous measurements being displayed, even though the sensor has become detached from the measurement point (e.g., as a result of the patient moving).

11.4.5 SpO2 Alarm Delay

The SpO2 alarm delay allows the user to adjust the time after which the Low SpO2 alarm will be escalated from a LOW priority alarm to a HIGH priority alarm, once a low SpO2 alarm condition has been initiated. Many desaturations are real, but transitory and, as such, may not require clinical intervention. Therefore, the priority of the desaturation alarm is escalated once the desaturation is determined not to be momentary, thus reducing the number of high priority alarm conditions.

The options available are NONE, 5 seconds, 10 seconds, and 15 seconds. If an SpO2 limit alarm occurs, and the time is less than the delay time, the SpO2 value is highlighted but the alarm is classified as LOW priority. If the alarm maintains its status for longer than the delay value, it is upgraded to HIGH priority. Selecting the option NONE removes the alarm delay and the priority of the Low SpO2 alarm will always be HIGH priority.

11.4.6 SpO2 Alarm Rapid Desat

If a very rapid desaturation occurs, a user would not want the alarm to be delayed. This is the reason for the SpO2 Alarm Rapid Desat feature. Options available are OFF, -5% and -10%. Percentages are in terms of the maximum 100% saturation possible, not a percentage of the user-set Low SpO2 Alarm Limit value. If the feature is set to OFF, the alarm delay time is always in effect.

The SpO2 Alarm Rapid Desat threshold is determined by taking the Low SpO2 Alarm Limit value and adding the value of the set point (either -5% or -10%). For example, imagine that the Alarm Delay is set to 15 seconds, the Low SpO2 Alarm Limit to 85% and Rapid Desat to -10%. Therefore, in this example, the SpO2 Alarm Rapid Desat threshold is 75%. As the SpO2 falls to 85%, the Low SpO2 alarm triggers at LOW priority and the following scenarios are possible:

If the SpO2 remains at 85%, then after 15 seconds the Low SpO2 alarm becomes HIGH priority.



- If the SpO2 falls below 85%, continues to fall, and after only 10 seconds reaches 75%, the HIGH priority alarm is triggered.
- If the SpO2 falls to 75% but then climbs to 76% in less than 15 seconds, then the Low SpO2 alarm does not
 return to Low priority. Once the Low SpO2 alarm becomes HIGH priority, the alarm stays HIGH priority until the
 low SpO2 alarm condition is no longer present.

11.4.7 PVI Display

Allows disabling or re-enabling display of the PVI measurement value depending on whether it is needed by the operator.

11.4.8 PVI Average Mode

The PVI Average Mode allows the clinician to select the desired level of visibility to subtle variations in the PVI value. This allows a clinician to fine tune PVI responsiveness to achieve the desired level of visibility to rapid variations in PVI values.

11.4.9 SpHb Arterial/Venous Mode

While monitoring Hb levels, there are two blood sample sources from which Hb readings can be obtained: arterial and venous. Arterial Hb measurements can be expected to be slightly lower than the Hb measurement derived from venous blood. This feature allows the clinician to tailor the SpHb measurement to their clinical practice and/or setting.

11.4.10 SpHb Average Mode

The SpHb Average Mode allows the clinician to select the desired level of visibility to subtle variations in the SpHb value. This allows a clinician to fine tune SpHb responsiveness to achieve the desired level of visibility to rapid variations in SpHb values.

11.4.11 ECG EMG Filter

When enabled (i.e., ON) this feature filters the ECG waveforms for EMG interference in the frequency range of 15-30 Hz.

11.4.12 Line Filter

Depending on the geographical region, the Line Filter setting must be configured to match the power line frequency used in that region. This setting is used by the ECG and Pulse CO-Oximeter modules to allow for cancellation of noise introduced by fluorescent lights and other sources which may improve signal quality.

11.4.13 Masimo Supported Parameters

Masimo parameters supported by MOVES[®] SLC™ are listed. More detailed Masimo information is supplied on the Info Screen.

11.5 WIFI SCREEN

11.5.1 Accessing the WIFI Screen

The WIFI Screen is used to connect to the remote screen tablet. The WIFI Screen is accessed from the Setup Screen. The user navigates to the options area using the Jog Wheel and then presses the Wheel to make it active. The user then navigates to the WIFI... option and presses the Jog Wheel again to proceed to the WIFI Screen.



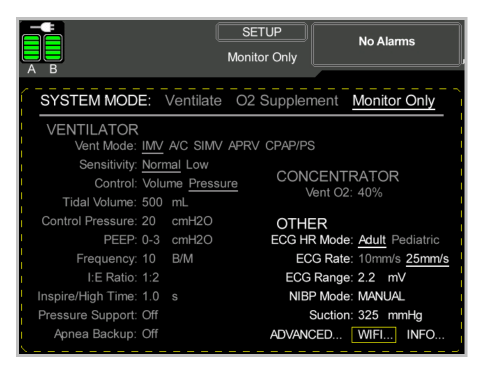


Figure 11-10: Accessing the WIFI Screen

11.5.2 The WIFI Screen

The WIFI Screen opens with the name of the most recently connected tablet in the Tablet field. On the following screen this is JFL127. From the WIFI Screen, you can connect wirelessly to a remote screen tablet. For details on connecting, see *Connecting to the Remote Screen Wirelessly from MOVES® SLC™ beginning on page 194*.



Figure 11-11: The WIFI Screen



11.6 INFO SCREEN

11.6.1 Accessing the Info Screen

The Info Screen is accessed from the Setup Screen. The user navigates to the System Mode area using the Jog Wheel and then presses the Wheel to make it active. The user then navigates to the INFO... option and presses the Jog Wheel again to proceed to the Info Screen.

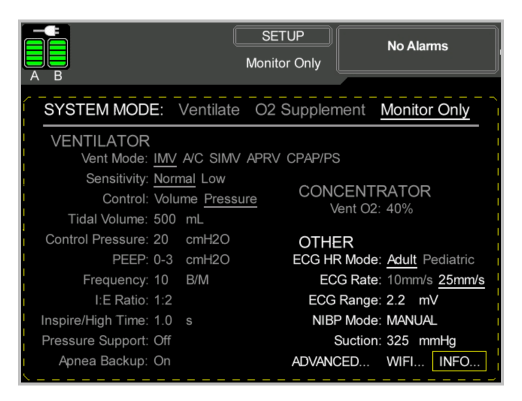


Figure 11-12: Accessing the Info Screen

11.6.2 The Info Screen

Only the Screen Button and Alarm View Button can be selected on the Info Screen. The remaining items simply list system information.

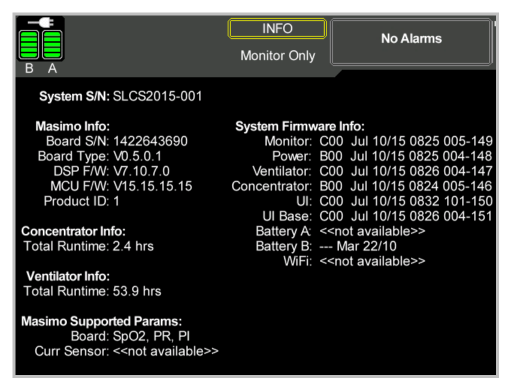


Figure 11-13: The Info Screen

11.6.3 Information Located on the Info Screen

Table 27: Info Screen Items

Info	Items
System S/N	System manufacturer serial number
Masimo Info	Board S/N (serial number)
	Board Type
	 DSP F/W (digital signal processor firmware version)
	MCU F/W (microcontroller firmware version)
	Product ID (identification)



Info	Items
Masimo Supported Params (parameters):	Board:
The screen provides a list of all supported parameters for the <i>Pulse CO-Oximeter (Board)</i> and the currently attached sensor (<i>Current Sensor</i>). A parameter must be supported by the Pulse CO-Oximeter and by the sensor in order for the parameter measurement to be displayed. Listed below are all possible parameters: SpO2 PR - Pulse Rate PI - Perfusion Index PVI – Pleth Variability Index SpCO SpMet SpHb SpOC NOTE: See Section 7.5 Pulse Oximeter Technology Overview beginning on page 54 for a description of each of these parameters.	 SpO2, PR and PI are always supported The extended parameters SpCO, SpMet, SpHb and SpOC are optional features which may or may not be present depending on the requested configuration when MOVES[®] SLC™ was manufactured. Current Sensor: All possible parameters the currently attached sensor can measure will be listed, even if the Pulse CO-Oximeter does not support the parameter. If no sensor is presently connected to the Pulse CO-Oximeter, then <<not style="list-style-type: square;">< not available>> is shown</not>
System Firmware Info (version date) NOTE: Upon completing system firmware updates, the system should be power cycled before verifying that correct versions are installed.	 Monitor (Patient Monitor) Power (Power Manager) Ventilator Concentrator UI (User Interface) UI Base Battery A Battery B WiFi
Concentrator Info	The cumulative number of hours of operation of the concentrator.
Ventilator Info	The cumulative number of hours of operation of the ventilator.



NOTE: If batteries are not inserted, the << not available >> notification is displayed.

11.7 MAIN SCREEN

The Main Screen is used to display patient status and monitored values. It is the primary screen of MOVES[®] SLCTM. The Main Screen is for status <u>only</u>. There are no functional settings to change.





NOTE: The values displayed on the Monitoring Screen are those that are actually measured from the patient and not how the system is configured (e.g., The Monitoring Screen displays the actual volume, pressure and respiratory rate measured from the ventilator.)

The available buttons are for the following:

- Choosing which chart to view
- Choosing the source of heart rate to view
- Zeroing the invasive pressures channels
- Labeling the invasive pressure channels

Any value that has an active alarm associated with it is drawn in reverse color (non black background).



Figure 11-14: Screen Items Drawn in Reverse

For all numeric or text display items, if no valid data is available, dashes are displayed.



Figure 11-15: Screen Items Displaying Dashes

For all numeric or text display items, if a fault prevents data display the fault icon (X) is displayed.

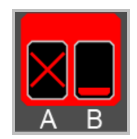




Figure 11-16: Screen Items Displaying the Fault Icon

For all display items, a trusted number is always shown or represented; otherwise, for values that are not possible – and therefore not trusted – dashes are displayed and the systems acts as if the data is not available. An alarm is usually generated for this untrusted number.

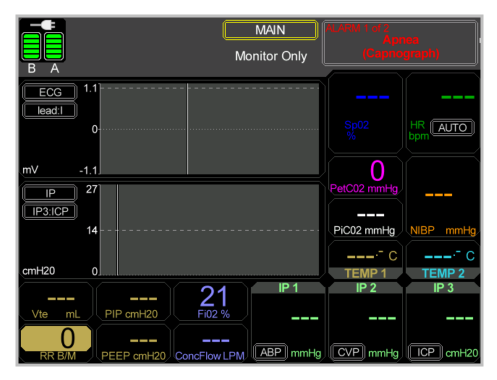


Figure 11-17: Main Screen with Dashes & Alarms

11.7.1 Selection Order for Options

If the Jog Wheel is turned clockwise, the selection order for Main Screen buttons follows a circular pattern around the screen as follows. If it is turned counterclockwise, the order is reversed.

- 1. Screen Button
- 2. Alarm View Button
- 3. HR Source
- 4. IP 3
- 5. IP 2
- 6. IP 1
- 7. Chart 2
- If Trends, followed by Data Source, Data Type & Time
- If ECG, followed by Data Source
- 8. Chart 1
- If Trends, followed by Data Source, Data Type & Time
- If ECG, followed by Data Source

- If IP, followed by Data Source
- All others followed by Chart 1
- If IP, followed by Data Source
- All others followed by Screen Button

SAMPLE CHARTS

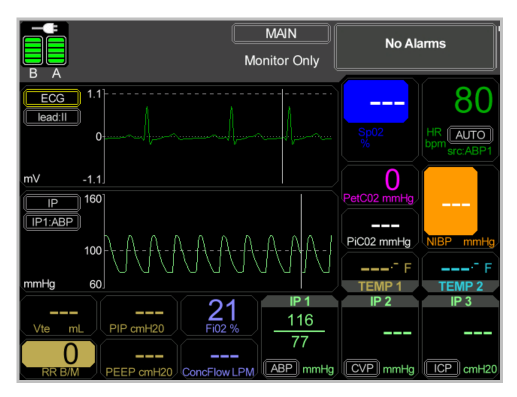


Figure 11-18: ECG & IP Graph Options & Sub-Options

11.7.2 Main Screen Items

Table 28: Main Screen Items and Descriptions

Label	Items on the Main Screen	Description
Vte	Expired Tidal volume	Numeric 1 to 2500 ml in steps of 1 ml, or a series of dashes () for unknown.
		 Only displayed when ventilating; otherwise, a series of dashes () is shown.
		• ">2.5 L" is shown if 2501 ml or higher.
Vti	Inspired Tidal volume	Vti is displayed in the location of Vte, alternating with Vte display every 2 seconds. This is only shown if:
		 "High expired tidal volume" alarm is active <u>OR</u>
		o "Leak detected" alarm is active.
		 Numeric 1 to 2500 ml in steps of 1 ml, or a series of dashes () for unknown.
		 Only displayed when ventilating; otherwise, series of dashes () is shown.
		• ">2.5 L" is shown if 2501 ml or higher.



Label	Items on the Main Screen	Description
RR	Respiratory Rate, calculated from the CO ₂ monitor	 Numeric 0–99 B/M in steps of 1, or a series of dashes () for unknown value. ">99" is shown if 100 B/M or higher.
PIP	Peak Inspiratory (Airway) Pressure	 Numeric 0–100 cmH20 in steps of 1, or a series of dashes () for unknown / out of range. Only displayed when ventilating, otherwise, or a series of dashes () is shown.
PEEP	Positive End Expiratory (Airway) Pressure	Same as PIP
FiO ₂	Fraction of Inspired Oxygen (O ₂)	 Numeric 0–100 % in steps of 1 or () for unknown value. If >100 and ≤105, shows 100, if >105 shows (). If unavailable due to calibration / warming-up shows "CAL".
ConcFlow	Concentrator Flow	 Volumetric flow of oxygen from the concentrator into the ventilator or to O2 Supplementation port (depending on system mode) at a nominal oxygen concentration of 90%. Numeric 0.0 to 4.0 LPM ">4.0 LPM" shown for any value over 4.0 LPM.
Temp1	Patient Temperature 1	 Numeric 82.4 to 108.0° F (or 28 to 42° C) in steps of 0.1 or () for unknown value. Values outside of human body temperature range are shown as follows: "<28.0" or ">42.0" (Celsius) "<82.4" or ">108" (Fahrenheit)
Temp2	Patient Temperature 2	Same as Temp1.
PetCO ₂	Partial Pressure of End-Tidal Carbon Dioxide	 Numeric 0–99 mmHg in steps of 1, or a series of dashes () for unknown value; otherwise, ">99" is shown. If unavailable due to calibration / warming-up, shows "CAL".
PiCO ₂	Partial Pressure of Inspired Carbon Dioxide	Same as PetCO ₂
SpO ₂	Saturation percentage of oxygen attached to hemoglobin. (The amount of oxygen being carried by the red blood cells in the blood.)	 Numeric 0–100% in steps of 1 or a series of dashes () for unknown value. If >100, shows ">100". If unavailable due to initialization shows "INIT".
	nd the appropriate Masimo sensor is present. The	onditionally displayed when the corresponding feature is hese five (5) values can only be found on the left of the pleth
PVI	Pleth Variability Index (PVI may help clinicians noninvasively assess fluid status of patients and predict fluid responsiveness.)	 Numeric 0–100% in steps of 1 or a series of dashes () for unknown value. If unavailable due to initialization shows "INIT".



Label	Items on the Main Screen	Description
SpCO	Saturation percentage of carbon monoxide attached to hemoglobin (i.e., carboxyhemoglobin). (The amount of carbon monoxide being carried by the red blood cells in the blood.)	 Numeric 0–100% in steps of 1 or a series of dashes () for unknown value. If unavailable due to initialization shows "INIT".
SpMet	Saturation percentage of methemoglobin. (Methemoglobin [MetHb] is an oxidized form of hemoglobin that is unable to carry oxygen.)	 Numeric 0.0–100.0% in steps of 0.1 or a series of dashes () for unknown value. If unavailable due to initialization shows "INIT".
SpHb	Total hemoglobin (Hb) concentration. (Hemoglobin is the part of a red blood cell that carries oxygen to the body. SpHb, which measures total hemoglobin, indicates the oxygen carrying capacity of the blood.)	System can be configured (by admin) to display in one of three different units: g / dL (grams hemoglobin / deciliter blood) Numeric 0.0–25.0 g/dL in steps of 0.1 or a series of dashes () for unknown value. mmol / L (millimoles hemoglobin / liter blood) Numeric 0.0–15.5 mmol/L in steps of 0.1 or a series of dashes () for unknown value. g / L (grams hemoglobin / liter blood) Numeric 0–250 g/L in steps of 1 or a series of dashes () for unknown value.x If unavailable due to initialization shows "INIT".
SpOC	Total oxygen content. (SpHb and SpO ₂ are used together to calculate the actual amount of oxygen in the blood.)	 Shown as a ratio: mL O₂ / dL blood (milliliters oxygen / deciliter blood) Numeric 0–35 mL/dL in steps of 1 or a series of dashes () for unknown value. If unavailable due to initialization shows "INIT".
n/a	Perfusion Index (Perfusion Index, or PI, is a relative assessment of the pulse strength at the monitoring site.)	 0–10.00% displayed as bar graph beside SpO2 Values between 10.001–20.000% display as a full bar graph. When there is no data, the graph is not drawn.



Label	Items on the Main Screen	Description
NIBP	Non-Invasive Blood Pressure The previous NIBP reading (if any), will be displayed while the new reading is being obtained. The current progress of the new reading will be displayed (if in progress) by showing the cuff pressure in a vertical bar graph beside the previous reading. The cuff pressure graph is from 0 to 260 mmHg (if over 260, bar graph is displayed as full). If the user cancels the current NIBP reading, the previous one will be removed from the screen. If the NIBP reading is invalid and/or cannot be obtained, the previous NIBP reading will be erased.	 If either the systolic or diastolic value is over the range of the sensor, ">300" is shown for that value. If the NIBP is one (1) or more minutes old, a message states "XXm ago" where XX is the number of minutes since the last reading. A NIBP measurement older than 15 minutes is discarded and no longer shown. If the NIBP reading was obtained using a cuff pressure that was close to the systolic value, which may make the reading lower than the actual systolic value of the patient, the systolic value is displayed with a series of dashes () to indicate the systolic pressure could not be obtained. The NIBP will automatically obtain another reading using a higher cuff pressure immediately after the reading. If neither the systolic or diastolic pressure is available, a single set of dashes () is shown and no divider line is drawn. While an NIBP reading is in progress, the current BP reading, if any, remains. As well, a bar graph representing cuff pressure from 0 to 260 mmHg is displayed to the right of the reading. The current cuff pressure is displayed below the cuff pressure bar.



Label	Items on the Main Screen	Description
ABP, CVP, or ICP	IP Invasive Pressure There are three IP inputs on MOVES [®] SLC™. They can be used to measure one or more of Arterial Blood Pressure (ABP), Central Venous Pressure (CVP) or Inter-Cranial Pressure (ICP)	 For ABP numeric, if either the systolic or diastolic value is over the range of the sensor, ">300" is shown for that value. If either the systolic or diastolic value is under the range of the sensor, "<-10" is shown for that value. For CVP numeric, Numeric -10 to 300 mmHg in steps of 1 is shown. If greater than 300 mmHg, shows ">300". If less than -10 mmHg, "<-10" is shown. For an unknown value, a series of dashes () is shown.
	ABP is displayed in a similar fashion to NIBP; that is, it is displayed showing systolic over diastolic in mmHg.	• For ICP numeric, Numeric –14 to 408 cmH ₂ O in steps of 1 is shown. If greater than 408 cmH ₂ O, shows ">408". If less than –14, "<–14" is shown. For an unknown value, a series of dashes () is shown.
	CVP or ICP is displayed as a single number, the mean pressure with up to 3 digits. CVP is displayed in mmHg, and ICP is displayed in cmH ₂ O. The software will be able to detect if a port is in use via hardware. There can be none, one, two, or three IPs in use at any time.	IP 1



Label	Items on the Main Screen	Description
HR	Heart Rate	The source of the HR is one of ECG, PulseOx, or IP in the following priority (if set to AUTO): APB1 APB2 APB3 SPO2 ECG HR Source: ABP or ECG Numeric 0–250 BPM in steps of 1 or a series of dashes () for an unknown value. ">250" is shown for values over 250 BPM HR Source: SPO2 (Pulse Oximeter) Numeric 0–239 BPM in steps of 1 or a series of dashes () for an unknown value. ">239" is shown for values over 239 and below 260 BPM The user can also set the source by pressing the source button below the HR and selecting the source from a list.
Chart Label 1	Chart area 1	Button to select which chart to view.Other buttons as required by the chart.
Chart Label 2	Chart area 2	Same as Chart 1



11.7.3 Additional Items Displayed with Masimo Sensors

As stated above, "PVI" and Rainbow SET® (SpMet, SpCO, etc) items are only displayed when the appropriate Masimo sensor is present. The values are found on the left of the pleth real-time graph. The following two screen captures illustrate the values obtained via two of the Masimo sensors.

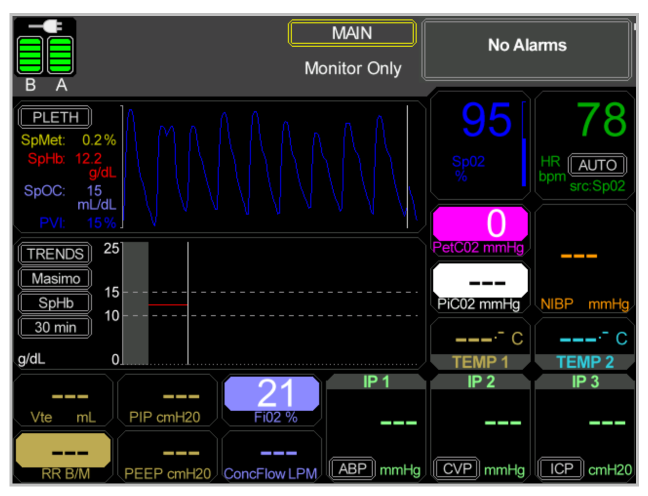


Figure 11-19: Values Obtained Via Masimo DC-3 Sensor



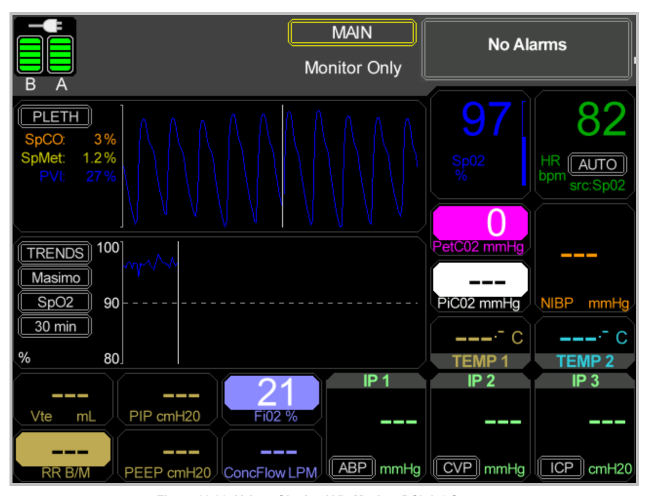


Figure 11-20: Values Obtained Via Masimo DCI-dc3 Sensor

11.7.4 Control Pressure, PEEP and PIP

It may be noticed that although an option for "Control Pressure" is present on the Setup Screen, only the items "PIP" and "PEEP" are displayed on the Main Screen.

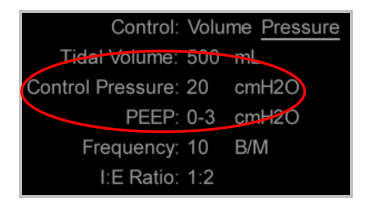


Figure 11-21: Control Pressure & PEEP on Setup Screen



Figure 11-22: PIP & PEEP on Main Screen



On ventilators with an active PEEP (Positive End Expiratory Pressure), such as MOVES[®] SLCTM, the user sets the Control Pressure, which is the pressure <u>above</u> PEEP, that the user wants maintained. If the user adjusts the PEEP value, the PIP (Peak Inspiratory Pressure) value goes up along with it.

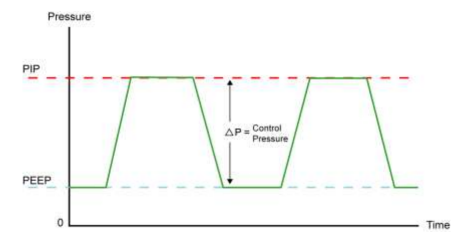


Figure 11-23: Relations of PEEP to PIP to Control Pressure

However, monitoring (as it is shown on the Main Screen) always displays the true airway pressures: PIP and PEEP.

11.7.5 Actual PEEP Higher than Set PEEP

Note that not all combinations of ventilator parameters are possible to obtain with all patients because of individual patient lung condition, resistance and compliance. If insufficient time is provided for the patient to exhale, the ventilator will delay the start of the next inhalation until the airway pressure has at least fallen to PEEP + 5 cmH₂O. It will also produce an "Expiratory obstruction" alarm. Whenever <u>set</u> PEEP is not being achieved due to this auto-PEEP scenario, the situation can be corrected by increasing the exhalation time, either by reducing the respiratory rate, reducing the inspire time or reducing the I:E ratio. When the ventilator is in an auto-PEEP scenario, the ventilator may under-report expired volumes.

11.7.6 Inverted Display of Patient Monitoring Values

Normally, patient monitoring values on the Main Screen are shown in their representative colors (e.g., Temp 1 in tan, Temp 2 in turquoise) against a black background (see Figure 11-24). However, when an active alarm is associated with monitoring value, the values are shown in either white or black against a background of their representative color (see Figure 11-25).



NOTE: The active state of the alarm will be indicated by inverted display regardless of whether the alarm is on or off. For example, if the patient temperature is high, causing an alarm condition, the patient temperature is drawn in reverse color, even if the patient temperature alarm has been turned off by the user. If an alarm is latched, meaning it must be acknowledged by the user even if the alarm condition no longer active, the monitoring display is only inverted while the alarm condition is active.







Figure 11-24: Regular Temperature Display

Figure 11-25: Inverted Display of Temperature (Alarm Condition)

11.7.7 Invasive Pressure (IP) Source Buttons

There are three (3) physical IP connections on the patient sensor panel. Each one maps to its corresponding IP channel box on the monitoring screen (labeled IP1, IP2 and IP3). Each channel is independently configurable, and can be set for ABP, ICP or CVP mode. The current mode is displayed on the config button and may be changed by using the Jog Wheel. An additional option is available, "ZERO", which does not change the mode, but rather forces an immediate rezeroing of the channel.

11.7.8 Zeroing a Channel

An IP port will auto detect the insertion of an IP probe. An inserted probe requires zeroing before data is available, and will show ZERO REQD until the channel config button – now titled "ZERO" – is clicked, zeroing the channel. The channel box will then immediately display data with format/units appropriate for the selected mode. The user can change modes without re-zeroing. The port will also detect a probe disconnect and raise an alarm to notify the operator. This alarm may be dismissed to acknowledge the condition.

11.8 ECG SCREEN

11.8.1 Overview

The ECG Screen will display up to 12 graphs (if all 10 ECG lead wires are used). The number of leads displayed depends on which lead wires are connected. If all lead wires are connected, all 12 graphs will be shown. The dashed reference line indicates a 1 mV signal amplitude.



NOTE: The timeout to the Monitoring Screen is disabled while the ECG Screen is displayed.





Figure 11-26: ECG Screen - Normal Mode

The only user interaction on the ECG Screen is the Pause / Resume button. Using the Jog Wheel, the user navigates to the button and presses the Wheel to toggle between Pause / Resume. If the selected button reads PAUSE when the Jog Wheel is pressed, then the ECG graphs will be frozen and can be examined more easily. Press the Jog Wheel when the selected button reads RESUME to restart normal graphing of the ECG waveform.



Figure 11-27: ECG Screen - Graphs Paused



11.9 ALARM LIMITS SCREEN

11.9.1 Overview

The Alarm Limits Screen allows for control of alarm limit values. When the Jog Wheel is rotated clockwise, the selection order for the buttons on the Alarm Limits screen is the following: Screen Button, Alarm View Button, Limit List (limits are selected from top to bottom with no wrap). When the Jog Wheel is rotated counterclockwise, the order is reversed.

A list of limits is displayed and the threshold name and value, along with the units of each limit, is shown. The list is scrollable once the Limit List part of the screen is activated.

If no limits have been changed from their default values, the phrase "All limits at default" is shown at right below the Status Bar. If any limits have been changed from their default values, an icon and a summary of the number of limits not at their default value is displayed (see *Figure 11-29: Alarm Limits Screen Active on page 173*).

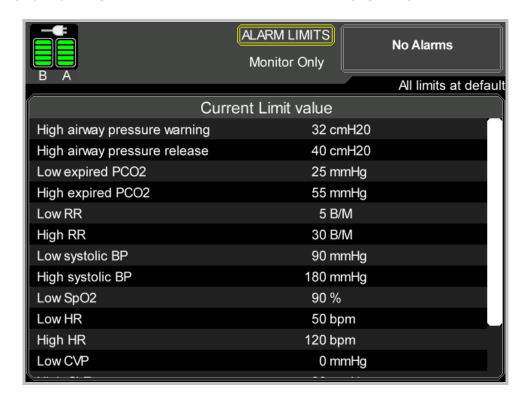


Figure 11-28: Alarm Limits Screen - All at Default



WARNING! WHEN SETTING ALARM LIMITS, DO NOT SET TO EXTREME VALUES THAT CAN RENDER THE ALARM SYSTEM USELESS.

11.9.2 Limit List Active

If the Limit List is active, a single limit is selected, as indicated by a yellow box around the limit (see Figure following). To modify the limit, the user presses the Jog Wheel to make the limit active and then uses the Wheel to select a value. The Jog Wheel is pressed once again to confirm the new value. The Back Button is used to abandon an actively selected limit and its new value and return to the previously selected limit value.



Limit values that have been changed from their default are shown with an asterisk. Also, an icon in the status bar indicates that limit values have been changed and accompanying text lists how many. As well, a message at the right below the status bar show the number of limits not at their default values.



NOTE: For more information on the number of limits changed icon, see <u>Table 23: MOVES® SLCTM Status</u> <u>Bar Items and Descriptions on page 139</u>.

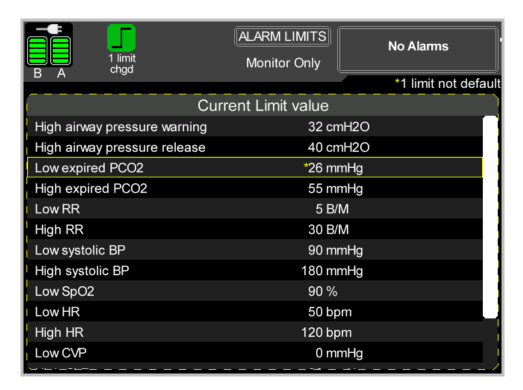


Figure 11-29: Alarm Limits Screen Active

The following table lists all the limits, their available values, and their defaults.

Table 29: Alarm Limits and Defaults

	Limits	Description
1.	High airway pressure warning	20 to 58 cmH ₂ O in steps of 2, default to 32
2.	High airway pressure release	40 to 58 cmH ₂ O in steps of 2, default to 40 NOTE: In addition to this being an alarm threshold, the ventilator uses this value as the pressure safety threshold at which the ventilator stops inspiration and switches to expiration.
3.	Low expired PCO ₂	15 to 35 mmHg in steps of 1, default to 25
4.	High expired PCO₂	50 to 60 mmHg in steps of 1, default to 55
5.	Low RR	5 to 15 B/M in steps of 1, default to 5
6.	High RR	20 to 42 B/M in steps of 1, default to 30



	Limits	Description
7.	Low systolic BP	70 to 140 mmHg in steps of 1, default to 90
8.	High systolic BP	140 to 200 mmHg in steps of 1, default to 180
9.	Low SpO ₂	85 to 95% in steps of 1, default to 90
10.	Low HR	30 to 150 BPM in steps of 1, default to 50
11.	High HR	100 to 250 BPM in steps of 1, default to 120
12.	Low CVP	-5 to 5 mmHg in steps of 1, default to 0
13.	High CVP	10 to 30 mmHg in steps of 1, default to 20
14.	Low ICP	-7 to 7 cmH ₂ O in steps of 1, default to 0
15.	High ICP	13 to 40 mmHg in steps of 1, default to 27
16.	Low PVI	1 to 98% in steps of 1, default to 5
17.	High PVI	2 to 99% in steps of 1, default to 40
18.	High SpMet	 1.0 to 2.0% in steps of 0.1 2.0 to 99.5% in steps of 0.5 Default to 3.0
19.	High SpCO	2 to 98% in steps of 1, default to 10
20.	Low SpHb	Alarm limit will be displayed in the SpHb units the system is configured to display, one of the following options:
		g / dL (grams hemoglobin / deciliter blood) 1.0 to 23.5 g/dL in steps of 0.1, default to 8.0
		mmol / L (millimoles hemoglobin / liter blood) 1.0 to 14.5 mmol/L in steps of 0.1, default to 5.0
		g / L (grams hemoglobin / liter blood) 10 to 235 g/L in steps of 1, default to 80
21.	High SpHb	Alarm limit will be displayed in the SpHb units the system is configured to display, one of the following options:
		g / dL (grams hemoglobin / deciliter blood) 2.0 to 24.5 g/dL in steps of 0.1, default to 17.0
		mmol / L (millimoles hemoglobin / liter blood) 2.0 to 15.0 mmol/L in steps of 0.1, default to 11.0
		g / L (grams hemoglobin / liter blood) 20 to 245 g/L in steps of 1, default to 170
22.	Low SpOC	1 to 33 mL O ₂ / dL blood in steps of 1, default to 10
23.	High SpOC	2 to 34 mL O ₂ / dL blood in steps of 1, default to 25



11.10 ALARM ON / OFF SCREEN

11.10.1 Overview

The Alarm On / Off screen allows for alarm On / Off control. When the Jog Wheel is rotated clockwise, the selection order for Alarm ON/OFF screen buttons is Screen Button, Alarm View Button, Alarm list (the Alarm list is selected from top to bottom with no wrap). When the Jog Wheel is rotated counterclockwise, the selection order is reversed.

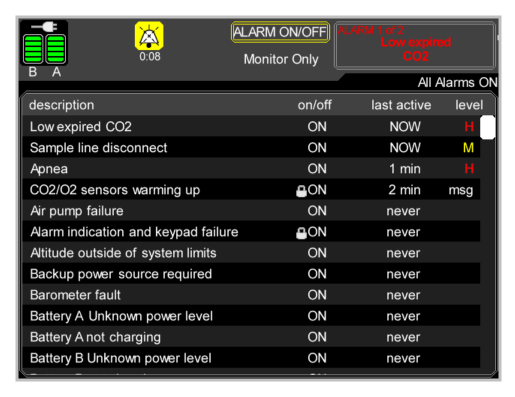


Figure 11-30: Alarm On / Off Screen

A summary of the alarms turned off is shown at top right as follows: All Alarms ON, 1 Alarm OFF, 2 Alarms OFF, etc.

The screen below presents a list of all alarms that can be turned off. The list is sorted when the user enters the screen according to the following:

- First according to last active, with alarms that are active now at the top.
- 2. Alphabetically according to the alarm message.

Alarm level is indicated to the right. A red "H" indicates a HIGH priority alarm; a yellow "M" indicates a MEDIUM priority alarm; a yellow "L" indicates a LOW priority alarm. The letters "msg" denote a "MESSAGE", not an alarm.

The last active column indicates the current state of the alarm. The text "NOW" indicates the alarm is currently active (even if it is turned off). The text "never" indicates the alarm has never been active since the device was turned on. Otherwise the time since the alarm was last active is given (first in seconds, then minutes and finally hours). In this case the priority level listed is the last active priority.



NOTE: Alarms and messages shown with a padlock beside them in the alarm list CANNOT be turned off or altered in any way, even by someone with Administrator privileges.



11.10.2 Alarm On/Off List Active

If the Alarm list is activated, one alarm in the list is selected. This is indicated by a yellow selection box. Turning the Jog Wheel clockwise or counter clockwise selects the next or previous item in the list, scrolling the list as needed. Note that there is no wrap around.

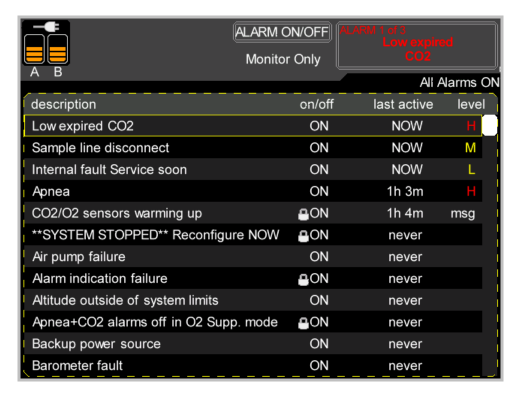


Figure 11-31: Alarm On / Off Screen Active

Pressing the Jog Wheel toggles the ON/OFF state of the selected alarm and updates the alarm summary. Pressing the Back Button exits the list active state.

12.0 Using the Remote Screen

MOVES[®] SLC[™] can also be used with the remote screen (P/N 126581) shown below. The remote screen is an optional accessory.



Figure 12-1: MOVES[®] SLC™ Remote Screen

12.1 OVERVIEW

The remote screen interface displays the same program screens (except for the System Test and WiFi screens) as the local user interface of $MOVES^{@}$ SLC^{TM} .

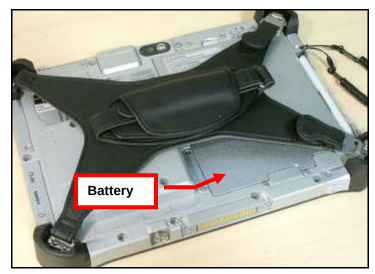
The remote screen offers the convenience of a secondary, and portable, monitor with a larger screen. It provides the user with the ability to remotely control the settings of MOVES[®] SLC[™]. There are two options for connection – wired and wireless (WiFi).



12.2 REPLACING / INSTALLING THE REMOTE SCREEN BATTERY

The MOVES[®] SLC[™] remote screen can be run on battery or AC power. The remote screen features a 10-hour battery life as well as a bridge battery configuration which makes batteries hot swappable. The second battery allows the user to change batteries without having to power off the device.

1. Turn over the remote screen. The X-strap on the back, shown in the photo on the right, must be removed or shunted to the left to access the battery.



2. To shunt the X-strap to the left, first undo the Velcro fasteners attaching the top and bottom right corners. Then pull the two straps to the left to loosen the entire X-strap.



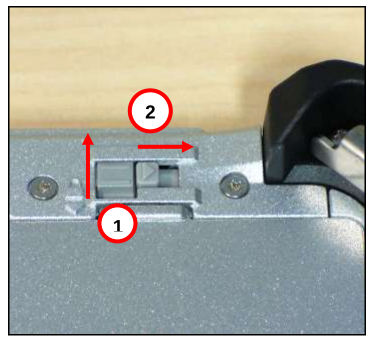
3. Undo the two hooks attaching the straps. Then shunt the X-strap to the left.



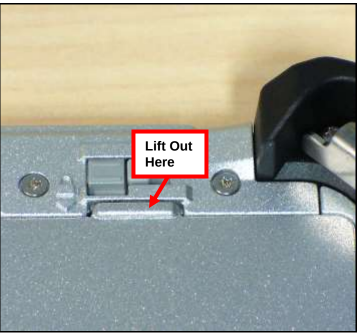
4. An inserted battery is shown in the photo at right.



5. To release and remove the battery, push the Lock button (#1) up to unlock the battery. Then push the Release button (#2) to the right to release the battery.



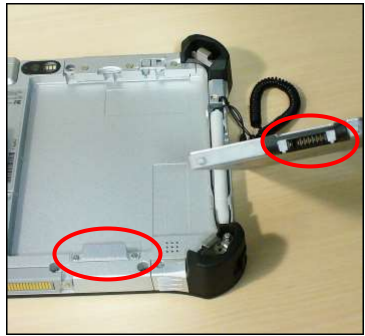
6. Lift out the battery by the handle indicated in the photo at the right.



7. The battery is shown removed (or uninstalled) in the photo at the right.



8. Do not touch the terminals of the battery pack or the computer. Doing so can make the terminals dirty or damage them which may cause the battery pack or the remote screen to malfunction.



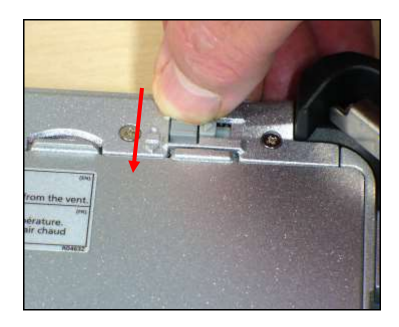
9. To install a battery, first align it as shown in the photo at the right.



10. Then press it down until it snaps into place.



11. Lastly, push the Lock button in the direction of the arrow to lock the battery in place.



12. Reinstall the X-strap (See Steps 3 - 1).

12.3 CONNECTING THE REMOTE SCREEN TO WALL POWER

 The cord shown at right, which includes an AC to DC adaptor, is used to connect wall power to the remote screen.



2. The DC connection for the remote screen is located under a plastic cap in the bottom left corner .



3. The plastic cap covering the DC connection is shown in the photo at right. To access the connection, lift the cap from the bottom.



4. The DC connection is shown in the photo at the right.



5. Power cord connected.

NOTE: The Installed battery is automatically recharged if it is below a full charge when wall power is connected to the remote screen.



12.4 REMOTE SCREEN INDICATORS

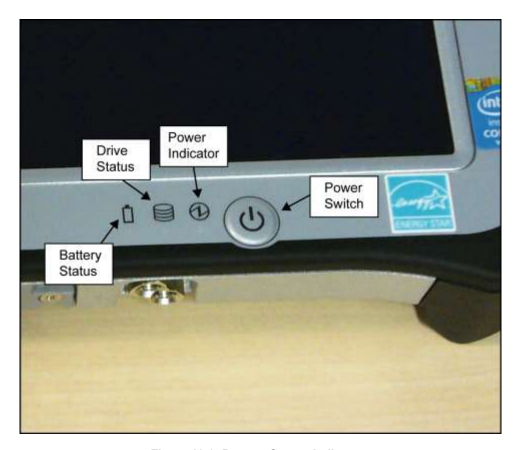


Figure 12-2: Remote Screen Indicators



Table 30: Remote Screen Indicators

Indicator	Function / Properties
Power Switch	To start, press and hold the power switch until the power indicator lights. To turn off the remote screen, press and hold the power switch for four (4) seconds.
Power Indicator	Off: Power off/Hibernation, Green: Power on, Blinking green: Sleep
Drive Status	Do not perform the following operations until the drive indicator turns off:
Battery Status	The battery indicator may not light even if the AC adaptor and the battery pack are correctly connected. The reason for this is that the AC adaptor's protection function may be working. If this is the case, pull out the AC cord and wait for more than one (1) minute before reconnecting the AC cord.

12.5 REMOTE SCREEN BATTERY STATUS

Battery Indicator	Battery Status
Not lit	The battery pack is not inserted or not being charged.
Orange	Charging is in progress.
Green	The battery is fully charged.
Blinking Green	When you close the battery cover with the battery pack already inserted, you can check the battery level by blinking times. Blinking times to Battery Level: • 5 times = 95 % to 100 %
	• 4 times = 50 % to 94%
	• 3 times = 25 % to 49%
	• 2 times = 5 % to 24%
	• 1 time = 0 % to 4 %
Red	The remaining battery is approximately 9% or less.
Blinking Red	When blinking approximately every 1 second:
	The battery pack or the charging circuit is not operating properly.
	When blinking approximately every 4 seconds:
	The battery cover is open. You can remove the battery pack in this case.
	When blinking approximately every 0.5 second:
	The battery cover is open. If you remove the battery pack in this case, the power supply is cut off and the computer is shut down. Close the battery cover immediately.



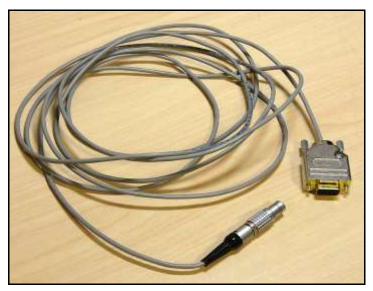
Battery Indicator	Battery Status	
Blinking Orange	The battery cannot be charged temporarily due to the following reasons:	
	 Its internal temperature is out of the acceptable range. 	
	 The power supply is not enough because software applications or peripheral devices are consuming a large amount of power. 	
Blinking Green and Orange Alternately	The temperature is low and the computer is warming up to prevent the hard disk drive or flash memory from malfunctioning. The computer will start automatically after warming-up.	

12.6 WIRED CONNECTION OF THE REMOTE SCREEN TO MOVES® SLC™

Two cables are required to make a wired connection of the remote screen to the auxiliary communications port on MOVES[®] SLC™: a short black cable that connects to the serial port on the remote screen and a long gray cable with a circular end that connects to the auxiliary communication port on MOVES[®] SLC™. The two cables connect to each other via two matching serial port connectors. The cables are shown below.

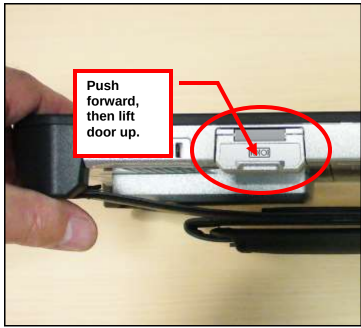


Remote Screen Serial Port Connection Cable



MOVES[®] SLC[™] Auxiliary Communication Port Connection Cable

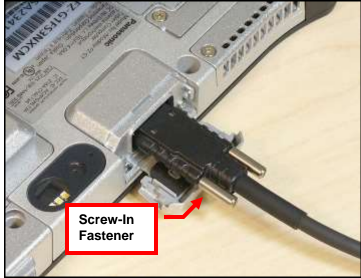
 Locate the serial port connection on the top side of the remote screen. The door indicated in the photo at the right is covering the serial port connection



2. Lift up the door to access the serial port connection. The connection shown at right.



3. Attach the small end of the short black cable to the remote screen by pushing it in and tightening the screw-in fasteners on either side. Make sure the cable is properly aligned. Do not force it. If the remote screen is lying face down, as in the photo, the wider side of the black cable connector should be facing up.



 Attach the large, pin-end of the black cable to the receiving serial port connection on the MOVES[®] SLC™ cable by pushing the two together.





5. Now connect the circular end of the MOVES[®] SLC[™] cable shown at the right to the auxiliary communication port on MOVES[®] SLC[™].



6. The auxiliary communication port on MOVES[®] SLC™ is shown in the photo at the right. To access the connection lift the rubber cover.



7. The auxiliary communication port connection is shown in the photo at the right.



8. Align the two slot keys on the remote screen cable connector with the two receiving channels in the auxiliary communication port connection and insert.



 The remote screen cable connector is shown connected to the auxiliary communication port on MOVES[®] SLC™.



12.6.1 Activating the Wired Connection

A typical connection scenario would be once the remote screen is physically connected to the auxiliary communication port on $MOVES^{@}$ SLC^{TM} , turn $MOVES^{@}$ SLC^{TM} on. Then turn on the remote screen. As soon as the remote screen is activated, it interfaces automatically with $MOVES^{@}$ SLC^{TM} and begins sharing information. No software program needs to be launched by the user.



NOTE: The Remote Screen, however, can be connected at anytime, and if MOVES[®] SLC™ is running, and the Remote Screen is running, then the communication link will start automatically.

12.7 WIRELESS CONNECTION OF THE REMOTE SCREEN TO MOVES® SLC™

Before MOVES[®] SLC[™] can be connected wirelessly to the remote screen, the remote screen must be enabled to accept a wireless connection (Wi-Fi).

12.7.1 Enabling the Remote Screen to Accept a Wireless Connection

The Administrator is the person responsible for enabling the remote screen to accept a wireless connection (Wi-Fi). The Administrator also configures the Wi-Fi security settings. The Administrator can also disable wireless connection. If the Administrator has disabled wireless connection, then the Wi-Fi control switch will be locked in the OFF position, and only a wired connection will be able to be made to the remote screen. Also, the words Wi-Fi and OFF will be grayed out on the switch.



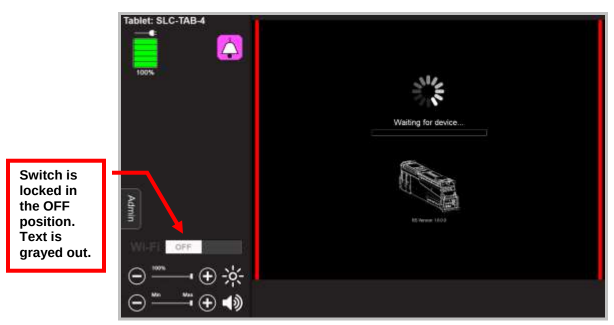


Figure 12 3: Wi-Fi Disabled on the Remote Screen

If the Administrator has enabled wireless connection on the remote screen, then the operator can control the use of a wireless connection using the on-screen Wi-Fi control switch.

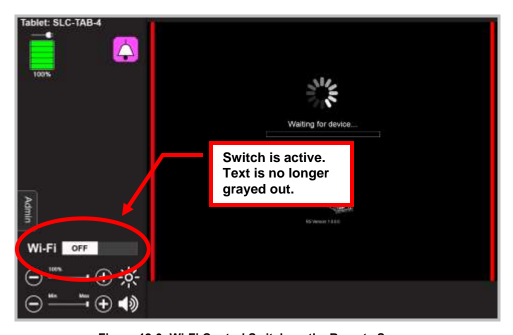


Figure 12-3: Wi-Fi Control Switch on the Remote Screen

Tapping or clicking the switch will toggle Wi-Fi OFF and ON. When the Wi-Fi Control Switch is set to ON then a wireless connection can be made with the remote screen.



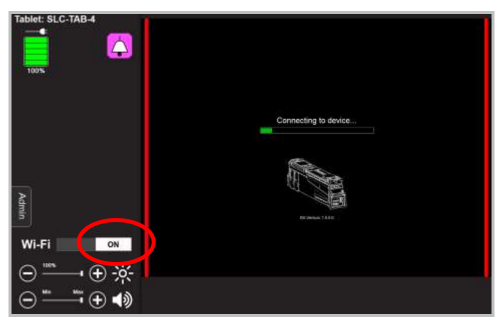


Figure 12-4: Wi-Fi Control Switch Set to ON

12.7.2 Connecting to the Remote Screen Wirelessly from MOVES[®] SLC™

Using the Jog Wheel, navigate to the options area of the Setup Screen, select WIFI... and press the Jog Wheel.

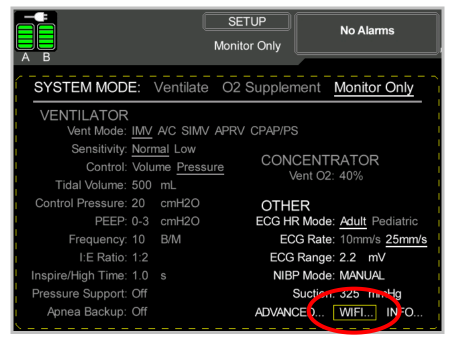


Figure 12-5: Accessing the WIFI Screen

The WIFI Screen opens with the name of the most recently connected tablet in the Tablet field. On the following screen this is JFL127. With the Tablet field selected, press the Jog Wheel to proceed to the WIFI TABLET Selection Screen.



Figure 12-6: The WIFI Connection Screen

A list of the available tablets that you can connect to is listed in order of signal strength with the first tablet selected. Signal strength is indicated by the amount of blue shown in the wireless icon (versus gray which indicates weakness).





Figure 12-7: WIFI Signal Strength Icons

To connect to a tablet, use the Jog Wheel to select it from the list (if it is not already selected). Then press the Jog Wheel to confirm the selection and return to the WIFI Screen.



Figure 12-8: Tablets Available for Connection

When you return to the WIFI Screen after choosing a tablet, the CONNECT button will already be highlighted. Press the Jog Wheel to connect.

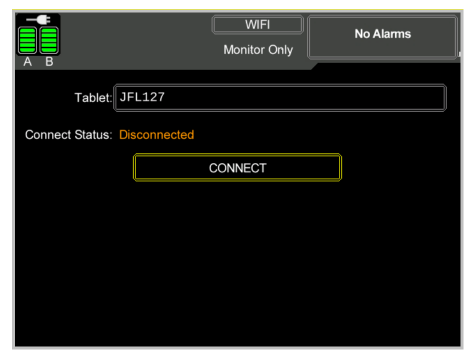


Figure 12-9: Ready to Connect

As you are connecting, the wireless symbol appears with gray bars and a pulsating dot that alternates between gray and colored (green if monitoring only, blue if the remote screen also has remote control capabilities).

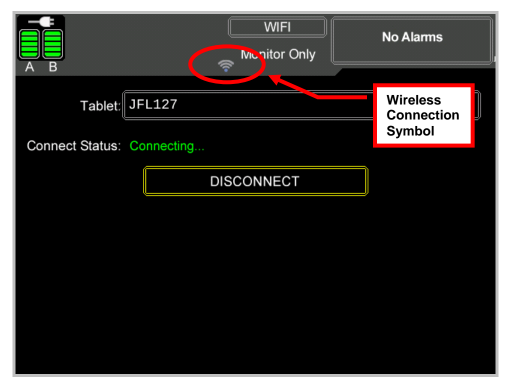


Figure 12-10: MOVES® SLC™ Connecting to Remote Screen

The wireless connection can be terminated at any time by pressing the DISCONNECT button while connecting or when connected. When a connection is initially being establishing between the MOVES[®] SLC[™] and the Remote Screen, the Connect Status will indicate "Connecting…" and will attempt to connect for 45 seconds. If a connection has been established and then lost, the MOVES[®] SLC[™] will try to reconnect for up to five (5) minutes, and the Connect Status will indicate "Reconnecting…". When a connection is made, the Connect Status will indicate the Tablet to which the MOVES[®] SLC[™] is connected (e.g., "Connected to: SLCRS-001").

12.8 THE REMOTE SCREEN USER INTERFACE (UI)

12.8.1 First Connecting

At first, the user will see the Initial screen. The information on the Initial screen may vary. What the Initial screen displays depends on whether the remote screen is starting from a total disconnect or from the hibernate/sleep state (i.e., whether software resources are loaded already or not). It also depends on whether the remote screen is connecting to MOVES[®] SLC™.

If the remote screen is starting from a total disconnect, and is NOT connecting to MOVES[®] SLC™, the user will see the information displayed in the screen capture that follows. A blue progress bar is shown while the program is loading. A "spinning dial" and the message, "Waiting for device…" indicate the screen is NOT connecting with MOVES[®] SLC™.



Figure 12-11: Initial Screen - Resources Loading

If the remote screen is starting from the hibernate/sleep state (or the software resources have finished loading), and the remote screen IS connecting to $MOVES^{\circledR}$ $SLC^{\intercal M}$, the user will see the information displayed in the following screen shot. A green progress bar is shown while the screen is connecting to $MOVES^{\circledR}$ $SLC^{\intercal M}$.



NOTE: The Dashboard now becomes visible on the left side of the screen once the resources have loaded.

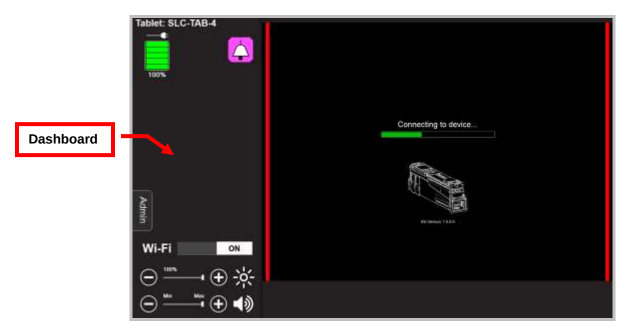


Figure 12-12: Main Remote Screen - Connecting and Resources Loaded



NOTE: In this second instance, sometimes the connection is so quick that the Initial screen is barely visible.

12.8.2 The Dashboard

The Dashboard contains the following:

- Battery charge level & wall power connection indicator
- Connection state icon
- Alarm audio pause or disable option
- Admin (Administrator) tab
- Wi-Fi On/Off switch
- Screen brightness slider
- Alarm volume slider

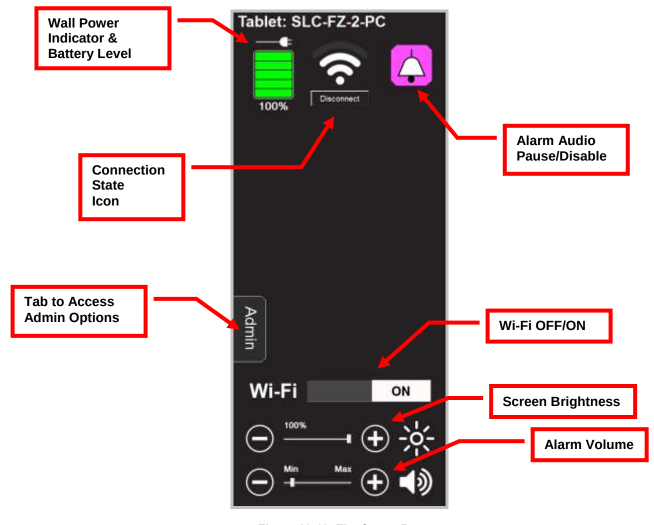


Figure 12-13: The Status Bar



BATTERY CHARGE LEVEL & WALL POWER CONNECTION INDICATOR

A wall plug icon is shown if the tablet is connected to wall power. When the tablet is running on AC, and the battery is charging, then the percent the battery is charged is displayed. If the tablet is running on battery power, then the estimated time remaining in the battery is displayed. In the illustration above, the tablet has a fully charged battery and is running on wall power.

CONNECTION STATE ICON

Indicates whether the tablet has established a connection with MOVES[®] SLC™. Clicking or tapping the Disconnect button breaks an established connection.

ALARM AUDIO PAUSE OR DISABLE OPTION

Click or tap the on-screen Alarm icon. A screen popup lets you choose to pause the alarm audio (for two [2] minutes) or disable the alarm audio.



Figure 12-14: Remote Screen Alarm Audio Options



NOTE: Disabling the alarm audio must be allowed by the Administrator. If it is not allowed, it will not appear as an option.

ADMIN (ADMINISTRATOR) TAB

The Admin tab is <u>not to be used</u> by the operator. The Administrator controls the following options:

- Allow/disallow alarm disable
- Minimum alarm volume
- Enable/disable Wi-Fi

If the Administrator has set the minimum alarm volume to maximum volume, then the alarm volume control is locked. This is shown by the lock icon and the word "Max" in the screen capture below.





Figure 12-15: Alarm Volume Locked at Maximum

WI-FI ON/OFF SWITCH

The WiFi control functions as a toggle switch. When it says ON, then WiFi is on; when it says OFF, WiFi off.

SCREEN BRIGHTNESS SLIDER & ALARM VOLUME SLIDER

Dragging the sliders or clicking the plus/minus icons will change the setting. Tapping the plus/minus icons will also change the setting.

12.8.3 No System Test Screen

The remote screen does not display the System Test screen. If tests have not been completed prior to connecting the remote screen, the remote screen will open on the Setup screen and display the status "Initializing". The Suction and NIBP icons will be shown at the top of the screen with the "No" symbol (circle with a diagonal line) indicating that they are not available. These two functions are not allowed until the startup tests have been passed (see Figure 12-16: Setup Screen – Initializing which follows). While the system is initializing (i.e., System Tests are not yet complete), the System Mode cannot be modified on the Setup screen; however, the settings on the Setup screen can be preconfigured in preparation for use after the System Tests are complete.



NOTE: The following screens are shown without the Dashboard for the sake of clarity.



Figure 12-16: Setup Screen - Initializing



Once the startup tests have been passed, the Setup screen SYSTEM MODE setting will unlock in Monitor Only mode and the not-available Suction and NIBP icons will disappear from the top of the screen.



Figure 12-17: Setup Screen - Monitor Only Mode

Just as on the display that is physically apart of MOVES[®] SLC™, the Setup screen on the remote screen "times out" or defaults to the Main screen after <u>one minute</u> of inactivity so that patient monitoring can be maintained. The only exception to this is the ECG screen.



Figure 12-18: Main Screen - On Remote Screen

12.8.4 Remote Screen Panel Buttons

NAVIGATING AND SELECTING VIA PANEL BUTTONS

Because the remote screen does not use a jog wheel, navigation is controlled by the use of buttons, shown below at the right. Only two of the buttons from the MOVES[®] SLC™ keypad are replicated on the remote screen – the NIBP button and the Suction button – which are shown at the left. They work in exactly the same way as they do on the physical keypad.



Figure 12-19: Remote Screen - Panel Buttons at Bottom

The remote screen is a touch-sensitive screen, and the user can navigate through the displayed user interface and select options with the touch-sensitive on-screen panel buttons. The way the buttons function is explained in the table that follows.



Table 31: Buttons on the Remote Screen

Buttons on the Remote Screen		
	Screen Button Press the Screen button to move to the next screen in sequence. If there are outstanding queries or parameters that need to be satisfied, the functionality of the Screen button will be inhibited until these queries or parameters are satisfied.	
	NOTE: This button also has quick navigation functionality. Pressing and holding the button will take you from any screen to the Monitoring screen. If you are already on the Monitoring screen, pressing and holding the button will take you to the Setup screen.	
83	Cancel Button Press this button to reject the current value that is being changed and return to the previous value. This button is also used to back out of an activated button (e.g., Alarm Queue button).	
	Check Button Press the Check button to start or end editing a currently selected item, or change the currently selected item. The Check button is also used to acknowledge messages.	
*	Next Button Selects the next item in a group or increases a numeric value.	
	Previous Button Selects the previous item in a group or decreases a numeric value.	

TOUCH NAVIGATING

Rather than navigating step-by-step through the screens, choices, and options as the user must do with the physical buttons on MOVES[®] SLC™, and still can do with the on-screen buttons when using the remote screen, the user can also directly select options with a finger press. Then, however, the Next and Previous arrow buttons and the Check button must be used to change options and confirm choices.

12.8.5 Graphs Independently Configurable

Although most of the functionality on the remote screen is linked and analogous to that on the display screen physically attached to MOVES[®] SLC™, the graphs are an exception. The graphs shown on the remote screen are NOT linked. Changing the graphs on the remote screen will not change the graphs on the display screen. This has been done to allow the user the option of displaying twice as much graph monitoring information.

12.8.6 Alarm Indicators

On the MOVES[®] SLCTM unit, there are four (4) System Visual Indicators, one at each top corner. These indicators display the alarm system status. On the remote screen, this functionality has been duplicated by Alarm Status Bars on either side of the screen (see Figures following).



The severity of the <u>highest priority active alarm</u> is shown. If the highest priority active alarm is turned off, the bars display the severity of the next highest priority active alarm which is turned on. The color-coding indicates the same alarm states.

Table 32: Alarm Side Bar States and Explanations

Status	Activity	Indicates
		System off or in Initializing mode (i.e., System Tests incomplete)
Green bars Solid green (no activity) No alarms active		No alarms active
Yellow bars	Solid (Low priority alarm) Flashing (Medium priority alarm) Low or Medium Priority Alarm(s	
Red bars Flashing (always) High Priority Ala		High Priority Alarm(s) active



Figure 12-20: Green Bars Indicating No Alarms Active





Figure 12-21: Red Bars Indicating High-Priority Alarm Active



Figure 12-22: Yellow Bars Indicating Low or Medium Priority Alarm Active

12.8.7 Remote Screen Software Version

The remote screen version can be determined by accessing the Info Screen. For information on accessing the Info Screen, see Section 11.6.1 Accessing the Info Screen on page 156.



Figure 12-23: Info Screen Showing Remote Screen Version



12.9 FEATURES DISABLED ON THE REMOTE SCREEN

Some of the features available on the UI screen that is physically a part of MOVES[®] SLC[™] may be disabled on a remote screen that is connected wirelessly. If this is the case, it has been done by the Administrator. If features are disabled, the Administrator has decided that wireless control of these feature(s) is unsafe in the current medical environment.

The following feature areas of the remote screen can be disabled by the administrator:

- Wireless Communication (no wireless remote screen connection allowed)
- Alarms (no changes to Alarm settings via the remote screen, includes dismissing alarms in alarm queue, changing alarm limits, and turning alarms on or off)
- Patient Monitor Settings (no changes to ECG/NIBP settings on Setup screen or any of the settings on the Advanced screen via the remote screen)
- Invasive Pressure Settings (no changes to IP channel mode [i.e., ABP, ICP or CVP] and no channel zeroing via the remote screen)
- Heart Rate Source (no selecting of HR source via the remote screen)
- Ventilator Settings (no changes to Ventilator settings via the remote screen)
- Suction (no changes to Suction pressure and no control of Suction on/off via remote screen)
- System Mode (no changes to System Mode [i.e., Ventilate, O2 Supplement, Monitor Only] via remote screen)

On the following Alarm Limits screen, access to Alarm settings has been disabled. This is shown by the padlock to the left of each item. As well, Suction has been disabled as can be seen by the Suction button at the bottom left being grayed-out and stroked-out.



Figure 12-24: Alarm Limits and Suction Disabled on the Remote Screen

On the following screen, the Ventilator and Patient Monitor settings have been disabled. This is shown by the padlock to the left of each item. As well, the NIBP and the Suction buttons have been disabled, as can be seen by their being grayed-out and stroked-out, due to Patient Monitor and Suction settings being disabled respectively.



Figure 12-25: Ventilator Settings and NIBP and Suction Disabled on the Remote Screen

On the following screen, the Heart Rate Source option (HR) is disabled. The double white lines that surround an option that is active (such as ABP) are missing from its display.



Figure 12-26: Heart Rate Source Option Disabled on the Remote Screen

On the following screen, the Invasive Pressure settings have been disabled. The double white lines that surround an option that is active (such as HR) are missing from the three IP channels set to ABP, CVP and ICP.



Figure 12-27: Invasive Pressure Settings Disabled on the Remote Screen

On the following screen, the System Mode option has been disabled. This is indicated by the small padlock beside the words SYSTEM MODE on screen. When the System Mode option has been disabled, the remote screen will only display in current System Mode.



Figure 12-28: SYSTEM MODE Disabled on the Remote Screen

13.0 Connecting the Patient

13.1 CONNECTION OVERVIEW

The patient may be intubated, in which case the patient will be connected to a ventilator breathing circuit. The patient may also be breathing spontaneously but be receiving supplemental oxygen via a nasal cannula (nasal prongs) or oxygen mask.



WARNING! DO NOT CONNECT A PATIENT TO MOVES[®] SLC™ UNTIL THE MOVES[®] SLC™ SYSTEM IS PROPERLY WARMED UP AND O₂ VALUES ARE DISPLAYED.



WARNING! BEFORE VENTILATING A PATIENT, ENSURE THAT A SPARE VENTILATOR CARTRIDGE AND BREATHING CIRCUIT ARE READILY AVAILABLE.

13.2 CONNECTING AN INTUBATED PATIENT

- Ensure that the ventilator cartridge and breathing circuit have been connected to MOVES[®] SLC™.
- 2. Ensure startup test procedures have been completed and all tests passed.
- 3. Configure MOVES[®] SLC[™] to operate in Ventilate mode.
- 4. Attach the elbow to the patient's endotracheal tube.



WARNING! WHEN VENTILATING PATIENTS UNDER 30 KG OR WITH TIDAL VOLUMES UNDER 150 ML, REPLACE THE AIRWAY FILTER WITH PEDIATRIC BREATHING SYSTEM FILTER (P/N 125245) TO REDUCE DEAD SPACE VENTILATION.

13.3 CONNECTING A SPONTANEOUSLY BREATHING PATIENT

- 1. Ensure startup test procedures have been completed and all tests passed.
- 2. Attach the O₂ sampling adaptor to the MOVES[®] SLC[™] O₂ Outlet port.
- 3. Attach the sampling line filters and tubing between the O₂ sampling adaptor and the Gas Sample port in the Patient Connections panel.
- Attach one end of the O₂ tubing to the O₂ sampling adaptor.
- Attach a fire suppression device (such as the BPR Firesafe™ Cannula Valve) to the other end of the O₂ tubing.
- 6. Attach the nasal cannula or mask tubing to the fire suppression device. (NOTE: Place the fire suppression device as close to the patient as possible.)
- 7. Configure MOVES[®] SLC[™] to operate in O2 Supplement mode.
- 8. Attach the nasal cannula or mask to the patient.



NOTE: For full procedures with accompanying photographs see <u>Delivering Supplementary Oxygen (O2)</u> <u>beginning on page 102</u>.



13.4 ATTACHING THE EAR OR FINGER CLIP PULSE CO-OXIMETER SENSOR



NOTE: It is recommended that a pulse oximeter is always used to ensure adequate patient oxygenation.

The $MOVES^{\circledR}$ SLCTM system comes with some or all of the pulse CO-oximeter sensors listed in the following table. Sensor position and whether the sensor is reusable or disposable is also indicated.

Table 33: CO-Oximeter Sensors

Se	nsor	Position	Reusable / Disposable
1.	Adult SpO ₂ (only) Finger Clip (M-LNCS DCI)	Any of the five (5) fingers on either hand (but not the toes)	Reusable
2.	Pediatric SpO2 (only) Finger Clip (M-LNCS DCIP)	Any of the five (5) fingers on either hand (but not the toes)	Reusable
3.	SpO ₂ Ear Clip (M-LNCS TC-I)	Either ear lobe NOTE: Ear lobe must NOT be pierced.	Reusable
4.	Adult SpO2 Disposable Ear Sensor (M-LNCS E1)	Either ear's Cavum Conchae	Disposable (i.e., single use)
5.	Adult SpO ₂ Adhesive Sensor (M-LNCS Adtx-3)	Any of the five (5) fingers on either hand (but not the toes)	Disposable (i.e., single use)
6.	Pediatric SpO ₂ Adhesive Sensor (M-LNCS Pdtx-3)	Any of the five (5) fingers on either hand (but not the toes)	Disposable (i.e., single use)
7.	Adult SpO ₂ , SpCO, & SpMet Finger Clip (Rainbow [®] DCI-dc3)	Any of the five (5) fingers on either hand (but not the toes)	Reusable
8.	Pediatric SpO ₂ , SpCO, & SpMet Finger Clip (Rainbow [®] DCIP-dc3)	Any of the five (5) fingers on either hand (but not the toes)	Reusable
9.	Adult SpO ₂ , SpHb and SpMet Adhesive Sensor (Rainbow [®] R1 25)	Any of the five (5) fingers on either hand (but not the toes)	Disposable (i.e., single use)
10.	Pediatric SpO ₂ , SpHb and SpMet Adhesive Sensor (Rainbow [®] R1 20)	Any of the five (5) fingers on either hand (but not the toes)	Disposable (i.e., single use)
11.	Adult SpO ₂ , SpCO and SpMet Adhesive Sensor (Rainbow [®] R25)	Any of the five (5) fingers on either hand (but not the toes)	Disposable (i.e., single use)
12.	Pediatric SpO ₂ , SpCO and SpMet Adhesive Sensor (Rainbow [®] R20)	Any of the five (5) fingers on either hand (but not the toes)	Disposable (i.e., single use)

The following photographs show correct placement of the finger clip and ear clip sensors.

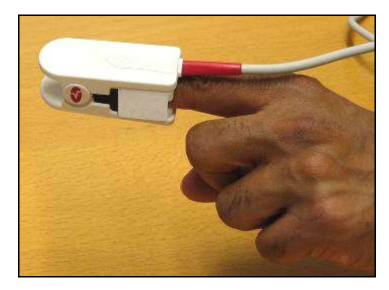


Figure 13-1: 1 of 5 Possible Sensor Finger Placements



Figure 13-2: Correct Ear Lobe Sensor Placement

AVOIDING INACCURATE READINGS

Pulse oximetry sensors work by transmitting red and infrared light through body tissue. Most light is absorbed by the tissue between the transmitting probe on one side and the actual sensor on the other. The small amount of light that is not absorbed is used to calculate oxygen saturation. Therefore, anything that disturbs the light flow can cause inaccurate readings. Note the following possible problems:

- Motion Artifact The most common cause of inaccurate SpO₂ readings is movement. Movement affects the ability of the light to travel from the light-emitting diode (LED) to the photo detector. Rhythmic movement such as tremors and seizure activity, as well as shivering, and vibrations caused by ground or air transport, can cause problems with detecting saturation and may measure false high pulse readings.
- **Ambient Light** Because pulse oximeters measure the amount of light transmitted through arterial blood, bright light that shines directly on the sensor whether from the sun or an overhead exam light can skew the readings. To fix this problem, move the sensor, or cover it with something opaque.
- **Light Absorbent Substances** Anything that absorbs light within the reading area may cause false-low readings (e.g., dried blood, polish, dyes [including intravenous]).

PULSE CO-OXIMETER SENSOR WARNINGS



WARNING! SPO₂ SENSOR APPLICATION SITES SHOULD BE INSPECTED AT LEAST EVERY FOUR (4) HOURS, OR AS DIRECTED IN THE SENSOR'S *DIRECTIONS FOR USE*, TO ENSURE CORRECT SENSOR ALIGNMENT AND SKIN INTEGRITY. IF CIRCULATION OR SKIN INTEGRITY IS COMPROMISED, THE SENSOR SHOULD BE APPLIED TO A DIFFERENT SITE. PATIENT SENSITIVITY MAY VARY DUE TO MEDICAL STATUS OR SKIN CONDITION. DISCONTINUE THE USE OF ADHESIVE TAPE STRIPS IF THE PATIENT EXHIBITS AN ALLERGIC REACTION TO THE ADHESIVE MATERIAL.



WARNING! OXIMETER READINGS MAY BE AFFECTED BY THE USE OF AN ELECTROSURGICAL UNIT (ESU).





WARNING! CIRCULATION DISTAL TO THE SENSOR SITE SHOULD BE CHECKED ROUTINELY.



WARNING! A FUNCTIONAL TESTER CANNOT BE USED TO ASSESS THE ACCURACY OF A PULSE OXIMETER MONITOR OR SENSOR.



WARNING! DO NOT USE DAMAGED SENSORS OR PATIENT CABLES. DO NOT USE A SENSOR OR PATIENT CABLE WITH EXPOSED OPTICAL OR ELECTRICAL COMPONENTS.

13.5 PLACEMENT OF THE HEART RATE ELECTRODES



WARNING! CONDUCTIVE PARTS OF ELECTRODES AND ASSOCIATED CONNECTORS FOR THE APPLIED PART, INCLUDING THE NEUTRAL ELECTRODE, SHOULD NOT CONTACT OTHER CONDUCTIVE PARTS AND EARTH.



CAUTION! ECG CABLES SHOULD BE DISCARDED AND REPLACED AFTER TWO (2) YEARS OF CONTINUOUS USE. CHECK CABLE USE BY RECORDING THE DATE THE CABLE WAS FIRST USED.

The snap-on ends of the MOVES[®] SLC™ ECG cable are color coded (see Figure following).

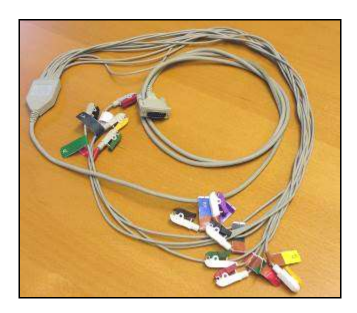


Figure 13-3: Color-Coded 12-Lead ECG Cable

The electrodes usually consist of a conducting gel, embedded in the middle of a self-adhesive pad onto which cables clip. It is important that the ECG electrodes to which the cable clips attach be placed properly to reduce motion artifact and receive the best signal possible.

13.5.1 ECG Cable Color Coding and Naming Conventions

The clamps of the ECG cables that are attached to the electrodes are color coded to ease connection for medical personnel. There are two definitions for these colors: AHA colors, used in the USA and Canada, and IEC colors used in all other countries (usually Europe). The following table explains the two systems.



Electrode	IEC Abbreviation	IEC Color	AHA Abbreviation	AHA Color
Right Arm	R	Red	RA	White
Left Arm	L	Yellow	LA	Black
Right Leg	N	Black	RL	Green
Left Leg	F	Green	LL	Red
Chest 1	C1	White-Red	V1	Brown-Red
Chest 2	C2	White-Yellow	V2	Brown-Yellow
Chest 3	C3	White-Green	V3	Brown-Green
Chest 4	C4	White-Brown	V4	Brown-Blue
Chest 5	C5	White-Black	V5	Brown-Orange
Chest 6	C6	White-Violet	V6	Brown-Violet

Table 34: ECG Cable Color Coding and Naming

13.5.2 Correct Electrode Placement

The following illustrations show correct electrode placement. The first illustration shows correct limb lead placement. The second illustration and accompanying table shows and explains the proper placement of the precordial leads V1–V6 (or C1–C6 in the IEC naming convention).

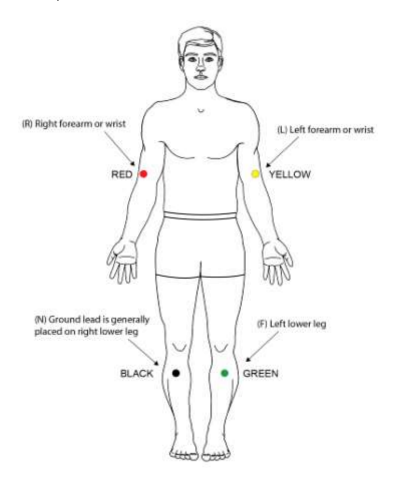


Figure 13-4: Proper Limb Lead Electrode Placement



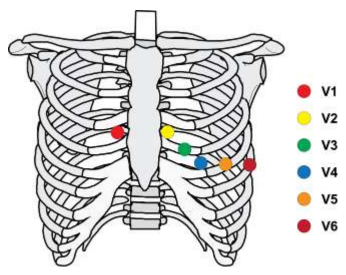


Figure 13-5: Precordial Leads: V1-V6

Table 35: Proper Placement of Precordial Leads

Electrode Identifier	Electrode Position on Body Surface
V1 / C1	Fourth intercostal space at right border of sternum
V2 / C2	Fourth intercostal space at left border of sternum
V3 / C3	Fifth rib between V2 / C2 and V4 / C4
V4 / C4	Fifth intercostal space on left mid clavicular line
V5 / C5	Left anterior axillary line at the horizontal level of V4 / C4
V6 / C6	Left midaxillary line at the horizontal level of V4 / C4

13.5.3 Reducing Artifacts

A common problem with 12-lead ECGs is that a lot of artifacts are generated if the patient is moving around and not fully relaxed. Here are a few guidelines on how to reduce artifacts and capture a good ECG:

- Place the patient in a supine or semi-Fowler's position. If the patient cannot tolerate being flat, you can do the ECG in a more upright position.
- Instruct the patient to place their arms down by their sides and to relax their shoulders.
- Make sure the patient's legs are uncrossed.
- Remove any electrical devices, such as cell phones, away from the patient as they may interfere with the machine.
- If you're getting artifacts in the limb leads, try having the patient sit on top of their hands.

13.5.4 Expiry Date of ECG Electrodes



CAUTION! CHECK THE EMBOSSED EXPIRY DATE ON THE ECG ELECTRODES PACKAGE BEFORE USING. ELECTRODES ARE GOOD FOR 45 DAYS ONCE PACKAGE IS OPENED.







Figure 13-6: Electrode Package

Figure 13-7: Electrode Use By Date

13.6 ZEROING THE PRESSURE IN THE IP TRANSDUCER

To zero the pressure in the invasive pressure transducer, turn the stopcock from the horizontal *running* position to the vertical *open-to-ambient-air* position.



Figure 13-8: Transducer Stopcock in Running Position (Horizontal)



Figure 13-9: Transducer Stopcock in Open-to-Ambient-Air Position (Vertical)

Zero Transducer

13.6.1 Zeroing the Transducer Channel

An IP port will auto detect the insertion of an IP probe. An inserted probe requires zeroing before data is available, and will show ZERO REQD until the onscreen channel config button -- now titled "ZERO" -- is clicked, zeroing the channel. The channel box will then immediately display data with format/units appropriate for the selected mode. The user can change modes without re-zeroing.



13.6.2 Transducer Warnings



EXTREME WARNING! WHEN USING FLUID FILLED PRESSURE TRANSDUCERS TO MONITOR INTRACRANIAL PRESSURE (ICP), MAKE SURE THAT THE TRANSDUCER AND THE LINE CONNECTING TO THE PATIENT'S DRAIN ARE FREE OF ANY AIR BUBBLES!



EXTREME WARNING! AFTER COMPLETING FILLING THE TRANSDUCER AND THE LINE, DISCONNECT THE FLUID BAG FROM THE TRANSDUCER, AND CAP THE END WITH THE STERILE CAP PRIOR TO CONNECTING THE TRANSDUCER TO THE PATIENT'S BRAIN!



EXTREME WARNING! NEVER FLUSH THE ICP TRANSDUCER WHILE CONNECTED TO THE PATIENT!



EXTREME WARNING! FAILURE TO OBSERVE THESE PRECAUTIONS MAY RESULT IN SERIOUS INJURY OR DEATH!

13.7 USING THE SUCTION FEATURE

To use the Suction feature:

- Ensure that the suction accessories are set up as shown in Section 9.10. Installing Suction Accessories beginning on page 109.
- 2. Activate the suction by pressing the Suction Control Button
- 3. Deactivate the suction at anytime by pressing the Suction Control Button a second time.



NOTE: The suction wand supplied with MOVES[®] SLC[™] has a control vent that must be occluded by the operator's thumb to allow suction to occur (See Figure following).

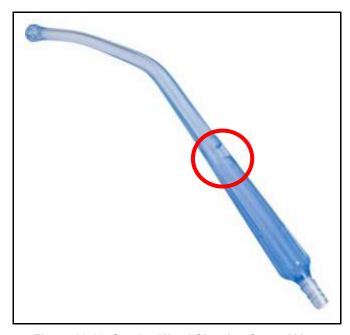


Figure 13-10: Suction Wand Showing Control Vent





NOTE: You can adjust suction pressure between 100 and 325 mmHg in increments of 25 mmHg. (For details, see the subsection <u>Changing Settings</u> in the section <u>Setup Screen beginning on page 144.</u>)

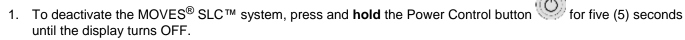


CAUTION! THE OPERATOR SHOULD ALWAYS HAVE AVAILABLE AN ALTERNATE MEANS OF SUCTION IN THE EVENT OF POWER FAILURE, MECHANICAL FAILURE, OR A SERIOUS OCCLUSION IN THE SUCTION SYSTEM.



WARNING! THE O2 CONCENTRATOR DOES NOT PRODUCE O2 WHEN THE SUCTION FEATURE IS IN USE. AN ALTERNATIVE O2 SUPPLY WILL BE NECESSARY IF HIGH O2 CONCENTRATION IS CRITICAL.

13.8 SHUTDOWN PROCEDURES



- 2. Detach all patient monitoring accessories from the patient connection panel.
- 3. Dispose of breathing circuit (but **NOT THE WATER TRAP**), sample lines (but **NOT THE NAFION TUBE**) and filters, breathing cartridge, ECG adhesive sensor pads, IP transducer, suction wand and tubing in a sanitary manner in accordance with local biohazard regulations.
- 4. Clean and sterilize all cables, NIBP cuffs and tubing with a disinfecting spray. Ensure that no moisture enters the tubing. Return items to storage in the accessories case.
- 5. Carefully remove the suction canister and suction filter and dispose of them and waste in a sanitary manner in accordance with local biohazard regulations.
- 6. Remove the MOVES[®] SLC[™] batteries from the system and store them in a dry area at room temperature.



WARNING! LEAVING BATTERIES INSTALLED IN A NON-OPERATIONAL UNIT MAY CAUSE THEM TO DRAIN TO AN UNRECHARGEABLE LEVEL.



CAUTION! THE SUCTION CANISTER AND SUCTION FILTER ARE INTENDED FOR <u>SINGLE USE ONLY</u> AND SHOULD BE DISPOSED OF IN ACCORDANCE WITH LOCAL BIOHAZARD REGULATIONS.



CAUTION! BREATHING CIRCUIT, SAMPLE LINES (BUT NOT THE NAFION TUBE) AND FILTERS, BREATHING CARTRIDGES, ECG ADHESIVE SENSOR PADS, IP TRANSDUCER AND SUCTION WAND AND TUBING ARE DISPOSABLE AND SHOULD BE DISPOSED OF IN ACCORDANCE WITH LOCAL BIOHAZARD REGULATIONS.



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14.0 Using System Graphs | Trends

14.1 OVERVIEW

Plotting appears on two graphs on the Main Screen. Either graph can be set to real-time waveforms or TRENDS, which plots historical values over a user-designated time frame (30 minutes, 1, 2, 4, 8, and 16 hours).

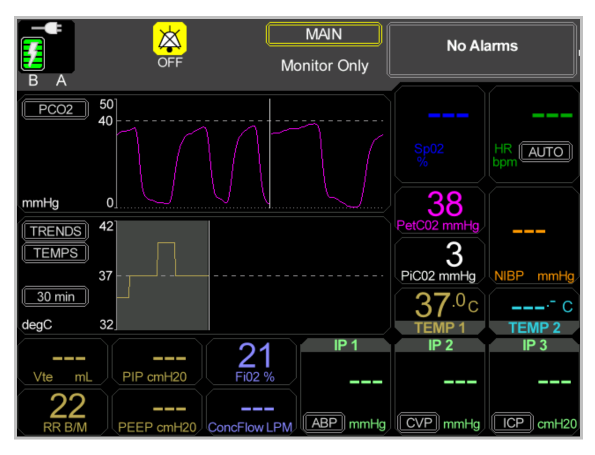


Figure 14-1: Plotting Graphs on Main Screen

The graphs have a vertical scale (from a minimum to a maximum value) for data of interest, and a horizontal time scale. Because plot data is stored, changing plots (e.g., from ECG to Airway) allows the part of the plot to be displayed immediately. Normally, patient data is displayed over time from left to right; however, if there is no data, the sweep continues to erase old data and plots new data when it is available.



NOTE: Data points above the top of the plot are plotted at the top of the plot. Data points below the bottom of the plot are plotted at the bottom of the plot. In other words, if the data is out of range, the display will only show either maximum or minimum values until the data comes back into range.



14.2 AVAILABLE SYSTEM GRAPHS / TRENDS

14.2.1 System Graphs

The following table lists all available System Graphs.

Table 36: System Graphs and Parameters

GRAPH	PARAMETERS			
	Minimum value: -1.1 mV			
	Maximum value: 1.1 mV			
	Vertical sensitivity:			
	Display	MAIN Screen	ECG Screen	
	MOVES® SLC™	10 mm/mV	5 mm/mV	
1. ECG – 2.2 mV	MOVES® SLC™ Remote Screen	14 mm/mV	7 mm/mV	
255 2.2	Plotting rates: 25 mm/sec, 10 mm/sec (MOVES [®] SLC™) 30 mm/sec, 15 mm/sec (MOVES [®] SLC™ Remote Screen)			
	Sampling rates: 150 Hz (25/30 mm/sec); 60 Hz (10/15 mm/sec)			
	Reference line: 0 mV, 1mV			
	Time scale: Real time			
	Minimum value: -1.5 mV			
	Maximum value: 1.5 mV			
	Vertical sensitivity:			
	Display	MAIN Screen	ECG Screen	
	MOVES [®] SLC™	8 mm/mV	4 mm/mV	
2. ECG – 3 mV	MOVES [®] SLC™ Remote Screen	10 mm/mV	5 mm/mV	
	Plotting rates: 25 mm/sec, 10 mm/sec (MOVES [®] SLC™) 30 mm/sec, 15 mm/sec (MOVES [®] SLC™ Remote Screen)			
	Sampling rates: 150 Hz (25/30 mm/sec); 60 Hz (10/15 mm/sec)			
	Reference line: 0 mV, 1mV			
	Time scale: Real time			



GRAPH	PARAMETERS			
	Minimum value: -3.0 mV			
	Maximum value: 3.0 mV			
	Vertical sensitivity:			
	Display	MAIN Screen	ECG Screen	
	MOVES [®] SLC™	4 mm/mV	2 mm/mV	
3. ECG – 6 mV	MOVES [®] SLC™ Remote Screen	5 mm/mV	3 mm/mV	
	Plotting rates: 25 mm/sec, 10 mm/sec (MOVES [®] 30 mm/sec, 15 mm/sec (MOVES [®]			
	Sampling rates: 150 Hz (25/30 mm	/sec); 60 Hz (10/15 mm/sec)		
	Reference line: 0 mV, 1mV			
	Time scale: Real time			
	Minimum value: -6.0 mV			
	Maximum value: 6.0 mV			
	Vertical Sensitivity:			
	Display	MAIN Screen	ECG Screen	
	MOVES [®] SLC™	2 mm/mV	1 mm/mV	
4. ECG – 12 mV	MOVES [®] SLC™ Remote Screen	3 mm/mV	1 mm/mV	
	Plotting rates: 25 mm/sec, 10 mm/sec (MOVES [®] SLC™) 30 mm/sec, 15 mm/sec (MOVES [®] SLC™ Remote Screen)			
	Sampling rates: 150 Hz (25/30 mm/sec); 60 Hz (10/15 mm/sec)			
	Reference line: 0 mV, 1mV			
	Time scale: Real time			
	Minimum value: 60 mmHg			
	Maximum value: 160 mmHg			
5. IP–ABP	Plotting rate: 7.7 mm/sec			
	Sampling rate: 20.8 Hz			
	Reference line: 100 mmHg			
	Minimum value: 0 mmHg			
	Maximum value: 20 mmHg			
6. IP–CVP	Plotting rate: 7.7 mm/sec			
	Sampling rate: 20.8 Hz			
	Reference line: 10 mmHg			



GRAPH	PARAMETERS	
	Minimum value: 0 mmHg	
	Maximum value: 27 mmHg	
7. IP–ICP	Plotting rate: 7.7 mm/sec	
	Sampling rate: 20.8 Hz	
	Reference line: 14 mmHg	
	Plotting rate: 7.7 mm/sec	
	Sampling rate: 20.8 Hz	
8. Pleth	Reference line: None	
(Plethysmograph)	NOTE: The pleth waveform is scaled to a fixed size for signal strengths above 10% or below 0.5% (i.e., all waveforms with a signal strength over 10% or less than 0.5% will have the same amplitude; however, the user will still see an oscillating waveform). Between the values indicated, the waveform is scaled according to signal strength.	
	Minimum value: 0 cmH ₂ O	
O Aimuov	Maximum value: 40 cmH ₂ O	
9. Airway	Plotting rate: 2.66 mm/sec	
	Sampling rate: 7.4 Hz	
	Reference line: 30 cmH ₂ O	
	Minimum value: 0 mmHg	
	Maximum value: 50 mmHg	
10. PCO ₂	Plotting rate: 4.00 mm/sec	
Sampling rate: 11.1 Hz		
	Reference line: 40 mmHg	
	Time scale: Real time	

14.2.2 System Trends

There are sixteen (16) trend charts:

•	FiO2 / SpO2	•	PetCO2
•	PIP	•	SpO2

• TEMPS (both TEMP1 and TEMP2) • PI

• HR • SpCO

ABP (available under IP1, IP2, or IP3)
 SpMet

CVP (available under IP1, IP2, or IP3) • SpHb

- ICP (available under IP1, IP2, or IP3)
- NIBP

- SpOC
- PVI

Trend plots can be set to display trend data for:

- 30 minutes
- 1 hour
- 2 hours
- 4 hours
- 8 hours
- 16 hours
- 24 hours

The following table lists all System Trends.

Table 37: System Trends and Parameters

PARAMETERS
Time: 30 min, 1, 2, 4, 8, 16, 24 hours
Range: 60–160 mmHg
Reference line: 100 mmHg
Time: 30 min, 1, 2, 4, 8, 16, 24 hours
Range: 0–20 mmHg
Reference line: 10 mmHg
Time: 30 min, 1, 2, 4, 8, 16, 24 hours
Range: 0–27 cmH ₂ O
Reference line: 14 cmH ₂ O
Time: 30 min, 1, 2, 4, 8, 16, 24 hours
Range: 60–160 mmHg
Reference line: 100 mmHg
Time: 30 min, 1, 2, 4, 8, 16, 24 hours
Range: 0–50 mmHg
Reference line: 40 mmHg
Time: 30 min, 1, 2, 4, 8, 16, 24 hours
Range: 0-100%
Reference line: 21%
_



TREND	PARAMETERS
	Time: 30 min, 1, 2, 4, 8, 16, 24 hours
7. Trend Plot - PIP	Range: 0–40 cmH ₂ O
	Reference line: 30 cmH ₂ O
	Time: 30 min, 1, 2, 4, 8, 16, 24 hours
8. Trend Plot – TEMP	Range: 89°F–108°F (32°C–42°C)
	Reference line: 98.6°F (37°C)
	Time: 30 min, 1, 2, 4, 8, 16, 24 hours
9. Trend Plot – HR	Range: 0–200 bpm
	Reference lines: 60, 100 bpm
	Time: 30 min, 1, 2, 4, 8, 16, 24 hours
10. Trend Plot – SpO2	Range: 80–100%
	Reference line: 90%
	Time: 30 min, 1, 2, 4, 8, 16, 24 hours
11. Trend Plot – Pl	Range: 0–20%
	Reference line: None
	Time: 30 min, 1, 2, 4, 8, 16, 24 hours
12. Trend Plot – SpCO	Range: 0–40%
	Reference line: 10%
	Time: 30 min, 1, 2, 4, 8, 16, 24 hours
13. Trend Plot – SpMet	Range: 0–70%
·	Reference line: 10%
	Time: 30 min, 1, 2, 4, 8, 16, 24 hours
14. Trend Plot – SpHb	Range: 0–25 g/dL (0–15.5 mmol/L, 0–250 g/L)
	Reference lines: 10, 15 g/dL (6.2, 9.3 mmol/L; 100, 150 g/L)
	Time: 30 min, 1, 2, 4, 8, 16, 24 hours
15. Trend Plot – SpOC	Range: 0–35 mL/dL
	Reference line: 13, 20 mL/dL
	Time: 30 min, 1, 2, 4, 8, 16, 24 hours
16. Trend Plot – PVI	Range: 0–50%
	Reference line: 15%



15.0 Alarms

Alarm conditions which require immediate or priority user attention can be distinguished from normal status alarms at a distance via their highlighted screen colors, audible annunciation, and alarm status LEDs. If a user is too far away from MOVES[®] SLCTM to see the contents of the display, the user will still be informed of the alarm condition by seeing that the status LEDs are <u>NOT green</u>.

15.1 ABOUT STATUS LEDS

There are four (4) status LEDs, each situated on a top corner of the MOVES[®] SLC[™]. They are placed away from the display so that they can be seen when the display cannot. Note the following:

- When the system is NOT in alarm, the status LEDs are solid green.
- When the system is IN alarm, the status LEDS are solid yellow, flashing yellow, or flashing red (from low to medium to high in order of priority).
- If alarms have been silenced, the status LEDs continue to indicate alarm level.

Alarms and messages are displayed in the top right of the screen's status area. When a new alarm condition becomes active, or an existing alarm becomes inactive, the system will update the visual and audible alarm state with the highest priority active alarm. To see the text of an alarm other than that of the highest-priority alarm, the user can navigate to the Alarm Queue on the status bar, press the Jog Wheel, and then turn the Jog Wheel to page through the alarms.

15.2 ALARM PRIORITIES AND CHARACTERISTICS

15.2.1 Standard Alarms

The following table details the MOVES® SLCTM standard alarms, their audio and visual characteristics, and their priorities.

Table 38: Alarm Types and Descriptions

ALARMS TABLE			
Alarm Priority	Description / Response	Visible Alarm	Audible Alarm
High	Immediate operator response required	Red LED 60% duty cycle flashing at 2.8 Hz	Three short notes of ascending pitch followed by two long notes of ascending pitch. The musical sequence is a common triad (C-E-G) followed by a perfect fourth (G-C). When measured with a microphone at a distance of one (1) meter, the sound pressure level of the High Priority alarm is 62 dB. (This was measured with an Aweighted background level of 27.2 dB which included any information signal or extraneous noise.)



	ALARMS TABLE				
Alarm Priority	Description / Response	Visible Alarm	Audible Alarm		
Medium	Prompt operator response required	Yellow LED 60% duty cycle at 0.8 Hz	Three notes of equal, medium length and ascending pitch. The musical sequence is a common triad (C-E-G). When measured with a microphone at a distance of one (1) meter, the sound pressure level of the Medium Priority alarm is 61.5 dB. (This was measured with an A-weighted background level of 27.2 dB which included any information signal or extraneous noise.)		
Low	Operator awareness required	Yellow LED Constant ON	None		
Information	Message	Green LED (Indicating no Alarm) Constant ON	None		



NOTE: Low priority alarms and/or messages will not be audibly indicated.

15.2.2 High Priority Communication Failure Alarm

The High Priority Communication Failure alarm is triggered when there is no longer communication between the MOVES[®] SLC™ user interface and its internal systems.

Alarm Priority Description		Visible Alarm	Audible Alarm	
High	System failure. Immediate operator response required.	Red LED 60% duty cycle flashing at 2.8 Hz	Three short notes of the same duration and pitch (the note "C").	

15.3 ALARM QUEUE

The Alarm Queue is displayed in the top right corner of the screen's Status Bar. Initially, only the text of the highest-priority alarm is displayed. To see the text of an alarm other than that of the highest-priority alarm, the user navigates to the Alarm Queue with the Jog Wheel, presses the Wheel, and then uses the Jog Wheel to page through the additional alarms. The Alarm Queue shows all active alarms, non-active latched alarms, and non-alarm messages. If there are no alarms, the message No Alarms is displayed.

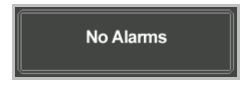


Figure 15-1: No Alarms Message in Alarm Queue

15.4 LOCKED ALARMS AND MESSAGES

Some alarms and messages CANNOT be turned off or altered in any way, even by someone with Administrator privileges. These alarms and messages are shown with a padlock beside them in the alarm list on the Alarm ON/OFF screen. Selecting them and pressing the Jog Wheel will have no effect on their status (unlike other alarms which can be toggled ON / OFF or dismissed).

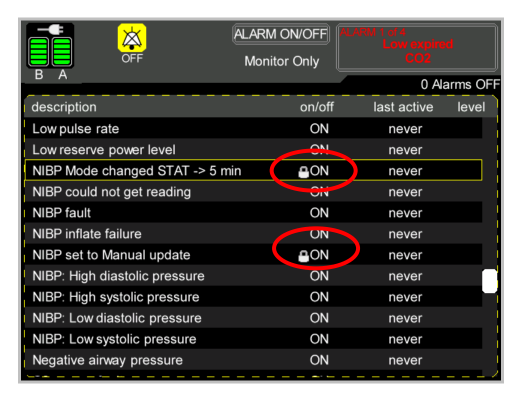


Figure 15-2: Two Locked Alarms

15.5 INHIBITABLE AND LATCHING ALARMS

Inhibitable and Latching alarms can be distinguished from General alarms by the "dismiss" option initially shown in the top right corner.

15.5.1 Inhibitable Alarms

Some, but not all, $MOVES^{\mathbb{R}}$ SLC^{TM} alarms are Inhibitable. An Inhibitable alarm can be dismissed and cleared even though the condition or event that triggered the alarm is still active.

An Inhibitable alarm is shown in two states in the images below.



Figure 15-3: Inhibitable Alarm Able To Be Dismissed



Figure 15-4: Inhibitable Alarm Cleared



To clear the Inhibitable alarm, select it in the Alarm Queue using the Jog Wheel and then press the Wheel. Once an alarm is cleared, it will be removed from the Alarm Queue.



NOTE: The alarm will, of course, also be cleared if the condition or event that triggered the alarm is addressed or corrected.

15.5.2 Latching Alarms

Some alarms (known as latching alarms) require operator acknowledgment before they are removed from the Alarm Queue (i.e., they will persist in the Alarm Queue even if the condition or event that triggered the alarm is addressed / corrected). This guarantees that the operator is made aware that the alarm condition occurred **even if the alarm condition has since disappeared**. A latching alarm is visually identical to an Inhibitable alarm. A latching alarm is shown is shown in two states in the images below.



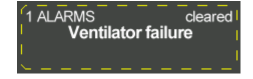


Figure 15-5: Latched Alarm Able To Be Dismissed

Figure 15-6: Latched Alarm Cleared

To clear the Latched alarm, select it in the Alarm Queue using the Jog Wheel and then press the Wheel. Once an alarm is cleared, it will be removed from the Alarm Queue. A Latched alarm will also not persist after MOVES[®] SLCTM is shut down.



NOTE: All alarms are logged in the log file.



NOTE: See <u>Alarm Conditions and Causes beginning on page 238</u> for identification of which alarms are Inhibitable, which are Latched, and which are General and neither.

15.5.3 Alarms That Have Been Turned Off

Alarms that are not Locked can be turned OFF using the Alarm ON/OFF screen. A white triangle with an "X" through it presented on a red chamfered rectangle indicates that some alarms have been turned off. This icon is shown in the top left corner of the Status Bar. The number of alarms turned off is indicated below the icon.



Figure 15-7: Latched Alarm Able To Be Dismissed

15.5.4 Auto Restoration of Alarm Settings Upon Power Loss

When power is lost for less than or equal to three (3) minutes, the alarm settings prior to the power loss are restored automatically. However, the latched or inhibited state of all alarms is lost immediately upon power loss and is NOT restored.

After three (3) minutes and no more than 30 minutes, the operator will be asked: "Configure system for new patient?". If the operator responds YES, then the alarms are returned to their default values; otherwise, the old alarm settings are used. After 30 minutes all alarms return to default values.

15.6 TESTING ALARMS

15.6.1 Testing Adjustable Alarms

Table 39: Adjustable Alarms, Test Procedures and Results

Alarm	Test Procedure	Result / Observation	
1. High Airway Pressure Warning	 Setup MOVES[®] SLC[™] breathing circuit with disposable ventilator cartridge, patient tubing, Y-piece, compliant reservoir (i.e., balloon) and sample line. Go to the "Alarm Limits" screen and change the "High airway pressure warning" to 20 cmH₂O. Start ventilating the reservoir in default ventilator mode, with Control Pressure = 30 cmH₂O. Wait for alarm. 	Alarm queue should display "High airway pressure airway warning" within 20 seconds.	
2. High Airway Pressure Release	 Setup MOVES[®] SLC[™] breathing circuit with disposable ventilator cartridge, patient tubing, Y-piece, compliant reservoir (i.e., balloon) and sample line. Go to the "Alarm Limits" screen and change the "High airway pressure release" to 40 cmH₂O. Start the ventilating reservoir in default ventilator mode, with Control Pressure = 45 cmH₂O. Wait for alarm. 	Alarm queue should display "High airway pressure release" within 20 seconds.	



Alarm	Test Procedure	Result / Observation
3. High Expired CO₂	 Ensure the O₂/CO₂ sensor is warmed up (values populate "Monitor" screen). Go to the "Alarm Limits" screen and change the "High expired pCO₂" to 50 mmHg. Quickly expose the sample line to gas with higher than ambient CO₂ concentration (should be > 7% CO₂ at sea level or greater for higher altitudes) until pCO₂ on "Monitor" screen goes > 50 mmHg. Wait for alarm. 	Alarm queue should display "High expired CO ₂ " within 20 seconds.
4. Low Expired CO2	 Ensure the O₂/CO₂ sensor is warmed up (values populate "Monitor" screen). Go to the "Alarm Limits" screen and change the "Low expired PCO₂" to 35 mmHg. Breathe lightly on sample line until pCO₂ on "Monitor" screen goes < 35mmHg. Wait for alarm. 	Alarm queue should display "Low expired CO2" within 20 seconds.
5. Low SpO2	 Plug an SpO₂ finger clip into the MOVES[®] SLC™ and place the finger clip on an SpO₂ simulator. Set the simulator for SpO₂ < 90%. Wait for alarm. (NOTE: M-LNCS SpO₂ sensors are compatible with more simulators.) 	Alarm queue should display "Low SpO2" within 20 seconds.
6. Low pulse rate	 Use one MOVES[®] SLC[™] heart rate source (ABP, SpO₂ or ECG) cable plugged into MOVES[®] SLC[™] and a patient simulator. Set heart rate to < 50 bpm. Wait for alarm. 	Alarm queue should display "Low pulse rate" within 20 seconds.
7. High pulse rate	 Use one MOVES[®] SLC[™] heart rate source (ABP, SpO₂ or ECG) cable plugged into MOVES[®] SLC[™] and a patient simulator. Set heart rate to > 120 bpm. Wait for alarm. 	Alarm queue should display "High Pulse rate" within 20 seconds.



Alarm	Test Procedure	Result / Observation	
8. IPx-ABP: High systolic pressure (where x is 1, 2 or 3)	 Plug the IP cable into one of the 3 MOVES[®] SLC™ IP ports and into a clinical blood pressure transducer. With the transducer open to air, select "Zero". Ensure that the channel in use is configured for ABP. Attach a syringe to the transducer and slowly increase the pressure on the transducer until the MOVES[®] SLC™ displays systolic pressure > 180 mmHg. Wait for alarm. 	Alarm queue should display "IPx-ABP: High systolic pressure" within 20 seconds.	
9. IPx-ABP: Low systolic pressure (where x is 1, 2 or 3)	 Plug the IP cable into one of the 3 MOVES[®] SLC™ IP ports and into a clinical blood pressure transducer. With the transducer open to air, select "Zero". Ensure that the channel in use is configured for ABP. Wait for alarm. 	Alarm queue should display "IPx-ABP: Low systolic pressure" within 20 seconds.	
10. IPx: Low CVP (where x is 1, 2 or 3)	 Plug the IP cable into one of the 3 MOVES[®] SLC™ IP ports and into a clinical blood pressure transducer. With the transducer open to air, select "Zero". Ensure that the channel in use is configured for CVP. Wait for alarm. 	Alarm queue should display "IPx: Low CVP" within 20 seconds.	
11. IPx: High CVP (where x is 1, 2 or 3)	 Plug the IP cable into one of the 3 MOVES® SLC™ IP ports and into a clinical blood pressure transducer. With the transducer open to air, select "Zero". Ensure that the channel in use is configured for CVP. Attach a syringe to the transducer and slowly increase the pressure on the transducer until the MOVES®SLC™ displays systolic pressure > 20 mmHg. Wait for alarm. 	Alarm queue should display "IPx: High CVP" within 20 seconds.	



Alarm	Test Procedure	Result / Observation	
12. IPx: Low ICP (where x is 1, 2 or 3)	 Plug the IP cable into one of the 3 MOVES[®] SLC™ IP ports and into a clinical blood pressure transducer. 	Alarm queue should display "IPx: Low ICP" within 20 seconds.	
	With the transducer open to air, select "Zero".		
	Ensure that the channel in use is configured for ICP.		
	4. Wait for alarm.		
13. IPx: High ICP (where x is 1, 2 or 3)	 Plug the IP cable into one of the 3 MOVES[®] SLC™ IP ports and into a clinical blood pressure transducer. 	Alarm queue should display "IPx: High ICP" within 20 seconds.	
	With the transducer open to air, select "Zero".		
	3. Ensure that the channel in use is configured for ICP.		
	 Attach a syringe to the transducer and slowly increase the pressure on the transducer until the MOVES[®] SLC™ displays systolic pressure > 27 cmH₂O. Wait for alarm. 		

15.6.2 Testing Non-Adjustable Alarms

Table 40: Non-Adjustable Alarms, Test Procedures and Results

	Alarm	Test P	rocedure	Result / Observation	
1.	Power Failure	 Turn MOVES[®] Unplug from the open both batter 	ne wall power and	Alarm visual indicator lights should be solid red for at least seven seconds.	
2.	High Oxygen Concentration FiO₂) 1. Setup MOVES [®] SLC™ breathic circuit with disposable ventilated cartridge, patient tubing, Y-piece compliant reservoir (i.e., balloo and sample line.		posable ventilator ent tubing, Y-piece, ervoir (i.e., balloon)	Check alarm queue for "Inspired O2 above target" alarm within 20 seconds.	
		ventilator mod Allow to ventila	g reservoir in default e. Set $O_2 = 70\%$. ate until ump stops $(O_2 >$		
		While still vent 30%. Wait for	ilating, set O ₂ = alarm.		



Alarm		Test Procedure	Result / Observation	
3.	Low Oxygen Concentration (FiO₂)	 Setup MOVES[®] SLC[™] for oxygen supplementation Redirect the sample line to ambient air. 	Check alarm queue for "Inspired O2 below target" alarm within 30 seconds.	
4.	Breathing Circuit Integrity	 Setup MOVES[®] SLC[™] breathing circuit with disposable ventilator cartridge, patient tubing, Y-piece, compliant reservoir (i.e., balloon) and sample line. Disconnect sample line from Y-piece. Start ventilating the reservoir on default ventilator settings. Wait for alarm. 	Check alarm queue for "Patient circuit disconnect" alarm within 20 seconds. May get "Leak detected" alarm within 30 seconds.	



15.7 ALARM CONDITIONS AND CAUSES



NOTE: If no values are given in the Locked / Latching or Inhibitable columns, these conditions do not apply to the listed alarm.

Table 41: Alarm Conditions and Causes

	Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
1.	Internal fault: Service soon	Low	Latching		Low priority fault indicating possible calibration issues with patient monitoring and/or system sensors, concentrator pump temperature sensor fault, real-time clock malfunction, or internal memory malfunction.	 Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
2.	Low Temperature System warming	Low			System temperature is less than 0°C (32°F), or concentrator pump temperature is less than 0°C (32°F) while system temperature is in the range of -26°C (-14.8°F) to 70°C (158°F). Once active alarm will stay active until both system temperature and concentrator pump temperature exceed 5°C (41°F).	 Wait for system to warm up or proceed to a higher temperature environment. If erroneous, return MOVES[®] SLC™ for servicing.
3.	System Temperature too high	Low			System temperature exceeds 70°C (158°F).	 Remove any contact between the patient and the system enclosure. Proceed to a lower temperature environment. If erroneous, return MOVES[®] SLC™ for servicing.

	Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
4.	System Temperature too low	Low			System temperature is less than -26°C (-14.8°F).	 Proceed to a higher temperature environment. If erroneous, return MOVES[®] SLC™ for servicing.
5.	TEMP1: High patient temperature	Low			Patient temperature is over 38°C (100.4°F) and no more than 42°C (107.6°F).	 Check integrity of interface with patient. Fully disconnect then reconnect temperature patient connection. Check for related alarms and follow actions if required. Clinical intervention may be required.
6.	TEMP1: Low patient temperature	Low			Patient temperature is no less than 28°C (82.4°F) and under 35°C (95.0°F).	 Check integrity of interface with patient. Fully disconnect then reconnect temperature probe patient connection. Check for related alarms and follow actions if required. Clinical intervention may be required.
7.	TEMP1: Temperature probe disconnect	Low		Inhibitable	Temperature probe patient connection is disconnected from either system or patient after previously being connected.	 Fully disconnect then reconnect temperature probe patient connection. Check integrity of interface with patient. Replace temperature probe patient connection. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.



	Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
8.	TEMP2: High patient	Low			Patient temperature is over 38°C (100.4°F) and no more than 42°C (107.6°F).	 Check integrity of interface with patient. Fully disconnect then reconnect temperature patient connection. Check for related alarms and follow actions if required. Clinical intervention may be required.
9.	TEMP2: Low patient	Low			Patient temperature is no less than 28°C (82.4°F) and under 35°C (95.0°F).	 Check integrity of interface with patient. Fully disconnect then reconnect temperature probe patient connection. Check for related alarms and follow actions if required. Clinical intervention may be required.
10	TEMP2: Temperature probe disconnect	Low		Inhibitable	Temperature probe patient connection is disconnected from either system or patient after previously being connected.	 Fully disconnect then reconnect temperature probe patient connection. Check integrity of interface with patient. Replace temperature probe patient connection. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
11	. Concentrator temperature sensor fault	Low			Redundant internal temperature sensor communication failure with concentrator.	 Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
12	. Concentrator failure	Medium			Concentrator communication failure with UI or concentrator reports concentrator pump fault.	 Turn suction on and off for 2 seconds. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.



Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
13. Concentrator in degraded mode	Medium			Concentrator reports partial failure due to unexpected data, including: product tank flow, product tank pressure and suction pressure. Oxygen flow or concentration may be reduced.	 Monitor inspired O₂ to ensure adequate oxygenation is being maintained in the ventilation circuit or O₂ supplement mask. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
14. Suction not available	Medium			Concentrator communication failure with UI or concentrator reports concentrator pump fault.	 Turn suction on and off for 2 seconds. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
15. Ventilator failure Use backup vent if attempting to ventilate	High, if vent problem while attempting to ventilate Low otherwise	Latching		Communication failure between ventilator module and UI, erroneous feedback from blower to ventilator module, blower air inlet occlusion, ventilator pressure sensor fault, or low supply voltage to ventilator module while in ventilation mode. NOTE: In the event of a communication failure, the ventilator will continue to operate with the last user-set parameters and will operate the air pump at least 9 LPM. NOTE: If a "Leak alarm" and a "Ventilator failure" alarm occur simultaneously, the leak is only substantial if you see alternating Ve/Vi values displayed.	 Ensure that the blower inlet is not occluded. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.



Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
16. Vent vol-flow fault Try PC-IMV mode if venting and not already in CPAP/PS or PC-IMV	Medium			Ventilator flow sensor communication failure with ventilator detected during system start-up tests.	 Power cycle MOVES[®] SLC[™]. Repeat system start-up tests. Revert to pressure control ventilation modes if necessary. Return MOVES[®] SLC[™] for servicing.
17. Air pump failure	Medium			Erroneous feedback from air pump is detected by ventilator.	 Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
18. Leak detected	Medium			Delivered inspired tidal volume is 25% or 100 mL more than exhaled tidal volume, or the internal ventilator bag has collapsed. NOTE: A high expiratory flow rate (over 80 LPM) can be misinterpreted as being lower than actual, resulting in an incorrect leak report. NOTE: If a "Leak alarm" and a "Ventilator failure" alarm occur simultaneously, the leak is only substantial if you see alternating Ve/Vi values displayed.	 Ensure ventilator cartridge door is closed and latched properly. Ensure all breathing circuit connections (including water trap, patient tubing, Y-piece and sample line, Nafion and sample line filters) are tight and no leaks are detected. Replace breathing circuit. Leak may be patient-related. Clinical intervention may be required. Return MOVES® SLC™ for servicing.
19. Internal power monitoring fault	Medium			Internal supply voltage mismatches between any or all of the ventilator, UI, concentrator, patient monitor, or power manager modules.	 Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
20. Power manager fault	Medium			Communication failure between power manager and UI.	 Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.



Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
21. Battery A: Unknown power level	Low			Power manager detects presence of battery but fails to communicate or power manager fault alarm is active.	 Replace affected battery or batteries. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
22. Battery B: Unknown power level	Low			Power manager detects presence of battery but fails to communicate or power manager fault alarm is active.	 Replace affected battery or batteries. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
23. Battery A: Not charging	Low			Battery charge level has not increased by more than 1% after one hour of charging and is less than 96%.	 Remove and replace problematic battery. Replace MOVES[®] SLC[™] power supply/charger. Power cycle MOVES[®] SLC[™]. Return battery and/or power supply/charger and/or MOVES[®] SLC[™] for servicing.
24. Battery B: Not charging	Low			Battery charge level has not increased by more than 1% after one hour of charging and is less than 96%.	 Remove and replace problematic battery. Replace MOVES[®] SLC[™] power supply/charger. Power cycle MOVES[®] SLC[™]. Return battery and/or power supply/charger and/or MOVES[®] SLC[™] for servicing.
25. Low battery	High	Locked		Running on battery power and all batteries are below 20%.	 Connect power supply/charger to MOVES[®] SLC™ to allow battery or batteries to charge. If running on a single battery then insert a second fully charged battery. If running on two batteries then replace each battery (one at a time) with a fully charged battery.



	Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
26	. Backup power source required	High			Only one valid power source (battery A, battery B or power supply/charger) is detected and the low battery alarm is not active. Battery charge level must be greater than or equal to 10% to be considered valid.	 Connect a second valid power source to MOVES[®] SLC™. If second valid power source is connected and alarm persists, ensure that battery doors are fully closed/latched and/or power supply/charger connection is secure.
27	. Low backup power level	Medium		Inhibitable	Backup battery charge level is less than 40% when two valid power sources (battery A, battery B or power supply/charger) are detected and the low battery alarm is not active. NOTE: Backup battery charge is all batteries present when connected to wall power or the battery with the lowest charge level when running on battery power.	 Connect power supply/charger to MOVES[®] SLC™ to allow battery or batteries to charge if not already connected. Replace battery with low charge level with a fully charged battery.
28	. Critically low backup power	High			Backup battery charge level is less than 20% when two valid power sources (battery A, battery B or power supply/charger) are detected and the low battery alarm is not active. NOTE: Backup battery charge is all batteries present when connected to wall power or the battery with the lowest charge level when running on battery power.	 Connect power supply/charger to MOVES® SLC™ to allow battery or batteries to charge if not already connected. Replace battery with critically low charge level with a fully charged battery.



Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
29. Patient monitor failure	Medium			Patient monitor module communication failure with MOVES [®] SLC™.	 Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
30. ECG lead off	Low		Inhibitable	ECG lead is disconnected from patient after previously being connected.	 Check integrity of interface with patient. Fully disconnect then reconnect ECG patient connections. Replace ECG patient connections. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
31. ECG fault	Medium			ECG module internal fault or ECG module communication failure with patient monitor.	 Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
32. Pulse Ox: Cable disconnect	Low		Inhibitable	Pulse oximeter extension cable or patient sensor cable (if no extension cable used) is disconnected from system after previously being connected.	 Fully disconnect then reconnect pulse oximeter extension cable or patient sensor to the system. Replace pulse oximeter extension cable and/or patient sensor. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
33. Pulse Ox: Replace cable	Low			Pulse oximeter extension cable is non-functional or the life of the cable has expired.	Replace pulse oximeter extension cable with another that has not expired.
34. Pulse Ox: Invalid cable	Low			Pulse oximeter extension cable is not compatible with system.	Replace pulse oximeter extension cable with another that is compatible with MOVES [®] SLC™.



Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
35. Pulse Ox: Sensor disconnect	Low		Inhibitable	Pulse oximeter patient sensor cable is disconnected from extension cable connected to system after previously being connected, or if using single-patient-use sensor the adhesive portion of the sensor has come loose after previously being connected. May be an incorrect sensor or a defective sensor or cable.	 Fully disconnect then reconnect pulse oximeter patient connection cable to extension cable, or if using single-patient-use sensor fully disconnect and reconnect adhesive portion of the sensor. Fully disconnect then reconnect pulse oximeter extension cable to system. Replace pulse oximeter extension cable and/or patient sensor. Power cycle MOVES® SLC™. Return MOVES® SLC™ for servicing.
36. Pulse Ox: Replace sensor	Low			Pulse oximeter patient sensor has expired and needs to be replaced.	Replace pulse oximeter patient sensor with another that has not expired.
37. Pulse Ox: Invalid sensor	Low			Pulse oximeter patient sensor is not compatible with system.	Replace pulse oximeter patient sensor with another that is compatible with MOVES [®] SLC [™] .
38. PulseOx: Patient disconnect	Low		Inhibitable	Pulse oximeter patient sensor is disconnected from patient. Sensor may not be connected to patient properly or sensor may be damaged.	 Check placement of sensor or if it is too tight. Confirm that the sensor is securely on the patient. Reapply sensor or select a new site. See the directions for use provided with your sensor. Reconnect the sensor to the MOVES® SLC™ or extension cable. If the sensor is damaged, replace the sensor. Power cycle MOVES® SLC™. Return MOVES® SLC™ for servicing.



Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
39. PulseOx: Check sensor	Low			Pulse oximeter sensor at system or patient is compromised.	 Check integrity of interface with patient. Fully disconnect then reconnect pulse oximeter patient sensor to system. Replace pulse oximeter patient sensor. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
40. PulseOx: Check cable and sensor	Low			Pulse oximeter cable and/or sensor at system or patient is compromised.	 Check integrity of interface with patient. Fully disconnect then reconnect pulse oximeter patient sensor to extension cable. Fully disconnect then reconnect pulse oximeter extension cable to system. Replace pulse oximeter patient sensor. Replace pulse oximeter extension cable. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
41. PulseOx: Interference detected	Low			High intensity light such as pulsating strobe lights, excessive ambient light sources such as surgical lights or direct sunlight, or other monitor displays. Incorrect Line Filter frequency setting (Hz).	 Shield the sensor from excessive or strobing light. Minimize or eliminate motion at the monitoring site. Adjust the Line Filter frequency to the correct setting (See Setup Screen on page 144 and following). Replace pulse oximeter patient sensor. Power cycle MOVES® SLC™. Return MOVES® SLC™ for servicing.



Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
42. PulseOx: Low perfusion	Low		Inhibitable	Pulse oximeter patient signal too small (related to SpO ₂ measurement).	 Check placement of sensor or if it is too tight. Confirm that the sensor is securely on the patient. Reapply sensor or select a better perfused site. See the directions for use provided with your sensor. Fully disconnect then reconnect pulse oximeter patient sensor to system. Replace pulse oximeter patient sensor. Clinical intervention may be required.
43. PulseOx: Low SpCO perfusion	Low		Inhibitable	Pulse oximeter detects low perfusion related to SpCO measurement.	 Check placement of sensor or if it is too tight. Confirm that the sensor is securely on the patient. Reapply sensor or select a better perfused site. See the directions for use provided with your sensor. Fully disconnect then reconnect pulse oximeter patient sensor to system. Replace pulse oximeter patient sensor. Clinical intervention may be required.



Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
44. PulseOx: Low SpMet perfusion	Low		Inhibitable	Pulse oximeter detects low perfusion related to SpMet measurement.	 Check placement of sensor or if it is too tight. Confirm that the sensor is securely on the patient. Reapply sensor or select a better perfused site. See the directions for use provided with your sensor. Fully disconnect then reconnect pulse oximeter patient connection to system. Replace pulse oximeter patient sensor. Clinical intervention may be required.
45. PulseOx: Low SpHb perfusion	Low		Inhibitable	Pulse oximeter detects low perfusion related to SpHb measurement.	 Check placement of sensor or if it is too tight. Confirm that the sensor is securely on the patient. Reapply sensor or select a better perfused site. See the directions for use provided with your sensor. Fully disconnect then reconnect pulse oximeter patient sensor to system. Replace pulse oximeter patient sensor. Clinical intervention may be required.



Al	larm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
	ulseOx: Low SpOC erfusion	Low		Inhibitable	Pulse oximeter detects low perfusion related to SpOC measurement.	 Check placement of sensor or if it is too tight. Confirm that the sensor is securely on the patient. Reapply sensor or select a better perfused site. See the directions for use provided with your sensor. Fully disconnect then reconnect pulse oximeter patient sensor to system. Replace pulse oximeter patient sensor. Clinical intervention may be required.
	ulseOx: SpO2 ading confidence oor	Low			Pulse oximeter indicating low confidence in SpO ₂ measurement value due to low signal quality. Low signal quality may be a result of excessive motion relative to perfusion, or sensor is damaged or not functioning.	 Check and see if blood flow to the monitoring site is restricted. Ensure proper sensor type and application. Minimize or eliminate motion at the monitoring site. Reapply sensor or select a better perfused site. Replace pulse oximeter patient sensor. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.



Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
48. PulseOx: HR reading confidence poor	Low			Pulse oximeter indicating low confidence in heart rate measurement value due to low signal quality when source of heart rate displayed by the system is the pulse oximeter. Low signal quality may be a result of excessive motion relative to perfusion, or sensor is damaged or not functioning.	 Check and see if blood flow to the monitoring site is restricted. Ensure proper sensor type and application. Minimize or eliminate motion at the monitoring site. Reapply sensor or select a better perfused site. Replace pulse oximeter patient sensor. If available, use alternate heart rate source on MOVES® SLC™. Power cycle MOVES® SLC™. Return MOVES® SLC™ for servicing.
49. PulseOx: PI reading confidence poor	Low			Pulse oximeter indicating low confidence in perfusion index measurement value due to low signal quality. Low signal quality may be a result of excessive motion relative to perfusion, or sensor is damaged or not functioning	 Check and see if blood flow to the monitoring site is restricted. Ensure proper sensor type and application. Minimize or eliminate motion at the monitoring site. Reapply sensor or select a better perfused site. Replace pulse oximeter patient sensor. Power cycle MOVES® SLC™. Return MOVES® SLC™ for servicing.



Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
50. PulseOx: SpCO reading confidence poor	Low			Pulse oximeter indicating low confidence in SpCO measurement value due to low signal quality. Low signal quality may be a result of excessive motion relative to perfusion, or sensor is damaged or not functioning	 Check and see if blood flow to the monitoring site is restricted. Ensure proper sensor type and application. Minimize or eliminate motion at the monitoring site. Reapply sensor or select a better perfused site. Replace pulse oximeter patient sensor. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
51. PulseOx: SpMet reading confidence poor	Low			Pulse oximeter indicating low confidence in SpMet measurement value due to low signal quality. Low signal quality may be a result of excessive motion relative to perfusion, or sensor is damaged or not functioning	 Check and see if blood flow to the monitoring site is restricted. Ensure proper sensor type and application. Minimize or eliminate motion at the monitoring site. Reapply sensor or select a better perfused site. Replace pulse oximeter patient sensor. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.



Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
52. PulseOx: SpHb reading confidence poor	Low			Pulse oximeter indicating low confidence in SpHb measurement value due to low signal quality. Low signal quality may be a result of excessive motion relative to perfusion, or sensor is damaged or not functioning.	 Check and see if blood flow to the monitoring site is restricted. Ensure proper sensor type and application. Minimize or eliminate motion at the monitoring site. Reapply sensor or select a better perfused site. Replace pulse oximeter patient sensor. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
53. PulseOx: SpOC reading confidence poor	Low			Pulse oximeter indicating low confidence in SpOC measurement value due to low signal quality. Low signal quality may be a result of excessive motion relative to perfusion, or sensor is damaged or not functioning.	 Check and see if blood flow to the monitoring site is restricted. Ensure proper sensor type and application. Minimize or eliminate motion at the monitoring site. Reapply sensor or select a better perfused site. Replace pulse oximeter patient sensor. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.



	Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
54.	PulseOx: PVI reading confidence poor	Low			Pulse oximeter indicating low confidence in PVI measurement value due to low signal quality. Low signal quality may be a result of excessive motion relative to perfusion, or sensor is damaged or not functioning.	 Check and see if blood flow to the monitoring site is restricted. Ensure proper sensor type and application. Minimize or eliminate motion at the monitoring site. Reapply sensor or select a better perfused site. Replace pulse oximeter patient sensor. Power cycle MOVES® SLC™. Return MOVES® SLC™ for servicing.
55.	PulseOx: Only SpO2 available	Low		Inhibitable	SpCO, SpMet, SpHb and SpOC measurements unavailable due to failed initialization. SpO2, HR and perfusion index (PI) measurements are still available. NOTE: Retry of SpCO, SpMet, SpHb and SpOC measurement initialization initiated by removing and then re-attaching patient sensor to MOVES® SLC™.	 Fully disconnect and reconnect pulse oximeter patient sensor to patient. Fully disconnect then reconnect pulse oximeter patient sensor to system. Ensure proper sensor type and application. Minimize or eliminate motion at the monitoring site. Reapply sensor or select a better perfused site. Replace pulse oximeter patient sensor. Power cycle MOVES® SLC™. Return MOVES® SLC™ for servicing.
56.	PulseOx fault	Low			Pulse oximeter module communication failure or self-diagnosed failure on pulse oximeter module.	 Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.



Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
57. PulseOx: Busy programming	Low	Locked		Pulse oximeter has been placed in special programming mode to allow software updates or feature activation. NOTE: Pulse oximeter measurement and monitoring are disabled when pulse oximeter is in programming mode.	 If intentionally updating software or enabling new features then let programming complete. If not intentionally updating software then ensure nothing is connected to SpO₂ port on Patient Connection Panel, wait 2 minutes to see if alarm clears, if it does not then power cycle MOVES[®] SLCTM. If alarm continues to persist after multiple power cycles then return MOVES[®] SLCTM for servicing.
58. Gas sensor pressure fault	Medium			Mismatch between O ₂ and CO ₂ sensor pressures, indicating one of the sensors is likely not operating correctly.	 Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
59. CO2 sensor fault	Medium			CO ₂ sensor internal fault, CO ₂ sensor communication failure with patient monitor or calibration issue.	 Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
60. O2 sensor fault	Medium			O ₂ sensor internal fault, O ₂ sensor communication failure with patient monitor or calibration issue.	 Power cycle MOVES[®] SLC™. Return MOVES[®] SLC™ for servicing.



Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
61. O2 reading may be biased low	Low			O2 reading determined to be less than 19.4% when sampling air (20.9% O2) during periodic O2 calibration check	 Ensure there are no obstructions to the gas calibration port on the patient monitor panel. Ensure environment in which MOVES[®] SLC™ is operating contains 20.9% O₂ and is free of impurities. Power cycle MOVES[®] SLC™ and wait for re-calibration. If unsuccessful, move MOVES[®] SLC™ into another area, power cycle MOVES[®] SLC™ and wait for re-calibration. Return MOVES[®] SLC™ for servicing
62. O2 reading may be biased high	High			O2 reading determined to be greater than 22.4% when sampling air (20.9% O2) during periodic O2 calibration check	 Ensure there are no obstructions to the gas calibration port on the patient monitor panel. Ensure environment in which MOVES® SLC™ is operating contains 20.9% O₂ and is free of impurities. Power cycle MOVES® SLC™ and wait for re-calibration. If unsuccessful, move MOVES® SLC™ into another area, power cycle MOVES® SLC™ and wait for re-calibration. Return MOVES® SLC™ for servicing.
63. Barometer fault	Low			Mismatch between O2 and CO2 cell pressures and barometric pressure when sample pump is not running.	 Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.



Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
64. Altitude above system limit	Medium			Barometric pressure is less than 45 kPa.	 Proceed to a lower altitude. If erroneous, return MOVES[®] SLC™ for servicing.
65. Altitude below system limit	Medium			Barometric pressure is greater than 110 kPa.	 Proceed to a higher altitude. If erroneous, return MOVES[®] SLC™ for servicing.
66. Inspired O2 low	High			Inspired O ₂ is less than 19% (while running in ventilate mode) OR Inspired O ₂ is less than 82% (while running in O ₂ supplement mode).	 Ventilate Mode Ensure sample line with Nafion and filters is connected to system and breathing circuit. Completely replace the sample line including Nafion and filters. Power cycle the system and run the sample line system tests and confirm that they pass. Return for servicing. Supplement Mode Ensure sample line with Nafion and filters is connected to system and O2 Outlet Sampling Adaptor. Ensure the O2 Outlet Sampling Adaptor is properly connected to the O2 Outlet port. Completely replace the sample line including Nafion and filters. Replace the O2 Outlet Sampling Adaptor. Power cycle the system and run the sample line system tests and confirm that they pass. Return for servicing.



Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
67. Inspired O2 above target	Low			Inspired O ₂ exceeds Vent O ₂ setting by 15% (ventilation mode only).	NOTE: If Vent O ₂ setting was recently lowered by more than 15% after breathing circuit reached Vent O ₂ setting level then this alarm will appear until patient consumes excess oxygen (O ₂). 1. Check for related alarms and follow actions if required. 2. Clinical intervention may be required.
68. Inspired O2 below target	Low			Inspired O ₂ is not within 5% of Vent O ₂ setting while running in ventilation mode.	 Check for related alarms and follow actions if required. Clinical intervention may be required. Return MOVES[®] SLC™ for servicing.
69. Low expired CO2	High if less than 15 mmHg Low otherwise			When the apnea (capnograph) alarm is not active, expired CO ₂ partial pressure is less than or equal to the operator set low expired CO ₂ alarm limit. NOTE: This alarm will not appear when operating in O ₂ supplement mode.	 Check for related alarms and follow actions if required. For patients under 30 kg or with tidal volumes under 150 mL, ensure that the pediatric breathing system filter is used (P/N 125245). Clinical intervention may be required.
70. High expired CO2	Medium			Expired CO ₂ partial pressure is greater than or equal to the operator set high expired PCO ₂ alarm limit. NOTE: This alarm will not appear when operating in O2 supplement mode.	 Check for related alarms and follow actions if required. For patients under 30 kg or with tidal volumes under 150 mL, ensure that the pediatric breathing system filter is used (P/N 125245). Clinical intervention may be required.



Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
71. High inspired CO2 Change cartridge	High if greater than or equal to 10 mmHg Low otherwise			While running in ventilate mode, inspired CO ₂ partial pressure is greater than 6 mmHg.	 Replace ventilator cartridge. Return MOVES[®] SLC™ for servicing.
72. Low SpO2	Low until SpO2 Alarm Delay time finished or SpO2 Alarm Rapid Desat threshold reached, then High			O ₂ saturation level is less than or equal to user-set low SpO ₂ alarm limit.	 Check for related alarms and follow actions if required. If in O2 supplementation mode, physically check for an oxygen flow leaving the port. Clinical intervention may be required.
73. Low Pleth Variability Index	Low			Pleth Variability Index (PVI) less than or equal to the operator set low PVI alarm limit. NOTE: This alarm will not appear if PVI Display is set to OFF on Advanced Screen.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
74. High Pleth Variability Index	Medium			Pleth Variability Index (PVI) greater than or equal to the operator set high PVI alarm limit. NOTE: This alarm will not appear if PVI Display is set to OFF on Advanced Screen.	 Check for related alarms and follow actions if required. Clinical intervention may be required.



Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
75. High SpMet	HIGH, if SpMet reading is greater than or equal to 6.0%, LOW otherwise			SpMet greater than or equal to the operator set high SpMet alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
76. High SpCO	High			SpCO greater than or equal to the operator set high SpCO alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
77. Low SpHb	High			SpHb less than or equal to the operator set low SpHb alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
78. High SpHb	Medium			SpHb greater than or equal to the operator set high SpHb alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
79. Low SpOC	High			SpOC less than or equal to the operator set low SpOC alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
80. High SpOC	Medium			SpOC greater than or equal to the operator set high SpOC alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
81. Apnea (Capnograph)	High			Respiratory rate is less than or equal to 4 breaths per minute and the sample line occlusion alarm is not active. NOTE: This alarm will not appear when operating in O2 supplement mode.	 Check for related alarms and follow actions if required. For patients under 30kg or with tidal volumes under 150mL, ensure that the pediatric breathing system filter is used (P/N 125245). Clinical intervention may be required.



Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
82. Apnea (Ventilator)	High			While running in CPAP+PS ventilator mode with Apnea Backup enabled, ventilator fails to see inspiratory efforts from patient within period defined by <i>Frequency</i> setting and is now delivering backup breaths based on the selected ventilator settings.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
83. Apnea (Ventilator and Capnograph)	High			Both Apnea (Capnograph) and Apnea (Ventilator) alarm causes are present at the same time. NOTE: This alarm will not appear when operating in O2 supplement mode.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
84. Low RR	Medium			Respiratory rate is less than or equal to operator set low RR alarm limit when apnea alarm is not active and respiratory rate is greater than zero. NOTE: This alarm will not appear when operating in O2 supplement mode.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
85. High RR	Medium			Respiratory rate is greater than or equal to operator set high RR alarm limit. NOTE: This alarm will not appear when operating in O2 supplement mode.	 Check for related alarms and follow actions if required. Clinical intervention may be required.



Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
86. High pulse rate	High, if heart rate is greater than or equal to 150 BPM, Low otherwise,			Heart rate is greater than operator set high HR alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
87. Low pulse rate	High			Heart rate is less than operator set low HR alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
88. Cannot reach set I/E ratio	Low		Inhibitable	Ventilator operational settings and/or patient parameters result in operator set I:E ratio not being achieved due to inadequate inspiratory time.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
89. High Time too long for Frequency	Low	Locked		In APRV ventilation mode High Time setting duration is too long for the chosen Frequency setting (i.e. insufficient low time). Frequency setting takes priority and the system uses the maximum high time which allows a minimum low time of 0.3 seconds.	 If Frequency setting is unintentionally set too high then reduce setting so that High Time setting properly fits inside cycle period while allowing a minimum low time of 0.3 seconds. If the High Time setting is unintentionally too long then reduce to appropriate set point.
90. Inspire Time too long for Frequency	Low	Locked		In SIMV ventilation mode Inspire Time setting duration is too long for the chosen Frequency setting (i.e. insufficient expire time). Frequency setting takes priority and the system uses the maximum inspire time which allows a minimum expire time of 0.3 seconds.	 If Frequency setting is unintentionally set too high then reduce setting so that Inspire Time setting properly fits inside breath period while allowing a minimum expire time of 0.3 seconds. If the Inspire Time setting is unintentionally too long then reduce to appropriate set point.



Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
91. Above target volume	Medium			Exhaled tidal volume is at least 15% greater than the current operator set point and patient circuit occlusion alarm is not active.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
92. Below target volume	Medium			Exhaled tidal volume is at least 15% less than the current operator set point and patient circuit occlusion alarm is not active.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
93. High expired tidal volume	Medium			Exhaled tidal volume is 30% greater than the delivered inspired tidal volume.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
94. NIBP: Low systolic pressure	High		Inhibitable	Systolic pressure of NIBP measurement is less than or equal to the operator set low systolic BP alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
95. NIBP: High systolic pressure	High, if systolic pressure is greater than or equal to operator set high systolic BP alarm limit plus 40 mmHg, Medium otherwise.		Inhibitable	Systolic pressure of NIBP measurement is greater than or equal to the operator set high systolic BP alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
96. NIBP: Low diastolic pressure	Medium		Inhibitable	Diastolic pressure of NIBP measurement is less than or equal to 40 mmHg.	 Check for related alarms and follow actions if required. Clinical intervention may be required.



Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
97. NIBP: High diastolic pressure	High, if diastolic pressure is greater than or equal to 120 mmHg, Medium otherwise.		Inhibitable	Diastolic pressure of NIBP measurement is greater than or equal to 100 mmHg.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
98. NIBP inflate failure	Low		Inhibitable	Pressure required for inflation cannot be generated in the NIBP patient arm cuff.	 Ensure NIBP patient arm cuff is full connected to MOVES[®] SLC™ and has no leaks. Replace NIBP patient arm cuff. Power cycle MOVES[®] SLC™. Return MOVES[®] SLC™ for servicing.
99. NIBP could not get reading	Low		Inhibitable	NIBP measurement was not obtained unrelated to NIBP patient arm cuff inflation failure.	 Attempt another NIBP reading. Ensure NIBP patient arm cuff is full connected to MOVES[®] SLC™ and has no leaks. Replace NIBP patient arm cuff. Power cycle MOVES[®] SLC™. Return MOVES[®] SLC™ for servicing.
100. NIBP set to: Manual update	Low	Locked & Latching	Inhibitable	NIBP measurement was not obtained three consecutive times when set to take automatic measurements.	 Attempt another NIBP reading. Ensure NIBP patient arm cuff is full connected to MOVES[®] SLC[™] and has no leaks. Replace NIBP patient arm cuff. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
101. NIBP fault	Low			NIBP module internal fault or NIBP module communication failure with patient monitor.	 Power cycle MOVES[®] SLC™. Attempt another NIBP reading. Return MOVES[®] SLC™ for servicing.



Α	larm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
102.	NIBP Mode changed STAT -> 5 min	Low	Locked & Latching		NIBP has been set to STAT mode for 15 minutes and has automatically changed to take readings every 5 minutes for long term monitoring.	No action required. NIBP mode setting can be changed back to STAT at any time if required.
103.	IP zero required	Low		Inhibitable	At least one invasive channel has the following set of conditions: a) invasive channel is connected b) zero is required c) channel not currently zeroing d) zero failed is not active on the channel	 Zero all invasive channels which require zeroing or disconnect IP cable(s) if not in use. Fully disconnect then reconnect IP patient connection and zero IP transducer as prompted. Replace IP patient connection. Check integrity of interface with patient. Power cycle MOVES® SLC™. Return MOVES® SLC™ for servicing.
104.	IP1: Measurement error	Low			IP measurement contains values that are outside of the measurement range of the sensor.	 Fully disconnect then reconnect IP patient connection and zero IP transducer as prompted. Replace IP patient connection. Check integrity of interface with patient. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
105.	IP1 disconnect	Low		Inhibitable	IP patient connection is disconnected from system after previously being connected.	 Fully disconnect then reconnect IP patient connection. Replace IP patient connection. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.



А	larm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
106.	IP1: Zero failed	Low		Inhibitable	IP one-point zero calibration has failed.	 Ensure transducer is ready to zero (i.e., not connected to patient). Fully disconnect then reconnect IP patient connection and re-zero IP transducer as prompted. Replace IP patient connection. If available, use a different IP channel on the MOVES[®] SLC[™]. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
107.	IP1-ABP: Low systolic pressure	High			Systolic pressure of ABP measurement is less than or equal to the operator set low systolic BP alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
108.	IP1-ABP: High systolic pressure	High, if systolic pressure is greater than or equal to operator set high systolic BP alarm limit plus 40 mmHg, Medium otherwise.			Systolic pressure of ABP measurement is greater than or equal to the operator set high systolic BP alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
109.	IP1-ABP: Low diastolic pressure	Medium			Diastolic pressure of ABP measurement is less than or equal to 40 mmHg.	 Check for related alarms and follow actions if required. Clinical intervention may be required.



Α	larm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
110.	IP1-ABP: High diastolic pressure	High, if diastolic pressure is greater than or equal to 120 mmHg, Medium otherwise.			Diastolic pressure of ABP measurement is greater than or equal to 100 mmHg.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
111.	IP1: Low CVP	High			CVP measurement is less than or equal to the operator set low CVP alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
112.	IP1: High CVP	Medium			CVP measurement is greater than or equal to the operator set high CVP alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
113.	IP1: Low ICP	High			ICP measurement is less than or equal to the operator set low ICP alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
114.	IP1: High ICP	Medium			ICP measurement is greater than or equal to the operator set high ICP alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
115.	IP2: Measurement error	Low			IP measurement contains values that are outside of the measurement range of the sensor.	 Fully disconnect then reconnect IP patient connection and zero IP transducer as prompted. Replace IP patient connection. Check integrity of interface with patient. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.



A	larm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
116.	IP2 disconnect	Low		Inhibitable	IP patient connection is disconnected from system after previously being connected.	 Fully disconnect then reconnect IP patient connection. Replace IP patient connection. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
117.	IP2: Zero failed	Low		Inhibitable	IP one-point zero calibration has failed.	 Ensure transducer is ready to zero (i.e., not connected to patient). Fully disconnect then reconnect IP patient connection and re-zero IP transducer as prompted. Replace IP patient connection. If available, use a different IP channel on the MOVES[®] SLC™. Power cycle MOVES[®] SLC™ for servicing.
118.	IP2-ABP: Low systolic pressure	High			Systolic pressure of ABP measurement is less than or equal to the operator set low systolic BP alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
119.	IP2-ABP: High systolic pressure	High, if systolic pressure is greater than or equal to operator set high systolic BP alarm limit plus 40 mmHg, Medium otherwise.			Systolic pressure of ABP measurement is greater than or equal to the operator set high systolic BP alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.



Α	larm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
120.	IP2-ABP: Low diastolic pressure	Medium			Diastolic pressure of ABP measurement is less than or equal to 40 mmHg.	Check for related alarms and follow actions if required. Clinical intervention may be required.
121.	IP2-ABP: High diastolic pressure	High, if diastolic pressure is greater than or equal to 120 mmHg, Medium otherwise.			Diastolic pressure of ABP measurement is greater than or equal to 100 mmHg.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
122.	IP2: Low CVP	High			CVP measurement is less than or equal to the operator set low CVP alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
123.	IP2: High CVP	Medium			CVP measurement is greater than or equal to the operator set high CVP alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
124.	IP2: Low ICP	High			ICP measurement is less than or equal to the operator set low ICP alarm limit.	Check for related alarms and follow actions if required. Clinical intervention may be required.
125.	IP2: High ICP	Medium			ICP measurement is greater than or equal to the operator set high ICP alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
126.	IP3: Measurement error	Low			IP measurement contains values that are outside of the measurement range of the sensor.	 Fully disconnect then reconnect IP patient connection and zero IP transducer as prompted. Replace IP patient connection. Check integrity of interface with patient. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.



Al	arm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
127.	IP3 disconnect	Low		Inhibitable	IP patient connection is disconnected from system after previously being connected.	 Fully disconnect then reconnect IP patient connection. Replace IP patient connection. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
128.	IP3: Zero failed	Low		Inhibitable	IP one-point zero calibration has failed.	 Ensure transducer is ready to zero (i.e., not connected to patient). Fully disconnect then reconnect IP patient connection and re-zero IP transducer as prompted. Replace IP patient connection. If available, use a different IP channel on the MOVES[®] SLC™. Power cycle MOVES[®] SLC™. Return MOVES[®] SLC™ for servicing.
129.	IP3-ABP: Low systolic pressure	High			Systolic pressure of ABP measurement is less than or equal to the operator set low systolic BP alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
130.	IP3-ABP: High systolic pressure	High, if systolic pressure is greater than or equal to operator set high systolic BP alarm limit plus 40 mmHg, Medium otherwise.			Systolic pressure of ABP measurement is greater than or equal to the operator set high systolic BP alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
131.	IP3-ABP: Low diastolic pressure	Medium			Diastolic pressure of ABP measurement is less than or equal to 40 mmHg.	 Check for related alarms and follow actions if required. Clinical intervention may be required.



Α	larm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
132.	IP3-ABP: High diastolic pressure	High, if diastolic pressure is greater than or equal to 120 mmHg, Medium otherwise.			Diastolic pressure of ABP measurement is greater than or equal to 100 mmHg.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
133.	IP3: Low CVP	High			CVP measurement is less than or equal to the operator set low CVP alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
134.	IP3: High CVP	Medium			CVP measurement is greater than or equal to the operator set high CVP alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
135.	IP3: Low ICP	High			ICP measurement is less than or equal to the operator set low ICP alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
136.	IP3: High ICP	Medium			ICP measurement is greater than or equal to the operator set high ICP alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
137.	High airway pressure warning	Medium			Peak inspiratory pressure (PIP) exceeds operator set high airway pressure warning alarm limit without reaching operator set high airway pressure release alarm limit.	 Check for related alarms and follow actions if required. Clinical intervention may be required.



А	larm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
138.	High airway pressure release	Medium	Latching		Peak inspiratory pressure (PIP) exceeds operator set high airway pressure release alarm limit.	 Ensure the high airway pressure release alarm limit is set appropriately for the patient's physiological condition and the ventilator configuration. Check for related alarms and follow actions if required. Clinical intervention may be required.
139.	Low pressure end of inspire	Medium	Latching		Peak inspiratory pressure (PIP) does not reach 10 cmH ₂ O.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
140.	Negative airway pressure	Low	Latching		Airway pressure is less than 0 cmH ₂ O when ventilating in any mode.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
141.	Target pressure not reached	Medium			Peak inspiratory pressure (PIP) is ± 5 cmH ₂ O or greater from target when ventilating in a pressure control mode other that APRV and CPAP+PS.	 Check for related alarms and follow actions if required. Clinical intervention may be required.
142.	Expiratory obstruction	High			Positive end expiratory pressure exceeds operator set positive end expiratory pressure (PEEP) by 5 cmH ₂ O or greater.	 Ensure breathing circuit patient tubing is not kinked or obstructed. Ensure expiratory pathway (ventilator cartridge and valve block) is not obstructed. Replace breathing circuit including ventilator cartridge. Return MOVES[®] SLC[™] for servicing.
143.	Insp and exp pressure similar	High			Positive end expiratory pressure is within 10 cmH ₂ O of peak inspiratory pressure (PIP) when ventilating in IMV, SIMV and A/C modes.	 Check for related alarms and follow actions if required. Clinical intervention may be required.



A	arm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
144.	Sample line occlusion	Medium			Pressure at O2/CO2 sensor cells is below expected value compared to barometer (above normal resistance when sampling).	 Ensure sample line has no kinks or obstructions. Replace sample line. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
145.	Sample line disconnected	Medium			Pressure at O ₂ /CO ₂ sensor cells is above expected value compared to barometer (below normal resistance when sampling).	 Ensure sample line is fully connected to MOVES[®] SLC[™] and has no leaks. Ensure that Nafion tubing and sample line filters are installed in the sample line. Replace sample line and filters. Replace Nafion tubing. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
146.	Patient circuit occlusion	Medium			Inspired tidal volume is at least 15% less than the current operator set point, and the peak inspiratory pressure (PIP) exceeds operator set high airway pressure release alarm limit.	 Ensure breathing circuit patient tubing is not kinked or obstructed. Ensure inspiratory pathway (ventilator cartridge and valve block) is not obstructed. Replace breathing circuit including ventilator cartridge. Return MOVES[®] SLC[™] for servicing.



Alarm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
147. Patient circuit disconnect	High			 While ventilating, any of the following conditions occur: a) Expiratory flow never exceeds inspiratory flow for 15 seconds. b) With ventilation modes which include mandatory volume controlled breaths, the following occurs for the most recent mandatory volume controlled breath: the inspired tidal volume is greater than the target tidal volume, and the expired tidal volume is less than 25 mL, and the difference between PIP and PEEP is less than 6 cmH₂O. c) With ventilation modes which include mandatory pressure controlled breaths, the following occurs for the most recent mandatory pressure controlled breath: the PIP was less than the target pressure by at least 1 cmH₂O, and the expired tidal volume is less than 25 mL, and the difference between PIP and PEEP is less than 6 cmH₂O. 	 Ensure all breathing circuit connections (including patient tubing, Y-piece and sample line) are tight and no leaks are detected. Ensure the ventilator cartridge door is properly closed and latched. Replace breathing circuit. Return MOVES® SLC™ for servicing.



Α	larm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
148.	Concentrator air intake filter blocked	High			Pressure at concentrator air intake filter exceeds expected value (above normal resistance when concentrator in use).	 Check for and remove any obstructions from intake filter (i.e., hydrocarbon filter). Replace intake filter. Return MOVES[®] SLC[™] for servicing.
149.	Concentrator air intake filter missing	Low			Pressure at concentrator air intake filter is below expected value (below normal resistance when concentrator in use).	 Ensure intake filter (i.e., hydrocarbon filter) is completely secured to MOVES[®] SLC[™]. Replace intake filter. Return MOVES[®] SLC[™] for servicing.
150.	Patient gas sampling system failure	Medium			Patient gas sampling pump flow is insufficient for proper patient gas monitoring.	 Check for related alarms and follow actions if required. Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
151.	Unexpected system reset	Medium	Locked	Inhibitable	System suffered unexpected power loss and power was restored within 3 minutes.	 Ensure MOVES[®] SLC[™] has at least two sources of power present at all times. Replace exhausted battery with spare charged battery.
152.	Safe Gas Mode	Low	Locked		System will revert to "Safe Gas Mode" if any of the following alarms are active while running in ventilation mode: 1. CO2/O2 sensors warming up 2. O2 sensor fault 3. CO2 sensor fault 4. Gas sensor pressure fault 5. Inspired O2 low 6. Sample line disconnect 7. Sample line occlusion 8. Patient monitor failure	 See actions for relevant alarms. NOTE: Do not continuously run MOVES[®] SLC™ in Safe Gas Mode. Safe Gas Mode is only intended for short term use to complete a transport.



Α	larm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
153.	Alarm indication failure	Medium	Locked		Communication failure with Alarm board, which controls alarm audio/visual indication.	 Power cycle MOVES[®] SLC[™]. Return MOVES[®] SLC[™] for servicing.
154.	CO2 / O2 sensors warming up	Message	Locked		O ₂ and CO ₂ sensors are warming up.	Wait up to 5 minutes for system to become ready for use. Warming time is dependent of system/environment temperature.
155.	**SYSTEM STOPPED** Reconfigure NOW	High	Locked	Inhibitable	System auto resumes (i.e., recovers from short power loss) and is unable to recover patient settings previously in use. Therefore, the system cannot resume patient treatment. Patient settings return to defaults and require immediate reconfiguration to resume patient treatment. NOTE: If previously ventilating, this alarm means that System Mode has become Monitor Only and ventilation has ceased. It is imperative that ventilator settings be re-configured and the System Mode set to Ventilate, or an alternative means of ventilation be utilized.	 Reconfigure patient settings immediately and resume patient treatment. Ensure adequate backup battery power is present. If occurs repeatedly return MOVES[®] SLC™ for servicing.



А	larm / Message	Priority	Locked / Latching	Inhibitable	Causes	Action(s)
156.	Remote Screen disconnect	Medium	Locked	Inhibitable	Remote Screen was connected and became disconnected. NOTE: When this alarm becomes active, alarm audio pause or alarm audio disable will clear (i.e., alarm audio will resume) on both MOVES® SLC™ and Remote Screen.	 Wired Connection Check cable connection if using wired connection. Confirm that both devices are still ON. Disconnect and reconnect cable connection. Replace connection cable. Try alternate Remote Screen with MOVES® SLC™. Try alternate MOVES® SLC™ with Remote Screen. Return faulty device(s) for servicing. Wireless Connection Confirm that both devices are still ON and that disconnect was not made intentionally from other side. Try to reconnect devices. Try alternate Remote Screen with MOVES® SLC™. Try alternate MOVES® SLC™ with Remote Screen. Try moving devices to another area with less radio interference. Return faulty device(s) for servicing.



15.8 SYSTEM TEST FAILURE MESSAGES AND CAUSES

Table 42: System Test Failure Messages and Causes

System Prompt	Causes/Actions
1. The ventilator is not responding correctly.	Displayed if ventilator fails to respond appropriately during initialization for testing.
	 Power cycle and repeat tests.
	If problem persists, return for servicing.
2. Refer unit for service.	Default "catch-all" for serious system malfunctions.
	 Power cycle and repeat tests.
	If problem persists, return for servicing.
3. Ensure that the Y-Piece is open and cartridge and	Failure of open Y-piece ventilator test.
tubing are not obstructed or damaged.	 Confirm that the breathing circuit is connected to the system correctly.
	Confirm Y-piece is open and circuit has no leaks or occlusions, then repeat tests.
	 Confirm that the ventilator blower inlet is not occluded in any way.
	 Replace breathing circuit, then repeat tests.
	Power cycle and repeat tests.
	6. If problem persists, return for servicing.
4. Ensure that hydrocarbon filter is not occluded. If problem persists, refer unit for service.	Concentrator detects occluded HC filter during open Y-piece test.
	 Confirm HC filter is not occluded, then repeat tests.
	 Remove HC filter and confirm there are no occlusions to concentrator pump inlet, then repeat tests.
	Replace HC filter and repeat tests.
	Power cycle and repeat tests.
	5. If problem persists, return for servicing.
5. Ensure that the hydrocarbon filter is present. If problem persists, refer unit for service.	Concentrator detects missing HC filter during open Y-piece test.
	 Confirm HC filter is installed and tightly secured, then repeat tests.
	Replace HC filter and repeat tests.
	Power cycle and repeat tests.
	4. If problem persists, return for servicing.



System Prompt	Causes/Actions
6. Ensure that Y-Piece is disconnected from the patient and kept stationary during test.	Failure of open Y-piece ventilator test, specifically the flow accuracy of the air pump.
	Confirm that the breathing circuit is connected to the system correctly.
	Confirm Y-piece is open and circuit has no leaks or occlusions, then repeat tests ensuring that the Y-piece and breathing tubes remain stationary during the test.
	Confirm that the air pump inlet is not occluded in any way.
	Replace breathing circuit, then repeat tests.
	Power cycle and repeat tests.
	If problem persists, return for servicing.
7. Ensure the test plug is inserted into the Y-Piece and the tubing is kept stationary during test.	Failure of closed Y-piece ventilator test, specifically the detection of flow when zeroing the flow meters and pressurizing the breathing circuit.
	 Confirm that the ventilator cartridge door is properly closed and latched.
	Confirm that the breathing circuit is connected to the system correctly.
	 Confirm Y-piece is securely blocked by the test plug, and that the circuit has no leaks, then repeat tests ensuring that the Y-piece and breathing tubes remain stationary during the test.
	Replace breathing circuit, then repeat tests.
	Power cycle and repeat tests.
	If problem persists, return for servicing.
8. Ensure the test plug is inserted into the Y-Piece, the ventilator cartridge door is closed and the circuit has	Failure of closed Y-piece ventilator test, specifically the pressurization of the ventilator and ventilator circuit.
no leaks.	 Confirm that the ventilator cartridge door is properly closed and latched.
	Confirm that the breathing circuit is connected to the system correctly.
	 Confirm that the test plug is securely inserted into the Y-piece and that the circuit has no leaks, then repeat tests.
	 Confirm that the ventilator driving gas inlet is not obstructed in any way.
	Replace breathing circuit, then repeat tests.
	6. Power cycle and repeat tests.
	7. If problem persists, return for servicing.
Communication problem with patient monitoring subsystem.	Displayed if there is a communication fault with patient monitoring subsystem.
	Power cycle and repeat tests.
	If problem persists, return for servicing.



System Prompt	Causes/Actions
10. Unable to verify clear sample line. Ensure sample line is not kinked or blocked, and occluding cap is	Displayed if unable to establish a stable clear-line condition during Sample Line testing.
removed.	 Check sample line for occlusions and repeat tests.
	Replace sample line and repeat tests.
	Remove sample line and repeat tests with sample port open.
	Power cycle and repeat tests.
	If problem persists, return for servicing.
11. Patient gas sensor fault occurred during test.	Displayed if O2 or CO2 sensor suddenly becomes not- ready during the execution of the test.
	 Power cycle and repeat tests.
	If problem persists, return for servicing.
12. Patient gas sample pump fault occurred during test.	Displayed if timed-out waiting for Sample Pump to enter the specific mode for occlusion-testing.
	 Power cycle and repeat tests.
	If problem persists, return for servicing.
13. Patient gas sensor fault. Unable to start test.	Displayed if unable to establish stable O2/CO2 sensor- readiness during setup for Sample Line testing.
	 Power cycle and repeat tests.
	If problem persists, return for servicing.
14. Ensure that the sample line is connected to the MOVES [®] SLC™ sample port and the cap on the	Displayed if unable to confirm stable occlusion condition during Sample Line testing.
Nafion [®] tubing is firmly connected to the end of the sample line.	 Confirm that sample line occlusion cap is in place.
	Check sample line connection to sample port.
	 Confirm that all sample line components are tightly connected and not leaking.
	 Replace sample line and repeat tests.
	Remove sample line and repeat tests with sample port occluded.
	Power cycle and repeat tests.
	7. If problem persists, return for servicing.



System Prompt	Causes/Actions
15. The system requires a backup source of power. Ensure that either wall power is present and a charged backup battery is installed or that two charged batteries are installed.	Displayed if the backup power source verification test fails to detect the presence of redundant power. 1. Confirm that at least 2 power sources are connected, and that any batteries installed have charge.
	 Confirm the presence of the power sources connected on the user interface. Ensure all battery doors are properly closed and latched.
	 4. Disconnect and reconnect any connected power sources and repeat tests. 5. Power cycle and repeat tests. 6. If problem persists, return for servicing.
16. The power source monitoring system is not operating correctly.	Displayed if there is a communication fault with the Power Manager. 1. Power cycle and repeat tests. 2. If problem persists, return for servicing.
17. Operator verification of alarm lights and/or audio failed.	Displayed if the lights/audio verification test fails (i.e., user responds 'No'). If alarm lights or audio is not functioning: 1. Select test and repeat. 2. Power cycle and repeat tests. 3. If problem persists, return for servicing.
18. The test timed-out.	Displayed if during ventilator open Y-Piece or closed Y-Piece tests, test progress stalls unexpectedly. 1. Select test and repeat. 2. Power cycle and repeat tests. 3. If problem persists, return for servicing.
19. You elected to skip this test, or the test cannot be performed due to other reported faults.	Displayed if the test is skipped either by the user explicitly or because of test dependency failures. 1. Select test and repeat. 2. Power cycle and repeat tests without skipping. 3. If problem persists, return for servicing.

15.9 SAFE GAS MODE



WARNING! THE MOVES® SLC $^{\text{TM}}$ SHOULD NOT BE RUN CONTINUOUSLY IN SAFE GAS MODE. SAFE GAS MODE IS INTENDED FOR SHORT TERM USE ONLY TO COMPLETE TRANSPORTS.



NOTE: The MOVES® SLCTM will enter Safe Gas Mode only when in <u>Ventilate</u> mode.



The system automatically enters SGM whenever the system cannot rely on the oxygen (O₂₎ and/or carbon dioxide (CO₂) value that it is sensing. It is also used (when ventilating) if inspired CO₂ becomes too high due to an exhausted CO₂ scrubber ventilator cartridge.



Figure 15-8: Status Bar Showing System in Safe Gas Mode

SGM can occur for several reasons (each of which generates a unique alarm message):

- Sample line is occluded or disconnected for more than 60 seconds.
- O2 or CO2 sensor faults and cannot get a reading for more than 60 seconds.
- Operating the system for more than 60 seconds while the O₂ or CO₂ sensor is still reporting that it is warming up.
- CO₂/O₂ pressure sensor fault for more than 60 seconds.
- Patient monitoring board fault for more than 60 seconds.
- Inspired O₂ is below 19%.
- System is ventilating, and a high PiCO₂ alarm is active at high priority (PiCO₂ ≥ 10 mmHg). System exits SGM when there is no high PiCO₂ warning (PiCO₂ ≤ 6 mmHg), and no other SGM conditions are active.
- O2 sensor readings are suspected of being biased high.

Standard Safe Gas Mode

If the Ventilator target O_2 setting is $\leq 40\%$:

• Safe Gas Mode will normally deliver approximately 5.5 to 6 LPM of 40–50% O2.

High O₂ Safe Gas Mode

If the Ventilator target O₂ setting is > 40%:

SGM delivers 2.5 LPM of 90% O2 (nominal).



NOTE: If the concentrator is in a degraded mode, or in a fault mode when operating in Safe Gas Mode, the system will deliver at least 9.5 LPM of air in addition to any oxygen it can produce.



16.0 Appendix A

16.1 SYSTEM DEFAULT SETTINGS

Table 43: System Default Settings

Setting	Default
Alarms	All alarms and audio enabled
System Mode	Monitor Only
Vent Mode	IMV
Vent Control	Pressure
Vt (not used by default vent mode PC-IMV)	500 ml
Frequency	10 B/M
PEEP	0–3 cmH ₂ O
I/E ratio	1:2
Inspire/High Time (not used by default vent mode PC-IMV)	1.0 second
Control Pressure	20 cmH₂O
Pressure Support (not used by default vent mode PC-IMV)	Off
Apnea Backup	Off
Vent O ₂	40%
NIBP Update	Manual
Suction	325 mmHg
ECG HR Mode	Adult or Pediatric depending on System Administration ECG HR Mode Default configuration
ECG Range	2.2 mV
ECG Sweep Speed	25 mm/sec
SpO2 Average Time	8s
SpO2 Sensitivity Mode	Normal
SpO2 Alarm Delay	15s
SpO2 Alarm Rapid Desat	-5%
PVI Display	On
PVI Average Mode	Long
SpHb Arterial/Venous Mode	Arterial
SpHb Average Mode	Long
ECG EMG Filter	On
High airway pressure warning alarm limit	32 cmH₂O
High airway pressure release alarm limit	40 cmH ₂ O
Low expired PCO₂ alarm limit	25 mmHg
High expired PCO ₂ alarm limit	55 mmHg



Setting	Default		
Low RR	5 B/M		
High RR	30 B/M		
Low systolic BP alarm limit	90 mmHg		
High systolic BP alarm limit	180 mmHg		
Low SpO₂ alarm limit	90%		
Low HR alarm limit	50 BPM		
High HR alarm limit	120 BPM		
Low CVP alarm limit	0 mmHg		
High CVP alarm limit	20 mmHg		
Low ICP alarm limit	0 cmH ₂ O		
High ICP alarm limit	27 cmH ₂ O		
Low PVI alarm limit	5%		
High PVI alarm limit	40%		
High SpMet alarm limit	3.0%		
High SpCO alarm limit	10%		
Low SpHb alarm limit	Dependent on System Administration SpHb units, either: • 8 g Hb / dL blood • 5.0 mmol Hb / L blood • 80 g Hb / L blood		
High SpHb alarm limit	Dependent on System Administration SpHb units, either: 17 g Hb / dL blood 11.0 mmol Hb / L blood 170 g Hb / L blood		
Low SpOC alarm limit	10 mL O ₂ / dL blood		
High SpOC alarm limit	25 mL O ₂ / dL blood		

16.2 SYSTEM CLEANING

The MOVES[®] SLC[™] is designed for easy maintenance. All exposed parts of MOVES[®] SLC[™] are corrosion resistant. The MOVES[®] SLC[™] device should be serviced after every three (3) months of use or after every twelve (12) months of storage.



CAUTION! DO NOT SUBMERGE THE MOVES $^{\otimes}$ SLC $^{\intercal}$ OR POUR CLEANING LIQUIDS OVER OR INTO THE MOVES $^{\otimes}$ SLC $^{\intercal}$.

The external body of the system can be cleaned using standard cleaning agents, excluding oxidizing agents. It is recommended that the external metal surfaces be wiped down with isopropyl alcohol during routine maintenance. Cables, NIBP cuffs and tubing can be cleaned with a disinfecting spray.

External surfaces of the MOVES[®] SLC™ can be wiped clean with one of the following:

- Isopropyl Alcohol
- Chlorine Compounds*
 - Maximum Concentration: 1:10



^{*}These compounds are diluted by volume in water.

For recommended MOVES[®] SLC[™] accessories not labeled as single use, refer to the cleaning instructions provided by the manufacturer.

16.3 SYSTEM MAINTENANCE

16.3.1 Replacing Filters

There are two user-serviceable components in the MOVES[®] SLC™ system:

- 1. Hydrocarbon Filter (P/N 100915)
- 2. Ventilator Filter (P/N 126504)

The hydrocarbon filter should be replaced when the system alarm indicates it is occluded. The ventilator filter should be replaced every three (3) months. For information on replacing the hydrocarbon filter see Section 9.7.3 Installing the Hydrocarbon Filter on page 98. For information on replacing the ventilator filter see Section 16.3.3 Replacing the Ventilator Filter on page 285.

16.3.2 End of Life

The MOVES[®] SLC™ ventilator has an expected service life in excess of 5 years and over 1000 operational hours when operated and serviced according to the manual. The minimum expected lifetime has been determined with the device at maximum oxygen concentration and default ventilation settings (Control Pressure=20 cmH₂O, Frequency=10 BPM, IE ratio of 1:2) which are considered a typical use case. Higher tidal volumes and ventilation at high pressures may reduce the service life of the device. In addition, any failure of the system self-test should result in servicing and replacement of components or determination of device end of life by manufacturer.

16.3.3 Replacing the Ventilator Filter

1. Locate the Ventilator Driving Gas Inlet.





2. Locate the lid release screw and unscrew it counterclockwise.



3. Lift the lid to access the filter.



4. The filter is shown in the photo at the right.



5. While holding the lid up, gently grip the filter and rotate it counterclockwise so that its projecting flange disengages from the filter clamps.



6. The flange is shown disengaged in the photo at the right.



7. Remove the filter.



8. Acquire a new filter. Note the three clearance cuts indicated in the photo at the right.



9. Place the filter down and align the clearance cuts on the filter so that the flanges are positioned to slide under the filter clamps.



10. Rotate the filter clockwise so that the flanges slide underneath the clamps.



A click will be felt when the filter is fully engaged.
 Visually, a tab will be touching each of the three
 clamps.

NOTE: If the filter is not properly seated the lid will not close. Do not force it. Unscrew the filter and then re-seat it.



16.3.4 General Maintenance



CAUTION! NO LUBRICANTS OTHER THAN THOSE RECOMMENDED BY THE MANUFACTURER SHALL BE USED ON THE MOVES[®] SLC™.

Regular maintenance and calibration should be carried out by authorized and qualified service personnel annually or after three (3) months of use. Systems should also be checked annually if not in use. In addition, if the MOVES[®] SLC™ system is subjected to extremely rough handling or environmental stress, or sustains damage, it should be referred to authorized and qualified service personnel for inspection and/or repair.





WARNING! ONLY AUTHORIZED SERVICE AND MAINTENANCE PERSONNEL SHOULD REMOVE ANY COVERS FROM MOVES[®]. UNAUTHORIZED REMOVAL OF COVERS FROM MOVES[®] SLC™ MAY RESULT IN ELECTRIC SHOCK AND POSSIBLY DEATH, AND MAY DAMAGE THE SYSTEM COMPONENTS.

16.4 ACCESSORIES MAINTENANCE

Reusable accessories should be regularly inspected for wear or damage. Inspect all cables and connections (especially the power cord) for signs of fraying or other damage. Keep accessories clean. Refer to original manufacturer's instructions for cleaning agents and procedures. Ensure that all gels and pastes are removed from electrode cables. Accessories that come in sanitary sealed packages should be inspected for damage to the sealed package before using. If the seal is broken, discard accessory. The label on the *package* of the Ventilator Cartridge contains an expiry date. Always check the expiry date on the Ventilator Cartridge package before using it to make sure that the cartridge has not expired. As well, monitor spare cartridges with regard to their remaining "shelf life".

16.5 CHECKING THE ACCURACY OF THE TEMPERATURE PROBE

To check the accuracy of the temperature probe:

- 1. Pour a glass of warm water. Using an external temperature probe, measure the temperature of the water. The water temperature needs to be between 28°C (82.4°F) and 42°C (107.6°F).
- 2. Place the MOVES[®] SLC[™] temperature probe in the water. Compare the temperature displayed by the MOVES[®] SLC[™] with temperature measured by external temperature probe (with at least 0.05°C accuracy).
- 3. The temperature displayed by the MOVES® SLC™ should not disagree with the temperature measured by the external probe by any more than the values listed in *Temperature Monitoring Specifications* on *page 302*.

16.6 MOVES® SLC™ SPECIFICATIONS

16.6.1 Model Number

The documentation in this manual is for MOVES® SLC™ model number 122752.



CAUTION! OPERATION OF MOVES[®] SLC™ OUTSIDE OF SPECIFIED LIMITS MAY CAUSE INACCURATE RESULTS.

16.6.2 Physical Properties

Table 44: Physical Properties of MOVES[®] SLC™

Property	Specification	Notes
Unit Weight (lbs)	37.45	Excludes batteries, accessories, options, cables, etc.
Battery Weight, ea. (lbs)	3.25	
Unit Length (in)	33	
Unit Width (in)	5.5-6.5	
Unit Height (in)	10.25	
Unit Exterior Housing Material	Aluminum	



Property	Specification	Notes
Operating Sound Level	< 70 dB	At a distance of 1 m level
Standards Compliance IEC 60601-1		
	IEC 60601-1-1	
	IEC 60601-1-2	
	IEC 60601-1-8	
	ISO 80601-2-12	
	ISO 80601-2-13	
	IEC 60601-2-27	
	IEC 80601-2-30	
	IEC 60601-2-34	
	IEC 60601-2-49	
	ISO 80601-2-55	
	ISO 80601-2-61	
	ISO 8359	
	ASTM E1112-00	
	BS EN 794-3 (2009)	
	MIL-STD-810G	
	JECETS	
WIFI Compliance	FCC (IDs are XF6-RS9113SB, XF6-RS9113DB)	
	IC (IDs are 8407A-RS9113SB, 8407A-RS9113DB)	
	CE/ETSI	
	TELEC	
	SRRC	
Device Classification	Class II, CF Defibrillation Proof	
Screen	115.2 mm (w) x 86.4 mm (h)	Resolution of 640 x 480 pixels



16.6.3 Oxygen Concentrator Specifications

Table 45: Oxygen Concentrator Specifications of MOVES[®] SLC™ in a Circle Circuit

Property	Specification			Notes
Time to FiO ₂	Temp (°C)	FiO ₂ (%)	Time (min.)	All figures measured with a 1 L test lung from
	22	50	2	cold start. With a patient, the FiO ₂ as measured on MOVES [®] SLC™ UI will depend
	22	70	4	on the patient's FRC (washout time) and their
	22	80	6	oxygen consumption rate.
	22	85	10	
	0	50	3.5	
	0	70	8	
	0	80	12	
	0	85	18	
	-10	50	5	
	-10	70	11	
	-10	80	15	
	-10	85	22	
Maximum Pressure	13.5 psi ± 10%	,		None

Table 46: Oxygen Concentrator Specifications of MOVES[®] SLC™ in O2 Supplement Mode

Property	Specification	1		Notes
Time to FiO ₂	Temp (°C)	FiO ₂ (%)	Time (min.)	None
	-25	87	36:36	
	-25	90	39:52	
	-10	87	10:32	
	-10	90	13:43	
	0	87	6:56	
	0	90	8:13	
	21	87	1:12	
	21	90	1:49	
	47	87	0:52	
	47	90	1:09	
	54	87	0:38	
	54	90	0:44	
Maximum Pressure	13.5 psi ± 10%	6		None
Output Concentration	> 87%			None



Property	Specification	Notes
Flow	@ 1 PSI (7 kPa) backpressure, O ₂ flow is measured as 2.4 LPM	None
	@ 0 PSI (0 kPa) backpressure, O ₂ flow is measured as 2.5 LPM	
Low O2 Alarm Threshold	< 82%	None

16.6.4 Ventilator Specifications and Vent Mode Definitions

IMV – INTERMITTENT MANDATORY VENTILATION

Pressure Controlled

Parameters: Frequency, I:E ratio, PEEP, Control Pressure

Triggers: Time only

<u>Description</u>: Produces a mandatory breath at the specified frequency, dividing the breath into an inspire and expire period as specified by the set I:E ratio. Inspiration starts at the specified PEEP pressure and linearly increases the pressure to the PEEP + Control Pressure over 90% of the inspire time. At the end of the inspire time, pressure is released down to the PEEP pressure, where it is maintained until the next start of inspiration.

Volume Controlled

Parameters: Frequency, I:E ratio, PEEP, Tidal Volume

Triggers: Time only

<u>Description</u>: Produces a mandatory breath at the specified frequency, dividing the breath into an inspire and expire period as specified by the set I:E ratio. Inspiration starts at the specified PEEP pressure and linearly increases the delivered volume to the specified tidal volume over the inspire time. At the end of the inspire time, pressure is released down to the PEEP pressure, where it is maintained until the next start of inspiration.

A/C - ASSIST/CONTROL

Pressure Controlled

Parameters: Frequency, I:E ratio, PEEP, Control Pressure, Trigger Sensitivity

<u>Triggers</u>: Time or pressure/flow

<u>Description</u>: Produces a mandatory breath at the specified frequency, or faster if triggered by patient pressure/flow. The breath is divided into an inspire and expire period as specified by the set I:E ratio. Inspiration starts at the specified PEEP pressure and linearly increases the pressure to the PEEP + Control Pressure over 90% of the inspire time. At the end of the inspire time, pressure is released down to the PEEP pressure, where it is maintained until the next start of inspiration, as determined by the set respiratory rate or a patient trigger.

<u>Triggering</u>: In normal sensitivity, a patient can trigger the start of inhalation with an inspiratory flow of at least 10 L/min or by causing a pressure drop of 3.0 cmH₂0 from PEEP. In low sensitivity, the flow must exceed 15 L/min or the pressure drops 6 cmH₂0 from PEEP.

Volume Controlled

Parameters: Frequency, I:E ratio, PEEP, Tidal Volume, Trigger Sensitivity



Triggers: Time or pressure/flow

<u>Description</u>: Produces a mandatory breath at the specified frequency, or faster if triggered by patient pressure/flow. The breath is divided into an inspire and expire period as specified by the set I:E ratio. Inspiration starts at the specified PEEP pressure and linearly increases the delivered volume to the specified tidal volume over the inspire time. At the end of the inspire time, pressure is released down to the PEEP pressure, where it is maintained until the next start of inspiration, as determined by the set respiratory rate or a patient trigger.

<u>Triggering</u>: In normal sensitivity, a patient can trigger the start of inspiration with an inspiratory flow of at least 10 L/min or by causing a pressure drop of 3.0 cmH₂0 from PEEP. In low sensitivity, the flow must exceed 15 L/min or the pressure drops 6 cmH₂0 from PEEP.

SIMV - SYNCHRONOUS INTERMITTENT MANDATORY VENTILATION

Pressure Controlled

Parameters: Frequency, Inspire Time, PEEP, Control Pressure, Trigger Sensitivity

Triggers: Time or pressure/flow

<u>Description</u>: Creates a timing window based on the specified frequency. The first patient trigger to occur in the time window will immediately initiate a mandatory breath, or if the timing window ends with no triggers, a mandatory breath will be taken. During a mandatory breath, inspiration starts at the specified PEEP pressure and linearly increases the pressure to the PEEP + Control Pressure over the specified inspire time. At the end of the inspire time, pressure is released down to the PEEP pressure, where it is maintained until the next start of inspiration. If additional patient triggers occur during the same timing window, the ventilator will allow the patient to breathe spontaneously, supporting the patient at the specified PEEP pressure.

<u>Triggering</u>: In normal sensitivity, a patient can trigger the start of inspiration with an inspiratory flow of at least 10 L/min or by causing a pressure drop of 3.0 cmH₂0 from PEEP. In low sensitivity, the flow must exceed 15 L/min or the pressure drops 6 cmH₂0 from PEEP.

Pressure Controlled + Pressure Support

Parameters: All the parameters of Pressure Controlled SIMV above plus Pressure Support.

<u>Description</u>: Operates similarly to Pressure Controlled SIMV, but additional patient triggered breaths in a time window will be supported to a pressure of PEEP + specified Pressure Support level.

Cycling (Pressure Support Breaths): In normal sensitivity, the breath will cycle back to PEEP level when the flow drops under 4 L/min. In low sensitivity, the breath will cycle to PEEP when the flow drops under 7 L/min.

Volume Controlled

Parameters: Frequency, Inspire Time, PEEP, Tidal Volume, Trigger Sensitivity

Triggers: Time or pressure/flow

<u>Description</u>: Creates a timing window based on the specified frequency. The first patient trigger to occur in the time window will immediately initiate a mandatory breath, or if the timing window ends with no triggers, a mandatory breath will be taken. During a mandatory breath, inspiration starts at the specified PEEP pressure and linearly increases the delivered volume to the specified tidal volume over the specified inspire time. At the end of the inspire time, pressure is released down to the PEEP pressure, where it is maintained until the next start of inspiration. If additional patient triggers occur during the same timing window, the ventilator will allow the patient to breathe spontaneously, supporting the patient at the specified PEEP pressure.



<u>Triggering</u>: In normal sensitivity, a patient can trigger the start of inspiration with an inspiratory flow of at least 10 L/min or by causing a pressure drop of 3.0 cmH₂0 from PEEP. In low sensitivity, the flow must exceed 15 L/min or the pressure drops 6 cmH₂0 from PEEP.

Volume Controlled + Pressure Support

Parameters: All the parameters of Volume Controlled SIMV above plus Pressure Support.

<u>Description</u>: Operates similarly to Pressure Controlled SIMV, but additional patient triggered breaths in a time window will be supported to a pressure of PEEP + specified Pressure Support level.

Cycling (Pressure Support Breaths): In normal sensitivity, the breath will cycle back to PEEP level when flow drops under 4 L/min. In low sensitivity, the breath will cycle to PEEP when flow drops under 7 L/min.

APRV – AIRWAY PRESSURE RELEASE VENTILATION

Parameters: Frequency, PEEP, Control Pressure, Inspire/High Time

Triggers: Time only

<u>Description</u>: Maintains a constant pressure at the specified PEEP + Control Pressure level for the specified High Time. At the end of the High Time, pressure is released down to the PEEP pressure, where it is maintained until the next start of inspiration as set by Frequency. The patient may breathe spontaneously at any time.

CPAP/PS - CONTINUOUS POSITIVE AIRWAY PRESSURE / PRESSURE SUPPORT

Parameters: PEEP, Pressure Support, Trigger Sensitivity, Apnea Backup

<u>Description</u>: Maintains a constant pressure at the specified PEEP level. When the patient inspiratory efforts exceed the flow trigger, increase the pressure to PEEP + Pressure Support until the inspiratory flow drops under the cycle level where the pressure will drop back to the specified PEEP level. If Pressure Support is set to Off, no triggering or cycling will occur.

<u>Triggering</u>: In normal sensitivity, a patient can trigger the start of inspiration with an inspiratory flow of at least 10 L/min or by causing a pressure drop of 3.0 cmH₂0 from PEEP. In low sensitivity, the flow must exceed 15 L/min or the pressure drops 6 cmH₂0 from PEEP.

Cycling (Pressure Support Breaths): In normal sensitivity, the breath will cycle back to PEEP level when flow drops under 4 L/min. In low sensitivity, the breath will cycle to PEEP when flow drops under 7 L/min.

Apnea Backup: When Pressure Support is on, the optional Apnea Backup mode will allow the additional parameters for IMV to be configured. The ventilator will take a mandatory IMV breath whenever the time period from the last trigger breath exceeds the time period as specified by 1 ÷ Frequency.

Table 47: Ventilator Specifications of MOVES® SLC™

Property	Specification
Tidal Volume	50–750 mL
Frequency	6–40 B/M
Inspiratory/Expiratory Ratio	1:1 to 1:3
Inspiratory Resistance	6 cmH ₂ O (at 60 LPM)
Expiratory Resistance	6 cmH ₂ O (at 60 LPM)



Property	Specification
Inspiratory Time	0.3–5.0 seconds
Peak Flow	60 LPM
Positive End Expiratory Pressure (PEEP)	0-20 cmH ₂ O
Positive Pressure Relief Valve (Mechanical)	70 cmH ₂ O
Control Pressure	10–55 cmH ₂ O (over PEEP). Control Pressure = PIP (Peak Inspiratory Pressure) – PEEP (Positive End Expiratory Pressure)
Pressure (Pw max)	Limited to 58 cmH ₂ O by software
Pressure Support Ventilation	Off, 5–40 cmH ₂ O
Apnea Backup (For CPAP+PS)	Off, On
Maximum Limited Pressure (Plim max)	100 cmH ₂ O (at 100 LPM)
Minimum (Sub atmospheric) Limited Pressure (Plim min)	- 6.0 cmH ₂ O (at 60 LPM)
Negative Pressure (Sub atmospheric) Available in the Expiratory Phase	None
Trigger Sensitivity (Normal)	10 LPM (flow) or 3 cmH ₂ O (pressure) below PEEP
Trigger Sensitivity (Low)	15 LPM (flow) or 6 cmH2O (pressure) below PEEP
External Oxygen Supply	15 LPM maximum (@ 2 psi minimum)
Modes	PC-IMV (default), VC-IMV, PC-A/C, VC-A/C, PC-SIMV, PC-SIMV+PS, VC-SIMV, VC-SIMV+PS, APRV, CPAP, CPAP+PS
Compressible Volume of Ventilator and Cartridge	1350 mL (NOTE: System compensates for compressible volume.)
Ventilator Circuit Compliance Including Cartridge	Approximately 0.7 mL/cmH ₂ O over the ventilator settable pressure range
Time to reach 90% FiO ₂ using external oxygen supplied into the O2 inlet port.	Tidal volume of 500 mL at 10 breaths/minute into a test lung with 5 cmH2O/L/s resistance and 50 mL/cmH2O compliance
	At oxygen flow of 5 LPM = 2:04 minutes
	At oxygen flow of 15 LPM = 1:36 minutes
	Tidal volume of 150 mL at 20 breaths/minute into a test lung with 20 cmH2O/L/s resistance and 20 mL/cmH2O compliance
	At oxygen flow of 5 LPM = 2:28 minutes
	At oxygen flow of 15 LPM = 2:00 minutes



Property	Specification
Ventilator Breathing System Filter characteristics (standard circuit with PALL Medical BB100E)	Internal Volume of 85 mL Resistance of 2.0 cmH2O at 60 LPM Airborne Bacterial/Viral Removal Efficiency of 99.999%
Ventilator Breathing System Filter characteristics (Pediatric filter P/N 125245)	Internal Volume of 35 mL Resistance of 3.6 cmH2O at 60 LPM Airborne Bacterial/Viral Removal Efficiency of 99.999%
Standards Compliance	ISO 80601-2-12, ISO 80601-2-13, EN 794-3

16.6.5 Suction Specifications

Property	Specification
Suction Vacuum	-100 to -325 mmHg
Free Flow Rate	20 LPM

16.6.6 Electrical Characteristics

Table 49: Electrical Specifications of MOVES[®] SLC™

Property	Specification	Notes
External Power	100-240 VAC, 50-60 Hz, 5.5 A max.	
Max Current Output	28 VDC, 14.3 A max.	Supplied by Power Supply / Battery Charger
Battery Type	25.9 V lithium polymer	
Charge Time (per set 2 batteries)	2.5 hrs when the system is idle	
Battery Life (per set 2 batteries)	Minimum 2.5 hrs on two fully charged new batteries at 101 kPa and 21°C. Typical > 4 hours with ventilator and monitors on, running concentrator with ¼ duty cycle.	Battery run time is highly dependent on use of the oxygen concentrator and / or suction.

16.6.7 WIFI Specifications

Property	Specification
WIFI Type	802.11n WIFI
Frequency Range	2402.0 – 2480.0 MHz
Output Watts	0.05188 W



16.6.8 Environmental Specifications

MOVES[®] SLC ™

MOVES[®] SLC[™] has been tested to function under the environmental conditions present during transport and battlefield use. These included mechanical testing for vibration, bump and shock as well as EMC and extreme temperature humidity and weather conditions. Details are provided in the table below:

Table 50: Environmental Specifications of MOVES $^{\circledR}$ SLC $^{\intercal M}$

Variable	Storage Condition	Operating Condition	
Temperature	-14°F to 140°F (-26°C to	-14°F to 129°F (-26°C to 54°C).	
	60°C) – system & batteries	Notes:	
		 MOVES[®] SLC[™] can be taken from any operating temperature into extreme cold of -26°C without affecting operation. It can be cold started at -26°C (- 14°F) on AC power. However, MOVES[®] SLC[™] cannot be cold started below -20°C (-4°F) on batteries. 	
		 When starting up MOVES[®] SLC™ in cold temperatures, the system may take a longer time to reach the required oxygen concentration. See Oxygen Concentrator Specifications on page 293. 	
		3. In modes of operation where the oxygen concentrator runs continuously (including O2 Supplement Mode and Ventilate Mode with oxygen set to Maximum), a degradation in concentrator oxygen purity may occur after operating in extreme high temperature environments for an extended period of time. Furthermore, in this situation it is recommended that alternate power sources are available (such as a power supply/charger) as the extreme heat and loading of the batteries may cause battery overheating and unexpected power loss.	
		4. NEVER CHARGE BATTERIES IN AMBIENT TEMPERATURES BELOW 32°F (0°C) OR ABOVE 104°F (40°C).	
Relative Humidity	15% to 95% non-condensing	Same	
Altitude	0–18,000 ft. (5,500m)	Same	
Water Resistance	MIL-STD-810G Method 506.4 for blowing rain	Same & IPX4 (Water splashing against the device)	
Blowing Sand and Dust Resistance	MIL-STD-810G Method 510.4 Procedure II	Same	
Radiated Immunity	N/A	30 V/m	
ESD	N/A	15 kV air discharge 8 kV contact discharge	



Variable	Storage Condition	Operating Condition
Mechanical Strength	Composite Wheeled Transport Vibration (JECETS)	 Vibration (sinusoidal) according to IEC 60068-2-6
		 Random vibration wide band – Reproducibility Medium according to IEC 60068-2-36
		Bump according to IEC 60068-2-29
		Drop Test: 1M attached to stretcher.
		Rotary Wing Combined (JECETS)
		Fixed Wing Jet Aircraft (JECETS)
		Fixed Wing Turbo-Propeller Aircraft



REMOTE SCREEN

Table 51: Environmental Specifications of the Remote Screen

Variable	Storage Condition	Operating Condition
Temperature	-4°F to 140°F (-20°C to 60°C)	14°F to 122°F (–10°C to 50°C).
Relative Humidity	30% to 90% non-condensing	30% to 80% non-condensing



NOTE: Even within the above temperature/humidity ranges, operation for a long time in extreme environments, smoking nearby, or operation in places where oil is used or where there is a lot of dust will result in the product deterioration and will shorten the product life.

16.7 PATIENT MONITORING SPECIFICATIONS

16.7.1 Heart Rate Monitoring Specifications

Table 52: Heart Rate Monitoring Specifications of MOVES[®] SLC™

Item	Specification
Source	Auto-detect the first available source in the priority sequence of ABP1, ABP2, ABP3, SpO ₂ , ECG; or manually select one of the available sources
Range	30–250 bpm for ABP and ECG, 30–239 bpm for SpO ₂
Accuracy	± 1% Full Scale (under stationary operation) ±5 BPM (under continuous vibration)
Filtering	ECG: In ECG pediatric heart rate mode, the detection range of QRS amplitude is 0.5 mV to 5 mV, for durations of the QRS wave ranging from 40 ms to 120 ms, up to a signal rate of 350 BPM. Otherwise, the detection range of QRS amplitudes is 0.5 mV to 5 mV, for durations of the QRS complex ranging from 50 ms to 120 ms, up to a signal rate of 300 BPM.
Pacemaker Pulse Rejection Capability	Pacemaker pulses may be detected by the ECG heart rate monitor and included in its calculation, depending on the type and model of pacemaker detected by the heart rate monitor.
Fixed Delays Due to Signal Processing	ABP: Pulse heart rate is calculated from the previous 6 beats. SpO2: Pulse heart rate is calculated based on the SpO2 average time selected on the Advanced screen (2–16 seconds, default 8 seconds). ECG: Heart rate is calculated from the previous 8 beats.
Alarm Condition Delay (onset of condition to internal realization) NOTE: "Alarm Condition Delay" derives its value from "Fixed Delays Due To Signal Processing" + 100 ms.	ABP: 6 beats + 100 ms SpO2: SpO ₂ average time + 100 ms ECG: 8 beats + 100 ms
Alarm Signal Generation Delay (realization to display)	Less than 200 ms



Item	Specification
Operating Mode That May Affect Alarm Generation	ECG: Incorrect setting of ECG pediatric mode.

16.7.2 Temperature Monitoring Specifications



NOTE: The thermometers conform to all of the requirements established in ASTM standard E1112.

Table 53: Temperature Monitoring Specifications of MOVES $^{\!@}$ SLC $^{\rm TM}$

Item	Specification	
Range	28°C to 42°C (82.4°F to 107.6°F)	
Accuracy	Range (Celsius)	Accuracy
	Less than 35.8°C:	± 0.3°C
	35.8°C to less than 37.0°C:	± 0.2°C
	37.0°C to 39.0°C:	± 0.1°C
	Greater than 39.0°C to 41.0°C:	± 0.2°C
	Greater than 41.0°C:	± 0.3°C
	Range (Fahrenheit)	Accuracy
	Less than 96.4°F:	± 0.5°F
	96.4°F to less than 98.0°F:	± 0.3°F
	98.0°F to 102.0°F:	± 0.2°F
	Greater than 102.0°F to 106.0°F:	± 0.3°F
	Greater than 106.0°F:	± 0.5°F
Standards Compliance	ASTM E1112-00	

16.7.3 Airflow Monitoring Specifications

Table 54: Airflow Monitoring Specifications of MOVES[®] SLC™

Item	Specification
Inspiratory/Expiratory Flow Range	- 60 to 60 LPM
Repeatability of Inspiratory/Expiratory Flow Measure	± 0.5% (% of reading)
Airway Pressure (Paw) Range	- 5 to 70 cmH ₂ O



Item	Specification
Accuracy of Airway Pressure Measure	± 2 cmH ₂ O + 4% of reading
Tidal Volume Accuracy	\pm (15% by Volume + 4mL)
Standards Compliance	ISO 80601-2-12, ISO 80601-2-13

16.7.4 CO₂ Monitoring Specifications

Table 55: CO₂ Monitoring Specifications of MOVES[®] SLC™

Item	Specification
Range	0 to 10% by Volume
Resolution	0.02%
Accuracy	± 1.0% Absolute
Rise Time	215 ms (10-90%) at 200 ml/min
Response time of gas sample readings	< 4 seconds
Flow Rate	250 ml/min ± 50 ml/min
Standards Compliance for CO ₂ Analyzer Used	ISO 80601-2-55: Medical electrical equipment – Particular requirements for the basic safety and essential performance of respiratory gas monitors

16.7.5 Respiratory Rate Monitoring Specifications

Table 56: Respiratory Rate Specifications of MOVES[®] SLC™

Item	Specification
Source	Capnograph (CO ₂)
Range	0 to 99 B/M
Accuracy	0 to 60 B/M: the greater of ± 2 B/M or ± 5% of actual value
Standards Compliance	ISO 80601-2-55: Medical electrical equipment – Particular requirements for the basic safety and essential performance of respiratory gas monitors

16.7.6 O₂ Monitoring Specifications

Table 57: O2 Monitoring Specifications of MOVES[®] SLC™

Item	Specification
Range	5 to 100% by Volume
Resolution	0.02%



Item	Specification
Accuracy	± 4% Absolute
Rise Time	150 ms (10-90%) at 150 ml/min
Response time of gas sample readings	< 4 seconds
Flow Rate	250 ml/min ± 50 ml/min
Operating Mode That May Affect Alarm Generation	None
Standards Compliance for O ₂ Analyzer Used	ISO 80601-2-55: Medical electrical equipment – Particular requirements for the basic safety and essential performance of respiratory gas monitors

16.7.7 ECG Specifications

Table 58: ECG Specifications of MOVES[®] SLC™

Item	Specification
Number of Leads	12
Lead View	Standard 12 lead with Wilson chest lead placement
Input Impedance	> 10 MOhm
Input Range	> 10 mVpp
Input Range (DC)	> 300 mV
Sensitivity	See ECG graphs of Section 14.2.1 System Graphs
Filtering	50 Hz, 60 Hz; optional 15–30 Hz EMG filter
Frequency Response	0.3–70 Hz
Pulse Detection	30–250 bpm ± 1%, ± 1 Digit, 8 beat averaging Adult detection – Will not respond to QRS amplitude of 0.15 mV or less or R-wave duration of 10 ms or less with an amplitude of 1 mV. Detection range of QRS amplitude is 0.5 mV to 5.0 mV, for duration of the QRS complex ranging from 50 ms to 120 ms, up to a signal rate of 300 BPM. Pediatric detection – Detection range of QRS amplitude is 0.5 mV to 5.0 mV, for duration of the QRS complex ranging from 40 ms to 120 ms, up to a signal rate of 350 BPM.
Defibrillator Protection	Yes
ST Analysis	None
Pacer Detection	None
Standard Complied to	IEC 60601-2-27



Item	Specification	
Maximum T-Wave Amplitude	0.6 mV	
T-Wave Rejection	The tall T-wave rejection is alwa rejection is 0.6 mV when R-wave	
Indicated Heart Rate (after a 20-second equipment		Rhythm
stabilization period)	A1 (80 BPM)	78
	A2 (60 BPM)	58
	A3 (120 BPM)	118
	A4 (90 BPM)	88
Response Time to Heart Rate Meter	Longest Time	< 10 seconds
	80 BPM-120 BPM	
	Longest Time	< 15 seconds
	80 BPM-40 BPM	
Time to Alarm for Tachycardia	B1 – Regular Amplitude	6 seconds
	B1 – Half Amplitude	6.2 seconds
	B1 – Double Amplitude	± 6 seconds
	B2 – Regular Amplitude (max)	5 seconds
	B2 – Half Amplitude (max)	4 seconds
	B2 – Double Amplitude (max)	4.2 seconds
Pacemaker Pulse Representation	Pacemaker pulses will be displa ECG waveform display and can estimation).	

16.7.8 NIBP Specifications

Table 59: NIBP Specifications of MOVES[®] SLC™

Item	Specification
Measurement Cycles	Stat, 1, 2, 3, 4, 5, 10, 15 minutes
Max Allowable Cuff Pressure	300 mmHg
Range	Systolic: 40–260 Diastolic: 20–200
Resolution	1 mmHg
Accuracy	± 5 mmHg Average with STD of 8 mmHg



Item	Specification
Calibration	The cuff pressure transducer should be verified once every 12 months.

16.7.9 Invasive Pressure Specifications

Table 60: Invasive Pressure Specifications of MOVES[®] SLC™

Item	Specification
Channels	3
Transducer Sites	ABP, CVP or ICP
Pressure Range	ABP: -10 to 300 mmHg CVP: -10 to 300 mmHg ICP: -14 to 408 cmH ₂ O
Temperature:	
Operating	15° to 40°C (57°F to 104°F)
Storage	-25° to 70°C (13°F to 158°F)
Accuracy	± 4 mmHg or 4% of reading whichever is greater

16.7.10 Pulse Oximetry Specifications

Table 61: Pulse Oximetry Specifications of MOVES[®] SLC™

Item	Specification
Method	Multiple visible and infrared LEDs (500 to 1400 nm)
Maximum Optical Power Output	≤ 25 mW
Fixed Delays Due to Signal Processing	Current SpO ₂ is calculated from previous 2–16 seconds depending on "SpO ₂ Average Time" setting in Advanced section of Setup Screen. Default: 8 seconds.
Alarm Condition Delay (onset of condition to internal realization)	Fixed delay due to signal processing + 100 ms
Alarm Signal Generation Delay (realization to display)	Less than 200 ms



16.7.11 Equipment Response Time

Table 62: Pulse Oximetry Equipment Response Time

SpO ₂ Values	Average	Latency
Standard/Fast Averaged SpO ₂	Choice of 2–4, 4–6, 8, 10, 12, 14, or 16 seconds. Default: 8 seconds	2 beats
Pulse Rate Values	Average	Latency
Standard/Fast Averaged Pulse Rate	Choice of 2–4, 4–6, 8, 10, 12, 14, or 16 seconds. Default: 8 seconds	2 beats

16.7.12 Drift in Sensing Accuracy

A drift of less than 0.4% in the oxygen reading can be expected over a 6 hour duration when external conditions are held constant.

A drift of less than 0.3% in the carbon dioxide reading can be expected over a 6 hour duration when external conditions are held constant.

16.7.13 Specifications of Masimo Rainbow SET® Pulse CO-Oximeter

Measurement Range

Measurement	Display Range
SpO2 (Oxygen Saturation)	0% to 100%
SpMet (Methemoglobin)	0.0% to 100.0%
SpCO (Carboxyhemoglobin)	0% to 100%
SpHb (Hemoglobin)	0.0 g/dL to 25.0 g/dL
	0.0 g/L to 250 g/L
	0.0 mmol/L to 15.5 mmol/L
SpOC (Oxygen Content)	0 ml of O2/dL to 35 ml of O2/dL of blood
PR (Pulse Rate)	25 bpm to 239 bpm, > 239 displayed when PR is 240–260 bpm
PI (Perfusion Index)	0% to 10% (bar graph beside SpO2)
PVI (Pleth Variability Index)	0% to 100%

Accuracy [7]

Oxygen Saturation Accuracy [1]	
No Motion	60% to 80%
Adults/Pediatrics	± 3%



No Motion [2]	70% to 100%
Adults/Pediatrics	± 2%
Motion [3]	70% to 100%
Adults/Pediatrics	± 3%
Low Perfusion [4]	70% to 100%
Adults/Pediatrics	± 2%
Pulse Rate Accuracy [5]	
Pulse rate range	25 bpm to 239 bpm
No Motion	L
Adults/Pediatrics	± 3 bpm
Motion	L
Adults/Pediatrics	± 5 bpm
Low Perfusion	
Adults/Pediatrics	± 5 bpm
Carboxyhemoglobin Accuracy [1]	
Adults/Pediatrics	1% to 40% ± 3%
Methemoglobin Saturation Accuracy [1]	'
Adults/Pediatrics	1% to 15% ± 1%
Total Hemoglobin Accuracy [6]	
Adults/Pediatrics	8 g/dL to 17 g/dL ±1 g/dL

Notes:

- 1. SpO₂, SpCO, and SpMet accuracy was determined by testing on healthy adult volunteers in the range 60% to 100% SpO₂, 0% to 40% SpCO, and 0% to 15% SpMet against a laboratory CO-Oximeter. SpO₂ and SpMet accuracy was determined on 16 neonatal NICU patients ranging in age from 7 days to 135 days old and weighting between 0.5 kgs and 4.25 kgs. Seventy-nine (79) data samples were collected over a range of 70% to 100% SaO₂ and 0.5% to 2.5% HbMet with a resultant accuracy of 2.9% SpO₂ and 0.9% SpMet. Contact Masimo for testing specifications.
- 2. The Masimo sensors have been validated for no motion accuracy in human blood studies on healthy adult male and female volunteers with light to dark skin pigmentation in induced hypoxia studies in the range of 70-100% SpO2 against a laboratory CO-Oximeter and ECG monitor. This variation equals plus or minus one standard deviation. Plus or minus one standard deviation encompasses 68% of the population.
- 3. The Masimo sensors have been validated for motion accuracy in human blood studies on healthy adult male and female volunteers with light to dark skin pigmentation in induced hypoxia studies in the range of 70-100% SpO2 against a laboratory CO-oximeter and ECG monitor. This variation equals plus or minus one standard deviation which encompasses 68% of the population.



- 4. The Masimo SET Technology has been validated for low perfusion accuracy in bench top testing against a Biotek Index 2 simulator and Masimo's simulator with signal strengths of greater than 0.02% and transmission of greater than 5% for saturations ranging from 70 to 100%. This variation equals plus or minus one standard deviation which encompasses 68% of the population.
- 5. The Masimo sensors have been validated for pulse rate accuracy for the range of 25-240 bpm in bench top testing against a Biotek Index 2 simulator. This variation equals plus or minus one standard deviation which encompasses 68% of the population.
- 6. SpHb accuracy has been validated on healthy adult male and female volunteers and on surgical patients with light to dark skin pigmentation in the range of 8 g/dL to 17 g/dL SpHb against a laboratory CO-Oximeter. The variation equals plus or minus one standard deviation which encompasses 68% of the population. The SpHb accuracy has not been validated with motion or low perfusion.
- 7. The following substances may interfere with pulse CO-Oximetry measurements:
 - Elevated levels of Methemoglobin (MetHb) may lead to inaccurate SpO2 and SpCO measurements.
 - Elevated levels of Carboxyhemoglobin (COHb) may lead to inaccurate SpO2 measurements.
 - Very low arterial Oxygen Saturation (SpO2) levels may cause inaccurate SpCO and SpMet measurements.
 - Severe anemia may cause erroneous SpO2 readings.
 - Dyes, or any substance containing dyes, that change usual blood pigmentation may cause erroneous readings.
 - Elevated levels of total bilirubin may lead to inaccurate SpO2, SpMet, SpCO and SpHb readings.

Resolution

Parameter	Step Size
%SpO ₂	1%
%SpCO	1%
%SpMet	0.1%
SpHb • g/dL • g/L • mmol/L	0.1 g/dL 1 g/L 0.1 mmol/L
SpOC mL/dL	1 mL O ₂ /dL blood
Pulse Rate	1 beats per minute
% PVI	1%



16.8 GENERAL ACCESSORIES SPECIFICATIONS

Table 63: General Accessories Specifications of MOVES[®] SLC™

Item	Category	Specification
Batteries	Operating Temperature:	-26°C to 54°C (-14°F to 129°F) (cold start above -20°C)
	Storage Temperature:	-26°C to 60°C (-14°F to 140°F)
	Charging Temperature:	0°C to 40°C (32° to 104°F)
Hydrocarbon/Particulate Filter (P/N 100915)	Effective Filtration Against:	GME Organic Vapor, Chlorine, Sulfur Dioxide, Chlorine Dioxide, Hydrogen Chloride, Hydrogen Sulfide, Ammonia, Methylamine, Formaldehyde, Hydrogen Fluoride: 99.97% effective against all particulate aerosols
NIBP Cuffs (ALL 4)	Temperature:	
	Operating	0°C to 40°C (32°F to 104°F)
	Storage	-34°C to 70°C (-29.2°F to 158°F)
Temperature Probe (Reusable/Autoclavable)	Temperature:	Interchangeable ± 0.1°C, 25°C to 45°C per EN 12470 Tested to ± 0.1°C, 0°C to 70°C for laboratory use
	Autoclave:	Withstands 100 autoclave cycles to 121°C Withstands 50 autoclave cycles to 134°C
Suction Canister		800 mL capacity

16.8.1 Approved Masimo Pulse Oximeter Accessories

The following Masimo accessories may be used in conjunction with the MOVES[®] SLC™ pulse oximeter. See the respective sensor instructions for detailed information regarding specified sensor use.

Table 64: Approved Masimo Pulse Oximeter Accessories

Accessory	Description
Rainbow [®] DCI-dc3	Adult Reusable Finger Sensor for SpCO, SpO ₂ & SpMet, 3 ft. NOTE: Can be used to measure only SpO ₂ if optional SpCO and/or SpMet features have not been activated on MOVES [®] SLC™ system.
Rainbow [®] DCIP-dc3	Pediatric Reusable Finger Sensor for SpCO, SpO ₂ & SpMet, 3 ft. NOTE: Can be used to measure only SpO ₂ if optional SpCO and/or SpMet features have not been activated on MOVES [®] SLC™ system.
Rainbow [®] RC-4	Rainbow [®] Patient Cable, 4 ft.



Accessory	Description
Rainbow [®] R1 25	Adult SpO ₂ , SpHb & SpMet Adhesive Sensor NOTE: Requires Rainbow [®] RC-4 Patient Cable. NOTE: Requires optional SpHb feature to be activated on MOVES [®] SLC [™] system in order for any SpO ₂ , SpMet or SpHb measurements to be made.
Rainbow [®] R1 20	Pediatric SpO ₂ , SpHb & SpMet Adhesive Finger Sensor NOTE: Requires Rainbow [®] RC-4 Patient Cable. NOTE: Requires optional SpHb feature to be activated on MOVES [®] SLC [™] system in order for any SpO ₂ , SpMet or SpHb measurements to be made.
Rainbow [®] R25	Adult SpO ₂ , SpCO & SpMet Adhesive Sensor NOTE: Requires Rainbow [®] RC-4 Patient Cable. NOTE: Can be used to measure only SpO ₂ if optional SpCO and/or SpMet features have not been activated on MOVES [®] SLC™ system.
Rainbow [®] R20	Pediatric SpO ₂ , SpCO & SpMet Adhesive Finger Sensor NOTE: Requires Rainbow [®] RC-4 Patient Cable. NOTE: Can be used to measure only SpO ₂ if optional SpCO and/or SpMet features have not been activated on MOVES [®] SLC™ system.
M-LNCS DCI [®]	Adult Reusable SpO ₂ Finger Sensor, 3 ft. NOTE: Requires Rainbow® RC-4 Patient Cable.
M-LNCS DCIP	Pediatric Reusable SpO ₂ Finger Sensor, 3 ft. NOTE: Requires Rainbow® RC-4 Patient Cable.
M-LNCS TC-I	Reusable SpO ₂ Ear Sensor, 3 ft NOTE: Requires Rainbow® RC-4 Patient Cable. NOTE: Sensor has not been validated under motion conditions. NOTE: Sensor is contraindicated for patients with pierced ears at the measuring site.
M-LNCS E1	Adult SpO2 disposable ear sensor, 3ft NOTE: Requires Rainbow® RC-4 Patient Cable. NOTE: Sensor has not been validated under motion conditions.
M-LNCS Adtx-3	Adult SpO ₂ Adhesive Sensor, 3 ft. NOTE: Requires Rainbow® RC-4 Patient Cable.
M-LNCS Pdtx-3	Pediatric SpO ₂ Adhesive Sensor, 3 ft. NOTE: Requires Rainbow® RC-4 Patient Cable.



16.8.2 Masimo Pulse Oximeter Accessories Specifications

Table 65: Masimo Pulse Oximeter Accessories Specifications

Sensor	Description	Preferred Application Site	Masimo P/N	Weight Range	Sp Accu	02 racy	P Accu	R iracy	Lo Perfu Accu	sion	SpCO/Hb Accuracy	SpMet Accuracy
					No Motion	Motion	No Motion	Motion	Sp02	PR	No Motion	No Motion
Rainbow DCI-dc3	Adult Reusable Finger Sensor for SpO2, SpCO, & SpMet, 3ft.	Finger	2201	>30 kg	60-80% ±3% 70- 100% ±2%	±3%	±3 bpm	±5 bpm	±2%	±3 bpm	(SpCO) ±3%	± 1%
Rainbow DCIP-dc3	Pediatric Reusable Finger Sensor for SpO2, SpCO, & SpMet, 3 ft.	Finger	2069	10-50 kg	60-80% ±3% 70- 100% ±2%	±3%	±3 bpm	±5 bpm	±2%	±3 bpm	(SpCO) ±3%	± 1%
M-LNCS DCI	Adult Reusable Finger Sensor	Finger	2501	>30 kg	±2%	±3%	±3 bpm	±5 bpm	±2%	±3 bpm	N/A.	N/A.
M-LNCS DCIP	Pediatric Reusable Finger Sensor	Finger	2502	10-50 kg	±2%	±3%	±3 bpm	±5 bpm	±2%	±3 bpm	N/A.	N/A.
M-LNCS TC-I	Adult Reusable Ear Sensor	Ear Lobe (Contraindicated for patients with pierced ears at the measuring site)	2503	>30 kg	±3.5%	N/A	±3 bpm	N/A	±3.5%	±3 bpm	N/A.	N/A.
M-LNCS E1	Adult Disposable Ear Sensor	Ear Concha	2919	>30 kg	±2.5%	N/A	±3 bpm	N/A	±2.5%	±3 bpm	N/A.	N/A.



Sensor	Description	Preferred Application Site	Masimo P/N	Weight Range	Sp Accu		P Accu		Lo Perfu Accu	sion	SpCO/Hb Accuracy	SpMet Accuracy
					No Motion	Motion	No Motion	Motion	Sp02	PR	No Motion	No Motion
M-LNCS Adtx-3	Adult SpO2 Adhesive Sensor, 3 ft.	Finger	2509	>30 kg	±2%	±3%	±3 bpm	±5 bpm	±2%	±3 bpm	N/A.	N/A.
M-LNCS Pdtx-3	Pediatric SpO2 Adhesive Sensor, 3 ft.	Finger	2511	10-50 kg	±2%	±3%	±3 bpm	±5 bpm	±2%	±3 bpm	N/A.	N/A.
Rainbow R1 25	Adult Adhesive Sensor for SpHb, SpO2, & SpMet	Finger	2416	>30 kg	60-80% ± 3% 70- 100% ± 2%	±3%	±3 bpm	±5 bpm	± 2%	±3 bpm	(SpHb) ±1 g/dL	± 1%
Rainbow R1 20	Pediatric Adhesive Finger Sensor for SpHb, SpO2, & SpMet	Finger	2417	10-50 kg	60-80% ± 3% 70- 100% ± 2%	±3%	±3 bpm	±5 bpm	± 2%	±3 bpm	(SpHb) ± 1 g/dL	± 1%
Rainbow R25	Adult Adhesive Sensor for SpO2, SpCO, & SpMet	Finger	2221	>30 kg	60-80% ± 3% 70- 100% ± 2%	±3%	±3 bpm	±5 bpm	± 2%	±3 bpm	(SpCO) ±3%	±1%
Rainbow R20	Pediatric Adhesive Finger Sensor for SpO2, SpCO, & SpMet	Finger	2222	10-50 kg	60-80% ± 3% 70- 100% ± 2%	±3%	±3 bpm	±5 bpm	± 2%	±3 bpm	(SpCO) ±3%	±1%



16.9 ELECTROMAGNETIC CONFORMITY INFORMATION

16.9.1 IEC 60601-1-2:2007 (Ed 3.0) Table 1 Requirements

MOVES[®] SLC[™] is intended for use in the electromagnetic environment specified below. The customer or the user of MOVES[®] SLC[™] should assure that it is used in such an environment.

Table 66: 5.2.2.1c IEC 60601-1-2:2007 (Ed 3.0) Table 1 Requirements

Emissions Test	Compliance	Electromagnetic Environment – Guidance
RF emissions CISPR 11	Group 1	MOVES [®] SLC [™] 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 emissions CISPR 11	Class A	MOVES [®] SLC™ is suitable for use in all establishments other
Harmonic emissions IEC 61000-3-2	Class A	than domestic, and may be used in domestic establishments and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic
Voltage fluctuations/ flicker emissions IEC 61000-3-3	Complies	purposes, provided the following warning is heeded: Warning: MOVES® SLC™ is intended for use by healthcare professionals only. MOVES® SLC™ may cause radio interference or may disrupt the operation of nearby equipment. It may be necessary to take mitigation measures, such as re-orienting or relocating MOVES® SLC™ or shielding the location.



16.9.2 IEC 60601-1-2:2007 (Ed 3.0) Table 2 Requirements

MOVES[®] SLC[™] is intended for use in the electromagnetic environment specified below. The customer or the user of MOVES[®] SLC[™] should assure that it is used in such an environment.

Table 67: 5.2.2.1f IEC 60601-1-2:2007 (Ed 3.0) Table 2 Requirements

Immunity Test	IEC 60601 Test Level	Compliance Level	Electromagnetic Environment – Guidance
Electrostatic discharge (ESD) IEC 61000-4-2	± 6 kV contact ± 8 kV air	± 6 kV contact ± 8 kV air	Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30%.
Electrical fast transient/burst IEC 61000-4-4	± 2 kV for power supply lines ± 1 kV for input/output lines	± 2 kV for power supply lines ± 1 kV for input/output lines	Mains power quality should be that of a typical commercial or hospital environment.
Surge IEC 610004-5	± 1 kV line(s) to line(s) ± 2 kV line(s) to earth	± 1 kV line(s) to line(s) ± 2 kV line(s) to earth	Mains power quality should be that of a typical commercial or hospital environment.
Voltage dips, short interruptions and voltage variations on power supply input lines IEC 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. End user shall ensure charged batteries are installed in the Equipment.
Power frequency (50 Hz) magnetic field IEC 61000-4-8	3 A/m	3 A/m	Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.



16.9.3 IEC 60601-1-2:2007 (Ed 3.0) Table 3 Requirements

 $\mathsf{MOVES}^{\circledR}$ SLCTM is intended for use in the electromagnetic environment specified below. The customer or the user of $\mathsf{MOVES}^{\circledR}$ SLCTM should assure that it is used in such an environment.

Table 68: 5.2.2.2 IEC 60601-1-2:2007 (Ed 3.0) Table 3 Requirements

Immunity Test	IEC 60601 Test Level	Compliance Level	Electromagnetic Environment – Guidance
Conducted RF IEC 61000-4-6	3 Vrms 150 kHz to 80 MHz Outside ISM band	3 Vrms	Portable and mobile RF communications equipment should be used no closer to any part of MOVES [®] SLC™, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter.
	10 Vrms ISM Frequencies: 6.78 MHz, 13.56 MHz, 27.12 MHz, 40.68 MHz	10 Vrms	Recommended separation distance: $d = [\frac{3.5}{V_1}]\sqrt{P}$
Radiated RF IEC 61000-4-3	10 V/m 80 MHz to 2.5 GHz	10 V/m	$d = \left[\frac{12}{V_2}\right]\sqrt{P}$
			$d = \left[\frac{12}{E_1}\right] \sqrt{P}$ 80 MHz to 800 MHz
			$d = [\frac{23}{E_1}]\sqrt{P}$ 800 MHz to 2,5 GHz
			Where <i>P</i> is the maximum output power rating of the transmitter in watts (<i>W</i>) according to the transmitter manufacturer and <i>d</i> is the recommended separation distance in meters (<i>m</i>).
			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 . Interference may occur in the vicinity of known RF transmitting devices and equipment marked with the following symbol:

Immunity Test IEC 60601 Test Level Compliance Level	Electromagnetic Environment – Guidance
---	--

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 equipment is used exceeds the applicable RF compliance level above, the equipment should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as re-orienting or relocating the equipment
- b) Over the frequency range 150 kHz to 80 MHz, field strengths should be less than 10 V/m.

16.9.4 IEC 60601-1-2:2007 (Ed 3.0) Table 5 Requirements

MOVES[®] SLC[™] is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or user of MOVES[®] SLC[™] can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and MOVES[®] SLC[™] as recommended below, according to the maximum output power of the communications equipment.

Table 69: 5.2.2.2 IEC 60601-1-2:2007 (Ed 3.0) Table 5 Requirements

Recommended Separation Distances Between Portable and Mobile RF Communications Equipment and MOVES [™] SLC [™]							
	Separation Distance According to Frequency of Transmitter (m)						
Rated Maximum Output Power of	150 kHz to 80 MHz outside ISM bands 150 kHz to 80 MHz in ISM bands		80 MHz to 800 MHz	800 MHz to 2.5 GHz			
Transmitter (W)	$d = \left[\frac{3,5}{V_1}\right]\sqrt{P}$	$d = \left[\frac{12}{V_2}\right] \sqrt{P}$	$d = \left[\frac{12}{E_1}\right] \sqrt{P}$	$d = \left[\frac{23}{E_1}\right] \sqrt{P}$			
0.01	0.12	0.12	0.12	0.23			
0.1	0.37	0.38	0.38	0.73			
1	1.17	1.20	1.20	2.30			
10	3.69	3.79	3.79	7.27			
100	11.67	12.00	12.00	23.00			

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

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

NOTE 2: The ISM (industrial, scientific and medical) bands between 150 kHz and 80 MHz are 6.765 MHz to 6.795 MHz; 13.553 MHz to 13.567 MHz; 26.957MHz to 27.283 MHz; and 40.66 to 40.70 MHz.

NOTE 3: An additional factor of 10/3 is used in calculating the recommended separation distance for transmitters in the ISM frequency bands between 150 kHz and 80 MHz and in the frequency range 80 MHz to 2.5 GHz to decrease the likelihood that mobile/portable communications equipment could cause interference if it is inadvertently brought into the patient area.

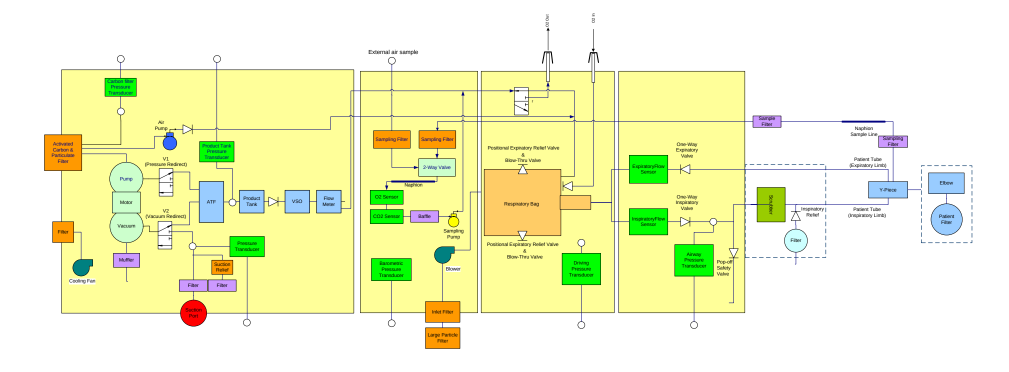
NOTE 4: These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.



MOVES® SLC™ Operator's Manual Thornhill Research Inc.

17.0 Appendix B – Pneumatic Diagram

17.1 MOVES[®] SLC™ PNEUMATIC DIAGRAM





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18.0 Appendix C – End-User License Agreement

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19.0 Appendix D - FCC and IC Declaration

This device complies with Part 15 of the FCC Rules. Operation is subject to the following two conditions:

- (1) This device may not cause harmful interference, and
- (2) This device must accept any interference received, including interference that may cause undesired operation.

NOTE: This equipment has been tested and found to comply with the limits for a Class B digital device, pursuant to part 15 of the FCC Rules. These limits are designed to provide reasonable protection against harmful interference when the equipment is operated in a commercial environment. This equipment generates, uses, and can radiate radio frequency energy and, if not installed and used in accordance with the instruction manual, may cause harmful interference to radio communications. Operation of this equipment in a residential area is likely to cause harmful interference in which case the user will be required to correct the interference at his own expense.

This Class B digital apparatus complies with Canadian ICES-003.

This device complies with Industry Canada license-exempt RSS standard(s). Operation is subject to the following two conditions:

- (1) this device may not cause interference, and
- (2) This device must accept any interference, including interference that may cause undesired operation of the device.

CAUTION: Any changes or modifications not expressly approved by the party responsible for compliance could void the user's authority to operate the equipment. This equipment should be installed and operated with minimum distance 20 cm between the radiator and your body.

FCC ID: XF6-RS9113DB

IC: 8407A-RS9113DB

CE/ETSI

TELEC

SRRC

IEEE 802-11n (2.4GHz)



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