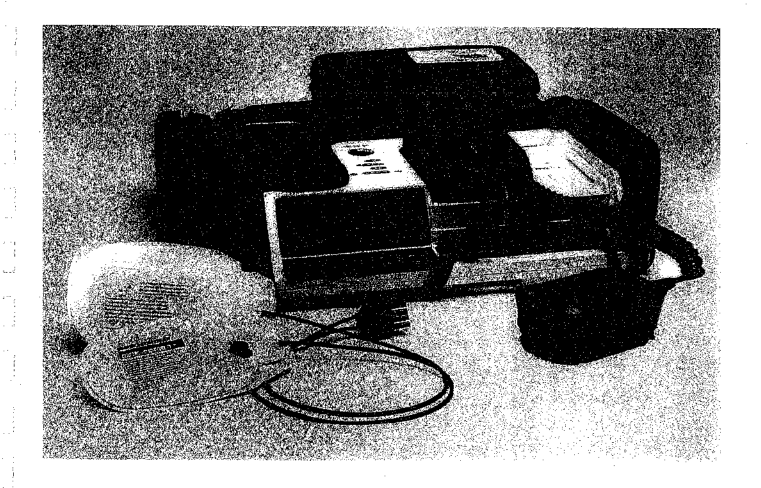
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LIFEPAK® 10C defibrillator/monitor/pacemaker



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OPERATING INSTRUCTIONS

LIFEPAK® 10C defibrillator/monitor/pacemaker

IMPORTANT

Federal (USA) law restricts this device to sale by or on the order of a physician.

This instrument is to be used by authorized medical personnel only.

Device Tracking

(USA only, including US government-owned units)
Under the Safe Medical Devices Act of 1990, defibrillator manufacturers and distributors are required to track the location of defibrillators. If your defibrillator has been sold, donated, lost, stolen, exported, or destroyed or if it was not obtained directly from Physio-Control, please notify Physio-Control at 1.800.442.1142, extension 4530.

Responsibility for Information

It is the responsibility of our customers to assure that the appropriate person(s) within their organization have access to this information.

Corporate Headquarters 11811 Willows Road Northeast Post Office Box 97005 Bedmond, WA 98073-9708 USA Tetephone: 256.867,4000 761 Free (USA only):600.426.8047 Fax: 205.867,4161 PRYSIO-CONTROL, LIFEPAK, FASTPAK, FAST-PATCH, QUIK-CHARGE, QUIK-PACE, DERMA JEL and LIFE-PATCH are registered tracemarks of Physio-Control Corporation.

CODE SUMMARY, PARTSLINE, and QUIK-COMBO are frademarks of Physio-Control Corporation.

Specifications subject to change without notice.

Lifto in USA.

**1935 Physio-Control Corporation.

P/N 3004087-000

TABLE OF CONTENTS

Pr	eface		
	About defibrillation	vii	
	Terms	Viji	
	General Warnings	viii	
	Symbols	ix	
1	Basic Operation		
	Introduction	1-2	
	Controls, Indicators, and Connectors	1-2	
	Setting the Clock	1-7	
2	Monitoring		
	Monitoring Warnings	2-2	
	Standard Paddles Monitoring Procedure	2-2	
	ECG Monitoring Procedure	2-3	
	ECG Electrode Requirements	2-4	
	QRS Detection	2-4	
	Monitoring Patients with Invasive Pacemakers	2-4	

3	Recording		
	Loading Paper	3-2	
	Using the Recorder	3-2	
	CODE SUMMARY Critical Event Record	3-3	
4	Standard Paddles Defibrillation		
	Defibrillation Warnings	4-2	
	Paddle Usage/Options	4-3	
	Standard Paddle Placement	4-3	
	Standard Paddles Defibrillation Procedure	4-4	
	Synchronized Cardioversion Procedure Using Patient ECG Cable	4-5	
 5	QUIK-COMBO Electrodes Procedures		
	About QUIK-COMBO Electrodes	5-2	
	QUIK-COMBO Electrode Placement	5-3	
	Connecting the QUIK-COMBO Electrode Cable	5-5	
	Monitoring Procedure	5-5	
	Defibrillation Procedure	5-6	
	Synchronized Cardioversion Procedure	5-7	
	Noninvasive Pacing	5-8	
	Defibrillation During Noninvasive Pacing	5-12	
	Patient Care Transfer to a Different Device	5-12	
_			
6	Maintenance and Testing		
	General Maintenance and Testing	6-2	
	Battery Maintenance and Testing	6-7	
	Troubleshooting	6-14	
	Service and Repair	6-18	
	Warranty	6-18	
	Supplies, Accessories, and Training Tools	6-19	•

Appendix A: Specifications

Appendix B: Basic Operation Checklist

LIST OF FIGURES

Figure 1-1	LIFEPAK 10C defibrillator/monitor/pacemaker controls, indicators, and connectors	1-2
Figure 1-2	Function buttons	1-4
Figure 1-3	Pacemaker buttons	1-5
Figure 1-4	Status display indicators	1-6
Figure 1-5	Paddle buttons	1-7
Figure 2-1	Electrode placement	2-3
Figure 3-1	Example of CODE SUMMARY report overview	3-4
Figure 3-2	Example of defibrillation format	3-4
Figure 3-3	Example of recorded ECG format	3-5
Figure 3-4	Example of noninvasive pacing format	3-5
Figure 4-1	Standard paddle anterior-lateral placement	4-3
Figure 5-1	Peeling the liner from the electrode	5-3
Figure 5-2	Anterior-lateral placement	5-3
Figure 5-3	Anterior-posterior placement	5-4
Figure 5-4	Removing cover and pushing connectors together	5-5
Figure 5-5	ECG recording strip of electrical capture	5-10
Figure 5-6	Electrical capture with intrinsic beat activity	5-10
Figure 5-7	ECG recording strip with distortion	5-11
Figure 5-8	Pacemaker refractory period	5-11
Figure 6-1	Reconditioning procedure form	6-10
Figure 6-2	Shelf Life Test form	6-11
Figure 6-3	Battery Maintenance Log	6-12

LIST OF TABLES

Table 1-1	Controls, indicators, and connectors	1-3
Table 1-2	Function button descriptions	1-4
Table 1-3	Pacemaker button descriptions	1-5
Table 1-4	Status display description	1-6
Table 1-5	Paddle button descriptions	1-7
Table 6-1	Recommended maintenance schedule for clinical personnel	6-2
Table 6-2	Inspection	6-3
Table 6-3	Recommended cleaning	6-3
Table 6-4	Troubleshooting the monitor and recorder	6-14
Table 6-5	Troubleshooting the defibrillator	6-16
Table 6-6	Troubleshooting the pacemaker	6-17
Table 6-7	Supplies, accessories, and training tools	6-19

PREFACE

About defibrillation

The LIFEPAK® 10C defibrillator/monitor/pacemaker is a therapeutic medical device intended for use by or under the direction or guidance of a physician. Direct current defibrillation is a recognized means of terminating certain potentially fatal cardiac dysrhythmias.

A direct current defibrillator applies a brief, high-energy pulse of electricity to the heart. This energy may be delivered either through external paddles or electrodes on the chest.

Defibrillation is only one aspect of the medical care required to resuscitate a patient in ventricular fibrillation. Depending on the situation, other supportive measures may include:

- · establishment and maintenance of a patent airway
- · ventilation, including administration of oxygen
- maintenance of blood circulation
- · pharmacologic measures.

Among other factors, it is recognized that the likelihood of successful resuscitation of a patient is related to the length of time between the onset of ventricular fibrillation and defibrillation. Rapid defibrillation and prompt follow-up care are essential. The physiological state of the patient may affect the likelihood of successful defibrillation or skeletal muscle contractility. Thus, failure to convert the dysrhythmia or to resuscitate a patient is not a reliable indicator of defibrillator performance. Similarly, the patient's muscular response to the defibrillator shock is not a reliable indicator of the energy delivered. Refer to the booklet *Defibrillation: What You Should Know* for further information (refer to page 6-19 for ordering information).

Daily inspection is important to determine the state of readiness of the equipment. In addition, the device must be kept in proper operating condition at all times through routine maintenance, testing, and repair by a qualified service technician. Refer to the *LIFEPAK 10C defibrillator/monitor/pacemaker Service Manual* for additional service information.

Terms

The following safety-related terms are used either in this manual or on the LIFEPAK 10C defibrillator/monitor/pacemaker:

Danger:

Immediate hazards which will result in serious personal injury or death.

Warning:

Hazards or unsafe practices which could result in serious personal injury or death.

Caution:

Hazards or unsafe practices which could result in minor personal injury or

product/property damage.

General Warnings

In addition to the following warnings, other warnings are provided near the beginning of each section.

△ WARNINGS

Possible loss of power during patient care.

Proper care and maintenance of batteries is vital to the performance of the LIFEPAK 10C defibrillator/monitor/pacemaker. Always carry a spare, fully-charged, properly-maintained battery.

When discharged, this defibrillator delivers up to 360 joules of electrical energy. Unless discharged properly as described in these Operating Instructions, this electrical energy may cause personal injury or death. Do not attempt to operate this device unless you are thoroughly familiar with these Operating Instructions and the function of all controls and indicators, as well as the connections and accessories.

Shock or fire hazard.

Do not immerse any portion of this device in water or other fluids. Avoid spilling any fluids on device or accessories. Do not autoclave this device or accessories unless otherwise specified.

Possible fire or explosion.

Do not use this device in the presence of flammable gases or anesthetics. Use care when operating this device close to oxygen sources (such as bag-valve-mask devices or ventilator tubing)

Possible improper device performance.

Use only Physic Control ECG and QUIK-COMBO : cables, electrodes, and Physic-Control batteries. Substitution of non-Physic-Control cables, electrodes, or batteries may cause the device to perform improperly.

Possible interference with implanted devices.

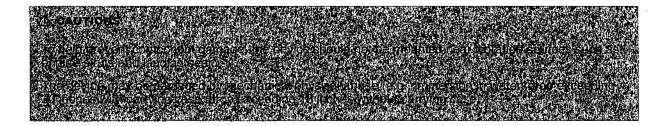
Magnets inside the standard defibrillation paddles may affect the function of an implanted device. Avoid placing standard paddles near an implanted device. Check function of implanted device after using standard paddles.

Possible electrical interference with ECG monitoring.

Equipment that emits certain radio frequency signals can cause electrical interference and distort the displayed or recorded ECG signal, thereby preventing accurate rhythm analysis. To minimize radio interference, move or reposition equipment away from defibrillator.

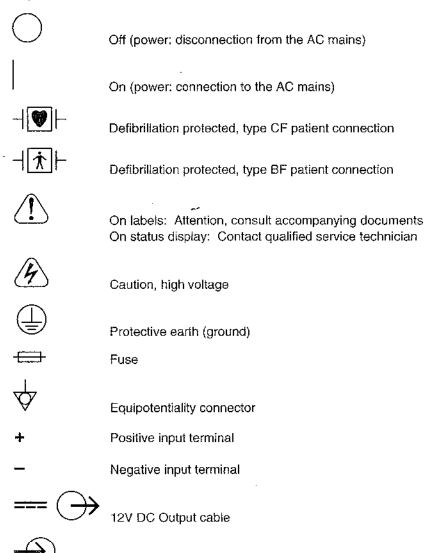
Safety risk and possible equipment damage.

Defibrillators, monitors, pacemakers, and their accessories (including electrodes and cables) contain ferromagnetic materials. As with all ferromagnetic equipment, these products must not be used in the presence of the high magnetic field created by a Magnetic Resonance Imaging (MRI) device. The high magnetic field created by an MRI device will attract the equipment with a force sufficient to cause death or serious personal injury to persons between the equipment and the MRI device. This magnetic attraction may also damage the equipment. Consult with the MRI manufacturer for more information.



Symbols

The symbols below may be found in this manual or on various configurations of the LIFEPAK 10C defibrillator/monitor/pacemaker and accessories.



12V DC Input cable



Paddles



Battery



Plug



Output



AC current



Output AC/Digital



Recycle battery



Recycle battery

BASIC OPERATION

This section describes the basic operation of the LIFEPAK 10C defibrillator/monitor/pacemaker. Topics include:

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Introduction

The LIFEPAK 10C defibrillator/monitor/pacemaker is a complete cardiac life support system used by paramedics, hospital staff, and other authorized healthcare providers. The defibrillator/monitor/pacemaker provides electrocardiogram (ECG) monitoring, defibrillation, synchronized cardioversion, and noninvasive pacing. It also allows single-cable, hands-free defibrillation, pacing, and monitoring therapy with the QUIK-COMBO™ pacing/defibrillation/ECG electrodes. The QUIK-COMBO electrodes allow patient transfer to other Physio-Control devices which use QUIK-COMBO electrodes.

The LIFEPAK 10C defibrillator/monitor/pacemaker includes the CODE SUMMARY™ critical event record which automatically stores critical events in memory. A 50mm thermal array printer provides printed copies of ECG monitoring and CODE SUMMARY reports. A cardioscope displays ECG monitoring. A Liquid Crystal Display (LCD) presents operating information such as heart rate, lead selection, and pacing current selection. Any one of three rechargeable NiCad batteries or an optional AC or DC Auxiliary Power Module provide power for the device.

Controls, Indicators, and Connectors

Figures 1-1 through 1-5 and Tables 1-1 through 1-5 provide an overview of the controls, indicators, and connectors for the LIFEPAK 10C defibrillator/monitor/pacemaker.

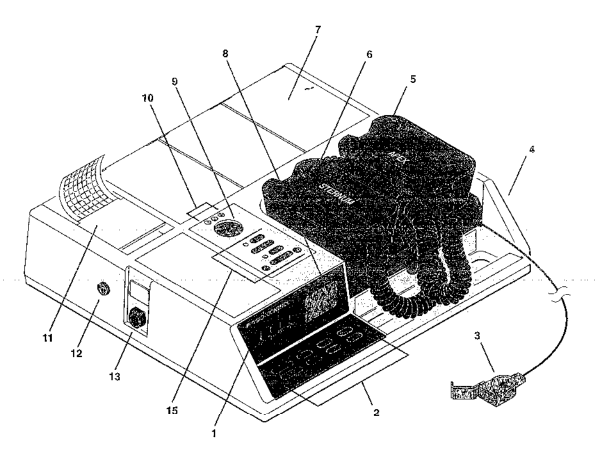


Figure 1-1 LIFEPAK 10C defibrillator/monitor/pacemaker controls, indicators, and connectors

Table 1-1	Controls, indicators, and con	nectors
1	Cardioscope	Non-fade display; ECG trace moves from right to left.
2	Function Buttons	Eight front-panel function buttons (see page 1-4).
3	QUIK-COMBO therapy cable	Allows hands-free defibrillation, pacing, or monitoring using QUIK-COMBO electrodes. Standard paddles must be stowed securely in paddle wells during use of QUIK-COMBO electrodes.
4	ELECTRICALLYISOLATED ECG Connector (far side, not shown)	Connection for 6-pin, 3-lead ECG cable (AHA or IEC version available).
5	APEX Paddle	QUIK-LOOKs, QUIK-CHARGEs defibrillation paddle with CHARGE, RECORD, and discharge button. Also serves as positive ECG electrode during standard paddle monitoring. (For button descriptions, see page 1-7.)
6	STERNUM Paddle	QUIK-LOOK defibriliation paddle with discharge button and ENERGY select dial. Also serves as negative ECG electrode during standard paddle monitoring. (For button descriptions, see page 1-7.)
7	Battery	Replaceable, rechargeable power source. Physio-Control® FASTPAK, LIFEPAK 5 FASTPAK or Battery Pak batteries may be used.
8	Status Display	Alphanumeric information indicates heart rate, AVAILABLE ENERGY, lead selected, SYNC mode, DIAG mode, pacing current and rate, pacing electrode connection message (LEADS), and service indicator (see page 1-6).
9	1 POWER	Rotary switch turns device power on or OFF. Select one of three batteries or, if available, auxiliary power source (AUX).
10	Low Battery Indicator	 When indicator is: Flashing - battery in use is nearly depleted; immediately switch to a charged battery. Continuously on - battery in use is depleted; replace depleted battery. The device may shut down with no low battery indication if the battery is damaged, improperly maintained, or depleted (e.g., if battery is very low on charge and operator attempts to charge defibrillator.)
11	Recorder	Thermal array recorder which prints ECG trace and annotations on 50mm thermal paper. Activated by RECORD button (see page 1-4).
12	MIEMSS Modulator Connector (optional)	Allows simultaneous output of one unmodulated and two modulated ECG signals.
13	AUX Connector	Allows connection to AC or DC Auxiliary Power Module to provide operating power. Also provides output (modulated or unmodulated) for ECG transmission (1V/mV ECG deflection).
14	Bail Incline (not shown)	Bail incline on bottom may be extended to tilt up device.
15	Pacemaker Controls	Controts for noninvasive pacemaker (see page 1-5).
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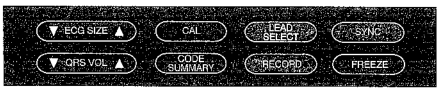


Figure 1-2 Function buttons

Table 1-2 Function bu	utton descriptions
V ECG SIZE A	Adjusts vertical size of ECG trace on cardioscope and recorder from 0.2 to 4.0cm/mV. Press ▲ to increase or ▼ to decrease ECG size.
(V ORSVOL ∕	Adjusts systole beeper volume. Press ▲ to increase, ▼ to decrease.
(CAL)	Superimposes 1mV calibration signal on cardioscope and recorder (not active in SYNC mode).
CODE SUMMARY.	Activates printing of report summarizing critical events (i.e., pre- and post-defibrillation/cardioversion events, pacing parameters, and selected monitored ECG segments).
TISELEON D	Selects ECG input: Paddles, Leads I, II, III. Press to change lead. Device may be programmed to power up in Paddles or Lead II.
(AREOGRA)	Activates thermal array recorder which prints time, date, ECG lead, ECG size, heart rate, SYNC (if activated), and pacing parameters. Press RECORD again to stop recorder. RECORD on APEX paddle performs identically.
	If device is programmed to enable diagnostic frequency response mode, holding RECORD down for more than 1 second selects diagnostic mode (DIAG) and starts recorder. Diagnostic mode must be reselected with each new recording. Recorder runs continuously in diagnostic mode.
(SYNCE)	Selects synchronized mode. To return to asynchronous mode, press SYNC again. Defibrillator automatically returns to asynchronous mode after discharge.
	Freezes cardioscope trace. Recorder continues to print delayed trace.

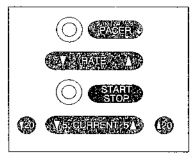
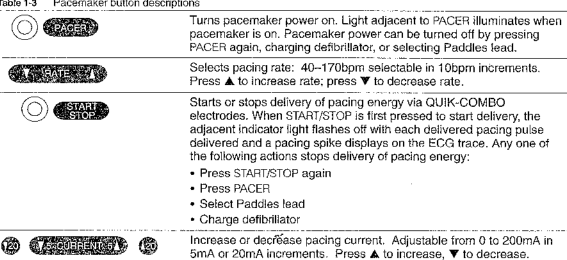


Figure 1-3 Pacemaker buttons

Pacemaker button descriptions Table 1-3



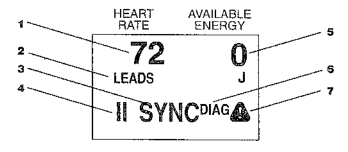


Figure 1-4 Status display indicators

1	Heart Rate	Displays two functions:									
		 PACER off-displays measured heart rate from ECG cable, QUIK-COMBO electrodes, or standard paddles; range: 20–295 beats per minute (bpm); symbol ~ ~ - indicates heart rate is outside the range of 20–295bpm. PACER on-displays selected (not measured) pacing rate from 									
2	LEADS Indicator	pacemaker control panel; range: 40–170 bpm. LEADS message displays when:									
		 Pacing is attempted without connecting the QUIK-COMBO therapy cable to the QUIK-COMBO electrodes. 									
		 QUIK-COMBO electrodes detach during pacing current delivery. During pacing, ECG monitoring is attempted in paddles lead . 									
3	SYNC Mode Indicator	SYNC message appears on status display indicating synchronized mode is enabled. Message blinks off with each detected QRS.									
ą	Lead Selection Indicator	Alphanumerics on status display identify lead selection:									
		Lead 1, II, or iii									
		Paddles lead.									
5	Available Energy	Displays two functions:									
		 PACER off-displays independent confirmation of energy level selected on ENERGY select dial (0-360J); a single tone sounds when charging is complete. 									
		 PACER on-displays pacing current (0-200mA) selected from pacemaker control panel. 									
3	DIAG Mode Indicator	DIAG message displays when diagnostic frequency response mode enabled.									
7	Service Indicator	Symbol indicates service is needed. If symbol displays continuously, have device promptly examined by a qualified service technician.									

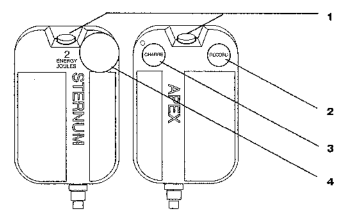


Figure 1-5 Paddle buttons

1	Discharge Buttons	Red buttons discharge the defibrillator. Both buttons must be pressed simultaneously to deliver energy. Energy is not delivered unless device is fully charged to selected energy level.
2	RECORD Button	Activates recorder. Functions identically to RECORD button on front panel.
3	3 CHARGE Button	Amber button initiates defibrillator charge cycle. Adjacent CHARGE indicator flashes when device is charging and glows steadily when fully charged. A single tone sounds when charging is complete.
4	2 ENERGY JOULES Dial	Rotary dial selects 1 of 9 discrete energy levels for defibrillation: 0, 5, 10, 20, 50, 100, 200, 300, or 360 joules.

Setting the Clock

To activate clock set-up mode:

- Turn the POWER switch to OFF. Then press and hold RECORD on the APEX paddle and turn the POWER switch to a power source. The heart rate section of the status display flashes the numbers 00. These numbers represent the hour digits of the 24-hour clock.
- 2 Press QRS VOL until the desired hour is displayed.
- 3 Press ▲ on ECG SIZE to scroll through the remaining clock settings in the heart rate display in the following order:
 - Minutes (0-59)
 - Month (1-12)
 - Day (1-31)
 - Year (0-99; the year 2000 shows as 00, 2001 as 01, etc.)
- 4 Press ▲ on QRS VOL to change any of the clock settings.
- 5 Turn the POWER switch to OFF to terminate the clock setting mode.
- To examine the clock setting, turn the POWER switch to a power source and press RECORD to start the recorder. Examine the printed strip and confirm the proper time and date is printed.

MONITORING

Patient ECG can be monitored with the standard paddles using the QUIK-LOOK defibrillation paddle feature, the 3-lead patient ECG cable, or the disposable QUIK-COMBO electrodes. For information about QUIK-COMBO electrodes refer to Section 5. Topics in this section include:

Monitoring War	nings	in di	J. 18			oage 2	-2
Standard Paddl	es Monit	oring P	rocedu	ıre		2	-2
ECG.Monitoring	Proced	ure				2	-3
ECG Electrode	Requirer	nents				. 2	-4
QRS Detection					Caltri	2	-4
Monitoring Pati	ents with	Invasiv	e Pac	emake	rs 🚶	2	-4

Monitoring Warnings

△ WARNINGS

Safety risk.

Use only Physio-Control patient ECG cables listed in this manual. Substitution of non-Physio-Control patient ECG cables may result in inaccurate ECG data.

Possible misinterpretation of cardioscope ECG data.

The cardioscope frequency response is only intended for rhythm identification. Use the recorder in DIAG mode for diagnostic interpretations.

Possible misinterpretation of ECG recordings.

When attempting to visually detect subtle ECG characteristics such as ST segment abnormalities, use the recorder only in diagnostic frequency response mode (DIAG). The monitor frequency response mode does not provide the resolution required for diagnostic and ST segment interpretation, and is intended only for basic ECG rhythm identification.

Possible electrical interference with ECG monitoring.

Do not operate this device in conjunction with electrocautery or diathermy equipment. Such equipment, as well as equipment which emits strong radio frequency signals, can cause electrical interference and distort the ECG signal displayed by the monitor, thereby preventing accurate rhythm analysis.

Standard Paddles Monitoring Procedure

To monitor with standard paddles:

- 1 Turn the 1 POWER switch to a power source. The device performs a 5-second, self-diagnostic test; all indicator lights and all status display messages illuminate momentarily.
 - The defibrillator may be programmed to power-on with Lead II selected (initial factory setting) or Paddles lead selected. For assistance in changing the power-on lead selection, contact a qualified service technician.
- 2 Press LEAD SELECT to select paddles lead (\mathbb{Q}).
- 3 Apply conductive gel over the entire paddle electrode surface.
- Place paddles firmly on patient's bare torso. The standard paddle electrode placement is STERNUM paddle on the patient's right upper torso below the clavicle and the APEX paddle lateral to the patient's left nipple in the midaxillary line.
- 5 Observe cardioscope to evaluate patient's rhythm.

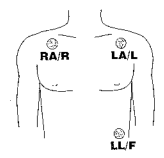
When the device is turned on, the ECG size is x1. The ECG size may need to be adjusted if the QRS complex is not clearly visible on cardioscope.

ECG monitoring after defibrillation is usually delayed by a defibrillation recovery time of a few seconds. During this time, it may not be possible to determine defibrillation results from the monitor trace.

ECG Monitoring Procedure

To perform 3-lead ECG monitoring:

- 1 Turn the 1 POWER switch to a power source.
- 2 Connect the patient ECG cable to the ELECTRICALLY ISOLATED ECG connector located on the right side panel.
- 3 Identify the appropriate electrodes sites on the patient:



AHA Labels
RA Right Arm
LA Left Arm
LL Left Leg

IEC Labels
R Right
L Left
F Foot

Figure 2-1 Electrode placement

- 4 Prepare patient's skin for electrode application:
 - Remove excessive hair at electrode site. Avoid locating electrodes over tendons and major muscle masses.
 - · For oily skin, clean skin with alcohol pad.
 - · Dry site with brisk rub.
- 5 Apply ECG electrodes:
 - Inspect electrode package and confirm package is sealed and date is not expired. Carefully tear open foil package and remove electrode carrier.
 - Attach an electrode to each of the lead wires.
 - Grasp electrode tab and peel electrode from carrier.
 - · Inspect electrode gel and make sure gel is intact (discard electrode if gel is not intact).
 - Hold electrode taut with both hands. Apply the electrode flat to the skin. Smooth tape outwardly in all directions. Do not press the center of the electrode.
- 6 Press LEAD SELECT to select desired lead (leads I, II, III, are available).
- 7 Adjust ECG SIZE if necessary. Size is automatically set to gain of x1 at power-on. To properly count heart rate during routine monitoring, the ECG size may need to be adjusted as follows:

 - Press ECG SIZE ▼ or ▲ until the systole beeper coincides with every QRS complex.
 - Adjust VOL ▼ or ▲ as desired.
- Secure the patient ECG cable with the cable clasp.
- To print an ECG strip, press RECORD. To stop the recorder, press RECORD again.

ECG Electrode Requirements

Electrode quality is critical for obtaining an undistorted ECG signal. Always check the date code on electrode packages for expiration date before patient use. Do not use electrodes with expired date codes.

For best ECG monitoring results, use silver/silver chloride (Ag/AgCl) electrodes such as Physio-Control LIFE•PATCH® ECG electrodes. The silver/silver chloride electrodes allow much faster post-defibrillation ECG display on the cardioscope than other electrode types.

Avoid using stainless steel electrodes; these electrodes can delay post-defibrillation ECG display on the cardioscope for 10 seconds or longer. If stainless steel electrodes must be used, perform careful patient evaluation including an extended period of cardioscope observation before pursuing further therapy.

QRS Detection

QRS detection is essential for use of the digital heart rate display, systole tone (QRS VOL), synchronized cardioversion, and noninvasive demand pacing.

The QRS defector in the LIFEPAK 10C defibrillator/monitor/pacemaker selectively detects QRS complexes. It discriminates against most noise, muscle artifact, T-waves, and other spurious signals.

Detection of QRS complexes and rejection of other signals depends on the proper setting of ECG size. If ECG size is set too low, QRS complexes will not be detected; no systole tones or sense (synchronizer) markers appear and the heart rate display is incorrect. If ECG size is set too high, systole tones and sense (synchronizer) markers may occur on spurious signals and the heart rate display may be incorrect.

The LIFEPAK 10C defibrillator/monitor/pacemaker displays a heart rate between 20 and 295bpm. Patient rates outside this range do not yield valid systole tones or heart rate display.

Monitoring Patients with Invasive Pacemakers

The LIFEPAK 10C defibrillator/monitor/pacemaker rejects most pacemaker impulses from internally implanted pacemakers. It does not use the pacemaker pulse for heart rate calculation or synchronization. Large amplitude pacemaker spikes can overload the QRS complex detector circuitry so that no paced QRS complexes are counted, resulting in blanking (heart rate displays - - -) of the heart rate display. To help minimize ECG pickup of large unipolar pacemaker pulses when monitoring patients with internal pacemakers, place ECG electrodes so the line between the positive and negative electrodes is perpendicular to the line between the pacemaker generator and the heart.

Smaller amplitude internal pacemaker pulses may not be distinguished clearly on the cardioscope and/or the recording strip in leads or paddles monitoring modes. To help distinguish internal pacemaker pulses on the recorder, try using the diagnostic mode. To help distinguish internal pacemaker pulses on the cardioscope and the recorded ECG strips, the leads monitoring mode can be programmed during set-up mode to agency frequency response. Refer to the LIFEPAK 10C defibrillator/monitor/pacemaker Service Manual or contact a qualified service technician for assistance.

RECORDING

This section describes how to record patient information. Topics include:

Loading Par)ér			page 3-2
Using the Re	ecorder.			: ∜∜3-2
CODE SUM	MARY Cri	ical Event	Record	3- 3

Loading Paper

The recorder is equipped with an out-of-paper sensor to protect the recorder printhead. The sensor automatically turns off the recorder if paper runs out or the recorder door is open.

To load the paper:

- Lift up the slotted edge of the front recorder door to open the recorder.
- 2 Remove empty paper roll.
- 3 Insert new paper roll, grid facing forward.
- 4 Pull out a short length of paper.
- 5 Pull the rear recorder door toward you and push down on the front recorder door to close.

⚠ CAUTION Possible equipment damage.

Use only-paper designed for thermal array recorders. Use of other types of paper may damage the print head.

Using the Recorder

To record:

- 1 Press RECORD.
- 2 Adjust ECG SIZE if necessary.
- 3 To stop printing, press RECORD again.

Recording can be performed with any lead selected.

Diagnostic Recording

If the diagnostic frequency response mode (DIAG) has been enabled during set-up, holding RECORD down for more than one second selects DIAG and turns on the recorder. The ECG signal now prints at a frequency response of 0.05 –100Hz (per AHA recommendations).

The DIAG mode must be reselected with each new recording. The recorder operates continuously when in DIAG mode.

Recorder Annotation

The recorder prints the time, date, ECG lead, ECG size, heart rate, defibrillation/synchronization parameters, pacing parameters, and CODE SUMMARY record. The beginning of each annotation is marked by an arrow symbol (>>).

While on, the recorder prints updated annotation information every 20 seconds. The recorder also updates the annotation if changes are made to lead selection, pacing parameters, or SYNC mode.

If FREEZE is pressed while the recorder is printing, the printing continues unaffected until FREEZE is released. At that time, frozen information is printed and annotated by ////ECG-FREEZE////. The recording function then returns to delayed mode.

If the recorder is on when the defibrillator is discharged, the recorder annotates the time, date, AVAILABLE ENERGY, and SYNC (if energy is transferred in SYNC mode) after discharge.

Handling Recordings

To help prevent the ECG annotation and tracing from fading or disappearing, follow these guidelines for thermal sensitive paper:

- Do not apply tape or other adhesives over printed information (adhesives may be applied to back of the paper).
- Store only in paper folders; do not store or file with plastics; avoid storing in temperatures exceeding 26.7°C (80°F) and relative humidity exceeding 70%.
- · Avoid extended exposure to sunlight.

CODE SUMMARY Critical Event Record

The CODE SUMMARY critical event record documents critical events during resuscitation. It records defibrillation and cardioversion details, operator-selected ECG segments, and pacing parameters in chronological order. Resuscitation details are prioritized for retention of the most critical events.

The CODE SUMMARY record does not store ECG data in DIAG mode. The CODE SUMMARY record stores ECG data at the monitoring frequency response (agency or domestic) selected in set-up.

Description of CODE SUMMARY Record

Critical events are retained in memory whenever the LIFEPAK 10C defibrillator/monitor/pacemaker is on. If power is removed, the CODE SUMMARY record may still be printed by applying power within five minutes and pressing CODE SUMMARY. After five minutes without power, CODE SUMMARY information may not be recovered.

Standard use of the recorder is available at any time by pressing RECORD once to interrupt CODE SUMMARY report printing, then pressing RECORD again to initiate recording. This does not delete information already stored in the CODE SUMMARY record.

If there is no paper in the recorder and the operator presses RECORD, additional ECG information is not stored in the CODE SUMMARY record. However, defibrillation, synchronized cardioversion, and pacing information is stored.

The CODE SUMMARY record only stores defibrillation and/or strip chart recording events if they are separated by at least a seven-second interval (i.e., if two defibrillation shocks are delivered within seven seconds, only the first shock is stored in the CODE SUMMARY record). The CODE SUMMARY record does not print whenever the defibrillator is charging. This helps prevent historical CODE SUMMARY data from being interpreted as real-time data.

Printing the CODE SUMMARY Report

To print the CODE SUMMARY report:

- Press CODE SUMMARY to initiate printing; unless interrupted, printing continues until the entire report is printed.
- 2 To interrupt printing, press CODE SUMMARY again.
 The CODE SUMMARY report printing is also interrupted if RECORD or CHARGE are pressed, power is turned off, or the recorder runs out of paper.
- 3 To resume printing, press CODE SUMMARY again. The recorder resumes printing beginning with the last event printed unless the interruption was caused by paper depletion. In this case, the recorder resumes printing beginning with the last three events printed.

Special Placement Situations

Implanted pacemaker patients. If possible, place paddles away from the internal pacemaker generator to help prevent damage to the pacemaker.

Patients with implanted defibrillators. Apply paddles in the preferred placement, APEX-STERNUM (anterior-lateral), and treat this patient like any other patient requiring emergency care. If defibrillation is unsuccessful, it may be necessary to increase the energy level or to use the alternate electrode placement (anterior-posterior) due to the insulation of implanted defibrillator electrodes.

Standard Paddles Defibrillation Procedure

To defibrillate using standard paddles:

- 1 Turn the POWER switch to a power source.
- a Apply defibrillation gel over entire paddle electrode surface.
- 3 Turn the ENERGY dial to the desired energy level. The defibrillator will not charge if the dial is between settings.
- 4 Place defibrillator paddles firmly on patient's chest.
- **5** Press CHARGE on APEX paddle. While the defibrillator is charging, the CHARGE indicator light flashes and the numbers increase in the AVAILABLE ENERGY display until the energy reached the selected level. A single tone sounds when the defibrillator is fully charged.
- **6** Make sure all personnel, including the operator, stand clear of the patient, bed, and any equipment connected to the patient.
- Discharge the defibrillator by simultaneously pressing both paddle discharge buttons. The defibrillator will not discharge until it completes charging to the selected energy level.
 If paddle discharge buttons are not pressed within 60 seconds, the stored energy is automatically removed within the defibrillator.
- **s** Observe patient and cardioscope to determine results. If additional countershock is necessary, repeat this procedure beginning at step 3.
- 9 To internally discharge an unwanted charge, rotate the ENERGY dial.
- 10 To turn off the defibrillator, turn POWER to OFF.
- Thoroughly clean defibrillator paddles and store them in the paddle storage area.

 If the ENERGY dial is rotated after charging is initiated, the AVAILABLE ENERGY display blanks, the charge indicator light goes out, and energy is internally removed. To reinitiate charging, press

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Synchronized Cardioversion Procedure Using Patient ECG Cable

Monitoring the ECG through the standard paddles (QUIK-LOOK monitoring) could introduce artifact and lead to improper synchronization during cardioversion. Always use the patient ECG cable or the QUIK-COMBO electrodes to monitor ECG during synchronized cardioversion.

There are two ways to monitor ECG for synchronized cardioversion:

- · Use the patient ECG cable with ECG electrodes and select Lead I, II, or III as described below.
- Use the QUIK-COMBO electrodes and select paddles lead as described on page 5-7.

To perform synchronized cardioversion when using the patient ECG cable:

- 1 Turn POWER switch to a power source.
- 2 Attach patient ECG cable and ECG electrodes. For proper placement of electrodes refer to ECG Monitoring Procedure on page 2-3.
- 3 Select lead with optimum QRS complex amplitude (positive or negative).
- 4 Press SYNC. Confirm the SYNC message on the status display blinks off with each detected QRS complex.
- 5 Observe the cardioscope. Confirm that a triangle sense marker appears near the middle of each QRS complex. If the sense markers do not appear or are displayed in the wrong locations, adjust ECG SIZE, select another lead, or reposition ECG electrodes. (It is normal for the sense marker location to vary slightly on each QRS complex.)
- 6 Rotate the ENERGY dial to select the desired energy. The defibrillator will not charge if the dial is between settings.
- 7 Prepare and place standard paddles on patient's chest.
- 8 Press CHARGE to charge defibrillator. Confirm the CHARGE indicator light flashes and the AVAILABLE ENERGY display indicates the energy level. When the single tone sounds the defibrillator is fully charged.
- Make certain all personnel, including operator, stand clear of the patient, bed, and any equipment connected to the patient.
- 10 Press and hold paddle discharge buttons until discharge occurs with next detected QRS complex. Release discharge buttons.
- 11 Observe patient and cardioscope. If synchronized cardioversion needs to be reattempted, press SYNC again and repeat procedure from step 5. (The defibrillator automatically returns to asynchronous mode after each discharge.)
- 12 To internally remove an unwanted charge, rotate the ENERGY select dial.
- 13 To turn off the defibrillator, turn POWER to OFF.
- 14 Thoroughly clean the paddles and store them in the paddle storage area.

The asynchronous defibrillation mode is automatically selected when the defibrillator powers on. The defibrillator automatically returns to asynchronous mode after each discharge.

Possible Causes of Pacing Interruption

If QUIK-COMBO electrodes become detached during pacing, the LEADS message is displayed and an audible alarm sounds. The pacing rate maintains its pre-alarm setting; however, the current resets to 0mA. Reattaching the QUIK-COMBO electrodes silences the audible alarm and removes the LEADS message. The pacing rate is maintained, but the current remains at 0mA unless increased by the operator.

Pacing therapy cannot be initiated or maintained in paddles lead. If the paddles lead is selected when cycling through leads during pacing, the current returns to 0mA and pacing therapy stops. If the paddles lead is selected and pacing is attempted, the LEADS message displays accompanied by an audible alarm.

Use of radio equipment while pacing may cause the current delivery to stop, the service message to appear, and an audible alarm to sound. To minimize radio interference, move radio equipment farther away from the defibrillator/monitor/pacemaker. If unable to move radio away, reorient the radio. Press PACER to stop the tones and erase the service message. To reinitiate pacing, follow the Pacing Procedure beginning with step 8 on page 5-9.

Defibrillation During Noninvasive Pacing

To defibrillate during noninvasive pacing:

- 1 Turn the ENERGY dial to the desired energy level. The defibrillator will not charge if the dial is between settings.
- 2 Press CHARGE on the APEX paddle to charge the defibrillator. A single tone sounds when the defibrillator is fully charged.
 - When CHARGE is pressed, pacing stops immediately (pacing control settings return to 40ppm and 0mA) and lights adjacent to PACER and START/STOP buttons go off. The HEART RATE display measures the patient's intrinsic rate in beats per minute and the AVAILABLE ENERGY display indicates the selected energy in joules.
- 3 Make certain all personnel, including the operator, stand clear of the patient, bed, and any equipment connected to the patient.
- 4 Discharge the defibrillator by simultaneously pressing both paddle discharge buttons.
- 5 Observe the patient and cardioscope. If additional countershock is necessary, repeat the procedure beginning at step 2.
- 6 To internally discharge an unwanted charge, rotate the ENERGY dial.

Patient Care Transfer to a Different Device

To transfer patient care between devices equipped to use QUIK-COMBO electrodes:

- Power off the device.
- 2 Disconnect the QUIK-COMBO electrode cable from the QUIK-COMBO therapy cable on the device. Leave the electrodes on the patient.
- 3 Connect the QUIK-COMBO electrode cable to the QUIK-COMBO therapy cable on the next device.
- 4 Follow instructions for the desired therapy.
- 5 Close the protective cover on the QUIK-COMBO therapy cable connector.

MAINTENANCE AND TESTING

This section describes how to perform operator-level maintenance, testing, and troubleshooting. Topics include:

General M	aintenance	and Tes	ing		page 6-2
Battery Ma	untenance	and Testi	ng:		6.7
Troublesh	ooting				6-14
Service an	d Repair				6-18
Warranty					6-18
Supplies, i	Accessorie	s, and Tra	aining To	ols .	6-19

Monitor/Recorder Test

Equipment Needed

- LIFEPAK 10C defibrillator/monitor/pacemaker
- Patient ECG cable
- QUIK-COMBO 3-Lead Patient Simulator

Test Procedure

- Turn defibrillator/monitor/pacemaker POWER switch to a power source. Confirm the device completes a self-test with no service indicator displayed.
- 2 Connect the patient ECG cable to the ECG connector and attach the leads to the patient simulator. The simulator power should remain off.
- 3 Select lead II on the defibrillator/monitor/pacemaker.
- 4 Press QRS VOL ▲ 5 times.
- 5 Press and release CAL. Confirm that a 1mV calibration pulse is displayed on the cardioscope.
- **6** Turn on simulator power and select NSR. Confirm the monitor displays a normal sinus rhythm with a heart rate of 72. Confirm that QRS tones sound with each beat.
- 7 Press FREEZE. Confirm the trace on the cardioscope stops. Release FREEZE.
- Press RECORD. Confirm the recorder operates and prints the ECG trace. Confirm that after approximately 3 seconds, the recorder annotates the time, date, lead II, ECG gain and heart rate on the paper.

⚠ CAUTION Possible equipment damage...

Recorder will not run without paper. Use only paper designed for thermal array recorders. Use of any other ECG paper may damage print head.

- 9 Press RECORD to turn off recorder.
- •• While in lead II, remove either the RA or LL lead from the simulator and confirm the NSR ECG trace is no longer displayed. Reconnect the lead.
- 11 Select lead I and remove either the RA or LA lead from the simulator. Confirm the NSR ECG trace is no longer displayed. Reconnect the lead. Confirm removing the reference lead, LL, does not affect the ECG display.
- 12 Select lead III and remove either the LA or LL lead from the simulator. Confirm the NSR ECG trace is no longer displayed. Reconnect the lead. Confirm removing the reference lead, RA, does not affect the ECG display.
- 12 To test the QUIK-COMBO therapy cable, connect the therapy cable to the QUIK-COMBO 3-Lead Patient Simulator.

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- 14 Confirm the standard paddles are securely stored in the paddle wells.
- 15 Select paddles lead ().
- Confirm the simulator power is on and NSR is selected. Confirm the monitor displays a normal sinus rhythm with a heart rate of 72. Confirm that QRS tones sound with each beat.

Defibrillator Test

Equipment Needed

- LIFEPAK 10C defibrillator/monitor/pacemaker
- · Battery Support System
- · QUIK-COMBO 3-lead patient simulator
- Timer

Test Procedure

- 1 Turn the defibrillator POWER switch to a power source.
- 2 Select 360 joules on the Battery Support System (refer to Battery Support System Operating Instructions if necessary).

riangle WARNING Possible paddle damage and patient burns.

Press paddles firmly onto the Battery Support System test plates when discharging to prevent formation of pits on paddle surfaces. Pitted or damaged paddle plates can cause patient skin burns during defibrillation.

- 3 Place the standard paddles on the Battery Support System test load plates. Make sure paddle surfaces do not contact your body or any other surface of the Battery Support System.
- 4 Rotate the ENERGY dial to select 360 joules.
- 5 Press CHARGE and start the timer.
- 6 Confirm that the tone indicating full charge sounds within 12 seconds or less.
- 7 Press RECORD.
- 8 Press only the APEX discharge button and confirm defibrillator does not discharge. Release the APEX discharge button.
- Press only the STERNUM discharge button and confirm defibrillator does not discharge. Release the STERNUM discharge button.
- Apply firm pressure with both paddles on the Battery Support System test load plates and press both paddle discharge buttons simultaneously. Confirm the Battery Support System displays the delivered energy. Confirm the recorder annotates the time, date, and energy selected on the ECG strip.
- 11 To test the QUIK-COMBO therapy cable, connect the therapy cable to the QUIK-COMBO 3-lead patient simulator.

⚠ WARNING Shock hazard.

During defibrilation cable testing, the discharged energy passes through the therapy cable connector. Be sure that the cable connector is securely attached to the testing device.

- 12 Rotate the ENERGY dial to select 360 joules.
- 13 Press CHARGE.
- 14 Discharge the defibrillator by pressing both paddle discharge buttons simultaneously while observing the DEFIB light on the simulator.
- 15 Confirm that the DEFIB light flashes briefly indicating discharge. If the indicator light does not flash, remove the defibrillator from active service and contact a qualified service technician.

Do not deliver more than 20 defibrillation pulses per hour at maximum energy, with no more than 45 coccurring in any 5-minute period. This helps prevent heat build-up in the defibrillator.

Synchronous Cardioversion Test

Equipment Needed

- LIFEPAK 10C defibrillator/monitor/pacemaker
- Battery Support System
- Patient ECG cable
- QUIK-COMBO 3-lead patient simulator

Test Procedure

- 1 Turn the defibrillator POWER switch to a power source.
- 2 Connect the patient ECG cable to the defibrillator and the QUIK-COMBO 3-lead patient simulator.
- 3 Connect the QUIK-COMBO therapy cable to the QUIK-COMBO 3-lead patient simulator.
- 4 Apply power to the simulator and select bradycardia rhythm.
- Solution of the cardioscope. Press LEAD SELECT as needed to select a lead with tall QRS complexes (positive or negative).
- e Press SYNC. Confirm the SYNC message is displayed. Adjust ECG SIZE until the sense markers appear on the upper portion of the QRS complexes. Confirm the SYNC message blinks off with each detected QRS complex and the heart rate is displayed.
- 7 Press RECORD to start the recorder.
- 8 Make sure the standard paddles are securely stored in the paddle wells.
- **9** Rotate the ENERGY dial on the STERNUM paddle to select 50 joules.
- 10 Press CHARGE to charge the defibrillator.
- After the tone sounds indicating full charge, simultaneously press and hold both paddle discharge buttons while observing the cardioscope.
- 12 Confirm the defibrillator discharges on the next QRS complex.
- 13 Confirm the defibrillator returns to asynchronous mode (SYNC message no longer displayed) and the recorder annotates the time, date, 50J, and SYNC.

Noninvasive Pacemaker Test

Equipment Needed

- LIFEPAK 10C defibrillator/monitor/pacemaker
- · Patient ECG cable
- QUIK-COMBO 3-lead patient simulator

Test Procedure

- Turn the defibrillator/monitor/pacemaker POWER switch to a power source.
- 2 Connect the QUIK-COMBO therapy cable to the QUIK-COMBO 3-lead patient simulator.
 - 3 Apply power to the simulator and select bradycardia rhythm.
- 4 Connect the 3-lead patient ECG cable to the defibrillator/monitor/pacemaker and the simulator.
 - 5 Press LEAD SELECT to select lead II.
- 6 Press PACER to apply power to the pacemaker. Press RATE & to select 35bpm (same rate as simulator bradycardia rhythm). Confirm the pacer output current is displayed as 0mA.
 - 7 Observe the cardioscope to confirm that the ECG signal from the simulator is displayed. Confirm that sense markers appear on each QRS complex. If sense markers do not appear, or appear elsewhere on the ECG, press ECG SIZE to adjust.
 - s Press RATE ▲ and ▼ and confirm that the selected rate changes on the status display. Select a rate of 60.
 - Press START/STOP and confirm the adjacent indicator light flashes off with each pacing spike. Confirm the pacing rate is 60. (Pacing energy is not delivered because the current is 0mA.)
 - to Press 20 ▲ and 20 ♥ and confirm the displayed current rate changes in 20mA increments.
 - 11 Press ▲ 5 CURRENT 5 ♥ and confirm the displayed current rate changes in 5mA increments.
 - 12 Increase the output current to 125mA.

- **13** Observe the cardioscope for captured complexes. Confirm the PACE indicator light on the simulator flashes with each delivered pacing pulse.
- 14 Disconnect the QUIK-COMBO therapy cable from the simulator. Confirm the pacemaker stops pacing, the LEADS message is displayed, and an audible alarm sounds.
- 15 Leave the QUIK-COMBO therapy cable disconnected and attempt to reinitiate pacing by pressing START/STOP. Confirm the LEADS message is displayed and an audible alarm sounds.
- 16 Reconnect the QUIK-COMBO therapy cable to the simulator.
- 17 Increase the output current to 125mA.
- **18** Press CHARGE to charge the defibrillator. Confirm the PACER indicator light goes off and the heart rate and available energy are displayed.
- 19 Turn the defibrillator POWER switch to OFF to internally discharge energy and remove power.
- 20 Press simulator OFF button to remove power.

Battery Maintenance and Testing

The LIFEPAK 10C defibrillator/monitor/pacemaker uses Nickel-Cadmium (NiCad) batteries. These NiCad batteries must be properly maintained using the Battery Support System to help maximize battery life and performance.

Use only Physio-Control batteries and battery chargers with Physio-Control devices. Use only the Battery Support System for battery maintenance.

△ WARNINGS

Possible loss of power during patient care.

Using an improperly maintained battery to power the defibrillator/monitor may cause premature power loss. Use only the Battery Support System to properly maintain batteries.

Possible loss of power during patient care.

Stored batteries lose charge. Failure to charge a stored battery before use may cause premature defibrillator/monitor power loss. Always charge a stored battery before returning it to active service.

Possible loss of power during patient care.

Physio-Control has no information regarding the performance or effectiveness of its LIFEPAK defibrillator/monitors if they are used with non-Physio-Control batteries or battery chargers. Using non-Physio-Control batteries or battery chargers may result in device failure and may void warranty. Use only Physio-Control batteries and the Battery Support System.

Fire or explosion hazard.

The two-well Battery Charger (Physio-Control Part Numbers 9-00284, 9-00288, or 801530) is not designed to charge FASTPAK batteries. Charging FASTPAK batteries in the two-well Battery Charger may reduce battery life and create risk of fire or explosion. Use only the Battery Support System to charge FASTPAK batteries.

When the low battery indicator appears, switch power to an alternate battery or auxiliary power module Do not continue using a discharged battery. Overdischarging can shorten battery life.

Battery Description

Physio-Control FASTPAK, LIFEPAK 5 FASTPAK, or Battery Pak batteries can power the LIFEPAK 10C defibrillator/monitor/pacemaker. The batteries perform similarly but require different charge times:

- Either FASTPAK battery charges in the Battery Support System in approximately 70 minutes.
- The Battery Pak battery charges in the Battery Support System in approximately 4 1/2 hours.

NiCad Battery Performance Factors

Three major factors affect the performance of NiCad batteries: temperature, voltage depression, and the self-discharge rate.

Temperature

Charging a battery at temperatures below 20°C (68°F) or above 25.5°C (78°F) prevents the battery from reaching its full capacity and may lead to irreversible cell damage.

Voltage Depression

Voltage depression is a condition which reduces battery performance, particularly when charging the defibrillator. This condition is often mistakenly called "memory." Voltage depression can usually be reversed by reconditioning the battery every 3 months as described on page 6-10. Voltage depression is caused by either:

- 1 Repeatedly attempting to add more charge to a fully charged or a nearly fully charged battery, or
- 2 Extended charging at temperatures above 25.5°C (78°F).

Self-Discharge Rate

Like most batteries, NiCad batteries self-discharge when not used. A new NiCad battery self-discharges approximately 1% of its capacity each day when stored at room temperature. In 10 days a new NiCad battery not installed in the defibrillator/monitor/pacemaker loses approximately 10% of its capacity. The self-discharge rate of the battery can be evaluated by performing a Shelf-Life Test as described on page 6-11. The actual battery self-discharge rate depends on:

- Battery age
- Temperature
- Frequency of use
- Length of time in storage
- Physical battery condition

These factors can combine to significantly increase the battery discharge rate. For example, an older battery stored in higher temperatures may have an accelerated self-discharge rate much greater than 1% a day.

Using the Battery Support System

Use only the Battery Support System to maintain FASTPAK, LIFEPAK 5 FASTPAK, and Battery Pak batteries. Refer to Battery Support System Operating Instructions for more information.

The AC and DC Auxiliary Power Modules do not perform all the procedures required to properly maintain or evaluate battery performance. Although the Power Modules supply a trickle-charge to any batteries installed in the defibrillator/monitor/pacemaker, their primary function is to supply external power to operate the device.

To properly maintain batteries, use only the Battery Support System with the following guidelines:

· Charge batteries at the proper temperature.

The optimum charging temperature is room temperature, or 20 to 25.5°C (68 to 78°F). Batteries charged outside room temperature may not reach full capacity even if the charge time is increased.

- · Properly locate the Battery Support System:
 - Place in a well-ventilated area.
 - Keep at room temperature.
 - Do not place in direct sunlight.
 - Do not place near a heat source or an air conditioner.
- Rotate batteries so all batteries in active service are used equally.
- · Recondition batteries every three months.
 - Reconditioning is a succession of discharge/charge cycles performed in the Battery Support System. Reconditioning a battery helps prevent or reverse effects of voltage depression and helps to keep track of battery capacity.
- Perform Shelf Life Test every six months (or alternate with the Reconditioning Procedure every 3 months).

The Shelf Life Test evaluates the self-discharge rate of a stored battery.

Installing and Removing a Battery

$\underline{\Lambda}$ WARNING Possible loss of power during patient care.

Battery pins (connectors) in the LIFEPAK 10C defibrillator/monitor/pacemaker and the Battery Support System may be damaged if batteries are dropped or forced into battery wells. Inspect pins routinely for signs of damage.

Do not drop or force a battery into the battery well.

To install a battery:

- Align the battery with battery well so the battery clip is toward connector pins.
- 2 Insert the end of the battery opposite the battery clip into the battery well.
- 3 Firmly press the other end of the battery into the battery well until it clicks into place.

To remove the battery, press the battery clip and lift.

Reconditioning Procedure

Reconditioning is a succession of discharge/charge cycles which may be performed on a battery inserted in the far right compartment of the Battery Support System. Reconditioning a battery helps prevent or reverse the effects of voltage depression and helps keep track of battery capacity.

Perform reconditioning *every three months* according to the Reconditioning Procedure in Figure 6-1. Discard any battery with a capacity reading of less than 80% on the third cycle. For information about ordering copies of the Reconditioning Procedure form, refer to page 6-19.

 R	CONDITIONING PROCEDURE		
For use with the Physio-Control® Battery Support System, FASTPAK® and Battery Pak batteries. – 80% or greater battery capacity is acceptable – Alternate every 90 days with Shelf Life Test – Use Battery Support System at 68–78° F – For Technical Support, call (800)442-1142 USA			
Test !	Date Battery ID		
Perfo	ermed by		
CHE	CKLIST (√ circle when done)		
○1 ○2 ○3 ○4 ○5 ○6	Charge battery until READY light appears Cycle #1: DISCHG-CHARGE-READY; disregard reading Cycle #2: DISCHG-CHARGE-READY; disregard reading Remove battery for 1 – 4 hrs Begin End Cycle #3: DISCHG-CHARGE-READY; bat. cap. =% Log Cycle #3 bat. cap.% on back of battery		
	Cycle #3 bat. cap. 80% or greater? (Yes-acceptable (No-unacceptable/discard battery		
	P/N 806017 -001 © 1993 Physio-Control Corporation		

Figure 6-1 Reconditioning procedure form

Shelf Life Test Procedure

The Shelf Life Test evaluates the self-discharge rate of a stored battery. Perform the Shelf Life Test described in Figure 6-2 every six months, or alternate it with the Reconditioning Procedure in Figure 6-1 every three months. Discard any battery with a Shelf Life Test value of more than 20. For information about ordering copies of the Shelf Life Test form, refer to page 6-19.

SHELF	LIFE TEST			
For use with the Physio-Control® Battery Support System, FASTPAK® and Battery Pak batteries. Shelf Life Test Value of 20 or less is acceptable. Alternate every 90 days with Reconditioning Procedure (Note: Steps 1–5 equals Reconditioning Procedure). Use Battery Support System at 68–78° F. For Technical Support, call (800)442-1142 USA.				
Test Date .	Battery ID			
Performed by				
CHECKLIST (√ circle when	n done)			
○ 1 Charge battery until	READY light appears			
○ 2 Cycle #1: DISCHG=0	CHARGE-READY; disregard reading			
○ 3 Cycle #2: DiSCHG-	CHARGE-READY; disregard reading			
4 Remove battery for f	I – 4 hrs Begin End			
○ 5 Cycle #3: DISCHG-	-CHARGE-READY; bat. cap. =%			
○ 6 Log Cycle #3 bat. ca	p.% on back of battery			
7 Remove battery for 7	7–8 days and store on shelf			
Begin <u>,,//</u>	End//			
○8 Cycle #4: DISCHG-	-CHARGE-READY; bat. cap. =%			
Record: Cycle #3 ba	ıt. cap %			
Subtract: Cycle #4 b	eat. cap. —%			
Result: Shelf Life Tes	st Value = %			
Shelf Life Test Value 20 o	or less?			
○ Yes-acceptable ○	No-unacceptable/discard battery			
P/N 806018~001 © 1992	Physio-Control Corporation			

Figure 6-2 Shelf Life Test form

Battery Maintenance Log

The Battery Maintenance Log shown in Figure 6-3 is available to help track battery maintenance procedures. For information about ordering copies of the Battery Maintenance Log, refer to page 6-19.

DATE i.D.	NUMBER BA	TTERY TEST PERFORMED	BATTERY TEST RESULTS	BATTERY A	CCEPTABL
		Faulty Reconditioning	Battery Capacity %	[]YES	☐ NO Discard
		Procedure Shelf Life Test Visual Inspection	Shelf Life Test Value		Battery
ļ		(case not cracked or broken)	Case OK Case not OK		
	, —	Faulty Reconditioning Procedure	Battery Capacity % Shelf Life Test Value	I∐ YES	□ NO Discard Battery
]	Sheff Life Test Visual Inspection (case not cracked or broken)	Case OK Case not OK		
		Faulty	Battery Capacity%	□ YES	□NO
•		Reconditioning Procedure Shelf Life Test	Shelf Life Test Value		Discard Battery
		Visual Inspection (case not cracked or broken)	Case OK ☐ Case not OK ☐	:	
	0.0	Faulty Reconditioning	Battery Capacity %	[]YES	☐ NO Discard
	-	Procedure Shelf Life Test	Shelf Life Test Value		Battery
		Visual Inspection (case not cracked or broken)	Case OK ☐ Case not OK ☐		
		Faulty Reconditioning	Battery Capacity %	☐ YES	□ NO Discard
		Procedure Shelf Life Test	Shetf Life Test Value		Battery
j	[.]	Visual Inspection (case not cracked or broken)	Case OK ☐ Case not OK ☐		
		Faulty Reconditioning	Battery Capacity %	YES	⊙ NO Discard
	l lo	Procedure Shelf Life Test	Shelf Life Test Value		Battery
		Visual Inspection (case not cracked or broken)	Case OK Case not OK		al neigh a
		Faulty Reconditioning	Battery Capacity %	. ☐ YES	☐ NO Discard
	.	Procedure Shelf Life Test	Shelf Life Test Value		Battery
1		Visual Inspection (case not cracked or broken)	Case OK ☐ Case not OK ☐		

Figure 6-3 Battery Maintenance Log

Receiving New Batteries

When newly-purchased batteries are received:

- Promptly label each new battery. Use a unique identification number so you can easily track the battery through all maintenance and rotation.
- Recondition each new battery. Because NiCad batteries self-discharge, a new battery may not be fully charged by the time it is received. Recondition a newly purchased battery according to the Reconditioning Procedure on page 6-10.

Storing Batteries

Store batteries in the Battery Support System or on a shelf. Batteries still require routine maintenance, even while in storage. When storing on a shelf:

- Store batteries between 4.4° and 26.7°C (40° and 80°F). Cooler temperatures reduce the battery self-discharge rate.
- Never freeze batteries.

Recycling Batteries at the End of Useful Life

When properly maintained, the Physio-Control NiCad batteries should have a battery life of approximately two years. A NiCad battery has reached the end of useful life if *one or more* of the following circumstances occur:

- Battery capacity is less than 80% after reconditioning
- There is a difference of greater than 20 after performing a battery Shelf Life Test
- · There is physical damage to the battery case
- The Battery Support System indicates FAULTY when you try to recharge the battery.

To promote awareness of battery recycling, Physio-Control NiCad batteries are marked with one of these symbols:





When a Physic-Control NiCad battery has reached the end of its useful life, recycle the battery as follows.

Battery Recycling in the USA

Recycle NiCad batteries by participating with Physio-Control Corporation in a national battery recycling program. Contact your Physio-Control representative to obtain shipping instructions and battery shipping containers. Do not return your batteries to the Physio-Control Corporate Headquarters in Redmond, Washington, unless instructed to do so by your Physio-Control representative.

Battery Recycling Outside the USA

Recycle NiCad batteries according to national and local regulations. Contact your local Physio-Control representative for assistance.

Troubleshooting

If a problem is detected during operation or testing, refer to the troubleshooting tips in the appropriate table:

Monitor/recorder problems

Table 6-4 on page 6-14

Defibrillator problems

Table 6-5 on page 6-16

Pacemaker problems

Table 6-6 on page 6-17

If the problem cannot be corrected, remove the monitor from active service and contact a qualified service technician for service and repair.

Table 6-4 Troubleshooting the monitor and recorder

when signal is applied or CAL is pressed.

Table	The state of the s	r
	servation	Corrective Action 4
s	Device does not function when POWER switch is turned to a power source. No race on cardioscope.	 Confirm batteries are fully charged and secured in battery wells. Check battery pins in selected battery well for signs of damage. (Power may be intermittent in some cases.) If using auxiliary power module, confirm it is
p	nterference on cardioscope when using patient ECG cable.	 connected to line power and to defibrillator/monitor. Check patient ECG cable connection to electrodes and patient. Check for damaged patient ECG cable. Check patient skin preparation, electrode contact, electrode placement and electrode expiration date. Check for presence of a strong radio frequency electrical field (such as diathermy, radio signals, etc.). If possible, turn off or move noise-generating equipment. Check whether paddles lead is selected. Select lead I, It, or III when using patient ECG cable. If excessive line frequency (50 or 60Hz) interference is suspected in DIAG, select notch frequency and enable the built-in notch filter via set-up menu. Contact a qualified service representative for assistance.
se (0	excessive interference (noise) on monitor creen with paddle monitoring QUIK-LOOK with standard paddles or NUIK-COMBO electrode monitoring).	 Check for paddle electrode surface dirt. Confirm paddles lead is selected. If using QUIK-COMBO electrodes, check for proper skin preparation, electrode contact, electrode placement, or expired electrodes. Confirm that material used between paddles and skin is appropriate for defibrillation. Confirm standard paddles are properly stowed in paddle wells if using the QUIK-COMBO electrodes. Confirm that paddle wells are clean.
u	Poor ECG signal on cardioscope when sing patient ECG cable. However, CAL oes provide a 1mV pulse on cardioscope.	 Confirm lead I, II, or III is selected (not paddles lead). Confirm electrodes are positioned correctly. Check for defective patient ECG cable.
5 S	Straight line on cardioscope and recorder	Increase ECG SIZE.

Table 6-4 Troubleshooting the monitor and recorde	
Observation	Corrective Action
 No ECG signal on cardioscope when using patient ECG cable. 	Confirm lead I, II, or III is selected (not paddles).
using patient EOG cable.	Check patient ECG cable. Confirm ECG electrodes are not positioned too sleep.
	 Confirm ECG electrodes are not positioned too close together
7 No ECG signal on monitor screen with	Confirm paddles lead is selected.
paddle monitoring (QUIK-LOOK with standard paddles or QUIK-COMBO	Discharge defibrillator into Battery Support System test lead to about and the and into a it.
electrode monitoring).	load to check paddle cord integrity. Discharge defibrillator into QUIK-COMBO Patient
	Simulator to check therapy cable integrity.
Recorder does not advance paper.	Replace battery with fully-charged battery.
	 Check/replace paper roll. Make sure paper is correctly loaded.
	 Recorder operating outside of specified operating temperature range. Allow device to cool down or warm up.
	Paper not loaded correctly.
9 ECG recording appears wrinkled.	 Check paper. ECG paper may be loaded improperly.
10 ECG recording appears smudged.	 Confirm correct ECG paper is in use. Use only paper designed for thermal array recorders.
11 No systole sound.	Increase QRS VOL (powers up at zero volume).
	 Increase ECG SIZE. Gain may be too low for proper QRS detection.
	 Select another lead or change electrode position. ECG amplitude may be too low in selected lead.
12 No SYNC marker on cardioscope when	Confirm SYNC selected.
sync mode is selected.	 Increase ECG SIZE. Gain may be too low for proper QRS defection.
	 Select another lead or change electrode position. ECG amplitude may be too low in that lead.
	 Reprep skin and apply new electrodes.
13 SYNC indicator does not blink.	 Increase ECG SIZE. Gain may be too low for proper QRS detection.
	 Select another lead or change electrode position. ECG amplitude may be too low in that lead.
14 SYNC marker not positioned within QRS complex.	 Adjust ECG SIZE until QRS indicator is properly positioned.
	 Select another lead or change electrode position. ECG amplitude may be too low in that lead.
15 Heart rate is not displayed.	 Increase ECG SIZE. Gain may be too low for proper QRS detection.
	 Select another lead or change electrode position. ECG amplitude may be too low in that lead.
·	 Noninvasive pacing in progress (heart rate display replaced by pacing rate).
	Patient's heart rate less than 20 bpm.
	 Reprep skin and apply new electrodes.
16 LOW BATTERY indicator remains flashing	Replace battery with fully-charged battery.
despite attempts to charge battery. However, device operates normally using auxiliary power module.	Use auxiliary power module.

Observation 17 Device shuts down with brief or no LOW BATTERY Indicator.	Corrective Action Battery is damaged, improperly maintained or depleted. (May occur if battery is very low on charge and defibrillation is attempted.) Switch to fully-charged battery or auxiliary power module.
18 Time or date on recorder incorrect (or 00ERR00 annotated).	Reset clock as described on page 1-7.
19 Service Indicator appears continuously on status display.	 Device requires service by qualified service technician. (It is normal for service indicator to appear while setting the clock.)

Table 6-5 Troubleshooting the defibrillator

	ble 6-5 Troubleshooting the defibrillator oservation	Corrective Action
1	Charge time to 360 joules exceeds 12 seconds.	 Replace battery with fully-charged battery. Use auxiliary power module. Allow device to warm up to 10°C (50°F).
	Energy is not delivered to patient when both paddle discharge buttons are pressed (using standard paddles or QUIK-COMBO electrodes).	 Device is in SYNC mode and no QRS complexes are detected. Defibrillator has not yet reached selected energy level (wait for tone indicating full charge). More than 60 seconds have elapsed since charge done tone. Energy has been internally removed. Energy has been internally removed because the ENERGY select dial was changed after charge was complete. Discharge defibrillator into Battery Support System test load to confirm paddle cord integrity. If using QUIK-COMBO electrodes: Confirm standard paddles are properly stored in paddle wells. Confirm therapy cable is properly connected. If cannot correct problem, try defibrillating patient through standard paddles as a backup.
3	AVAILABLE ENERGY does not match energy selected when defibrillator is fully charged.	Defibrillator is out of calibration. Contact a qualified service technician.
··4	Numbers do not appear or scroll very slowly in AVAILABLE ENERGY window in status display when CHARGE pressed.	 Replace battery with fully-charged battery. Connect device to auxiliary power module if available.
5	AVAILABLE ENERGY flashes and scrolls to zero after defibrillator discharge.	 Paddles discharged into open air. Confirm proper paddle pressure and contact is maintained during discharge. Possible failure in defibrillator discharge pathway (connectors, cables, etc.).
6	Patient didn't "jump" (no muscle response) during defibrillator discharge.	 No action specified. Patient muscle response is variable and depends on patient condition. Lack of visible response to defibrillation does not necessarily mean the discharge did not occur.

6-16

Table 6-6 Houbleshooting the pacernake	Table 6-6	 Troubleshooting the pacemake
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	Troubleshooting the pacema	aker
Observatio		Corrective Action
	does not function when is pressed.	 Replace battery with fully-charged battery. Use auxiliary power module if available.
	light on, but START/STOP es not illuminate when i.	 Pacing lead off. Check for LEADS message displayed. Inspect QUIK-COMBO cable and electrode connections. Paddles lead selected; select another lead.
3 Pacing	stops spontaneously.	 PACER power off. Press PACER to apply power. Detection of an internal failure has occurred. The pacemaker is inoperative and requires service by a qualified service technician. QUIK-COMBO electrode off. Check for LEADS message. Check QUIK-COMBO cable and electrode connections. Paddles lead selected. Select lead I, II, or III and reinitiate pacing. CHARGE has been pressed. Use of radio equipment while pacing may cause current delivery to stop and the service message to appear accompanied by tones. Press PACER to turn off pacemaker and discontinue service message and tones. To reinitiate pacing, follow steps as outlined in Pacing Procedure. To minimize radio interference, move radio farther away from defibrillator/monitor. If unable to move radio away, reorient the radio. Replace battery with fully-charged battery or use auxiliary power module if available.
4 No ECC	G trace on monitor.	 Confirm ECG leads are connected and lead I, II, or III is selected (not paddles lead). Check ECG cable and patient/electrode connections. Check proper power source is selected.
	scope displays ence while pacing.	 ECG electrodes not optimally placed with respect to pacing electrodes. ECG signal may be difficult to interpret at higher pacing rates. Select another lead (I, II, or III). Patient response to pacing is highly variable with respect to capture threshold and ECG distortion. Consider changing pacing rate. Consider moving ECG electrodes away from pacing electrodes to optimize patient response and ECG signal integrity.
	e does not occur with stimulus.	 Increase pacing current level. (Administer sedation/analgesia as needed.) Check pacing electrode placement. Consider invasive pacing. Patient response to pacing therapy (noninvasive and invasive) is dependent upon many factors. Perform Noninvasive Pacemaker Test to confirm pacemaker is delivering energy.
7 LEADS	message appears.	 Check for proper use of patient ECG cable during pacing. Select leads I, II, or III. Inspect QUIK-COMBO cable and electrode connections.
	QRS complexes not when pacing.	 Adjust ECG SIZE until sense markers are properly positioned. Amplitude of ECG signal too low in that lead. Select another lead (I, II, or III) or move ECG electrodes. Intrinsic QRS complexes are occurring during pacemaker's refractory period.

Service and Repair

/ WARNING

Possible shock

Do not attempt to remove the instrument cover to service or repair this instrument. Contact qualified service personnel for service or repair.

If the LIFEPAK 10C defibrillator/monitor/pacemaker requires service as indicated by testing, troubleshooting, or the service indicator, contact the local Physio-Control service representative. In the USA, call Physio-Control Technical Services at 1-800-442-1142.

When calling Physio-Control to request service, identify model and serial number and describe the observation. If the device must be shipped to a service center or the factory, pack the device in the original shipping container, if possible, or in protective packing to prevent shipping damage.

The LIFEPAK 10C defibrillator/monitor/pacemaker Service Manual provides detailed technical information to support service and repair by qualified service personnel.

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Warranty

Refer to the warranty statement which is included in the accessory kit shipped with the product. For duplicate copies, contact the local Physio-Control representative. In the USA call 1-800-442-1142.

Supplies, Accessories, and Training Tools

Supplies, accessories, and training tools for the LIFEPAK 10C defibrillator/monitor/pacemaker are listed in Table 6-7. For information about ordering, contact the local Physio-Control representative. In the USA, call 1-800-442-1142.

Table 6-7 Supplies, accessories, and training tools

Table 6-7 Supplies, accessories, and harring tools	
Description	Part Number
FASTPAK battery	09-10424
Recorder paper, 50mm	804700
LIFE◆PATCH ECG electrodes	800139
QUIK-COMBO pacing/defibrillation/ECG electrodes	806086
Pediatric paddle, external (2 required)	800418
Posterior paddle	802461
Battery Support System	801807
Battery Support System wall bracket assembly	802562
QUIK-COMBO Therapy Cable Tester	805550
QUIK-COMBO 3-lead patient simulator	806223
QUIK-COMBO 12-lead patient simulator	806395
12-Lead ECG Adapter	805600
Cables:	
Patient ECG Cable, 3-Lead (AHA, 90-degree angle connector)	805400
Patient ECG Cable, 3-Lead (IEC)	800947
Literature:	
LIFEPAK 10C defibrillator/monitor/pacemaker Operating Instructions	3004087
LIFEPAK 10C defibrillator/monitor/pacemaker Service Manual	3005330
Booklet Noninvasive Pacing: What You Should Know	805074
Booklet Defibrillation: What You Should Know	805662
Battery Reconditioning Procedure check sheet	806017
Battery Shelf Life Test check sheet	806018
Battery Maintenance Log check sheet	806019

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APPENDIX A: SPECIFICATIONS

Table A-1 lists the specifications for the LIFEPAK 10C defibrillator/monitor/pacemaker.

Table A-1	LIFEPAK	10C defibrillator/monitor/pacemake	r Specifications
E00 NA0	ヘトロナヘロ		

ECG	MON	III C	Ж
	-00		~ c

ECG Lead Selection

Paddles, I, II, or III

Input

Isolated ECG via QUIK-LOOK defibrillation paddles, QUIK-COMBO

electrodes, or 3-lead patient ECG cable

Electrical Isolation and

Shielding

Input protected against high voltage defibrillator pulses and radio frequency interference per FDA Standard MDS-201-0004. RF interference depends

on distance from RF source, radio output power, radiating efficiency,

vehicle environment, etc.

3-Lead ECG Cable Length

QUIK-COMBO Therapy

Cable Length

4.0m (13ft) total length; 3.0m cable (10ft) with 0.9m leads (3ft)

3.0m (10ft)

Common Mode Rejection

Minimum 100dB with respect to chassis ground at 60Hz, 65dB minimum with respect to isolated ground when using 3-lead patient ECG cable

CARDIOSCOPE DISPLAY

Size

72.5mm (2.85 in) x 43.5mm (1.7 in)

Sweep Speed

Frequency Response

25mm/sec

Monitor (domestic): 1 to 30Hz, -3dB Monitor (agency): 0.5 to 25Hz, -1.4dB

Expanded freq. response while recorder in DIAG mode: 0.05 to 30Hz, -3dB

Paddles: 2 to 20Hz, -3dB

STRIP CHART RECORDER

Paper Size

50mm x 30m (100 ft)

Paper Speed

25mm/sec

Frequency response:

Monitor (domestic): 1 to 30Hz, -3dB

Monitor (agency): 0.5 (-1.4dB) to 40 (-3dB) Hz

Diagnostic: 0.05 to 100Hz, -3dB Paddles: 2 to 20Hz, -3dB

CODE SUMMARY frequency response:

Domestic: 1 to 30Hz Agency: 0.5 to 40Hz

Annotation:

Includes time, date, lead, gain, heart rate, defibrillation and/or pacing

arameters

CODE SUMMARY critical

event record:

Digitally stored record of critical ECG and device parameters

STATUS DISPLAY

Heart Rate (bpm):

3-digit readout displays rates from 20 to 295 bpm

Available Energy: Pacing Rate:

0-360 joules 40-170bpm

Pacing Current:

0-200mA

DIAG message:

Indicates recorder frequency response is 0.05 to 100Hz, -3dB

MONITOR CONTROLS

ECG SIZE

Adjusts ECG gain

QRS VOL

CAL

Adjusts loudness of QRS beeper Sends calibration pulse to monitor input

CODE SUMMARY

Activates CODE SUMMARY printout

LEAD SELECT

Selects ECG Input: Paddles, I, II, or III

RECORD

Activates strip chart recorder; activates diagnostic mode if held for more

than 1 second (when enabled)

SYNC FREEZE

Triggers energy delivery to patient's QRS complex Momentarily halts ECG trace on the cardioscope

Table A-1	LIFEPAK 10	C defibrillator/monitor/	pacemaker S	pecifications ((cont.)	
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1V/mV at x1 gain .		
1V/mV at x1 gain .		
1400Hz ±2% center frequency, 1Vrms ±10%		
Matches strip chart recorder		
5msec monophasic pulse (Edmark) per AAMI spec		
0, 5, 10, 20, 50, 100, 200, 300, 360 joules		
360 joules in less than 12 seconds above 0°C (32°F)		
Adult: 82cm²		
Pediatric: 16cm²		
2.3m (7.5 ft)		
Energy discharge within 20msec of sync marker on cardioscope (triggers to patient-generated QRS complex)		
Both paddles: energy discharge buttons		
STERNUM: ENERGY select dial rotates to select 0 to 360 joules		
APEX: CHARGE (with indicator light) initiates charging RECORD activates strip chart recorder		
40 to 170 bpm		
±1.5% over entire range		
Monophasic, truncated, exponential current pulse 20 ±1msec duration measured at output current ≥10mA peak.		
0 to 200mA ±10% or 5mA (whichever is greater) for a load of 0 to 800 ohms		
Pacing rates Refractory period		
40-90 340msec ± 3%		
100 300msec ± 3%		
110–120 250msec ± 3%		
130-140 220msec ± 3%		
150-170 200msec ± 3%		
Standby: 5 to 55°C (41 to 131°F)		
Operating: -10 to 55°C (14 to 131°F) after minimum 2-hour storage at standby temperature		
Storage (exclusive of batteries): -30 to 65°C (-22 to 149°F)		
0 to 95% (non-condensing) from 0 to 34°C (32 to 93.2°F)		
0 to 80% (non-condensing) from 35 to 55°C (95 to 131°F)		
797 to 439mmHg (~570 to +15,000 ft)		
Helicopter Aircraft: MIL-STD-810D, method 514.3 (category 6). Test levels		
per US Army Aeromedical Research Laboratory Report no. 91-14, section 2.6.3 (March 1991), (UH-1 helicopter, floor under co-pilot's seat).		
Fixed-Wing, Turboprop Transport: (take off and climb). Test level of 0.0016g²/Hz, the maximum level per figure 32(31) of ECRI Report, contract		
no. 223-77-5035, prepared for FDA (April 1979).		
With carrying case (soft case), passes drops of 43 inches from the handle (30 inches from case). This exceeds test levels per ECRI report, contract no. 223-77-5035, prepared for FDA (April 1979).		
MIL-STD-108E and IEC 601-2-4		
6; / 1 1 5 /		

Table A-1 LIFEPAK	10C defibrillator/monitor/p	pacemaker Specifications	(cont.)
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BATTERY	3 NiCad batteries, 12V, 1.0 amp hours each. A single new battery registering at least 100% capacity on the Battery Support System will provide at a minimum:	
	 45 minutes of monitoring, or 	
	• 20 minutes of pacing, or	
	 25 discharges at 360 joules per battery. 	
SIZE		
Height	13.3cm (5.3in)	
Width	40.6cm (16in)	
Depth	37cm (14.6in)	
Weight	10kg (22lbs)	

All specifications at 25°C unless otherwise stated.

△: CAUTION

(CAUTION Possible equipment damage).
If a help prevent component damage, the device should not be mounted near vibration sources such as engine struts and landing gear.

APPENDIX B: BASIC OPERATION CHECKLIST

After reviewing this manual, you may use the checklists in Tables B-1 through B-4 to familiarize yourself with the use of QUIK-COMBO electrodes. If you cannot perform any of these steps, review the appropriate sections of this manual. For other operation support, refer to your local Physio-Control Corporation representative. To properly complete all of these steps, you will need the following equipment:

- LIFEPAK 10C defibrillator/monitor/pacemaker with fully-charged battery
- 3-Lead Patient ECG Cable
- QUIK-COMBO electrodes
- Physio-Control QUIK-COMBO 3-lead patient simulator

Table 8-1 Basic Operation Checklist-Monitoring Procedure

13-60	
	Completed Completed
1	Identify the QUIK-COMBO electrode placement for monitoring from the electrode
	package label.
2	Remove electrodes from package and connect to QUIK-COMBO therapy cable.
3	Identify which electrode should be placed in lateral position on patient's chest, and
	which electrode should be placed in anterior position on chest.
4	Disconnect QUIK-COMBO therapy cable from QUIK-COMBO electrodes and connect
•	to simulator.
5	Press simulator ON button and confirm the NSR indicator light is on.
6	Turn the defibrillator POWER switch to the power source.
7	Confirm the standard paddles are securely stored in paddle wells before monitoring.
8	Press LEAD SELECT to select paddles lead.
9	Confirm the cardioscope displays a normal sinus rhythm.
	Turn the defibrillator POWER switch to OFF.
10	Turn the delibriliator FOWEN Switch to OPF.
11	Press simulator OFF button to remove power.

Table B-2 Basic Operation Checklist-Defibrillation Procedure

Tab.	ie B-2 Basic Operation Checklist-Defibrillation Procedure
Şt	P Completed S
1	Identify the two possible QUIK-COMBO electrode placements for defibrillation from the electrode package label.
2	Connect QUIK-COMBO therapy cable to simulator.
3	Press simulator ON button to apply power.
4	Press simulator VF button and confirm the VF indicator light is on.
5	Turn the defibrillator POWER switch to the power source.
8	Press LEAD SELECT to select paddles lead and monitor ECG through the QUIK-COMBO therapy cable.
7	Rotate the ENERGY dial on the STERNUM paddle to select 200 joules.
8	Press CHARGE on the APEX paddle to charge the defibrillator.
9	Confirm the standard paddles are securely stored in paddle wells before discharge.
10	Confirm the cardioscope displays a shockable rhythm (VF).
11	After the defibrillator is fully charged (tone sounds), loudly announce to anyone else present "Stand clear!," then simultaneously press both paddle discharge buttons to discharge energy.
12	Turn the defibrillator POWER switch to OFF. (Unwanted charge may be internally discharged by turning POWER to OFF or rotating the ENERGY dial on the STERNUM paddle.)
13	Press simulator OFF button to remove power.

Table B-3 Basic Operation Checklist-Synchronized Cardioversion Procedure

St		Completed
1	Identify the two possible QUIK-COMBO electrode placements for cardioversion from the electrode package label.	
2	Turn the defibrillator POWER switch to the power source.	
3	Select one of the electrode placements:	
	A Anterior-Lateral	
	B Anterior-Posterior	
4	Connect QUIK-COMBO therapy cable to simulator.	
5	Select ECG monitoring method:	
	Anterior-Lateral: Select paddles lead; press simulator ON button and confirm NSR is selected (NSR light is on), or press V-TACH button to select V-TACH (V-TACH light is on).	
	Anterior-Posterior: Connect 3-lead patient ECG cable to defibrillator and simulator; press LEAD SELECT to select lead i, II, or III; press simulator ON button and confirm NSR is selected (NSR light is on), or press V-TACH button to select V-TACH (V-TACH light is on).	
6	Press SYNC and confirm the SYNC message flashes off with each detected QRS complex.	
7	Observe the cardioscope and confirm that sense markers appear on each QRS complex. If sense markers do not appear, or appear elsewhere on the ECG, press ECG SIZE as needed to properly adjust (or select another lead if using 3-lead cable). If using QUIK-COMBO electrode placement A and still cannot obtain proper sense markers, connect 3-lead patient ECG cable to defibrillator and simulator; press LEAD SELECT to select lead I, II, or III, and adjust ECG SIZE.	Weblan
8	Rotate the ENERGY dial on the STERNUM paddle to select 100 joules.	
9	Press CHARGE on the APEX paddle to charge the defibrillator.	
10	Confirm the standard paddles are securely stored in paddle wells before discharge.	
11	After the defibrillator is fully charged (tone sounds), loudly announce to anyone else present "Stand clear!"	<u></u>
	Simultaneously press and hold both paddle discharge buttons until the defibrillator discharges on the next QRS complex. Release the discharge buttons. (Unwanted charge may be internally discharged by turning POWER to OFF or rotating the ENERGY dial on the STERNUM paddle.)	
13	Turn the defibrillator POWER switch to OFF.	
14	Press simulator OFF button to remove power (and disconnect 3-lead patient ECG cable if used).	

Table B-4 Basic Operation Checklist-Noninvasive Pacing Procedure

labi	le B-4 Basic Operation Checklist-Noninvasive Pacing Procedure	
1	ep Identify the two possible QUIK-COMBO electrode placements for pacing from the electrode package label.	Completed:
2	Turn the defibrillator POWER switch to the power source.	
3	Connect the 3-lead patient ECG cable to defibrillator and simulator.	
4	Press LEAD SELECT to select lead I, II, or III.	
5	Connect QUIK-COMBO therapy cable to simulator.	
6	Press simulator ON button, then press BRADY and confirm the BRADY indicator light is on.	
7	Confirm the cardioscope displays a bradycardia rhythm.	
8	Press PACER and confirm the PACER indicator light is on.	
9	Press RATE to select a pacing rate of 80.	
10	Observe the cardioscope and confirm that sense markers appear on each QRS complex. If sense markers do not appear, or appear elsewhere on the ECG, press ECG SIZE to properly adjust or select another lead.	
11	Confirm the standard paddles are securely stored in paddle wells before pacing.	
12	Press START/STOP to start pacing.	
13	Press CURRENT to increase current until electrical capture is observed.	
14	Press START/STOP to stop pacing.	
15	Turn the defibrillator POWER switch to OFF.	
16	Press simulator OFF button to remove power; disconnect cables.	

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INDEX

A

Accessories, 6-19

Anterior-Lateral Placement QUIK-COMBO electrodes, 5-3 Standard paddles, 4-3

Anterior-Posterior Placement QUIK-COMBO electrodes, 5-4 Standard paddles, 4-3

APEX paddle definition, 1-3

Applying ECG electrodes, 2-3

Applying QUIK-COMBO electrodes, 5-5, 5-6

AUX connector, 1-3

11

Basic Operation Checklist, B-2

Batteries. See Battery Maintenance and Testing

Battery Maintenance and Testing, 6-7
Battery description, 6-8
Battery Maintenance Log, 6-12
Battery Support System, 6-8
Installing a battery, 6-9
NiCad battery performance factors, 6-8
Self-discharge, 6-8
Voltage depression, 6-8

Receiving new batteries, 6-13

Reconditioning Procedure, 6-10 Recycling batteries, 6-13 Shelf Life Test, 6-11 Storing batteries, 6-13 Battery Maintenance Log, 6-12

Battery Support System, 6-8

CAL button definition, 1-4

Cardioversion. See Synchronized cardioversion

Checklist for basic operation, B-2

Cleaning, 6-3

Clock setting, 1-7

CODE SUMMARY Record, 3-3
Description, 3-3
Event storage priority, 3-6
Format of reports, 3-4
Printing procedure, 3-3

Controls, indicators, connectors, 1-2

Date/time setting, 1-7

Defibrillation
Pacing and defibrillation, 5-12
Paddle placement, 4-3
Pediatric paddles, 4-3

Posterior paddles, 4-3
QUIK-COMBO electrodes, 5-6
Standard paddles procedure, 4-4
Synchronized cardioversion with
QUIK-COMBO electrodes,
5-7

Synchronized cardioversion with standard paddles, 4-5

Troubleshooting, 6-16 Warnings, 4-2

Defibrillator Test. 6-4

DIAG message description, 1-6

DIAG Mode, 3-2

DIAG mode activation. See RECORD button definition

Diagnostic Mode, Recording, 3-2

글 그는 건강하다 하셨다는 하네요. 한

ECG electrode placement, 2-3 Interference from invasive pacemakers, 2-4

ECG electrodes

Applying, 2-3

Required type, 2-4

ECG Monitoring Procedure 3-Lead ECG, 2-3

QUIK-COMBO electrodes, 5-5 Standard paddles, 2-2

ECG output. See AUX connector

Electrode placement

3-Lead ECG Monitoring, 2-3 QUIK-COMBO electrodes, 5-3

Implanted defibrillator, Standard paddle placement for defibrillation, 4-4

Inspection, 6-3

Invasive pacemakers Damage from defibrillation, 4-4 Interference with monitoring, 2-4

LEADS message description, 1-6

Low battery indicator, 1-3

M

Maintenance and testing Battery maintenance, 6-7

Cleaning, 6-3

Inspection, 6-3

Overview, 6-2

Schedule for maintenance, 6-2 Troubleshooting, 6-14

MIEMSS modulator connector, 1-3

Monitor/Recorder Test, 6-4

Monitoring Procedure 3-Lead ECG, 2-3

QUIK-COMBO electrodes, 5-5 Standard paddles, 2-2

N

Noninvasive Pacemaker Test, 6-6

Noninvasive Pacing, 5-8

Assessing for capture, 5-10

Defibrillation during pacing, 5-12

ECG distortion during pacing, 5-11

ECG monitoring during pacing, 5-9

Pacemaker refractory period, 5-11

Pacing interruption, 5-12

Patient response, 5-9

Procedure, 5-8

Troubleshooting, 6-17.

Operation Checklist, B-2

Ordering replacement parts, 6-19

Pacemakers

Damage from defibrillation, 4-4 Interference with monitoring, 2-4 Pacing, 5-8

See also Noninvasive Pacing

Procedure, 5-8

Paddle Placement, 4-3

Anterior-Lateral, 4-3

Anterior-Posterior, 4-3

Near implanted defibrillator, 4-4

Near implanted pacemaker, 4-4

Defibrillation with standard paddles, 4-4

Monitoring, 2-2

Paper loading in recorder, 3-2

Patient transfer and QUIK-COMBO electrodes, 5-12

Periodic maintenance. See Maintenance and testing

Preparing skin for electrodes

3-Lead ECG Monitoring, 2-3

QUIK-COMBO electrodes

Defibrillation, 5-6

Monitoring, 5-5

Printer

See also Recorder

Paper loading, 3-2

Q

QRS Detection ***

Adjusting for ECG monitoring, 2-3

Summary description, 2-4

QUIK-COMBO Electrodes

Applying electrodes, 5-5, 5-6

Cable connection, 5-5

Defibrillation procedure, 5-6

Electrode placement, 5-3

Anterior-Lateral, 5-3

Anterior-Posterior, 5-4

Near implanted defibrillators, 5-4 Near implanted pacemakers, 5-4

Obese patients, 5-4

Thin patients, 5-4

Monitoring procedure, 5-5

Noninvasive pacing, 5-8

Assessing for capture, 5-10

Defibrillation during pacing, 5-12

ECG distortion during pacing, 5-11

ECG monitoring during pacing,

5-9

Pacemaker refractory period,

5-11

Pacing interruption, 5-12

Patient response, 5-9

Patient transfer, 5-12

Removing and replacing electrodes, 5-4

Summary description, 5-2

Synchronized cardioversion procedure, 5-7

Warnings, 5-2

R

Reconditioning Procedure, 6-10 RECORD button definition, 1-4

Recorder

Annotations, 3-2

Handling recordings, 3-3

Paper loading, 3-2

Recorder paper, 6-19

Recording, Diagnostic mode, 3-2

Recording Procedure, 3-2

Recycling batteries, 6-13

Refractory Period, Noninvasive pacing, 5-11

Repair, 6-18

Replacement parts, 6-19

S

Schedule for maintenance, 6-2

Self-Discharge for batteries, 6-8

Sense markers, 4-5

Service and repair, 6-18

Service indicator, 1-6, 6-18

Setting the clock, 1-7

Shelf Life Test, 6-11

Skin preparation

3-Lead ECG monitoring, 2-3

QUIK-COMBO electrodes

Defibrillation, 5-6 Monitoring, 5-5

Specifications, A-2

Standard paddles

Defibrillation, 4-4

Monitoring, 2-2

STERNUM paddle definition, 1-3

Supplies and accessories, 6-19

Symbol definitions, ix

Sync markers. See Sense markers

Synchronized Cardioversion

QUIK-COMBO electrodes, 5-7

Standard paddles, 4-5

Synchronous Cardioversion Test, 6-6

Testing

Defibrillator test, 6-4

Monitor/Recorder test, 6-4

Noninvasive pacemaker test, 6-6

Schedule, 6-2

Synchronous cardioversion test,

6-6

Time/date setting, 1-7

Troubleshooting
Defibrillator, 6-16
Monitor and recorder, 6-14
Pacemaker, 6-17

·V

Voltage depression, 6-8

Warnings
Defibrillation with paddles, 4-2

General warnings, viii

Monitoring, 2-2

QUIK-COMBO electrodes, 5-2

Warranty, 6-18

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