The LIFEPAK 20 defibrillator/monitor is the ideal crashcart defibrillator putting early, effective defibrillation into the hands of BLS responders while transitioning easily to full therapy and monitoring capabilities for the ALS-trained code team.

The world leader in defibrillation technology brings you the improved LIFEPAK 20 defibrillator/monitor, created especially for hospitals and clinics. Designed with extensive input from clinicians around the world, the 20 is simple yet sophisticated, providing a flexible and effective tool to meet diverse defibrillation and monitoring needs.

**User Enhancements**

The 20 is highly intuitive, making it easy for infrequent AED-trained responders to quickly understand and use. Our proven Shock Advisory System™ guides the user with voice and visual prompts through each step and simple 1-2-3 operation. With an easy push of a latch, the 20 automatically converts to a manual defibrillator, including waveform displays. In manual mode, all of the advanced cardiovascular life support tools become available. The 20 offers noninvasive pacing, Masimo SET pulse oximetry ($\text{SpO}_2$), ECG monitoring (3- or 5-wire), and synchronized cardioversion.

Designed for indoor use, the 20 is compact, lightweight, and easy to rush to the scene or use during transport. A convenient, optional docking station enables the device to be firmly attached to the crash cart for safe and rapid transport, or easily released—whatever the situation requires. The docking station swivels, for multiple viewing angles.

Superb color graphics, easily viewed from many angles, enable clinicians to clearly see color matched waveforms and values for efficient and informed patient care decisions.
The LIFEPAK 20 defibrillator/monitor has seven main operating modes:

**Manual Mode:** Provides a normal operating capability for ALS users. Allows access to manual mode energy selections up to 360J, synchronous cardioversion and pacing. ECG waveform is displayed.

**AED Mode:** (Consistent with 2005 AHA Guidelines for CPR and ECC and ERC Guidelines for Resuscitation 2005.) Provides a normal operating capability for BLS users. All user features are available except manual defibrillation, synchronous cardioversion, pacing, and access to archived patient records. Provides shock energy defaults up to 360J. User selectable option to display ECG waveforms and/or visual AED prompts.

**Setup Mode:** Allows the operator to configure the device settings.

**Service Mode:** Allows the operator to execute diagnostic tests and calibrations, to display device module software and hardware versions, and to display and print the diagnostic code log.

**Inservice Mode:** Simulated waveforms are available for demonstration purposes. The waveforms consist of short segments of realistic data, which are repeated to form a continuous waveform.

**Auto Test Mode:** Performs daily self tests.

**Daily Auto Test:** Each day at approximately 0300 (3:00 am), the 20 automatically completes the following tasks:

- Turns itself on.
- Performs self-tests.
- Changes to a low energy level and then discharges through a test load.
- Tests the pacing circuitry if noninvasive pacing installed.
- Turns itself off.

### POWER

The device is an AC line operated device with an internal battery as backup.

**AC Powered:** 90–132 VAC 50/60 Hz, 198–264 VAC 50/60 Hz, total power draw less than 120 volt-amperes (VA).

**Internal Battery Backup:** NiMh. Batteries charge while device operates from AC Power.

**Operating Time:** A new fully charged internal backup battery will provide the following prior to shutdown:

<table>
<thead>
<tr>
<th>Task</th>
<th>Time (Minutes)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Monitoring</td>
<td>120</td>
</tr>
<tr>
<td>Monitoring in device without</td>
<td>135</td>
</tr>
<tr>
<td>Pulse Oximeter (minutes)</td>
<td>90</td>
</tr>
<tr>
<td>Defibrillation (360J)</td>
<td>70</td>
</tr>
<tr>
<td>Typical Battery Charge Time</td>
<td>&lt;2 hours</td>
</tr>
</tbody>
</table>

### SPECIFICATIONS

**Low Battery Indication and Message:** When the device is unplugged from AC power, it switches to battery. When battery gets low, the battery detection icon is indicated with a low battery message in the status area, and a warning tone occurs.

**Service Indicator:** When error detected.

### PHYSICAL CHARACTERISTICS

**Weight:**
- Fully featured defibrillator/monitor (pacing and SpO₂): 5.58 kg (12.3 lbs)
- QUIK-COMBO cable: .20 kg (.43 lbs)
- Standard (hard) paddles: .88 kg (1.95 lbs)

**Height:** 21.3 cm (8.4 in)

**Width:** 26.2 cm (10.3 in)

**Depth:** 26.2 cm (10.3 in)

### DISPLAY

**Size (active viewing area):** 115.18 mm (4.53 in) wide x 86.38 mm (3.4 in) high

**Resolution:** 320 x 240 dot color active LCD.

**Display symbol:** ---

**Out of Range Indication:** +/− 3 digits (during motion conditions)

**Pulse Rate Accuracy:** +/− 2 digits (during no motion conditions)

**Pulse Rate Range:** 20–350 BPM digital display

**SpO₂ Waveform Display sweep speed:** 25 mm/sec for ECG

### DATA MANAGEMENT

The device can easily print a CODE SUMMARY™ report, including an introduction with patient information and critical event record. The summary report also includes event and vital signs log, and waveforms associated with certain events. The device can print archived patient records and has two data communication ports—infared (IRDA) and a direct serial port, which supports a serial data cable.

### COMMUNICATIONS

The device is capable of transferring data records by serial connection:

- EIA/TIA-232E compatible at 9600, 19200, 38400, 57600 and 115200 bps

### ALARMS

**Quick Set:** Activates alarms for all parameters

**VF/VT Alarm:** Activates continuous CPSS monitoring in Manual Mode

**ECG**

ECG can be monitored through 3-wire or 5-wire ECG cables.

Standard paddles or therapy electrodes (QUIK-COMBO pacing/defibrillation/ECG electrodes or FAST-PATCH® disposable defibrillation/ECG electrodes) are used for paddles lead monitoring.

Compatible with LIFEPAK 12 ECG and therapy cables.

**Lead Selection:**
- Leads I, II and III, (3-wire ECG cable)
- Leads I, II, III, AVR, AVL, and AVF, V (c) acquired simultaneously, (5-wire ECG cable)

**ECG size:** 4, 3, 2.5, 2, 1.5, 1, 0.5, 0.25 cm/mV

**Heart Rate Display:** 20–350 BPM digital display

**Out of Range indication:** Display symbol “---”

**Heart symbol flash** for each QRS detection.

**Continuous Patient Surveillance System (CPSS):** In AED mode, while Shock Advisory System is not active, CPSS monitors the patient, via QUIK-COMBO paddles or lead II ECG, for potentially shockable rhythms.

**Voice Prompts:** Used for selected warnings and alarms (Configurable On/Off)

**Analog ECG Output:** 1V/mV X 1.0 gain < 30 ms delay

**Common Mode Rejection:** 90db at 50/60 Hz

**SpO₂**

Masimo SET Sensors

**Suction Range:** 1 to 100%

**Saturation Accuracy:** (70–100%) (0–6% unspecified)

**Adults/Pediatrics:** +/- 2 digits (during no motion conditions)

**Neonates:** +/- 3 digits (during motion conditions)

Dynamic signal strength bar graph

Pulse tone at the onset of the pleth waveform

**SpO₂ Update Average Rate:** User selectable 4, 8, 12 or 16 seconds

**SpO₂ Measurement:** Functional SpO₂ values are displayed and stored

**Pulse Rate Range:** 25 to 240 pulses per minute

**Pulse Rate Accuracy:** (Adults/Pediatrics/Neonates) +/- 3 digits (during no motion conditions)

**SpO₂ waveforms with autogain control**
**FREQUENCY RESPONSE**

**Diagnostic:** 0.05 to 0.15 Hz or 0.05 to 40 Hz (user configurable)

**Monitor:** 0.67 to 40 Hz or 1 to 30 Hz (user configurable)

**Paddles:** 2.5 to 30 Hz

**Analog ECG Output:** 0.67 to 32 Hz (except 2.5 to 30 Hz for paddles ECG)

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**DEFIBRILLATOR**

**Waveform:** Biphasic Truncated Exponential, The following specifications apply from 25 to 200 ohms, unless otherwise specified.

**Energy Accuracy:** ±1 joule or 10% of setting, whichever is greater, into 50 ohms ±1 joule or 15% of setting, whichever is greater, into any impedance from 25–100 ohms.

**Voltage Compensation:** Active when disposable therapy electrodes are attached. Energy output within ± 5% or ± 1 joule, whichever is greater, of 50 ohm value, limited to the available energy which results in the delivery of 360 joules into 50 ohms.

**Patient Impedance**

<table>
<thead>
<tr>
<th>Patient Impedance</th>
<th>Phase 1 Duration (ms)</th>
<th>Phase 2 Duration (ms)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Min.</td>
<td>Max.</td>
</tr>
<tr>
<td>25</td>
<td>5.1</td>
<td>6.0</td>
</tr>
<tr>
<td>50</td>
<td>6.8</td>
<td>7.9</td>
</tr>
<tr>
<td>100</td>
<td>8.7</td>
<td>10.6</td>
</tr>
<tr>
<td>125</td>
<td>9.5</td>
<td>11.2</td>
</tr>
</tbody>
</table>

**Paddle Options**

- QUIK-COMBO pacing/defibrillation/ECG electrodes (standard)
- Standard adult paddles with embedded pediatric paddles (optional)
- Internal handles with discharge control (optional)
- External sterilizable paddles (optional)
- FAST-PATCH disposable defibrillation/ECG electrodes (optional)

**Cable length:** 2.44 meter (8-foot) long QUIK-COMBO cable (not including electrode assembly)

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**AED**

**Shock Advisory System (SAS):** An ECG analysis system that advises the operator if the algorithm detects a shockable or nonshockable ECG rhythm. SAS acquires ECG via therapy electrodes only.

**Shock Ready Time:** Using a fully charged battery at normal room temperature, the device is ready to shock within 16 seconds of power on, if initial rhythm finding is “Shock Advised.”

The AED mode of the LIFEPAK 20 defibrillator/monitor is not intended for use on children less than 8 years of age.

**cprMAX technology Setup Options** (items marked with * are default settings):

- **Stacked Shocks:** Off*, On
- **Initial CPR:** Off*, Analyze First, CPR First
- **Preshock CPR:** Off*, 15, 30 seconds
- **Auto Analyze:** Off*, After 1st Shock, On
- **Pulse Check:** Never*, After Second No Shock Advised, After Every No Shock Advised, Always
- **CPR Time 1 & 2:** 15, 30, 45, 60, 90, 120*, 180 seconds, 30 minutes

Users should refer to the LIFEPAK 20 defibrillator/monitor operating instructions for details on how to customize the configuration of their devices to hospital protocols.

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**PACER**

**Pacing Mode:** Demand or nondemand

**Rate and current defaults (user configurable)**

**Pacing Rate:** 40 to 170 ppm

**Rate Accuracy:** +/- 1.5% over entire range

**Output Waveform:** Monophasic, amplitude stable to +/- 5% relative to leading edge for currents greater than or equal to 40 mA, Duration 20 +/- 1 ms, Rise/Fall times <= 1 ms [10–90% levels]

**Output Current:** 0 to 200 mA

**Pause:** Pacing pulse frequency reduced by a factor of 4 when activated

**Refractory Period:** 200 to 300 ms +/- 3% (function of rate)

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**ENVIRONMENTAL**

**Temperature, Operating:** 5 to 45° C (41 to 113° F)

**Temperature, Nonoperating:** -20 to +60° C (-4 to +140° F) except therapy electrodes

**Relative Humidity:** 5 to 95%, noncondensing

**Atmospheric Pressure, Operating:** Ambient to 429 mmHg (0 to 4,572 meters) (0 to 15,000 feet)

**Water Resistance, Operating (without accessories except for ECG Cable and hard paddles):** IPX1 (spillage) per IEC 60601-1 clause 44.6 (1995)

**Vibration:** MIL-STD-810E Method 514.4, Cat1

**Shock (Drop):** 1 drop on each side from 457.2 mm (18 in.) onto a steel surface

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**EMC**

IEC 60601-2-4:2002; Clause 36, Particular Requirements for the Safety of Cardiac Defibrillators and Cardiac Defibrillator monitors

All specifications are at 20° C (68° F) unless otherwise stated.
**ADAPTIV Biphasic Technology**

The 20 is equipped with ADAPTIV biphasic technology, which adjusts the shock waveform duration and voltage based on the patient’s impedance level. In AED mode, the device can provide escalating shocks (200–360 joules), depending on the patient’s needs. In manual mode, the ALS clinician can give energy shocks according to established protocols.

**cprMAX Technology**

The 20 in AED mode is also equipped with cprMAX technology, which supports the 2005 American Heart Association (AHA) Guidelines for Cardiopulmonary Resuscitation (CPR) and Emergency Cardiovascular Care (ECC) and European Resuscitation Council (ERC) Guidelines for Resuscitation 2005. cprMAX technology is aimed at optimizing the timing and sequencing of CPR and defibrillation. The 20 with cprMAX technology is highly flexible to accommodate various patient and CPR protocol requirements. cprMAX empowers users to customize the interaction of defibrillation and CPR, with options to:

- Provide a specified CPR interval before delivering the first shock.
- Provide CPR while the device is charging.
- Prompt for CPR after each single shock.
- Customize the option for and timing of pulse checks.

**Use and Maintenance**

QUIK-COMBO pacing/defibrillation/ECG electrodes are compatible with the entire LIFEPAK family of products, for standardization and continuity of care. Standard adult paddles with embedded pediatric electrodes, sterilizable adult paddles, and internal paddles provide flexible therapy options for response in various cardiac emergencies. The 20 is easy to maintain and service, comes with a five-year, in-hospital warranty, and is equipped with AC power and a backup internal nickel metal hydride (NiMh) battery.

The LIFEPAK 20 defibrillator/monitor extends and enhances the LIFEPAK family of products, providing flexible, compatible and standardized defibrillation solutions for patients across a range of hospital settings.