

CPRmeter 2

User Guide



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Intended Use

The CPRmeter 2 with Q-CPR[®] technology is a small, lightweight device powered by replaceable batteries. The device is intended for use by responders who have been trained in CPR and use of the CPRmeter 2.

When attached to the bare chest of a suspected victim of sudden cardiac arrest (SCA), the CPRmeter 2 provides real-time feedback on CPR compressions in accordance with current CPR guidelines. It displays CPR feedback indicators for depth, release, and rate of chest compressions. It also counts the number of compressions in a series, and provides notification of lack of expected CPR activity.

If in doubt about the appropriateness for use, perform CPR without using the CPRmeter 2.

Indication for Use

The CPRmeter 2 is used as a guide in administering cardio-pulmonary resuscitation (CPR) to a suspected SCA victim at least 1 years old.

Important Information



Read this User Guide and become familiar with the operation of the product prior to use. Use the product only as described in this User Guide.

Warnings and Cautions

A Warning states a condition, hazard, or unsafe practice that can result in serious personal injury or death.

A Caution states a condition, hazard, or unsafe practice that can result in minor personal injury or damage to the product.

Notes

A note states important information about the product or its operation.

Warning

The CPRmeter 2 is not intended for use on SCA victims under 1 years old.

Note

CPR cannot assure survival, no matter how well it is performed. In some patients, the underlying problem causing the cardiac arrest is not survivable despite any available care.

Items Included



CPRmeter 2

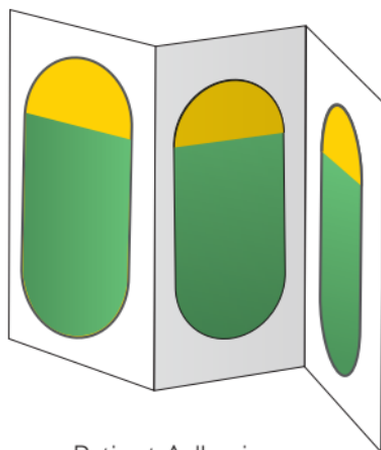


AAA Batteries

Items Included



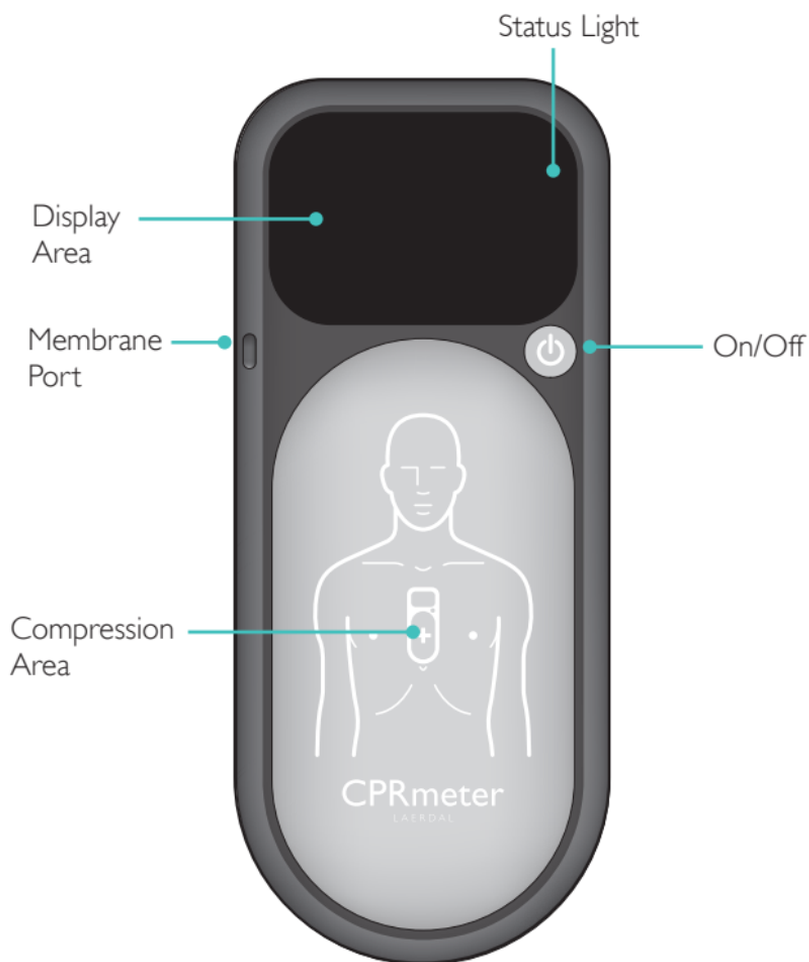
Protective Sleeve



Patient Adhesives

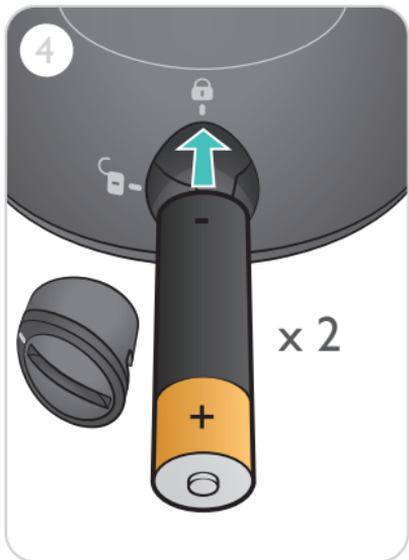
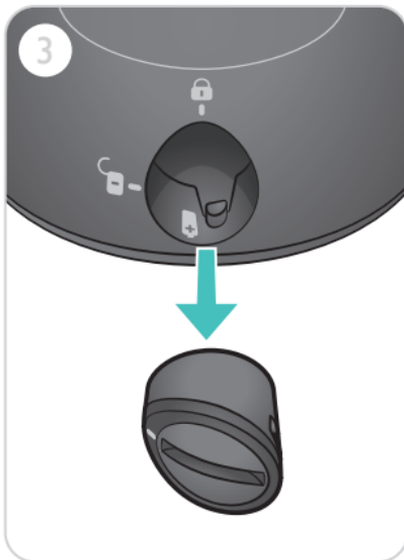
Items included may vary in appearance and are subject to change. Visit www.laerdal.com for more information including latest product downloads, spare parts and accessories.

Overview

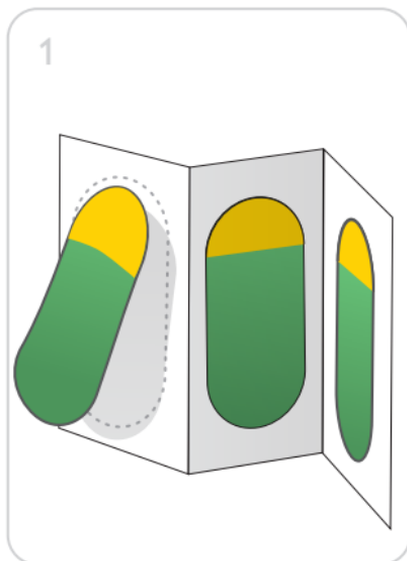




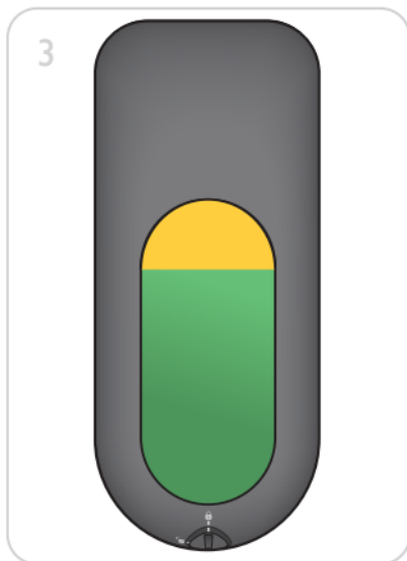
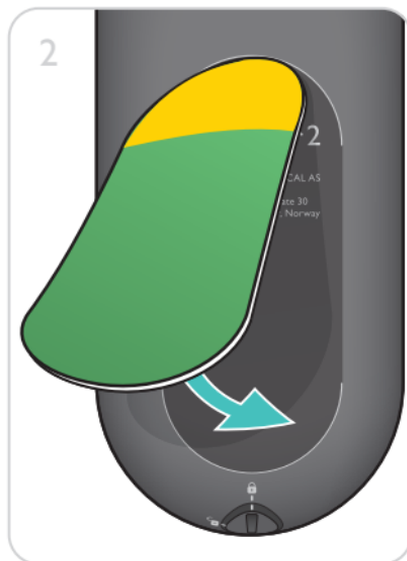
Setup - Insert Batteries



Setup - Apply Patient Adhesive



 **Caution**
Ensure Patient Adhesives are within their expiration date. Adhesives should be removed from the device and disposed of after 2 years.

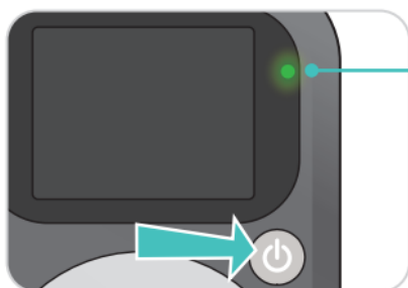


Getting Started

Notes

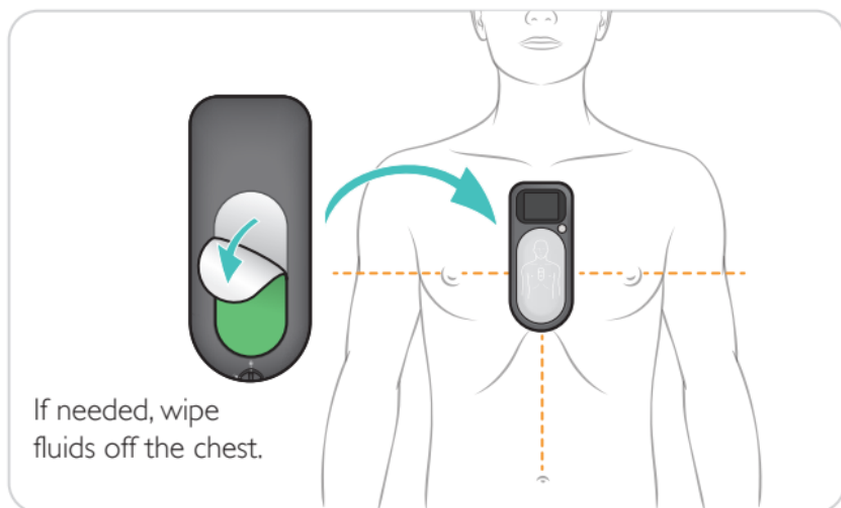
- Remove the device from its protective cover.
- Ensure the patient is on a firm surface.
- Remove clothing from the patient's chest.

Turn On



Status Light turns green for a few seconds, when CPRmeter 2 is turned on.

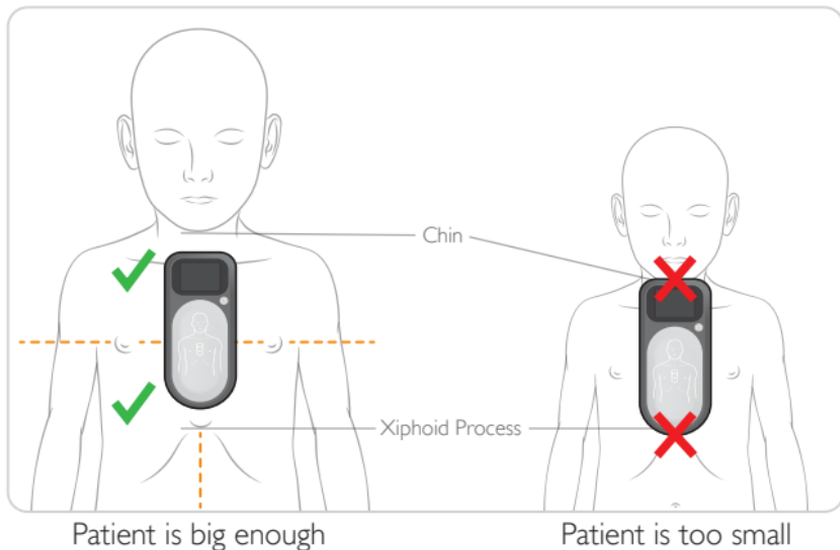
Place CPRmeter 2



If needed, wipe fluids off the chest.

Evaluate Fit

Place the CPRmeter 2 with the centre of the compression area aligned with the nipples. If the device touches the chin or xiphoid process, the patient is too small. Remove the CPRmeter 2 and perform CPR without it.



⚠ Warnings

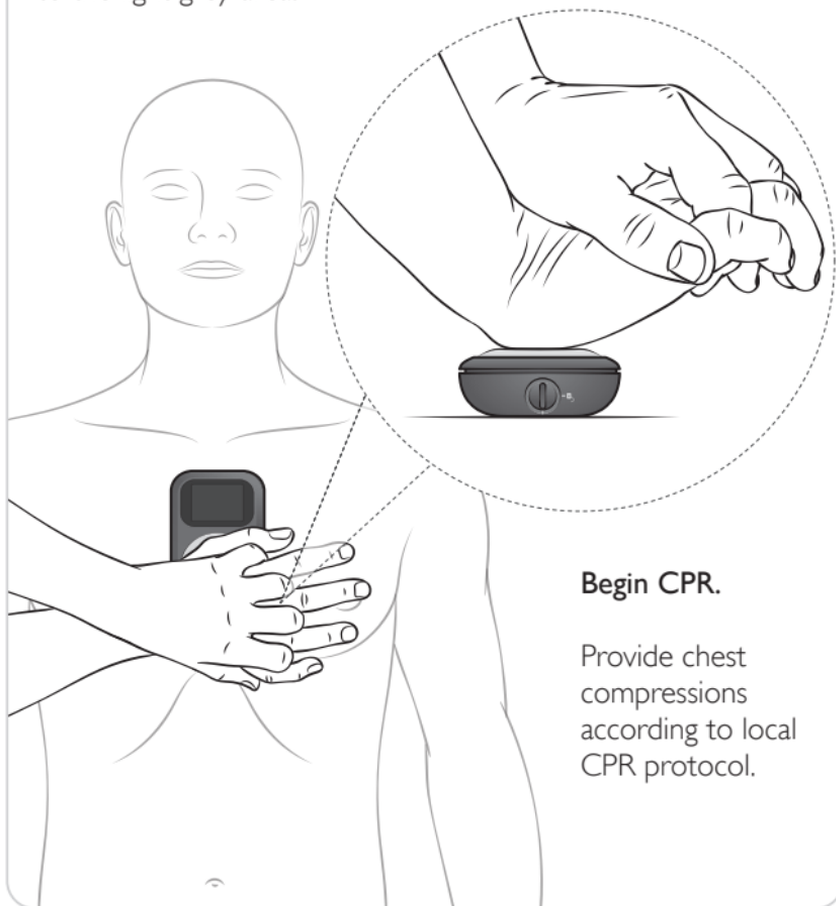
- Do not use the CPRmeter 2 on patients where the CPRmeter 2 comes into contact with the airway or the xiphoid process.
- Ensure that the compression point is in the middle of the chest.



Do not use the CPRmeter 2 on children under the age of 1 years.

Getting Started

Use the heel of the hand and apply pressure to the light grey area.



Begin CPR.

Provide chest compressions according to local CPR protocol.

 **Caution**

If the CPRmeter 2 moves during use, re-position it to the centre of the chest, as shown.

Warnings and Cautions

Warnings

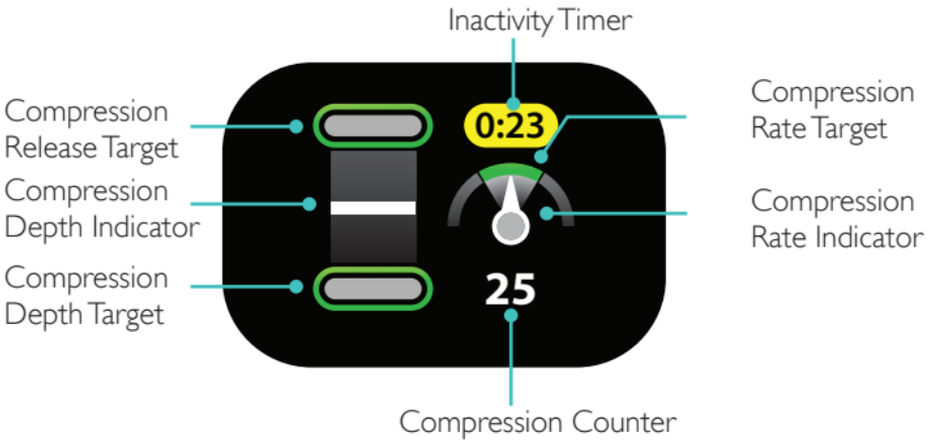
- *Do not use the device in conjunction with any mechanical or automated compression device.*
- *Do not use the device on top of defibrillation pads, unless the manufacturer of the defibrillator and the defibrillation pads has explicitly stated that the device can be used in such manner.*
- *Do not delay CPR. If there are any problems using the device, continue CPR without it.*
- *If the device appears to be damaged e.g. develops cracks or sharp edges, do not use.*

Cautions

- *If difficulty is encountered in applying the device, do not delay initiation of CPR. Remove the device and begin compressions.*
- *An orange status light indicates a technical error. If this occurs, stop using the CPRmeter 2 and continue CPR. Contact Laerdal technical support after the event.*

Contact a local Laerdal representative for further assistance, or visit www.laerdal.com for more information and Frequently Asked Questions (FAQ).

Feedback Display Overview



Compression Feedback

Depth

Adequate depth



Too shallow



Release

Adequate release



Incomplete release



Note

Release pressure fully between compressions.

Compression Feedback

On Soft Surface

If the CPRmeter 2 detects a compression that exceeds 60 mm (2.36"), it will show the depth indicator below the target area. If a specific CPR event requires CPR to be performed on a patient lying on a mattress, slide a backboard under the patient and compensate for the mattress softness by ensuring that for each chest compression the area below the compression depth target lights up.



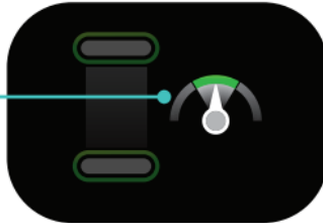
Warning

When performing CPR on a patient lying on a mattress, a backboard must be used to limit the amount of compressed depth which is absorbed by the mattress. Depending on characteristics of the mattress, backboard and patient, the depth compensation does not guarantee that the patient chest is compressed by 50 mm (2").

Compression Feedback

Rate

Adequate rate



Too slow



Too fast



Compression Counter

When compressions are started a counter will display grey up to 25 compressions.



Compression Counter

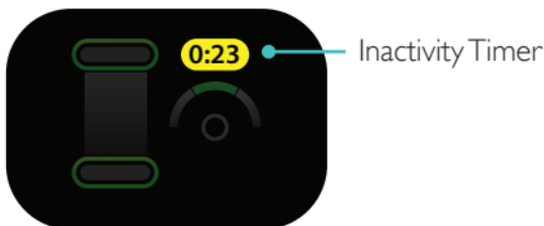
Compression Feedback



During a cycle of 30 compressions, the counter changes to solid white between 25 and 30 compressions. Beyond 30 compressions, the counter digits flash solid white for every tenth compression.

Without a compression the counter is reset after 3 seconds.

Inactivity

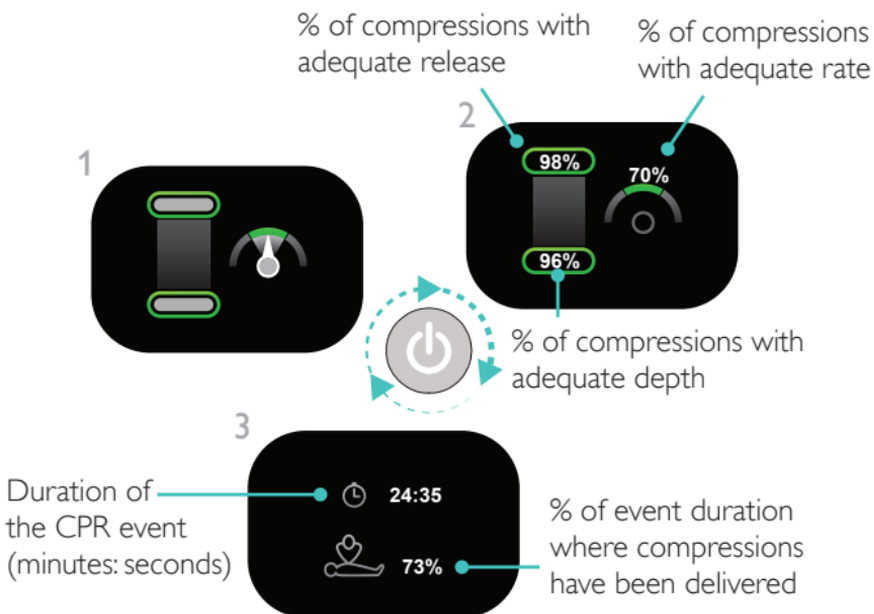


- After 3 seconds the CPRmeter 2 displays an inactivity timer which counts the seconds since the last compression.
- After 20 seconds since the last compression, the inactivity timer starts flashing.
- To conserve battery power the CPRmeter 2 display fades down after 1 minute. The display is restored when a new compression is delivered. The device shuts down automatically after 10 minutes without compressions performed.

QCPR Quick Review

The CPRmeter 2 can display CPR performance statistics for the last CPR event. After the device is turned on, press the On/Off button briefly to activate QCPR Quick Review. The statistics are shown over two displays.

Press the On/Off button briefly to cycle between the Compression Feedback and QCPR Quick Review screens.



The CPRmeter 2 reverts to Compression Feedback mode if a compression is delivered.

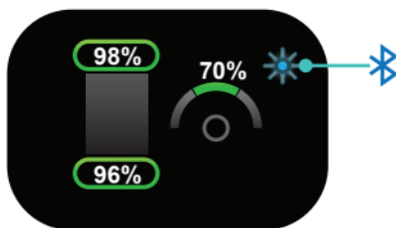
Debriefing

Notes

- The CPR event statistics are stored when the CPRmeter 2 is turned off. When turned on again, the statistics from the last stored CPR event can be reviewed.
- When the CPRmeter 2 is used in a new CPR event, the QCPR Quick Review will display the current event's statistics.
- CPR performance statistics are only calculated if at least 10 compressions have been delivered.

Wireless Data Transfer

The CPRmeter 2 has Bluetooth Low Energy functionality for uploading complete event data to an external device.



To connect a device, go to the QCPR Quick Review screen by pushing the On/Off button. The status light will flash blue indicating that Bluetooth is on and available for connection. When connected to a device, the flashing blue light turns steady. The CPRmeter 2 is now ready to transfer CPR performance data.

Note

Ensure Bluetooth connectivity is disabled during clinical use.

Battery Monitoring

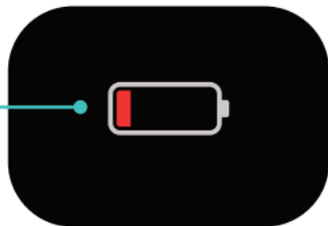
The CPRmeter 2 continuously monitors the power of its batteries. On a routine basis, particularly after periods of non-use, check the CPRmeter 2 battery status by turning on and checking if the low battery icon is displayed.

If the remaining power is estimated to be less than that required for a 30 minute CPR event, the visual indicators signal that the batteries should be replaced before the next use.



A low-battery icon appears when the CPRmeter 2 is being turned on.

A low-battery icon appears when the CPRmeter 2 is being turned off.



Routine Maintenance

1. On a routine basis check the battery status (as described).
2. Replace the batteries at least every 2 years.
3. On a routine basis, check that the CPRmeter 2 has a Patient Adhesive in place and that the liner remains on it. Replace the Patient Adhesive at least every 2 years if it is not used.

Maintenance and Cleaning

Warning

Do not interrupt CPR to replace the battery. Continue CPR without feedback from the CPRmeter 2.

Notes

- *If the remaining battery power during use becomes too low to sustain further operation, the low-battery icon is shown for 10 seconds and then the CPRmeter 2 turns itself off.*
- *After removal of low-batteries, wait 10 seconds before inserting new batteries.*

After Each Use

After use on a patient, the CPRmeter 2 may be contaminated and should be handled appropriately.

1. Place the contaminated CPRmeter 2 in a plastic bag until it can be cleaned (do not insert a contaminated CPRmeter 2 into its casing).
2. If it is visibly soiled, wipe the CPRmeter 2 with a soft cloth or paper towel to remove as much contamination as possible.
3. Remove the Patient Adhesive from the back of the CPRmeter 2.
4. Clean the CPRmeter 2 as described under Cleaning and Disinfection. Proper cleaning is required to achieve disinfection.
5. Check the exterior of the CPRmeter 2 for signs of damage. Contact Laerdal to arrange for replacement if needed.
6. Apply a new Patient Adhesive to the device as described in Setup- Apply Patient Adhesive.

Maintenance and Cleaning

Cleaning after Training on Manikin

If the CPRmeter 2 has been used for training on a manikin, it can be wiped using an alcohol wipe with minimum 70% ethanol.

Cleaning and Disinfection after Clinical Use

If the device has been used in a clinical situation, the device must be cleaned and disinfected.

Cleaning

1. Prepare a cleaning solution by mixing 5 ml of a mild dishwashing liquid in 4 l of warm tap water (40 °C – 50 °C).
2. Dip a small brush (e.g. toothbrush) in the cleaning solution and scrub the device for a minimum of 2 minutes.
3. If the membrane port is clogged, use the wet brush to remove any obstruction.
4. Wipe the exterior with a soft cloth dampened in lukewarm water (22 °C - 40 °C).

Disinfection

1. The exterior can be disinfected in a 0.55% solution of ortho-phthalaldehyde, or a 70% solution of isopropyl alcohol (isopropanol). Spray the exterior surfaces with the solution and ensure it remains wet for a minimum of 12 minutes. Repeat spraying as necessary to account for evaporation.
2. Wipe the exterior at least 3 times with a clean damp cloth to remove all traces of the disinfectant.
3. Allow to dry completely.

Maintenance and Cleaning

Caution

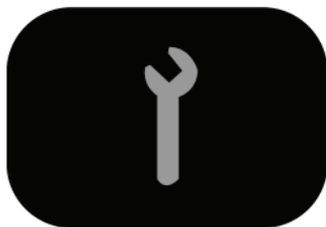
Do not immerse the CPRmeter 2 in water, hold it under running water, or allow moisture to penetrate it. Do not sterilize the CPRmeter 2.

Storing the CPRmeter 2 between use

Store the CPRmeter 2 in its protective cover to shield the display screen from scratches and to protect the patient adhesive from damage. Ensure that the On/Off button can not be inadvertently activated during storage.

Customer Service Indicator

If the Customer Service Indicator appears on the CPRmeter 2 at shutdown, contact a local Laerdal representative for further support.



CPR Targets

Category	Specification
Compression Depth Target	> 50 mm (2") Depth accuracy: $\pm 10\%$
Compression Release Target Force	< 2.5 kg (5.5 lbs) Force accuracy: +1.5 kg to - 2.0 kg (+3.3 lbs to - 4.4 lbs)
Compression Rate Target	100 to 120/min ± 3 /min

CPRmeter 2 [REF 801-002xx]

The CPRmeter 2 meets the performance requirements of IEC 60601-1, 3rd edition.

Category	Specification
Dimensions	153 mm x 64 mm x 25 mm (6.0" x 2.5" x 1.0")
Weight	163 g (5.7 oz) (excluding batteries)
Batteries	2 x 1.5V Size AAA (LR03)
Temperature	Transport and Storage: -20 °C to 70 °C (-4 °F to 158 °F) Operation: 0 °C to 50 °C (32 °F to 122 °F) Transient temperature: -20 °C to 70 °C (- 4 °F to 158 °F)
The time required for CPRmeter 2 to warm from the minimum storage temperature between uses until it is ready for intended use is minimum 15 minutes at room temperature.	
The time required for CPRmeter 2 to cool from the maximum storage temperature between uses until it is ready for intended use is minimum 15 minutes at room temperature.	
Sealing	Meets IP55 rating as defined by standard ISO/IEC 60529
Relative Humidity	Transport: 5% to 95% Storage: 5% to 75% Operation: 5% to 95%
Atmospheric Pressure (Atm.p.)	Transport, Storage and Operation: 1060 to 617 mbar (1060 to 617 hPa)

Specifications

Electromagnetic Compatibility	Meets IEC 60601-1-2 and RTCA/DO-160F Section 21 Category M
Defibrillation Protection	The CPRmeter 2 is defibrillation protected, type BF patient connection.

CPRmeter Patient Adhesives [REF 801-10850]

These CPRmeter 2 Patient Adhesives are disposable and are for single patient use only. Do not re-use. Re-use will lead to increased risk of cross contamination, and/or degradation of adhesive performance.

Category	Specification
Dimensions	39 mm x 90 mm (1.5" x 3.5")
Temperature	Transport and Storage: -20 °C to 70 °C (-4 °F to 158 °F) Operation: 0 °C to 50 °C (32 °F to 122 °F)
Relative Humidity	Transport: 0% to 75% Storage: 0% to 75% Operation: 0% to 95%
Material	Foam pad with biocompatible adhesive on each side.
Shelf Life	2 years when applied to the CPRmeter 2 or 4 years in unopened package. Do not exceed the expiration date on the packaging.

Environmental Considerations

Product	Information
CPRmeter 2	The CPRmeter 2 contains electronic components. Dispose of it at an appropriate recycling facility in accordance with local regulations.
CPRmeter Patient Adhesive	The used Patient Adhesive may be contaminated with body tissue, fluid, or blood. Dispose of it as infectious waste.

Symbol Glossary

Symbol	Definition
	CE Marking
	Do not re-use
	Defibrillation Protection
	Manufacturer
	Waste Electrical and Electronic Equipment (WEEE). Dispose of in accordance with your country's requirements.
	Reference order number
	Ingress Protection rating - protected from limited dust ingress and low pressure water jets from any direction.
	Expiration date
	Not made with natural rubber latex
	Warning/Caution
	Note
	Temperature limitations for transport/storage
	Contains number of CPRmeter 2 Patient Adhesives shown as "#."
	Consult Directions for Use

Symbol Glossary

	Peel symbol
	Do not use on children under 1
	Australian RCM mark
	Bluetooth symbol
	CSA Certification Mark

Warnings

- When the CPRmeter 2 is used together with a defibrillator, make sure to follow the defibrillator manufacturer's instructions. Stop compressions, remove hands from the CPRmeter 2 and remain clear of all patient contact during defibrillation or when otherwise required, in accordance with a proper defibrillation protocol.
- The CPRmeter 2 is not intended for use in a moving environment, such as an air, sea or road ambulance. If used during patient transport, the device may provide inaccurate feedback. If CPR is indicated in a moving environment, do not rely on the depth feedback during such conditions. It is not necessary to remove the device from the patient.
- Do not practice CPR by using the CPRmeter 2 on a person. It may be used with a training manikin or simply on a compliant surface for practice.
- Properly performed CPR may result in fracturing of the patient's ribs.¹ If rib integrity has been compromised, continue to provide CPR in accordance with your local protocol.
- Properly performed CPR may result in chest injuries¹ e.g. external chest wall bruising or abrasion.
- Do not rely on CPRmeter 2 feedback during aircraft ascent and descent, as its accuracy is reduced in such conditions.

Cautions

- Do not apply the CPRmeter 2 to an open wound or recent incision site.
- The device is designed to be used only with Laerdal-approved accessories and may perform improperly if non-approved accessories are used. Do not attempt to modify the device in any way.
- Use only model 801-10850 Patient Adhesives with the CPRmeter 2.
- The CPRmeter 2 is not a serviceable device. If experiencing technical difficulties contact a local Laerdal representative for support.

¹ Black CJ, Busuttill A, Robertson C. Chest wall injuries following cardiopulmonary resuscitation. Resuscitation. 2004;63:339 –343.

Regulatory Information



Note

Changes or modifications not expressly approved by Laerdal Medical could void the user's authority to operate the equipment.

Recommendation

Responders should receive training, including regular refresher training, in use of the CPRmeter 2. When training with the device on a CPR manikin, disable or ignore feedback from the manikin.



WEEE

This appliance is marked according to the European directive 2012/19/EU on Waste Electrical and Electronic Equipment (WEEE).

By ensuring this product is disposed of correctly, you will help prevent potential negative consequences for the environment and human health, which could otherwise be caused by inappropriate waste handling of this product.

The symbol on the product indicates that this appliance may not be treated as household waste. Instead it shall be handed over to the applicable collection point for the recycling of electrical and electronic equipment. Disposal must be carried out in accordance with local environmental regulations for waste disposal.

For more detailed information about treatment, recovery and recycling of this product, please contact your local city office, your household waste disposal service or the Laerdal representative where you purchased the product.



The product is in compliance with the essential requirements of Council Directive 93/42/EEC as amended by Council Directive 2007/47/EC and Council Directive 2014/53/EU.

Electromagnetic Conformity

Guidance and manufacturer's declaration: The CPRmeter 2 is intended for use in the electromagnetic environment specified in the tables below. The user of the CPRmeter 2 should assure that it is used in such an environment.

Electromagnetic Emissions

Emissions Test	Compliance	Electromagnetic Environment Guidance
RF CISPR 11	Group 1 Class B	<p>The CPRmeter 2 uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.</p> <p>The CPRmeter 2 is suitable for use in all establishments, including domestic establishments and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.</p>

Regulatory Information

Electromagnetic Immunity

Immunity Test	IEC 60601 Test Level	Compliance Level	Electromagnetic Environment - Guidance
Electrostatic Discharge (ESD) IEC 61000-4-2	± 6 kV contact ± 8 kV air	± 6 kV contact ± 8 kV air	There are no special requirements with respect to electrostatic discharge.
Power Frequency (50/60/400 Hz) Magnetic Field IEC 61000-4-8	3 A/m	3 A/m	Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial/hospital environment. There are no special requirements for non-commercial/non-hospital environments.
Radiated RF IEC 61000-4-3	3 V/m 80 MHz to 2.5 GHz	3 V/m	Portable and mobile RF communications equipment should be used no closer to any part of the CPRmeter 2, than is absolutely necessary. †, ‡ The recommended separation distances for various transmitters and the CPRmeter 2 are shown in the following table. Interference may occur in the vicinity of equipment marked with the following symbol: 

† The ISM (industrial, scientific and medical) bands between 150 kHz and 80 MHz are 6.765 MHz to 6.795 MHz; 13.553 MHz to 13.567 MHz; 26.957 MHz to 27.283 MHz; and 40.660 MHz to 40.700 MHz.

‡ Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast, and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the CPRmeter 2 is used exceeds the applicable RF compliance level above, the CPRmeter 2 should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as re-orienting or relocating the CPRmeter 2.

Recommended separation distances between portable and mobile RF communications equipment and the CPRmeter 2

The CPRmeter 2 is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the CPRmeter 2 can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the CPRmeter 2 as recommended below, according to the maximum output power of the communications equipment.

Regulatory Information

Electromagnetic Emissions

Rated maximum output power of transmitter [W]	Separation distance according to frequency of transmitter [m]		
	150 kHz to 80 MHz	80 MHz to 800 MHz $d=1.2\sqrt{P}$	800 MHz to 2.5 GHz $d=2.3\sqrt{P}$
0.01	NA	0.12	0.23
0.1	NA	0.38	0.72
1	NA	1.2	2.3
10	NA	3.8	7.28
100	NA	12	23

For transmitters rated at a maximum output power not listed above, the recommended separation distance d in meters (m) can be determined using the equation applicable to the frequency of the transmitter; where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer:

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

NOTE 2. The ISM (industrial, scientific and medical) bands between 150 kHz and 80 MHz are 6.765 MHz to 6.795 MHz; 13.553 MHz to 13.567 MHz; 26.957 MHz to 27.283 MHz; and 40.660 MHz to 40.700 MHz.

NOTE 3. An additional factor of $10/3$ is used in calculating the recommended separation distance for transmitters in the ISM frequency bands between 150 kHz and 80 MHz and in the frequency range 80 MHz to 2.5 GHz to decrease the likelihood that mobile/portable communications equipment could cause interference if it is inadvertently brought into patient areas.

NOTE 4. These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

NOTE 5. Transmitters/antennas of this power-level are most likely mounted on an emergency vehicle chassis. The distances cited here are for open field. For an external antenna, the separation distance is most likely shorter.

Warranty

The Laerdal CPRmeter 2 has a one-year limited Warranty. Refer to the Laerdal Global Warranty for terms and conditions.

About this edition

The information in this applies to the model 801-002xx CPRmeter 2. This information is subject to change.

The CPRmeter™ with Q-CPR® is protected by U.S. and International registered patents. The design of CPRmeter™ and its feedback symbols are protected in several jurisdictions under international design registrations.

CPRmeter™ and Q-CPR® are trademarks or registered trademark of Laerdal Medical AS.

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