

MEDUCORE Standard²

Monitor/Defibrillator

Instructions for use of devices from software version 3.9





Read these instructions for use before using the product. Ignoring the instructions for use may lead to serious injury or death.

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1 Introduction

1.1 About this document

This document describes all possible versions of the device.

Depending on the version purchased, it may be that you do not have available all the functions, components, and accessories described in this document and shown in the illustrations.

If functions of the device can only be enabled by purchasing a specific option, this document makes this clear by appending "(optional)" and "(only with option XXXX)" to the text.

Illustrations in these instructions for use are for general understanding, and may differ from the actual version. No claims can be brought on the basis of any deviations.

Texts shown in the device display appear in this document in bold. Example: Press the **Charge** function button.

The voice prompts of the device are shown in italics in this document. Example: *Stand clear of the patient*.

1.2 Explanation of warnings

DANGER

Danger!

DANGER indicates a dangerous situation which will result in death or serious injury if not prevented.

Warning!

WARNING indicates a dangerous situation which may result in death or serious injury if not prevented.

Caution!

CAUTION indicates a dangerous situation which may result in minor injury if not prevented.



Notice!

NOTICE indicates risks which may possibly cause material or environmental damage.



Designates useful tips relating to a particular sequence of actions.

1.3 Description of function

The device offers the following monitoring and diagnostic functions:

- 6-lead monitoring ECG: The electrical activity of the heart is derived and displayed. This allows the user to interpret cardiac rhythms and heart rate. To do so, the 6-lead monitoring ECG derives the limb leads according to Einthoven (I, II, III) and according to Goldberger (aVR, aVL, aVF) and displays these in a curve view.
- 12-lead ECG: The electrical activity of the heart is derived and displayed. This allows the user to carry out differentiated ECG diagnosis. The 12-lead ECG derives the following leads for this purpose:
 - Limb leads acc. to Einthoven (I, II, III)
 - Limb leads acc. to Goldberger (aVR, aVL, aVF)
 - Chest wall leads acc. to Wilson (V1 to V6)

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1 Introduction

- Pulse oximetry: Pulse oximetry monitoring allows continuous, non-invasive measurement of arterial oxygen saturation with the aid of different pulse oximetry sensors for different application sites. At the same time, a photosensor in the pulse oximetry sensor registers the percentage of oxygenated hemoglobin in the arterial blood (SpO₂) using different light wavelengths. In addition, the pulse oximetry sensor registers pulse rate. The values for SpO₂ and pulse rate are shown in the display numerically, the plethysmogram in the form of a curve.
- Non-invasive blood pressure measurement (NIBP monitoring): NIBP monitoring allows measurement of blood pressure on a limb in adults, children, and infants. Measurement is based on oscillometric blood pressure measurement technology. After the measurement has been carried out, systolic and diastolic blood pressures are shown numerically in the display in mmHg.

The device has the following therapy functions:

- Manual defibrillation and cardioversion: Based on the information displayed by the ECG, the user decides whether it is necessary to deliver a shock. If a shock is necessary, the user can select the shock energy, charge the device for shock delivery, and trigger the shock manually.
- Semi-automatic defibrillation: In AED mode, the device guides the user through resuscitation by means of audio and visual instructions. The device determines the resuscitation sequence. The device automatically carries out an ECG analysis and, if necessary, charges for electric shock delivery. The shock is triggered manually by the user.

2 Safety

The instructions for use are part of the device. If the instructions for use and the following safety instructions are not followed in their entirety, therapy may fail or be put at risk. This may cause serious or life-threatening injury or death to the patient, the user, and bystanders.

- \Rightarrow Follow instructions for use in their entirety.
- \Rightarrow Keep the instructions for use accessible and near the device at all times.
- \Rightarrow Use the device only for the intended purpose (see "2.1 Intended purpose", page 11).
- \Rightarrow Do **not** use the device if it is contra-indicated.
- ⇒ Follow the instructions for use of the components and accessories.

2.1 Intended purpose

MEDUCORE Standard² is used as a mobile external defibrillator with monitoring functions on patients who are exhibiting symptoms of a cardiac arrest or cardiac arrhythmias. MEDUCORE Standard² is also used when measured values of the measuring functions integrated in the device are to be recorded.

Patient target groups

MEDUCORE Standard² is suitable for use with emergency patients of any age (exception: semi-automatic defibrillation only from 1 year old upwards).

Intended users

Qualified medical personnel

Contra-indications

- Defibrillation may not be carried out on patients with a body temperature < 27 °C.
- Defibrillation may not be carried out on patients suffering ventricular tachycardia with a pulse.
- Cardioversion may not be carried out on patients who have overdosed on digitalis.

2.2 Intended areas of application

- Mobile use in emergency medicine or primary care in emergency response
- Transport between hospital rooms and departments
- Transport between hospitals and other premises by ambulance, aircraft, helicopter or ship

2.3 Possible side effects/complications

- Electrostimulation may cause burns, skin irritation and arrhythmias.
- Electrostimulation may damage implanted medical devices or those connected to the patient externally.

2.4 Exclusions and limitations of the intended purpose

The device has not been approved for the following applications:

- Operation in hyperbaric chambers
- Operation in conjunction with magnetic resonance imaging machines (MRI, MRT, NMR)
- Operation in conjunction with high-frequency electrosurgical equipment
- Electrostimulation in an electrically conductive and damp surroundings
- Electrostimulation in an oxygen-enriched atmosphere with simultaneous presence of combustible materials
- Electrostimulation in an atmosphere with flammable gases

2.5 Requirements on the user

The user must meet the following requirements:

- The user is medically trained and has the necessary technical knowledge and experience in the emergency medical treatment of patients.
- On the basis of this technical knowledge and experience, the user is in a position to perform the tasks assigned to him or her safely, and independently to recognize, assess and avoid potential risks to him or herself, to the patient or to bystanders.
- The user has been trained and has received instruction in the use of the device.
- The user is trained in extended measures to treat emergency patients and is able to use manual mode, cardioversion and interpretation of 12-lead ECGs as a function of the options purchased.

2.6 Safety information

2.6.1 Qualification

Warning

Risk of injury due to lack of knowledge and failure to follow guidelines!

Use of the device by users without medical qualifications and training in defibrillation/cardioversion and/or the failure to follow guidelines may result in injury to the patient, user or bystanders.

- \Rightarrow Only use the device if the user meets the requirements on the user (see "2.5 Requirements on the user", page 13).
- ⇒ Only use the device if the user is familiar with defibrillation/ cardioversion and the operation of the device.
- ⇒ Only use the device if the user is familiar with the enabled options.
- ⇒ Follow currently applicable guidelines on defibrillation/ cardioversion.
- ⇒ Observe national and regional provisions on defibrillation/ cardioversion.
- ⇒ Observe organizational guidelines on defibrillation/ cardioversion.
- \Rightarrow Include the patient condition when deciding on treatment.

2.6.2 How to use the device

Warning

Risk of injury if the device, components, and accessories are used in damp or electrically conductive surroundings!

Using the device, components, and accessories in damp or electrically conductive surroundings may result in an electric shock and injure the patient, user, and bystanders.

- ⇒ Only connect the device, components, and accessories in dry surroundings.
- ⇒ Only use the device, components, and accessories in surroundings that are not electrically conductive.
- ⇒ Keep conductive parts of the electrodes and plug connections away from other conductive parts and the ground.

Risk of injury due to ingress of liquids!

The device is only protected from water jets according to IP55 when the battery is inserted, the water jet protection for the SD card slot is closed, the cables and the NIBP connecting tube including NIBP cuff are connected. Ingress of liquids and dust may damage the device, components, and accessories.

- \Rightarrow Do not immerse the device, components, and accessories in liquids.
- \Rightarrow Only operate the device with the battery inserted.
- \Rightarrow Always close the water jet protection of the SD card slot.
- \Rightarrow Always leave cables connected.
- ⇒ Leave the NIBP connecting tube including NIBP cuff connected during use.

Risk of injury from malfunctions of the device, the components, and the accessories!

A damaged device, damaged components or damaged accessories may injure the patient, user, and bystanders.

- ⇒ Only operate the device, components, and accessories if they have no external damage.
- ⇒ Only operate the device, components, and accessories if the function check has been passed.
- \Rightarrow Do not leave the device and patient unsupervised.
- ⇒ In the event of device failure during resuscitation: Carry out cardiopulmonary resuscitation in accordance with currently applicable guidelines and obtain a replacement device.
- ⇒ In the event of device failure during a monitoring session: Monitor patient by monitoring breathing and taking pulse and obtain a replacement device if required.

Risk of injury due to device being inaccessible or alarm being hidden!

An inaccessible device or hidden alarm (alarm light, loudspeaker and display) mean that the user may not notice alarms and consequently react belatedly to hazardous situations. This may delay therapy and injure the patient.

- \Rightarrow Keep the device accessible at all times.
- \Rightarrow Always keep the alarm (alarm light, loudspeaker and display) clear.
- \Rightarrow Do not operate the device in a closed bag if this conceals the alarms.

Risk of injury due to alarm limits which are too high or too low!

Alarm limits which are either too high or too low may prevent the device triggering an alarm, thereby putting the patient at risk.

⇒ Always set alarm limits which have been adapted to the patient.

Risk of injury due to incorrectly set parameters or too few/too many enabled functions in the operator menu!

Incorrectly set parameters or too few/too many enabled functions in the operator menu may result in incorrect settings in the user menu or too limited/too extensive device functions. This may cause critical operating situations and injure the patient.

- ⇒ The operator menu should only be used by operators familiar with the settings in the operator menu and their impact on the user menu and device functions. Otherwise use the device with factory settings.
- \Rightarrow Adapt the device functions to the user's know-how.
- ⇒ Protect the operator menu with a secure access code (at least one letter and one number).

Risk of injury from incorrectly secured access to functions!

Too simple an access code gives the user access to functions with which he or she is not familiar or in which he or she is not trained. This may cause the patient or the user serious or life-threatening injury.

⇒ Protect functions with a secure access code (at least one letter and one number).

Risk of injury from operating the device, accessories, and components outside the prescribed ambient conditions!

Use of the device, accessories, and components outside the specified ambient conditions may lead to incorrect results to the extent that the device fails, injuring the patient.

- ⇒ Only operate the device within the prescribed ambient conditions (see "18 Technical data", page 385).
- ⇒ Allow the device, components, and accessories to acclimatize to operating temperature.

Disrupted and failed therapy due to incorrect use of disposables!

Reusing and reprocessing disposables might induce unpredictable reactions as a result of aging, embrittlement, wear, thermal stress, and chemical action. This may put the functionality and safety of the device at risk, and cause the patient and user serious or lifethreatening injury.

- \Rightarrow Do not reuse disposables.
- \Rightarrow Do not subject disposables to hygienic reprocessing.

Risk of injury from using third-party accessories!

Accessories which have not been approved by

WEINMANN Emergency may result in explosions, electric shocks, incorrect monitoring, impaired functions and a negative impact on interference immunity and interference emission or lead to material damage and injure the patient.

⇒ Only use accessories which have been approved by WEINMANN Emergency (see "17.2 Accessories and other parts", page 378).

Delay in treatment due to overly loud audio output!

When the defibrillator is used in conjunction with devices with audio output (e.g. alarm tones, voice prompts), overly loud audio output from one device may drown out audio output from the other device and thus delay treatment.

⇒ When using multiple devices with audio output at the same time, set the volume on the devices to the same level.

Risk of injury and treatment delay due to imperceptible alarm signals!

Alarm signals which are quieter than the ambient noise level prevent alarm situations from being detected. This may result in delayed treatment and thus in injury to the patient.

- \Rightarrow Always set device volume to be louder than ambient noise level.
- \Rightarrow Do not stack devices.

Therapy at risk due to inadequate patient monitoring!

If the patient and the device are not observed and monitored during therapy, delayed response by medical personnel to alarms and faults may result in serious or life-threatening injury to the patient and incorrect therapy.

⇒ Continuously observe and monitor the patient and device during therapy.

Disrupted or failed therapy due to modifications to the design of the device or accessories!

Modifications to the design of the device may result in disrupted or failed therapy. This may cause serious or life-threatening injury to the patient.

⇒ Do not make any modifications to the design of the device, components or accessories.

Risk of injury and delayed treatment from connecting the device, the components, and the accessories to several patients!

Connecting the device to several patients simultaneously may lead to measured values and shock delivery being incorrectly assigned. This may injure the patient.

⇒ Only connect the device, components, and accessories to one patient.

2.6.3 Power supply

DangerRisk of injury due to electric shock when the device is opened!The device contains a shock capacitor. Opening the device leads

to serious injuries or death from electric shock.

- \Rightarrow Do not open the device.
- ⇒ The device should only be opened by WEINMANN Emergency or by technicians authorized by WEINMANN Emergency.
- ⇒ Measures such as repairs and maintenance should only be carried out by the manufacturer or by a technician expressly authorized by it.

Warning Risk of injury due to electric shock when connecting an incorrect power supply unit and charger to line power!

The power supply unit and charger contains a safety device to prevent electric shock. The use of an unsuitable power supply unit and charger may result in injury to the user.

⇒ Only operate the device on line power using the power supply unit and charger recommended by WEINMANN Emergency.

Risk of injury due to missing, discharged or defective battery! A missing, discharged or defective battery impedes the therapy functions.

- \Rightarrow Carry out a function check before each use in order to check the battery.
- \Rightarrow Always have a charged, ready-to-use spare battery on hand.

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No therapy due to extended storage of the battery without recharging!

Storing the battery for an extended period of time without recharging may result in the rapid shutdown of and irreparable damage to the battery.

- ⇒ When the battery is stored in the device without a power supply: Charge battery every 3 months (see "15.2 Storing the battery", page 373).
- ⇒ If the battery is not stored in the device: Charge batteries with serial numbers < 20,000 every 5 months and batteries with serial numbers ≥ 20,000 every 9 months (see "15.2 Storing the battery", page 373).

Therapy fails due to battery being used for too long!

The battery has a lifetime of 2 years. Exceeding battery lifetime to a considerable extent may lead to sudden failure of the battery at low temperatures and when power consumption is high. \Rightarrow Replace battery after 2 years.

Risk of injury from damaged power supply unit and charger! When used in emergency vehicles, the power supply unit and charger may be damaged and fail to charge the battery properly. This may injure the patient.

- \Rightarrow Do not use the power supply unit and charger in emergency vehicles.
- \Rightarrow Use the power supply unit and charger only in indoor spaces.
- ⇒ Observe the technical data for the power supply unit and charger (see "18.6 Power supply unit and charger", page 390).

Disrupted or failed therapy due to removal of the battery during shock delivery!

Removing the battery during shock delivery may cause damage to the device and thus put therapy at risk.

⇒ Always leave the battery in the device while the device is delivering a shock.

Risk of injury due to inaccessible power plug!

An obstructed power plug cannot be pulled out in an emergency and may thus result in injury.

 \Rightarrow Keep the power plug and line power accessible at all times.

Risk of injury due to using the charging station in damp or electrically conductive surroundings!

Using the charging station in damp or electrically conductive surroundings or outside indoor areas may lead to electric shock and injure the user.

- \Rightarrow Only use the charging station in dry surroundings.
- \Rightarrow Only use the charging station in surroundings that are not electrically conductive.
- \Rightarrow Protect the charging station from ingress of liquids.
- \Rightarrow Only use the charging station indoors.

Impaired readiness for use as a result of a defective power cord or defective power supply unit and charger!

A defective power cord or a defective power supply unit and charger prevent the battery charging in the charging station and thus the readiness of devices operated with the battery.

⇒ Check the power cord and the power supply unit and charger regularly.

Risk of injury from electric shock!

The contacts in the battery compartment of the charging station are live. Touching the contacts may lead to electric shock and injure the user.

 \Rightarrow Do not touch the contacts in the battery compartment of the charging station.

Caution Risk of injury from touching the contacts in the battery compartment and the patient at the same time!

The contacts in the battery compartment are live. Touching the contacts and the patient at the same time may injure the user or the patient.

 \Rightarrow Only operate the device with the battery inserted.

Risk of injury due to trailing connecting cable!

Trailing connecting cables are a trip hazard, which may hinder operation and cause injuries.

- ⇒ During operation with line power, position the power cord so that it does not present a hindrance.
- ⇒ During 12 V operation, position the connecting cable so that it does not present a hindrance.

Risk of injury as a result of falsified measurement results when the line filter is not set correctly!

An incorrectly set line filter may falsify the measurement results of the ECG and lead to misinterpretations. This may injure the patient.

- \Rightarrow Adapt the line filter to suit the regional power supply network.
- \Rightarrow When assessing the ECG, take account of an activated line filter.

2.6.4 Defibrillation/cardioversion

Warning Risk of injury due to sparks during defibrillation/cardioversion in combination with oxygen/flammable gases and combustible materials!

During defibrillation/cardioversion in an oxygen-enriched atmosphere/an atmosphere containing flammable gases and in the presence of combustible materials (e.g. textiles), sparks generated by defibrillation/cardioversion may cause explosion and fire which may injure the patient, user or bystanders.

- ⇒ When treating patients with oxygen masks, nasal tubes or nasal cannulas: Switch off the oxygen supply or place the inhalation points at least 1 meter away from the patient during defibrillation, and ensure that the flow of the oxygen/air mixture can escape away from the torso.
- ⇒ When treating patients with a bag-valve mask: Leave the bag-valve mask securely in place on the patient tube or place it at least 1 meter away from the patient, and ensure that the flow of the oxygen/air mixture can escape away from the torso.
- ⇒ When connecting patients to a ventilator: Ensure that the flow of the oxygen/air mixture coming from the exhalation valve can escape away from the torso.
- ⇒ When carrying out defibrillation/cardioversion in cramped spaces with an oxygen-enriched atmosphere, ensure that there is adequate ventilation.

Risk of injury due to missing battery!

Operation with line power without a battery prevents the device being fully ready for use as the shock capacitor in the device cannot charge. This prevents shock delivery and delays treatment. \Rightarrow Only operate the device with the battery inserted.

Risk of injury due to unsuitable AED analysis algorithm in children below one year of age!

The device's AED analysis algorithm is not designed for children below one year of age and may result in injury to the child. \Rightarrow Do not use AED mode on children below one year of age.

Risk of injury during resuscitation due to incorrect settings in the operator menu!

Incorrect settings in the operator menu may result in undesirable effects during resuscitation as well as injure the patient.

- ⇒ Only allow persons with specialist knowledge of the latest resuscitation recommendations to make settings in the operator menu.
- ⇒ If you are unaware of the most recent recommendations for resuscitation: Use the factory settings.

Delay in treatment due to movement artifacts during ECG analysis!

Movement artifacts may falsify the ECG analysis in AED mode and the ECG in manual mode. They may result in the user or the device interpreting the ECG incorrectly, delaying treatment.

During cardiac rhythm analysis:

- \Rightarrow Keep the patient still.
- \Rightarrow Stand clear of the patient.
- \Rightarrow Do not carry out chest compressions.
- \Rightarrow Do not ventilate the patient.
- \Rightarrow Do not transport the patient.

Risk of injury and delay in treatment due to incorrectly attached defibrillation electrodes!

Incorrectly attached defibrillation electrodes may falsify the ECG and result in the user triggering an unnecessary shock, not triggering a necessary shock or in unsuccessful defibrillation due to incorrect interpretation of an ECG.

- ⇒ Attach the defibrillation electrodes correctly as per the instructions for use.
- ⇒ Always place defibrillation electrodes together on only one person.
- \Rightarrow Prevent contact with the defibrillation electrodes.
- \Rightarrow Keep the defibrillation electrodes away from other electrodes and parts in contact with the patient.

Risk of injury due to air/moisture between defibrillation electrodes and the patient's skin!

Air (e.g. in the case of hirsute patients) or moisture between the defibrillation electrodes and the patient's skin prevent correct shock delivery and may result in burns to the skin and unsuccessful defibrillation/cardioversion.

- \Rightarrow Remove hair from hirsute patients.
- \Rightarrow Rub the patient's skin dry.
- \Rightarrow Wipe down oily skin with an alcohol pad.
- \Rightarrow Press the defibrillation electrodes on firmly.

Risk of injury due to air/moisture between the paddles and the patient's skin!

Air (e.g. in the case of hirsute patients) or moisture between the paddles and the patient's skin prevent correct shock delivery and may result in burns to the skin and unsuccessful defibrillation/ cardioversion.

- \Rightarrow Remove hair from hirsute patients.
- \Rightarrow Rub the patient's skin dry.
- \Rightarrow Wipe down oily skin with an alcohol pad.
- \Rightarrow Always use electrode gel with paddles.
- ⇒ Always press paddles on firmly with a contact pressure of about 8 kg.

Risk of injury due to non-functional defibrillation electrodes!

Non-functional defibrillation electrodes may result in injury and in unsuccessful defibrillation/cardioversion.

- \Rightarrow Only use defibrillation electrodes with undamaged packaging.
- ⇒ Do not use defibrillation electrodes with a dried-on layer of gel, damage or detached protective film.
- \Rightarrow Replace damaged defibrillation electrodes.
- ⇒ Observe the expiry date of the defibrillation electrodes and, if necessary, replace the defibrillation electrodes.
- \Rightarrow Dispose of defibrillation electrodes after use and do not reuse them.
- ⇒ Only use defibrillation electrodes approved by WEINMANN Emergency for the device.

Risk of injury due to incorrect handling of electrode gel!

Incorrect handling of electrode gel may lead to electric shock, to ineffective shock delivery, and to burns, injuring the patient, user, and bystanders.

 \Rightarrow Always use electrode gel with paddles.

- ⇒ Do not allow any electrode gel to get between the surface of the electrode and the handle in order to prevent electric shock.
- \Rightarrow Do not use too much electrode gel to prevent a gel bridge and thus burns on the chest.
- ⇒ Do not use too little electrode gel to keep the resistance to the patient low for effective shock delivery and to prevent burns on the chest.

Risk of injury and delay in treatment due to implanted cardiac pacemakers!

Pulses from implanted cardiac pacemakers may affect the detection of cardiac rhythms which can be defibrillated, delaying treatment. Carrying out defibrillation on patients with implanted cardiac pacemakers may irreversibly damage the myocardium.

- ⇒ Position defibrillation electrodes at least 8 cm away from cardiac pacemakers.
- ⇒ Consider alternative positions (e.g. anterior-lateral, anteriorposterior) for the defibrillation electrodes.

Risk of injury due to shock being delivered at the wrong time during cardioversion!

A cardioversion carried out at the wrong time may lead to cardiac arrhythmias and cause the patient serious or life-threatening injury.

- \Rightarrow Check whether the ECG is stable.
- \Rightarrow Check whether the device is detecting and marking R waves correctly in the ECG.

Risk of injury due to ECG misinterpretation if ECG is derived from the defibrillation electrodes!

If the ECG is derived from the defibrillation electrodes, the device shows a non-diagnostic ECG curve. This ECG curve is designed to detect shockable cardiac rhythms and is not suitable for differential diagnosis. This may result in ECG misinterpretation, and thus in injury to the patient.

⇒ Do not use ECGs derived from defibrillation electrodes for differential diagnosis.

Failure of therapy due to defibrillation electrodes accidentally coming loose during shock delivery!

Defibrillation electrodes accidentally coming loose during shock delivery may lead to damage to the device and thus to the failure of treatment. This may injure the patient.

⇒ Ensure that the defibrillation electrodes are always connected to the device during shock delivery.

Malfunction or failure of other electrical devices due to delivery of shock energy!

Charging and then delivery of shock energy may impair the function of other electrical devices or damage devices connected to the patient or located in the vicinity of the defibrillator. This may injure the patient.

- ⇒ Disconnect from the patient any electrical devices without defibrillation protection.
- ⇒ After using the defibrillator, check the function of electrical devices in its vicinity.
- ⇒ Maintain separation distances between the defibrillator and portable and mobile high-frequency communications devices.

Caution Delay in treatment due to simultaneous voice prompts from defibrillator and ventilator!

If the defibrillator in AED mode is used in conjunction with a ventilator (MEDUMAT Easy CPR) which also guides the user through CPR by means of voice prompts, the simultaneous voice prompts from the defibrillator and ventilator may confuse the user and delay treatment.

⇒ When using the defibrillator in AED mode and a ventilator at the same time: Switch off the ventilator voice prompts.

Risk of injury due to incorrectly selected size of defibrillation electrodes!

If the wrong size of defibrillation electrodes is selected, this may result in sub-optimal defibrillation results or in burns.

⇒ Select the correct size of defibrillation electrodes in line with currently applicable guidelines and not based on the weight specifications given on the packaging.

2.6.5 Pulse oximetry monitoring

Warning

Risk of injury due to incorrect use of the pulse oximetry sensor!

Incorrect use of the pulse oximetry sensor may falsify measurement results and lead to patient injury.

- ⇒ Keep the pulse oximetry sensor away from strong electromagnetic sources (e.g. electrosurgical devices).
- \Rightarrow Do not use the pulse oximetry sensor in areas subject to X-ray (e.g. with MRI devices).

- ⇒ Keep the pulse oximetry sensor away from strong and fluctuating ambient light (including infrared and UV light). If necessary: Cover the pulse oximetry sensor.
- ⇒ Avoid strong movement of the pulse oximetry sensor. If necessary: To relieve strain, loop the pulse oximetry sensor cable and the pulse oximetry sensor connecting cable and fix to the patient with a plaster.
- ⇒ Do not attach the pulse oximetry sensor to a limb on which there is already an NIBP cuff or catheter port.
- ⇒ Keep the pulse oximetry sensor away from nail polish and artificial fingernails.
- \Rightarrow Keep the pulse oximetry sensor away from intravascular dyes.
- ⇒ Be aware of deviations from the measurement result with a high proportion of dysfunctional hemoglobins.
- ⇒ Be aware of deviations from the measurement result in the case of severe anemia, venous pulsation and high total bilirubin values.
- ⇒ Be aware of deviations in pulse rate with an intra-aortic balloon pump or certain arrhythmias.

If necessary: Compare the pulse rate with the heart rate determined by ECG monitoring.

- ⇒ Be aware of deviations from the measurement result during defibrillation/cardioversion.
- \Rightarrow Only use undamaged pulse oximetry sensors.
- ⇒ Only use the pulse oximetry sensors and pulse oximetry sensor connecting cables quoted in the scope of supply and in the accessories.

Risk of injury from unsuitable pulse oximetry monitoring in children with a body weight of less than 10 kg!

The device's pulse oximetry monitoring is not designed for children with a body weight of less than 10 kg and may result in injury to the child.

⇒ Do not use pulse oximetry monitoring in children with a body weight of less than 10 kg.

2.6.6 6-lead ECG monitoring/12-lead ECG recording and assessment

Warning Risk of injury from incorrect, expired or damaged ECG electrodes!

Incorrect, expired or damaged ECG electrodes impair the quality of the ECG signal and falsify measurement results. This may injure the patient.

- \Rightarrow Only use ECG electrodes which satisfy all of the points listed here.
- \Rightarrow Only use AAMI EC 12-certified ECG electrodes.
- \Rightarrow Only use high-quality ECG electrodes.
- \Rightarrow Observe the expiry date of the ECG electrodes and, if necessary, replace the ECG electrodes.
- \Rightarrow Only use ECG electrodes with undamaged packaging.
- \Rightarrow Do not use ECG electrodes with a dried-out layer of gel layer, damage or detached protective film.
- \Rightarrow Do not remove ECG electrodes from the packaging until immediately before the session.
- \Rightarrow Replace ECG electrodes damaged during the session.
- \Rightarrow Do not use ECG electrodes for defibrillation/cardioversion.
- \Rightarrow Dispose of ECG electrodes after use and do not reuse them.

Risk of injury from using the 6-lead ECG for differential diagnosis!

The ECG curve of the 6-lead ECG is not suitable for differential diagnosis (e.g. diagnosis of cardiac infarction). This may result in ECG misinterpretation, and thus in injury to the patient.

- \Rightarrow Do not use the 6-lead ECG for differential diagnosis.
- \Rightarrow For differential diagnosis, use the 12-lead ECG option of the device or use a different 12-lead ECG device.

Risk of injury and delay in treatment due to implanted cardiac pacemakers!

In the case of patients with cardiac pacemakers, the device detects the pacemaker pulses and suppresses the heart rate display and heart rate alarms. This may result in injury to the patient.

 \Rightarrow Monitor patients with pacemakers very closely.

Risk of injury and delay in treatment due to implanted cardiac pacemakers!

On patients with cardiac pacemakers, heart rate displays may count pacemaker pulses during a cardiac arrest or in the case of some cardiac arrhythmias. This may result in injury to the patient.

- \Rightarrow Do not rely entirely on the device's alarm signals.
- \Rightarrow Monitor patients with pacemakers very closely.
- ⇒ See information on suppressing pacemaker pulses (see "18.10 6-lead ECG monitoring/12-lead ECG recording and assessment", page 393).

Risk of injury due to ECG malfunction in the vicinity of electrosurgical devices!

ECG functions may be affected by electrosurgical devices and result in injury to the patient.

 \Rightarrow Only use approved ECG cables.

Risk of injury due to delayed delivery!

The user can deliver a 12-lead ECG to a teleconsultation service for further diagnosis. Delivering a 12-lead ECG is purely to obtain a second opinion, and does not replace adequately trained specialists at the session location. Specialist staff at the session location are responsible for treatment within the scope of their training and in accordance with the specifications of the operator. If the 12-lead ECG reaches the teleconsultation service very late or not at all for technical reasons, delayed treatment may result in a potentially life-threatening situation. The operator is responsible for ensuring that the necessary configuration for 12-lead ECG delivery has been carried out correctly.

- ⇒ When configuring e-mail delivery 12-lead ECG: Test e-mail delivery.
- ⇒ If the IT infrastructure permits it: Configure several networks for various devices or session locations so that if one network is unavailable, it is possible to select another network.
- ⇒ Check the configuration of the networks at regular intervals and update them if necessary.
- \Rightarrow Ensure that the e-mail addresses are correct and complete.
- ⇒ Check e-mail addresses at regular intervals and update them if necessary.

Caution Risk of injury from burns during defibrillation/cardioversion!

ECG cables without defibrillation protection may result in injury to the patient.

 \Rightarrow Only use approved ECG cables.

Therapy put at risk by incorrectly set low-pass filter for the 12-lead ECG!

An incorrectly set low-pass filter may lead to incorrect assessment of the 12-lead ECG and injure the patient.

 \Rightarrow Select a suitable low-pass filter (50 Hz or 150 Hz).

⇒ When assessing the 12-lead ECG, take account of an activated low-pass filter.

2.6.7 Non-invasive blood pressure measurement (NIBP monitoring)

Caution Risk of injury if NIBP cuff selected or put on incorrectly! An NIBP cuff selected or put on incorrectly may interrupt the blood supply or falsify measurement results due to excessive contact

pressure. This may result in injury to the patient.

- ⇒ Always use the NIBP cuff best suited to the patient's limb. Selecting the right NIBP cuff is the key to ensuring a goodquality measured value.
- \Rightarrow Attach the NIBP cuff level with the heart.
- \Rightarrow Attach the NIBP cuff so that the blood supply is not stopped.
- \Rightarrow Avoid moving the NIBP cuff during NIBP measurement.
- \Rightarrow When NIBP measurement is for an extended period: Check the position of the NIBP cuff regularly and, if necessary, reposition the NIBP cuff.
- ⇒ Repeat the NIBP measurement if measurement results are implausible. If the repeat measurement is still implausible, select an alternative method.
- \Rightarrow Do not bend or crush the NIBP cuff tube or the NIBP connecting tube.
- \Rightarrow Do not attach the NIBP cuff to a limb with poor circulation.
- \Rightarrow Do not attach the NIBP cuff to a limb with an intravenous infusion.
- \Rightarrow Do not attach the NIBP cuff to a limb on which there is already a pulse oximetry sensor or another monitoring device.
- \Rightarrow Do not attach the NIBP cuff to a limb with a shunt.
- \Rightarrow Do not attach the NIBP cuff to a limb with open wounds or burns.
- \Rightarrow In the case of patients who have undergone a mastectomy, do not attach the NIBP cuff to the affected side. In the case of patients who have undergone double mastectomies, attach the NIBP cuff to the non-dominant arm.
- \Rightarrow Only use an undamaged NIBP cuff.
- \Rightarrow Only use the NIBP cuffs and NIBP connecting tubes quoted in the scope of supply and in the accessories.
- \Rightarrow Follow the instructions for use of the NIBP cuff.

Risk of injury from falsified measured values during noninvasive blood pressure measurement!

Cardiac rhythm disorders, arteriosclerosis, reduced perfusion, diabetes, pregnancy, pre-eclampsia, arrhythmias (in newborns up to 28 days), kidney problems, shaking, shivering or the use of a cardiac pacemaker may impair the ability of the non-invasive blood pressure measuring module to record correct measured values. Evidence of the safety and efficacy of non-invasive blood pressure measurement has not been obtained for patient groups with these characteristics. This may injure the patient.

- \Rightarrow Include the status of these patient groups when evaluating the measured values of non-invasive blood pressure measurement.
- ⇒ Only use non-invasive blood pressure measurement on patient groups for whom evidence of the safety and efficacy of noninvasive blood pressure measurement has been provided.

Risk of injury from overly frequent measurements!

Overly frequent measurements may lead to circulation problems and patient injury.

- \Rightarrow Select the measurement intervals to guarantee sufficient perfusion.
- ⇒ With extended NIBP measurements, check the position of the NIBP cuff regularly and, if necessary, reposition it.

2.6.8 Printing

Warning Risk of injury due to electric shock when the printer is touched!

The printer's USB port is intended only for service purposes. A connection between the printer and another device via the USB port may lead to an electric shock if the printer is touched. This may injure the patient or the user.

 \Rightarrow Do not connect the printer to another device via the USB port. \Rightarrow Only use the USB port for service purposes.

Risk of injury from electric shock when charging the printer battery in the printer!

If the power supply unit and charger is defective, charging the printer battery in the printer may lead to an electric shock if the printer is touched. This may injure the patient or the user.

⇒ Only charge the printer battery in the charging station for the printer battery or the quadruple charging station for the printer battery.

- ⇒ Only use the power supply unit and charger for the printer battery for charging the printer battery in the charging station for the printer battery.
- \Rightarrow Do not use the power supply unit and charger in the patient's vicinity.

Caution Malfunction or failure of the printer due to electromagnetic radiation in airplane mode!

Electromagnetic radiation in airplane mode may lead to special characters appearing in the printout or to failure of the printer. \Rightarrow Do not use the printer in airplane mode.

Delayed therapy due to unsuitable printer paper!

Unsuitable printer paper may lead to poor print quality on the printout or to paper jams and delay treatment.

- ⇒ Only use printer paper approved by WEINMANN Emergency. If this is not possible: Only use printer paper which satisfies the specifications listed (see "18.8 Printer and printer paper", page 391).
- \Rightarrow Do not used printer paper where the paper is glued to the roll.
- \Rightarrow Use printer paper that has an end marking.

Delayed therapy due to incorrect handling of printer paper! Incorrect handling of the printer paper may lead to poor print quality on the printout and delay treatment.

- \Rightarrow To prevent the inks on the printer paper fading, do not use chemicals or oil.
- ⇒ Do not expose printer paper to significant heat, moisture or sunlight (see "18.8 Printer and printer paper", page 391).
- \Rightarrow Keep printer paper away from sharp or hard objects (e.g. fingernails or metal) in order not to scratch the printer paper.
- \Rightarrow Store printer paper in a cool, dry, dark place.
- \Rightarrow Do not use chemical adhesive.
- \Rightarrow Keep printer paper away from plastics which emit vapors.
- \Rightarrow Always use clean printer paper.

Malfunction or failure of the printer due to external influences and incorrect use!

The printer may be damaged by external influences and incorrect use.

- \Rightarrow Always use the printer in the printer case.
- \Rightarrow Do not drop the printer.
- \Rightarrow Do not expose the printer to severe shocks.
- \Rightarrow Do not expose the printer to direct sunlight or similar conditions.

- ⇒ Keep the paper compartment closed during the printing process.
- \Rightarrow Switch off the printer when it is not in use.
- \Rightarrow Keep the printer away from powerful electromagnetic sources.
- \Rightarrow Keep all the openings of the printer closed during printing.
- \Rightarrow Only use the products quoted in the cleaning and disinfection plan (see "8.7 Cleaning and disinfection plan", page 248).

2.6.9 Electromagnetic compatibility

Warning

Risk of injury from mutual influence of medical electrical devices!

Medical electrical devices which are operated directly next to or on top of one other may cause mutual interference to functionality and thus patient injury.

- \Rightarrow Do not stack the device with other medical electrical devices.
- ⇒ Do not operate the device in the direct vicinity of other medical electrical devices (exception: Combinations of devices approved for MEDUCORE Standard² on the portable units from WEINMANN Emergency).
- ⇒ If stacking or operation in the immediate vicinity cannot be avoided: Closely monitor the functioning of all affected medical electrical devices and do not use if functions are disrupted.

Risk of injury from overly powerful high-frequency interference signals or electromagnetic fields!

Overly strong high-frequency interference signals from portable high-frequency communications equipment (e.g. radios) or electromagnetic fields may lead to incorrect analysis, incorrect measurement results, and incorrect alarms, and thus impair the functioning of the device and injure the patient.

- ⇒ Maintain separation distances (see "18.17 Electromagnetic compatibility (EMC)", page 400).
- ⇒ With portable high-frequency communications devices, maintain a minimum distance of 30 cm from the device, components, and accessories.

Delay in treatment due to power supply network disruption!

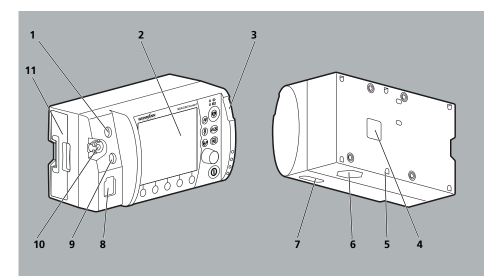
Disruptions to the power supply network may cause visible disruptions to the ECG. This may lead to incorrect measurement results and alarms and in AED mode, lead to shock delivery at the wrong time. This may delay therapy and injure the patient. \Rightarrow If there is major disruption to the power supply network, only

operate the device with a battery.

2.7 General instructions

- If third-party items are used, functional failures and restricted fitness for use may result. Biocompatibility requirements may also not be met. Please note that in such cases, any warranty claim and liability will be voided if neither the accessories recommended in the instructions for use nor original replacement parts are used. Third-party items may increase radiation output or reduce interference immunity.
- The manufacturer, WEINMANN Emergency, guarantees the compatibility of the device and of all components or accessories connected to the patient prior to the session. Have modifications to the device or accessories (exception: software update) carried out only by the manufacturer, WEINMANN Emergency, or by a technician expressly authorized by it. Do not use any articles from third parties.
- Follow the section on hygienic reprocessing in order to avoid infection or bacterial contamination (see "8 Hygienic reprocessing", page 238).
- As the user, always stay in the immediate vicinity of the device and the patient.
- Risks due to software errors have been minimized by means of extensive qualification measures.
- This device's software contains code which is subject to the General Public License (GPL). We will send you the source code and the GPL on request.

3 Description



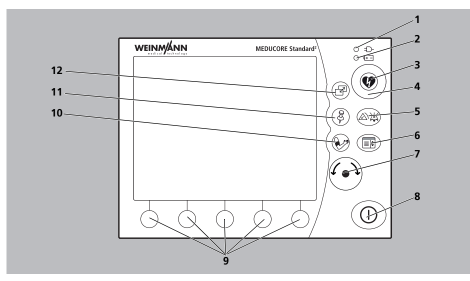
3.1 Overview

3-1 Device

No.	Designation	Description
1	ECG connection for ECG cable	Connects the device to an ECG cable.
2	Display	Displays settings and current values (see "3.4 Symbols in the display", page 54).
3	Alarm light	Indicates high-priority alarms visually.
4	Power supply connection	Connects the device to the power supply.
5	Security seal	Indicates whether the device has been opened without authorization.
6	Loudspeaker	Emits audible voice prompts, alarms and heart rate tones/pulse tones.
7	SD card slot	Takes an SD card.

No.	Designation	Description
8	Pad connection for master cable	Connects the device to one of the following components via the master cable: • Defibrillation electrodes • Paddles • Function test resistor
9	SpO ₂ port for pulse oximetry sensor connecting cable	Connects the device to a pulse oximetry sensor via the pulse oximetry sensor connecting cable.
10	NIBP connection for NIBP connecting tube	Connects the device to an NIBP cuff via the NIBP connecting tube.
11	Battery compartment with battery	Houses the battery.

3.2 Control panel



3-2 Control panel

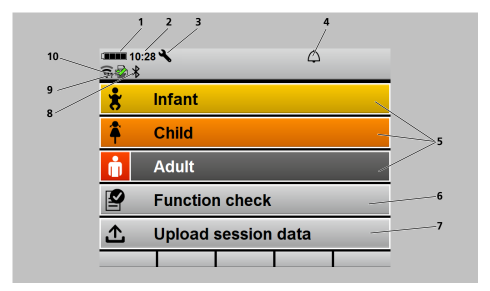
No.	Designation	Description
1	Line power indicator	Indicates that the device is being supplied by line power.
2	Battery status indicator	 Green light on: The battery is full or is not being charged because it is outside charging temperature range. Green light flashing: The battery is being charged. Red light on: The battery is defective or not in the device. No light: The device is not connected to line power.
3	Shock button	Triggers an electric shock for defibrillation/ cardioversion.
4	Shock standby indicator	Red light flashes when the device is ready to deliver a shock.
5	Alarm button	 Pauses audio alarm output for a certain length of time. Mutes audio alarm output. Cancels audio alarm output. Deactivates muting of audio alarm output and alarm cancellation.
6	Menu button	 In the start menu: Provides access to the operator menu. In a mode: Provides access to the user menu. In the button function test during the function check: Cancels the button function test.
7	Navigation knob	 Allows values to be selected (turn). Confirms selected values (press). In a mode: Provides access to the application menu (press).
8	On/Off button	 With device switched off: Press briefly to switch the device on. With device switched on: Press and hold to switch the device off.
9	Function buttons	 Provide access to the mode shown in the display. Activate/deactivate the functions shown in the display.
10	NIBP button	 Activates NIBP function mode (press NIBP button < 2 s). Starts an NIBP measurement (press NIBP button for > 2 s).

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No.	Designation	Description
11	Event button	Opens the events list.
12	View button	Switches between the following views: Parameter view Curve view Switches between the following views (only in 12-lead ECG function mode): 1-curve view 3-curve view 6-curve view

3.3 Display

3.3.1 Start menu

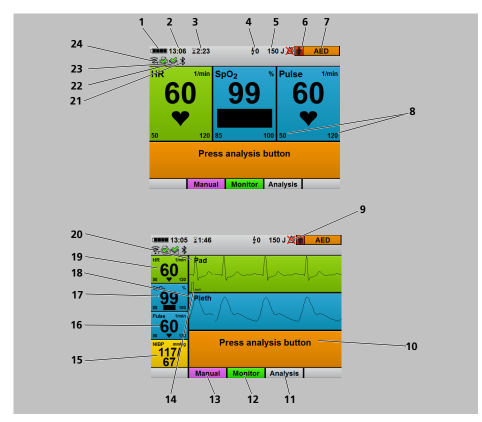


3-3 Start menu display

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No.	Designation	Description
1	Battery status	Displays the charge level of the battery.
2	Time	Displays the time.
3	Service reminder (if activated)	Displayed when the service interval is \leq 30 days.

No.	Designation	Description
4	Alarm indicator	 Indicates the status of audio alarm output: Audio alarm output active Audio alarm output muted/paused Audio alarm output canceled
5	Patient groups	Starts the device with the presets specific to the patient groups.
6	Function check	Provides access to the function check.
7	Upload session data (only with Session data upload option)	Allows device session data to be uploaded to WEINMANN Connect.
8	Bluetooth [®] symbol (only with Bluetooth [®] data transmission option)	 Black: Indicates that the device is ready for Bluetooth[®] data transmission to an external device. Blue: Indicates that a Bluetooth[®] connection to an external device has been set up.
9	Printer symbol (only with Printing option)	Indicates whether the printer is connected to the device.
10	WiFi symbol	 Indicates whether the default network can be reached. Displays the signal strength of the WiFi connection: The more bars displayed, the stronger the signal.



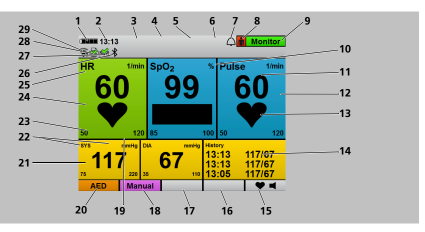
3.3.2 AED mode

3-4 Display in AED mode: Parameter view (top) and curve view (bottom)

No.	Designation	Description
1	Battery status	Displays the charge level of the battery.
2	Time	Displays the time.
3	Session duration	Displays the duration of the current session.
4	Number of shocks delivered	Displays the number of shocks delivered during the current session.
5	Shock energy	Displays the shock energy selected for the next shock.

No.	Designation	Description
6	Patient group	Displays the selected patient group: • Adult • Child
7	Mode indicator	Indicates the mode currently selected.
8	Alarm limits	Displays the set alarm limits.
9	Alarm off indicator	Indicates whether alarm output is deactivated in AED mode.
10	AED instructions	Give instructions on carrying out CPR.
11	Analysis	 Allows you to start a cardiac rhythm analysis manually. Only displayed if Start analysis automatically is deactivated in the operator menu (see "13.3 AED settings", page 325).
12	Monitor mode	Provides access to monitor mode.
13	Manual mode (only with Manual defibrillation option)	Provides access to manual mode.
14	ECG calibration mark	Displays the section corresponding to 1 mV of the ECG signal.
15	NIBP	Displays blood pressure.
16	Pulse	Displays pulse rate.
17	SpO ₂	Displays oxygen saturation.
18	Center curve field	Displays the plethysmogram.
19	HR	Displays heart rate.
20	Top curve field	Displays the ECG lead (Pad, II).
21	Bluetooth [®] symbol (only with Bluetooth [®] data transmission option)	 Black: Indicates that the device is ready for Bluetooth[®] data transmission to an external device. Blue: Indicates that a Bluetooth[®] connection to an external device has been set up.

No.	Designation	Description
22	Upload symbol	 Indicates that e-mail delivery 12-lead ECG has been started (only with E-mail delivery 12-lead ECG option). Indicates whether the last e-mail delivery started was successful (only with E-mail delivery 12-lead ECG option). Indicates whether the function check has been uploaded to WEINMANN Connect successfully. Indicates whether the session data have been uploaded to WEINMANN Connect successfully (only with Session data upload option). Indicates whether the service data have been uploaded to WEINMANN Connect successfully (only with Session data upload option).
23	Printer symbol (only with Printing option)	Indicates whether the printer is connected to the device.
24	WiFi symbol	 Indicates whether the default network can be reached. Displays the signal strength of the WiFi connection: The more bars displayed, the stronger the signal.



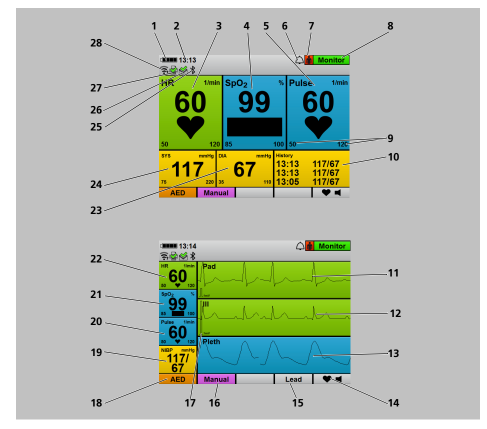
3.3.3 Manual mode (only with Manual defibrillation option)

3-5 Display in manual mode

No.	Designation	Description
1	Battery status	Displays the charge level of the battery.
2	Time	Displays the time.
3	Session duration	Displays the duration of the current session.
4	Operating time since last shock delivery	Displays device operating time since the last shock delivery.
5	Number of shocks delivered	Displays the number of shocks delivered during the current session.
6	Shock energy	Displays the shock energy selected for the next shock.
7	Alarm indicator	 Indicates the status of audio alarm output: Audio alarm output active Audio alarm output muted/paused Audio alarm output canceled
8	Patient group	Displays the selected patient group: • Adult • Child • Infant
9	Mode indicator	Indicates the mode currently selected.
10	SYNC (only with Cardioversion option)	Indicates that delivery of the following shock will be synchronized with the R wave.

No.	Designation	Description
11	Top curve field	Displays the ECG lead via the defibrillation electrodes/ paddles (Pad, II).
12	R wave marking (only with Cardioversion option)	Marks the R waves detected by the device in the ECG if SYNC has been activated in the application menu.
13	Center curve field	 Displays the selected ECG lead: I, II, III, aVR, aVL or aVF V1 to V6 (only with 12-lead ECG extension cable and only with 12-lead ECG option)
14	Bottom curve field	Displays the plethysmogram.
15	Energy	Allows shock energy to be adjusted.
16	ECG lead selection	 Allows switching between the ECG leads displayed in the center curve field: Limb leads acc. to Einthoven (I, II, III) Limb leads acc. to Goldberger (aVR, aVL, aVF) Chest wall leads acc. to Wilson (V1 to V6) (only with 12-lead ECG extension cable and only with 12-lead ECG option)
17	Monitor mode	Provides access to monitor mode.
18	Charge	Charges the shock capacitor.
19	ECG calibration mark	Displays the section corresponding to 1 mV of the ECG signal.
20	AED mode	Provides access to AED mode.
21	NIBP	Displays blood pressure.
22	Alarm limits	Displays the set alarm limits.
23	Pulse	Displays pulse rate.
24	SpO ₂	Displays oxygen saturation.
25	HR	Displays heart rate.
26	Bluetooth [®] symbol (only with Bluetooth [®] data transmission option)	 Black: Indicates that the device is ready for Bluetooth[®] data transmission to an external device. Blue: Indicates that a Bluetooth[®] connection to an external device has been set up.

No.	Designation	Description
27	Upload symbol	 Indicates that e-mail delivery 12-lead ECG has been started (only with E-mail delivery 12-lead ECG option). Indicates whether the last e-mail delivery started was successful (only with E-mail delivery 12-lead ECG option). Indicates whether the function check has been uploaded to WEINMANN Connect successfully. Indicates whether the session data have been uploaded to WEINMANN Connect successfully (only with Session data upload option). Indicates whether the service data have been uploaded to WEINMANN Emergency successfully.
28	Printer symbol (only with Printing option)	Indicates whether the printer is connected to the device.
29	WiFi symbol	 Indicates whether the default network can be reached. Displays the signal strength of the WiFi connection: The more bars displayed, the stronger the signal.



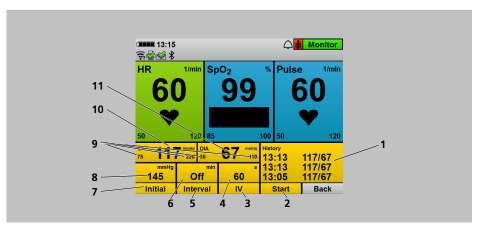
3.3.4 Monitor mode

3-6 Display in monitor mode: Parameter view (top) and curve view (bottom)

No.	Designation	Description
1	Battery status	Displays the charge level of the battery.
2	Time	Displays the time.
3	HR	Displays heart rate.
4	SpO ₂	Displays oxygen saturation.
5	Pulse	Displays pulse rate.

No.	Designation	Description
6	Alarm indicator	 Indicates the status of audio alarm output: Audio alarm output active Audio alarm output muted/paused Audio alarm output canceled
7	Patient group	Displays the selected patient group: • Adult • Child • Infant
8	Mode indicator	Indicates the mode currently selected.
9	Alarm limits	Displays the set alarm limits.
10	History	Shows the time and values of the last three NIBP measurements.
11	Top curve field	Displays the ECG lead via the defibrillation electrodes/ paddles (Pad, II).
12	Center curve field	 Displays the selected ECG lead: I, II, III, aVR, aVL or aVF V1 to V6 (only with 12-lead ECG extension cable and only with 12-lead ECG option)
13	Bottom curve field	Displays the plethysmogram.
14	Heart rate tone/pulse tone	Switches the heart rate tone/pulse tone on and off.
15	ECG lead selection	 Allows switching between the ECG leads displayed in the center curve field: Limb leads acc. to Einthoven (I, II, III) Limb leads acc. to Goldberger (aVR, aVL, aVF) Chest wall leads acc. to Wilson (V1 to V6) (only with 12-lead ECG extension cable and only with 12-lead ECG option)
16	Manual mode (only with Manual defibrillation option)	Provides access to manual mode.
17	ECG calibration mark	Displays the section corresponding to 1 mV of the ECG signal.
18	AED mode	Provides access to AED mode.
19	NIBP	Displays blood pressure.
20	Pulse	Displays pulse rate.
21	SpO ₂	Displays oxygen saturation.
22	HR	Displays heart rate.
23	DIA	Displays the diastolic measured value for an NIBP measurement.
24	SYS	Displays the systolic measured value for an NIBP measurement.

No.	Designation	Description
25	Bluetooth [®] symbol (only with Bluetooth [®] data transmission option)	 Black: Indicates that the device is ready for Bluetooth[®] data transmission to an external device. Blue: Indicates that a Bluetooth[®] connection to an external device has been set up.
26	Upload symbol	 Indicates that e-mail delivery 12-lead ECG has been started (only with E-mail delivery 12-lead ECG option). Indicates whether the last e-mail delivery started was successful (only with E-mail delivery 12-lead ECG option). Indicates whether the function check has been uploaded to WEINMANN Connect successfully. Indicates whether the session data have been uploaded to WEINMANN Connect successfully (only with Session data upload option). Indicates whether the service data have been uploaded to WEINMANN Emergency successfully.
27	Printer symbol (only with Printing option)	Indicates whether the printer is connected to the device.
28	WiFi symbol	 Indicates whether the default network can be reached. Displays the signal strength of the WiFi connection: The more bars displayed, the stronger the signal.



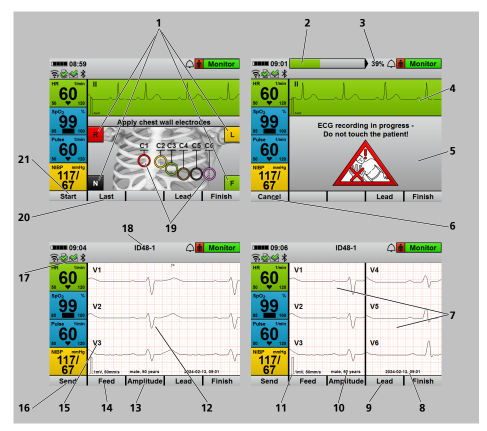
3.3.5 NIBP function mode

3-7 Display in monitor mode with NIBP function mode superimposed

No.	Designation	Description
1	History	Shows the time and values of the last three NIBP measurements.
2	Start/Stop	 Starts or stops an NIBP measurement. Starts or stops an interval measurement. Stops venous stasis.
3	IV	Starts venous stasis.
4	Duration of venous stasis	Displays the time for which the NIBP cuff is maintaining venous stasis.
5	Interval	 Specifies whether the NIBP measurement is an individual NIBP measurement or an interval measurement. Specifies the time between two consecutive NIBP measurements for interval measurement.
6	Interval duration	Displays the time between two consecutive NIBP measurements for interval measurement.
7	Initial	Allows initial NIBP cuff pressure to be changed.
8	Initial cuff pressure	Displays the pressure to which the device will inflate the NIBP cuff at the next NIBP measurement.
9	Alarm limits	Displays the alarm limits for systolic and diastolic measured values.

No.	Designation	Description
10	SYS	Displays the systolic measured value for an NIBP measurement.
11	DIA	Displays the diastolic measured value for an NIBP measurement.

3.3.6 12-lead ECG function mode (only with 12-lead ECG option)

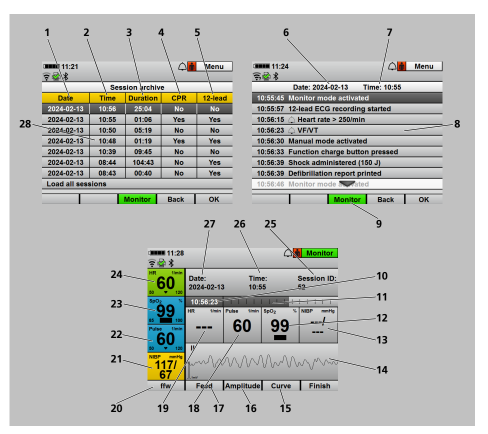


3-8 Display in monitor mode with superimposed 12-lead ECG function mode

No.	Designation	Description
1	Limb electrodes	Indicates that the limb electrodes are connected. The names of the limb electrodes change depending on the set electrode marking (ERC or AHA) (see "13.10 12-lead ECG settings (only with 12-lead ECG option)", page 350).
2	Progress display	Indicates how far the 12-lead ECG has already recorded.
3	Progress display in percent	Indicates how many percent of the 12-lead ECG has already been recorded.
4	Top curve field	Displays the selected ECG lead: I, II, III, aVR, aVL, aVF, V1, V2, V3, V4, V5, V6
5	Instruction field	Gives instructions on what to do.
6	Cancel	Cancels a 12-lead ECG recording.
7	6-curve view	Displays 6 ECG leads simultaneously.
8	Date/time	Displays the date and time of the 12-lead ECG recording.
9	ECG lead selection	 Allows you to switch between ECG leads: Limb leads acc. to Einthoven (I, II, III) Limb leads acc. to Goldberger (aVR, aVL, aVF) Chest wall leads acc. to Wilson (V1 to V6)
10	Patient data	Display the age and gender of the patient.
11	ECG calibration mark	Displays the section corresponding to 1 mV of the ECG signal.
12	3-curve view	Displays 3 ECG leads simultaneously.
13	Amplitude scaling	Here you can set the amplification of the ECG signal and thus the height of the ECG curve.
14	Feed rate	Here you can set the feed rate of the ECG curve display and so change the time resolution.
15	ECG lead	Displays the selected ECG lead: I, II, III, aVR, aVL, aVF, V1, V2, V3, V4, V5, V6
16	E-mail delivery (only with E-mail delivery 12-lead ECG option)	Here you can deliver the 12-lead ECG by e-mail.
17	Upload symbol (only with E-mail delivery 12-lead ECG option)	 Indicates that e-mail delivery 12-lead ECG has been started. Indicates whether the last e-mail delivery started was successful.
18	Patient ID	Displays the ID of the 12-lead ECG recorded. This ID also appears in the PDF and in the printout and allows the 12-lead ECG to be identified.

No.	Designation	Description
19	Chest wall electrodes	Indicates whether the chest wall electrodes are connected. The names of the chest wall electrodes change depending on which electrode marking is set (ERC or AHA) (see "13.10 12-lead ECG settings (only with 12-lead ECG option)", page 350). Connected chest wall electrodes are indicated by a check mark.
20	Last 12-lead ECG recording	Here you can call up the last 12-lead ECG of the current session.
21	Start	Starts the 12-lead ECG recording.





3-9 Display in replay view

No.	Designation	Description
1	Date	Displays the date on which the session took place.
2	Time	Displays the time at which the session began.
3	Duration	Displays the duration of the session.
4	CPR	Indicates whether the session involved resuscitation. Resuscitation took place if a shock was delivered or a resuscitation mode (AED mode or manual mode) was activated for > 2 min.
5	12-lead	Indicates whether one or more 12-lead ECGs were recorded during the session.

No.	Designation	Description
6	Date	Displays the date on which the session took place.
7	Time	Displays the time at which the session began.
8	Events	Displays events during a session.
9	Monitor mode	 Provides access to monitor mode. Depending on the mode from which the session archive was opened, other function buttons may be possible.
10	Recording point	Displays the exact time at which a momentary recording in an event took place.
11	Scroll bar	Indicates how far time was fast-forwarded within an event.
12	NIBP	Displays blood pressure at the time of recording.
13	SpO ₂	Displays oxygen saturation at the time of recording.
14	Bottom curve field	 Displays the selected ECG lead or plethysmogram at the time of recording: I, II, III, aVR, aVL or aVF ECG lead via the defibrillation electrodes/paddles (Pad, II)
15	Curve	Allows you to select the ECG lead or plethysmogram at the time of recording.
16	Amplitude scaling	Allows you to set the amplification of the ECG signal and thus the height of the ECG curve.
17	Feed rate	Allows you to set the feed rate of the ECG curve display and so change the time resolution.
18	Pulse	Displays pulse rate at the time of recording.
19	HR	Displays heart rate at the time of recording.
20	Scroll speed	Allows you to select the resolution of scrolling.
21	NIBP	Displays current blood pressure.
22	Pulse	Displays current pulse rate.
23	SpO ₂	Displays current oxygen saturation.
24	HR	Displays current heart rate.
25	Session ID:	Displays the session ID.
26	Time:	Displays the time at which the session began.
27	Date:	Displays the date on which the session took place.
28	Session archive	Displays a list of all the device sessions.

3.4 Symbols in the display

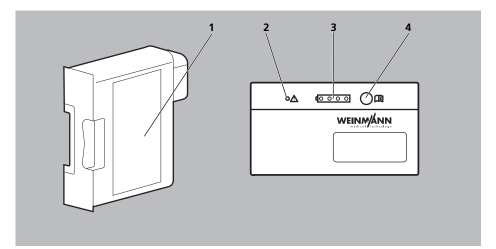
Symbol	Designation	Description
	Battery status symbol	Battery status (see "3.5 Battery and battery status indicator", page 57)
	Printer symbol (only with	Printer connected
	Printing option)	No printer connected
	Upload symbol: Only displayed if a 12-lead ECG has already been delivered in the current session (only with E-mail delivery 12-lead ECG option) or when data have been uploaded. Bluetooth [®] symbol (only with Bluetooth [®] data transmission option)	 Last e-mail delivery 12-lead ECG successful (only with E-mail delivery 12- lead ECG option) Function check uploaded to WEINMANN Connect successfully Session data uploaded to WEINMANN Connect successfully (only with Session data upload option) Service data uploaded to WEINMANN Emergency successfully
		 Symbol moves: Sending e-mail with 12-lead ECG (only with E-mail delivery 12-lead ECG option) Uploading data to WEINMANN Connect
*		 Last e-mail delivery failed (only with E-mail delivery 12-lead ECG option) Upload to WEINMANN Connect failed
*		Device is ready for Bluetooth [®] data transmission to an external device
*		$Bluetooth^{\textcircled{\text{$\mathbb 8$}}}$ connection to an external device set up

Symbol	Designation	Description
((î°:	_ WiFi symbol	 Indicates whether the default network can be reached. Displays the signal strength of the WiFi connection: The more bars displayed, the stronger the signal.
((•		Indicates that there is currently a connection to WEINMANN Connect and data are being uploaded to or with the aid of WEINMANN Connect.
→	Airplane mode symbol	Indicates that all wireless connections are switched off.
\bigtriangleup		Audio alarm output activeDesignates an alarm in the Replay view.
		Audio alarm output canceled
	Alarm symbol	Audio alarm output paused for the time set in the operator menu
X		Audio alarm output muted with no time limit
×		Alarm output deactivated in AED mode
*		Infant patient group
₹	Patient group symbol	Child patient group
İ		Adult patient group
	Heart rate tone/pulse tone	Heart rate tone/pulse tone on
• ×	function button	Heart rate tone/pulse tone off

Symbol	Designation	Description
		Displays access to the function check.
		Requirements for function check met
	- Function check symbols	Requirements for function check not met
	runction check symbols	Fault found during function check
(ji		Consult instructions for use
4		Service due in \leq 30 days or service interval exceeded
•	Heart symbol	 In the HR parameter field: Flashes at the measured heart rate. In the Pulse parameter field: Flashes at the measured pulse rate.
	R wave marking (only with cardioversion option and only in manual mode)	Marks the R waves detected by the device for cardioversion.
	Signal bar	Indicates the signal quality of SpO ₂ measurement.
	You may now touch the patient (in AED mode with the Pictograms in parameter view menu item activated (see "13.3 AED settings", page 325))	You may now touch the patient
	 Stand clear of the patient: In AED mode with the Pictograms in parameter view menu item activated (see "13.3 AED settings", page 325) In 12-lead ECG function mode (only with 12-lead ECG option) 	Stand clear of the patient

Symbol	Designation	Description
	Progress display of 12-lead ECG (only with 12-lead ECG option)	Indicates how far the 12-lead ECG has already recorded.
	Progress display of shock capacitor charging	Indicates how far the shock capacitor has already been charged.
曹	Symbol for venous stasis	Indicates that venous stasis has been carried out.
	Timeline (only with Replay view option)	Displays the duration of an event in replay view.
Ţ	Upload session data (only with Session data upload option)	Displays access to session data upload.
4	Service reminder	Service due in \leq 30 days or service interval exceeded.

3.5 Battery and battery status indicator



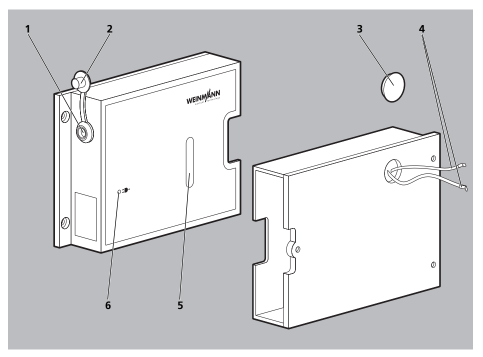
3-10 Battery and battery status indicator

ſ	No.	Designation	Description
	1	Battery	Supplies power to the device.

No.	Designation	Description	
2	Fault indicator (red)	On if the battery is defective.	
3	Status LEDs (green)	Indicate battery status.	
4	Status button	Activated by pressing the status LEDs.	

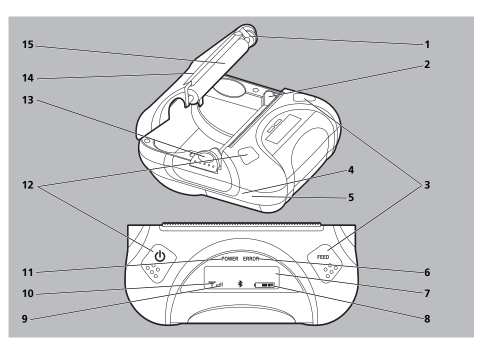
Status indicator on the battery	Status indicator on the device display	Meaning
		Battery status > 90 %
		Battery status approx. 60 % - 90 %
		Battery status approx. 40 % - 60 %
0000		Battery status approx. 10 % - 40 %
4 <u>000-0</u>		 Battery status < 10 % In the display: The last remaining segment in the battery status symbol is red. Battery weak appears in the display. In AED mode, the device outputs: <i>Battery weak</i>.
0000		 Battery is deeply discharged. Charge battery in the device for 24 h. After 24 hours: Green LED is on: Battery fully charged and ready for use Red LED or no LED on: Battery defective. Replace battery.
		Battery is empty Battery empty appears in the display and in AED mode, the device outputs: <i>Battery empty</i> . The device can be used for about another 15 minutes.
		Battery is defective. Replace battery.
		 Battery is defective. or No battery. or Battery not at suitable temperature.
		Green arrow: Battery charging





3-11 Charging station

No.	Designation	Description
1	Line power connection	Connects the charging station to line power.
2	Сар	Seals off the line power connection.
3	Cover cap	Covers the 12 V supply cable.
4	12 V supply cable	Connects the charging station to the 12 V on-board power supply in an emergency vehicle.
5	Window for viewing battery status indicator	Allows the battery status indicator to be viewed.
6	Line power indicator	Indicates that the charging station is connected to line power.



3.7 Printer and printer display

3-12 Printer and printer display

No.	Designation	Description	
1	Feed mechanism	Drives the feed roller.	
2	Print head	Prints the printer paper.	
3	Feed button (FEED)	Press and hold: Feeds the printer paper.	
4	USB port (hidden)	For service purposes only.	
5	Charging interface (hidden)	Not in use.	
6	Error indicator (ERROR)	Flashes when an error has occurred.Flashes red during switching on/switching off.	
7	LCD display	Indicates printer status.	
8	Battery status indicator	Indicates the status of the printer battery.	
9	Strength indicator	Indicates the strength of the connection.	
10	Connection indicator	Indicates that a connection has been set up between the printer and the device.	

No.	Designation	Description	
11	Status indicator (POWER)	 Flashes when the printer is switched on. Green on when the printer is switched on and ready to use. 	
12	On/Off button	 Press and hold (> 5 s): Switches the printer on or off. Press briefly (< 5 s): When the printer is switched on, displays the printer's MAC address. 	
13	Latch to release paper compartment cover	Opens the paper compartment.	
14	Paper compartment cover	Protects the paper compartment.	
15	Feed roller	Feeds the paper forwards.	

3.8 Components and accessories

3.8.1 Power supply

Accessory	Designation	Description
	Accu-Pack battery	Supplies power to the device when it is operating.
	Power supply unit and charger	Supplies the device or the charging station with power.

3 Description

Accessory	Designation	Description
	Charging station for battery WM 45045	Allows external battery charging.
	Charging adapter	Connects the power supply connection on the device to the power supply unit and charger or to the adapter cable for 12 V on- board power supply/circular connector.
	Adapter cable for 12 V on-board power supply/circular connector	Connects the device to the 12 V on-board power supply of a vehicle via the charging adapter.

3.8.2 Defibrillation/cardioversion

Accessory	Designation	Description
P. S.	MCS2-Connect master cable	Connects the following parts to the device: • Defibrillation electrodes • Paddles • Function test resistor
	MCS2-Softpads defibrillation electrodes for adults	 Conduct electrical cardiac activity to the device and shock energy to the patient. Connected to the master cable via the Pad connector. Allow defibrillation/ cardioversion of adults.
	MCS2-Softpads defibrillation electrodes for children	 Conduct electrical cardiac activity to the device and shock energy to the patient. Connected to the master cable via the Pad connector. Allow defibrillation/ cardioversion of children.
	MCS2-Hardpads paddles	 Conduct electrical cardiac activity to the device and shock energy to the patient. Connected to the master cable via the Pad connector. Allow defibrillation/ cardioversion of adults and children.

Accessory	Designation	Description
	Electrode gel	Improves electrical conductivity between the paddles and the patient's skin.

3.8.3 Pulse oximetry monitoring

Accessory	Designation	Description
	MCS2-Adapt pulse oximetry sensor connecting cable	Connects the pulse oximetry sensor to the device via the SpO ₂ connector.
	MCS2-SoftTip pulse oximetry sensor	 Measures oxygen saturation. Available in various sizes (see "17.2 Accessories and other parts", page 378).
	MCS2-Wrap pulse oximetry sensor	Measures oxygen saturation.

Accessory	Designation	Description
R	MCS2-Earclip pulse oximetry sensor with ear clip	Measures oxygen saturation.
	MCS2-Wrap pulse oximetry sensor (disposable)	 Measures oxygen saturation. Available in various sizes (see "17.2 Accessories and other parts", page 378).

3.8.4 6-lead ECG monitoring/12-lead ECG recording and assessment

Accessory	Designation	Description
	MCS2-Line ECG cable	 Conducts electrical cardiac activity to the device via the ECG connector. Can only be used for 6-lead ECG monitoring. Available in various versions (see "17.2 Accessories and other parts", page 378).

Accessory	Designation	Description
	MCS2-Line ECG cable with connection for MCS2-Line 12-lead ECG extension cable	 Conducts electrical cardiac activity to the device via the ECG connector. Can be used for 6-lead ECG monitoring and 12-lead ECG recording and assessment. Optionally connects the 12-lead ECG extension cable to the device (only with 12-lead ECG option). Available in various versions (see "17.2 Accessories and other parts", page 378).
	MCS2-Line 12-lead ECG extension cable (only with 12-lead ECG option)	 Conducts electrical cardiac activity to the device. Allows a 12-lead ECG to be derived. Available in various versions (see "17.2 Accessories and other parts", page 378).

3.8.5 Non-invasive blood pressure measurement (NIBP monitoring)

Accessory	Designation	Description
	NIBP cuff	 Measures blood pressure. Available in various sizes and versions (see "17.2 Accessories and other parts", page 378).
	NIBP connecting tube	Connects the NIBP cuff to the device.
	Adapter tube for connecting NIBP disposable cuffs for neonates	Connects NIBP cuffs for neonates (disposable).



Accessory	Designation	Description
	Printer	 Allows different ECGs and reports to be printed (see "6.16 Printing ECGs and reports (only with Printing option)", page 203). May only be used in the vicinity of the patient if it is in the printer case.
	Printer battery	 Supplies power to the printer when it is operating. May only be used in the vicinity of the patient if it is in the printer case.
	Printer case	 Protects the printer from damage and contamination. Allows the printer to be transported.
	Printer paper	Allows different ECGs and reports to be printed (see "6.16 Printing ECGs and reports (only with Printing option)", page 203).

Accessory	Designation	Description
	Power supply unit and charger for charging station for printer battery	 Charges the printer battery in the charging station for the printer battery (see "4.7.1 Charging the printer battery", page 94). Not suitable for use in the vicinity of the patient.
	Charging station for the printer battery	 Allows a printer battery to be charged (see "4.7.1 Charging the printer battery", page 94). Not suitable for use in the vicinity of the patient.
	Power supply unit and charger for quadruple charging station for the printer battery	 Charges the printer battery in the quadruple charging station for the printer battery (see "4.7.1 Charging the printer battery", page 94). Not suitable for use in the vicinity of the patient.
	Quadruple charging station for the printer battery	 Allows 4 printer batteries to be charged simultaneously (see "4.7.1 Charging the printer battery", page 94). Not suitable for use in the vicinity of the patient.



Accessory	Designation	Description
	Function test resistor	Allows a device function check to be carried out.
	SD card	Records session data and service data.
	DEFIview PC software	Facilitates the read-out and analysis of session data.
	ECG simulator	 For training purposes and for demonstrating the device. Allows ECGs to be simulated. Available in various versions (see "17.2 Accessories and other parts", page 378). Not suitable for use in the vicinity of the patient.

3.9 Transport options

In order to transport the device, carry accessories, provide a power supply for charging, and attach to a wall mounting, you can mount the device on one of the following portable units:

- LIFE-BASE 1 NG XS
- LIFE-BASE 1 NG XL
- LIFE-BASE 3 NG
- LIFE-BASE 4 NG

You can also transport the device in one of the following protective transport bags:

- MCS2-Bag protective transport bag
- MCS2-Bag protective transport bag, large

Protective transport bags cannot be attached to a wall mounting.

3.10 Options

You can tailor the range of functions on the device to your needs using the options. You can enable these options using a devicespecific enable code (see "12.2 Enabling options", page 301). The following options are available:

Option	Description	Parts required/conditions
Manual defibrillation	This option enables manual mode.	-
Cardioversion	This option allows cardioversion to be carried out in manual mode.	Manual defibrillation option enabled and activated
Printing	 This option allows various reports and printouts to be printed: Function check report Live printout Defibrillation report 12-lead ECG printout (only with 12-lead ECG option) Replay printout (only with Replay view option) Session report 	 Printer Printer paper Printer battery Charging station for printer battery with power supply unit and charger Printer case
12-lead ECG	This option enables 12-lead ECG.	 ECG cable with connection for 12-lead ECG extension cable 12-lead ECG extension cable
E-mail delivery 12-lead ECG	This option allows you to deliver a 12-lead ECG to an e-mail address.	 12-lead ECG option enabled and activated Network connected to the Internet Device registered with WEINMANN Connect (see "12.4 Registering the device with WEINMANN Connect", page 308)

Option	Description	Parts required/conditions
Replay view	This option allows an event to be reproduced and analyzed on the device.	-
Bluetooth [®] data transmission	This option allows data to be made available to a compatible system for digital patient data recording via the Bluetooth [®] interface.	Compatible system for digital patient data recording with Bluetooth [®] interface
Session data upload	This option allows session data to be uploaded to WEINMANN Connect.	 Network connected to the Internet Device registered with WEINMANN Connect (see "12.4 Registering the device with WEINMANN Connect", page 308)

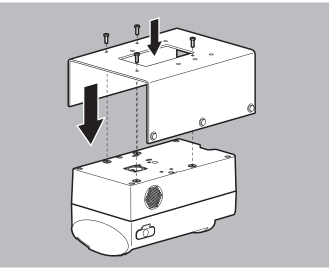
4 Preparation

4.1 Mounting the device

4.1.1 Mounting the device in the protective transport bag

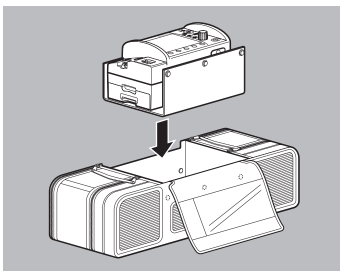
Required auxiliary equipment Phillips screwdriver, size PH1

1. Place the device on a smooth, firm surface with the control panel membrane facing down.

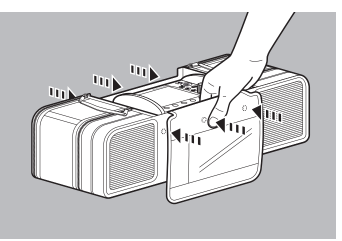


- Put the holding plate on the device. When doing so, please note: The holes of device and holding plate must line up.
- 3. Screw the device to the holding plate using screws.
- 4. Place the protective transport bag with its back on a level, firm surface.
- If the protective transport bag is closed: Release the magnetic catch of the protective transport bag and open the viewing window.

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6. Introduce the device on the holding plate into the protective transport bag from above until the snap fasteners on the protective transport bag and the holding plate line up.



- 7. Use both hands simultaneously to push opposing snap fasteners together until you hear them engage.
- *Result* The device is mounted in the protective transport bag.

4.1.2 Mounting the device on the portable unit

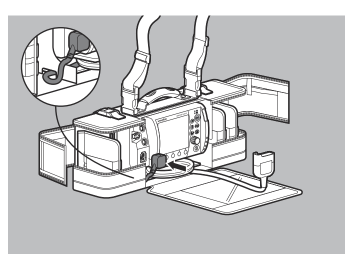
Follow the instructions for use of the portable unit.

4.2 Stowing components and accessories

4.2.1 Stowing components and accessories in the protective transport bag

Requirement The device is mounted in the protective transport bag (see "4.1.1 Mounting the device in the protective transport bag", page 74).

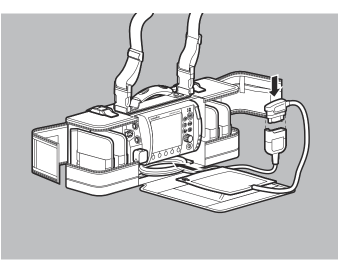
- 1. Release the magnetic catch of the protective transport bag and open the viewing window.
- 2. Open the side compartments.
- 3. If there is a divider in the left-hand side compartment: Remove the divider.



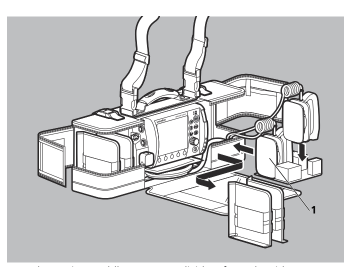
- Guide the master cable through the opening between the lefthand side compartment and the device compartment of the protective transport bag.
- 5. Connect the master cable connector to the Pad connection for master cable on the device.

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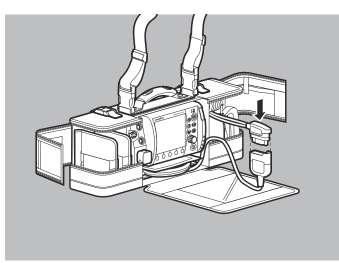
6. Put the divider back into the left-hand side compartment.



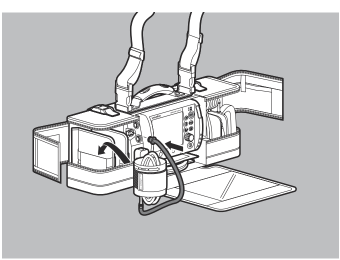
- When using defibrillation electrodes: Attach the Pad connector of the defibrillation electrodes to the master cable.
 When doing so, please note: The Pad connector must be plugged in firmly.
- 8. Wind up the master cable and stow under the device with the defibrillation electrodes.



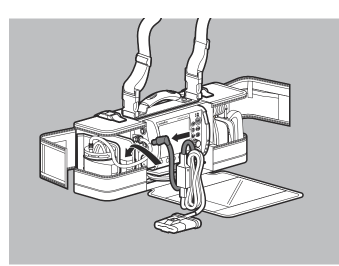
- When using paddles: Remove dividers from the side compartment.
 When doing so, please note: The paddles can only be stowed in the large versions of the protective transport bag and the protective bags for the portable units.
- 10. Insert the insert for accommodating paddles in the side compartment.
- 11. Insert the paddles one after the other. When doing so, please note:
 - The cables for the paddles must point upwards.
 - The paddle on the left (1) must be inserted first.



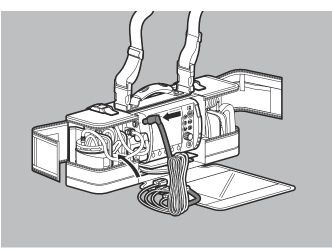
- 12. Connect the Pad connector of the paddles to the master cable. When doing so, please note: The Pad connector must be plugged in firmly.
- 13. Route the master cable in the groove between the device compartment and the side compartment of the protective transport bag.
- 14. Wind up the master cable and stow under the device.



- Connect the NIBP connecting tube to the device and the NIBP cuff (see "6.12.1 Preparing non-invasive blood pressure measurement (NIBP measurement)", page 189).
- 16. Wind up the NIBP connecting tube and the tube for the NIBP cuff and surround them with the NIBP cuff.
- 17. Stow the NIBP cuff with the tubes inside in the side compartment.



- 18. Connect the pulse oximetry sensor connecting cable to the device and to the pulse oximetry sensor (see "6.9.1 Preparing pulse oximetry monitoring", page 164).When doing so, please note: MCS2-SoftTip pulse oximetry sensors which have been turned inside out during hygienic reprocessing must be turned the right way out again.
- 19. Wind up the pulse oximetry sensor connecting cable and the cable for the pulse oximetry sensor and stow them in the side compartment.



- 20. Connect the selected ECG cable to the device (see "6.10.1 Preparing 6-lead ECG monitoring", page 171).
- 21. Wind up the ECG cable and stow it in the side compartment.
- 22. Stow ECG electrodes in their packaging in the side compartment.
- 23. If a printer is in use (only with Printing option): Stow the printer case in the protective transport bag.

Alternatively:

Mount the printer case on the protective transport bag (see "4.7.5 Mounting the printer case on the protective transport bag or the protective bag of the portable unit", page 101).

24. Stow further components and accessories in the side compartments.When doing so, please note:

- Components and accessories must be stowed so that they do not interfere with one another when removed.
- Components and accessories must be stowed so that the viewing window and the side compartments can be closed easily and securely.

25. Close side compartments and viewing window.

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Result Components and accessories are stowed in the protective transport bag.

4.2.2 Stowing components and accessories on the portable unit

Follow the instructions for use of the portable unit.

4.3 Connecting a power supply

A WARNING

Risk of injury due to missing battery!

Operation with line power without a battery prevents the device being fully ready for use as the shock capacitor in the device cannot charge. This prevents shock delivery and delays the patient's treatment.

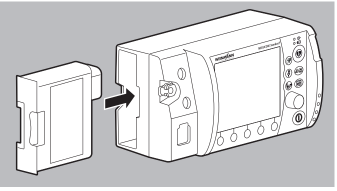
 \Rightarrow Only operate the device with the battery inserted.

- 1. Check battery status (see "3.5 Battery and battery status indicator", page 57).
- 2. If necessary: Charge battery (see "4.4.2 Charging the battery in the device", page 86).

Risk of injury from battery being incorrectly inserted!

An incorrectly inserted battery may lead to device failure and thus injure the patient.

⇒ Always push the battery into the battery compartment until you hear it engage and it is flush with the housing.



3. Slide the fully charged battery into the battery compartment until you hear it engage.

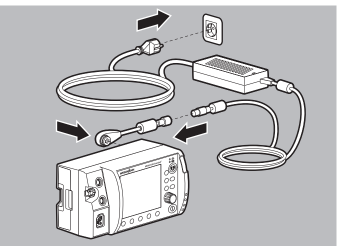
When doing so, please note: If the battery is not inserted properly, or not inserted at all, defibrillation/cardioversion is not possible and the following message appears:



4. If necessary:

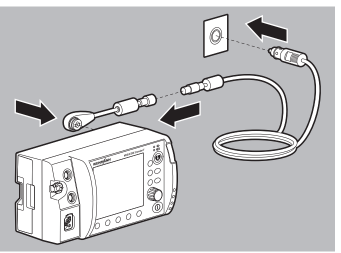
When operating on the portable unit, suspend the portable unit in a wall mounting with a charging interface.

Alternatively:



Connect the device to line power using the charging adapter and power supply unit and charger.

Alternatively:



Connect the device to a 12 V on-board power supply with the charging adapter and the adapter cable for 12 V on-board power supply/circular connector.

Result The power supply is connected.

4.4 Using the battery

4.4.1 General instructions

- Always operate the device with a battery.
- Note the methods of storing the battery and the charging intervals for extended storage (see "15.2 Storing the battery", page 373).
- The expected lifetime of the battery is 2 years. Recommendation: Replace the battery after 2 years. If battery operating time has substantially dropped before then, replace the battery earlier.
- If you receive a replacement battery, you need to charge it fully before the first use.

4.4.2 Charging the battery in the device

Requirement The portable unit is suspended in a wall mounting with a charging interface.

Alternatively:

The device is connected to line power with the charging adapter and power supply unit and charger.

Alternatively:

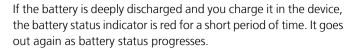
The device is connected to a 12 V on-board power supply with the charging adapter and the adapter cable for 12 V on-board power supply/circular connector.

- 1. Insert battery into the battery compartment. When doing so, please note:
 - Charging starts automatically if the following conditions are met:

Specification	Description	
External voltage	12 V to 15.1 V	
Battery status	< 95 % charged	
Battery temperature	Between 0 °C and 45 °C	

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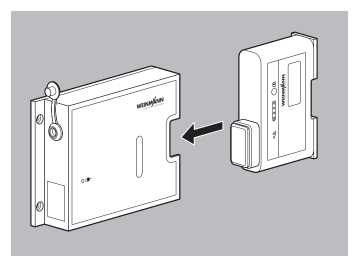
- If the device is switched on, the green arrow appears in the battery status symbol in the display (example: 4).
 and the battery status indicator on the device flashes green.
- If the device is switched off, only the battery status indicator flashes green.
- The device remains fully ready for use.



- 2. When the battery status indicator is green and/or the symbol appears in the display: Disconnect the device from the charging interface or from the power supply unit and charger.
- *Result* The battery is fully charged.

i

4.4.3 Charging the battery in the charging station



- Push the battery into the charging station until you hear it engage. Charging starts automatically if the following conditions are met:
 - External voltage of at least 12 V to 15.1 V connected

- Battery not fully charged (< 95 % charge)
- Battery temperature between 0 °C and 45 °C

The battery status LEDs flash green during the charging process.

- 2. Once the status LEDs go permanently green: Release the battery latch and take the battery out of the charging station.
- *Result* The battery is fully charged.

4.4.4 Changing the battery

Requirement The replacement battery is fully charged.

- 1. If the device is not connected to line power: Switch off the device (see "6.2 Switching the device off", page 126).
- 2. Take battery out of the battery compartment.

WARNING

Risk of injury from battery being incorrectly inserted!

An incorrectly inserted battery may lead to device failure and thus injure the patient.

⇒ Always push the battery into the battery compartment until you hear it engage and it is flush with the housing.

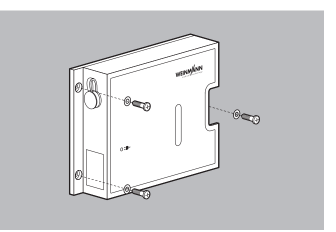
- 3. Slide the replacement battery into the battery compartment until you hear it engage.
- If necessary: Switch on the device (see "6.1 Switching on the device", page 125).
 The symbol appears in the display.
- *Result* The battery is changed.

4.5 Using the charging station

4.5.1 Mounting the charging station on the wall

Risk of injury from incorrectly mounted charging station!

If a charging station is mounted incorrectly, the charging station may come off the wall or the battery may fall out. \Rightarrow Position and attach the charging station correctly.



- 1. Position the charging station with its rear on a wall so that the compartment for inserting the battery is facing the right or left-hand side.
- Screw the charging station firmly to the wall using 3 screws and 3 plain washers (maximum screw-in depth 6 mm). When doing so, please note: The screws must be firmly tightened.
- *Result* The charging station is firmly attached to the wall.

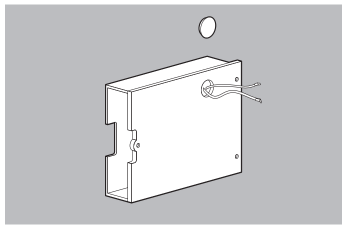
4.5.2 Operating the charging station on a mobile basis

- 1. Glue the rubber feet to the rear of the charging station.
- 2. Place the charging station with the rear on a level surface.
- *Result* The charging station is used on a mobile basis (e.g. as a desktop device).

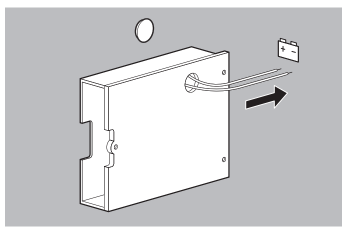
4.5.3 Connecting the charging station to line power

- 1. Remove the protective cap from the line power connection.
- 2. Connect power supply unit and charger WM 28937 to the line power connection.
- 3. Connect the power supply unit and charger to line power.
- *Result* The charging station is connected to line power. The line power indicator on the charging station is green.

4.5.4 Connecting the charging station to a 12 V on-board power supply



- 1. Remove the cover cap on the rear of the charging station.
- 2. Take the 12 V supply cable out of the charging station.
- 3. Remove the electrical insulation.
- Seal off the cable openings with liquid adhesive to stop the 12 V supply cable slipping back into the charging station and to stop dirt or water penetrating the charging station.



- 5. Connect the red wire of the 12 V supply cable to the positive terminal of the 12 V on-board power supply in the vehicle.
- 6. Connect the black wire of the 12 V supply cable to the negative terminal of the 12 V on-board power supply in the vehicle.
- *Result* The charging station is connected to the 12 V on-board power supply in the vehicle. The line power indicator of the charging station is green.

4.5.5 Charging the battery in the charging station

- 1. Charge the battery in the charging station (see "4.4.3 Charging the battery in the charging station", page 87).
- *Result* The battery is fully charged.

4.6 Using an SD card

NOTICE

Loss of data due to incorrect SD card!

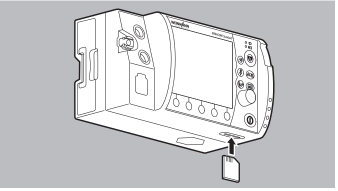
Functionality may be restricted or data may be lost in the case of SD cards not ordered via WEINMANN Emergency.

- ⇒ Only use SD cards approved by WEINMANN Emergency for use with this device.
- \Rightarrow Do not use the SD card for third-party files.

4.6.1 Inserting an SD card

The device has only a limited internal memory. To record session data over an extended period of time, you must insert an SD card:

1. Open the water jet protection of the SD card slot.



- Push the SD card into the SD card slot until you hear it engage. When doing so, please note: The beveled corner of the SD card must be at the front on the right during insertion.
- 3. Close the water jet protection to protect the device from ingress of dust and water.
- *Result* The SD card is inserted in the device and ready for use.

4.6.2 Removing the SD card

Requirement There is an SD card in the SD card slot.

1. Open the water jet protection of the SD card slot.

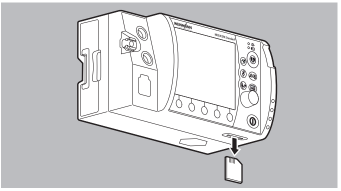


Incorrect use may result in loss of data!

If you remove the SD card while data is being written to it, data may be lost or the SD card damaged.

 \Rightarrow Only remove the SD card with the device switched off.

2. Briefly push in the SD card. The SD card is ejected slightly.



- 3. Remove the SD card.
- 4. Close the water jet protection to protect the device from ingress of dust and water.

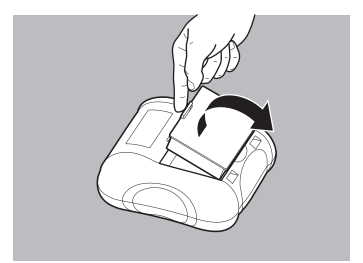
Result The SD card is removed.

4.7 Preparing printer (only with Printing option)

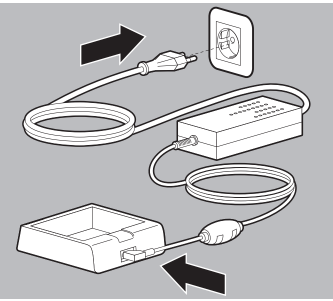
4.7.1 Charging the printer battery

Requirement

- The printer has been removed from the printer case.
 - The printer is switched off.
 - 1. Turn over the printer.

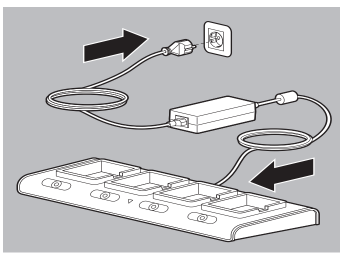


- 2. Release the printer battery latch.
- 3. Remove the printer battery from the printer.

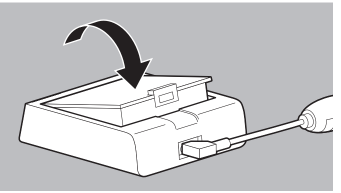


4. Connect the charging station for the printer battery to line power.

Alternatively:

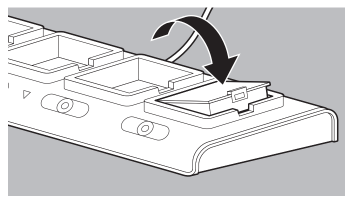


Connect the quadruple charging station for the printer battery to line power.



5. Insert the printer battery in the charging station for the printer battery until you hear it engage.

Alternatively:



Insert the printer battery in the quadruple charging station for the printer battery until you hear it engage. When doing so, please note:

- The printer battery and the charging stations heat up during charging.
- With the charging station for the printer battery: The LED on the power supply unit and charger is red when the battery is being charged.
- On the quadruple charging station for the printer battery: The CHARGING-LED is red when the battery is being charged.

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6. When the LED on the power supply unit and charger is green: Take the printer battery out of the charging station for the printer battery.

Alternatively:

If the FULL CHARGED LED is green: Take the printer battery out of the quadruple charging station for the printer battery.

Result The printer battery is fully charged.

4.7.2 Inserting the printer battery

- The printer battery is fully charged (see "4.7.1 Charging the printer battery", page 94).
 - The printer is switched off.

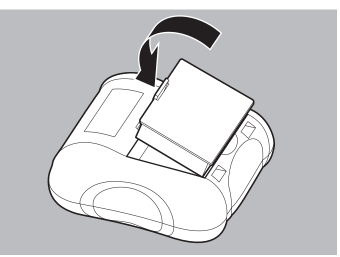


Risk of injury from electric shock when simultaneously touching a printer and the patient!

Simultaneously touching a printer and the patient may lead to an electric shock and injure the patient.

 \Rightarrow Stand clear of the patient if the printer is out of its case.

1. Turn over the printer.



2. Insert the printer battery in the battery compartment of the printer until you hear it engage.

3. Set the printer the right way up.

Result The printer battery is inserted in the printer.

4.7.3 Inserting printer paper

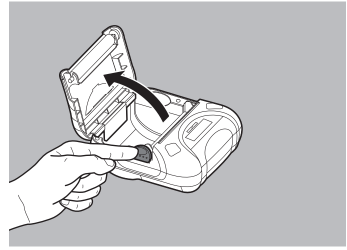
- *Requirement* The printer is not printing.
 - The printer paper provided by WEINMANN Emergency is in use.
 - The printer has been removed from the printer case.

A WARNING

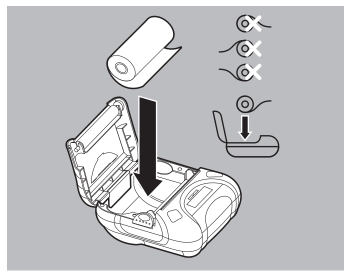
Risk of injury from electric shock when simultaneously touching a printer and the patient!

Simultaneously touching a printer and the patient may lead to an electric shock and injure the patient.

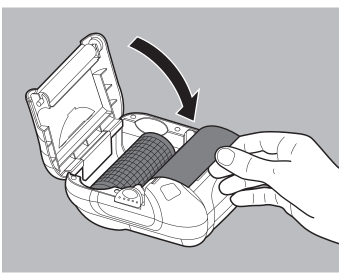
 \Rightarrow Stand clear of the patient if the printer is out of its case.



- 1. Push down the latch to release the paper compartment cover.
- 2. Open the paper compartment cover.



3. Insert printer paper in the paper compartment. The start of the printer paper must be at the bottom of the printer.



- 4. Guide the printer paper over the tear-off edge.
- 5. Close the paper compartment cover.

- 6. Press and hold the feed button (FEED) on the printer for a few seconds to feed printer paper about another 10 cm.
- *Result* The printer paper is inserted.

4.7.4 Stowing the printer in the printer case

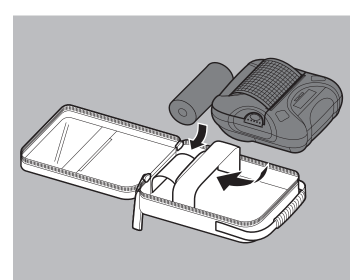
- The printer battery is inserted (see "4.7.2 Inserting the printer battery", page 97).
 - The printer paper is inserted (see "4.7.3 Inserting printer paper", page 98).

A WARNING

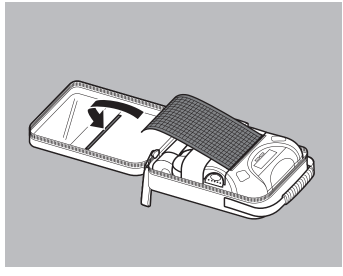
Risk of injury from contaminated printer!

A contaminated printer cannot be subjected to complete hygienic reprocessing. This may injure the patient and the user. \Rightarrow Only ever use the printer in the printer case.

1. Open the zipper on the printer case.



- 2. Stow the printer in the holder provided for it in the printer case.
- 3. If required: Stow spare printer paper in the holder provided for it in the printer case.



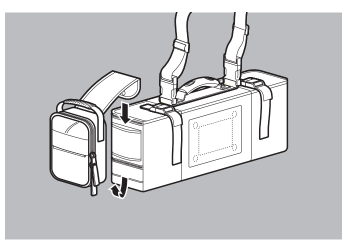
- 4. Guide printer paper through the slot in the printer case to the outside.
- 5. Close the zipper on the printer case.
- *Result* The printer is stowed in the printer case ready for printing.

4.7.5 Mounting the printer case on the protective transport bag or the protective bag of the portable unit

- The printer battery is inserted (see "4.7.2 Inserting the printer battery", page 97).
 - The printer paper is inserted (see "4.7.3 Inserting printer paper", page 98).
 - The printer is stowed in the printer case (see "4.7.4 Stowing the printer in the printer case", page 100).
 - A protective transport bag with a side holder for a printer case is being used.

Alternatively:

A protective bag of the portable unit with a side holder for a printer case is being used.



1. Push the tab of the printer case through the side holder on a protective transport bag or a protective bag of the portable unit.

When doing so, please note: Not all protective bags for the portable unit have a tab for mounting the printer case. For these portable units, the printer needs to be stowed in the accessories bag of the portable unit.

Result The printer case is mounted on the protective transport bag or a protective bag of the portable unit.

4.8 Configuring a default wireless network

A WARNING

Risk of injury due to delayed delivery!

The user can deliver a 12-lead ECG to a teleconsultation service for further diagnosis. Delivering a 12-lead ECG is purely to obtain a second opinion, and does not replace adequately trained specialists at the session location. Specialist staff at the session location are responsible for treatment within the scope of their training and in accordance with the specifications of the operator. If the 12-lead ECG reaches the teleconsultation service very late or not at all for technical reasons, delayed treatment may result in a potentially life-threatening situation. The operator is responsible for ensuring that the necessary configuration for 12-lead ECG delivery has been carried out correctly.

- ⇒ When configuring e-mail delivery 12-lead ECG: Test e-mail delivery.
- ⇒ If the IT infrastructure permits: Configure several wireless networks for various devices or session locations so that if one wireless network is unavailable, it is possible to select another wireless network.
- ⇒ Check the configuration of the wireless networks at regular intervals and update them if necessary.
- \Rightarrow Ensure that the e-mail addresses are correct and complete.
- ⇒ Check e-mail addresses at regular intervals and update them if necessary.

4.8.1 Adding a wireless network automatically

Prerequisite
 The WiFi interface of the device has been activated by the operator (see "12.3.1 Activating the WiFi interface", page 303).

- The **Allow wireless network configuration** menu item in the operator menu has been activated by the operator (see "13.11 Communication settings", page 353).
- The user menu is activated (see "10.1 Navigating the user menu", page 260).
- 1. Select Communication settings | Configure wireless network.

14:19		\triangle	Menu	
(;; 🖓 🖇				
Configure wireless network				
Add wireless network automatically				
Add wireless networ	k manually			
Edit wireless networ	k			
Back				
	N A a a i i a a	Deals	01/	
	Monitor	Back	OK	

 Select the Add wireless network automatically menu item. When doing so, please note: Some smartphones automatically switch off the WiFi hotspot. In order to ensure that the device can use the smartphone's hotspot, the dialog for switching on the WiFi hotspot must be open in the smartphone whilst the device sets up the WiFi connection.

All the wireless networks in range of the device are displayed.

A WARNING

New risks from integrating the device into an IT network! When the device is integrated into an IT network, new risks may arise for patients, users or third parties, and these must be determined, analyzed, and managed.

 \Rightarrow As the operator of the IT network, reassess changes to the IT network. When doing so, please note:

- Modified IT network configuration
- Connection of additional elements to the IT network
- Removal of additional elements from the IT network
- Updates of devices connected to the IT network
- Upgrades of devices connected to the IT network

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A WARNING

Delayed therapy due to integrating the device into a wireless network with a login page!

If the device is integrated into a wireless network with a login page, it is not possible to upload a 12-lead ECG by e-mail. \Rightarrow Do not use a wireless network with a login page.

- Select the desired wireless network. When doing so, please note: The device must only be connected to an encrypted wireless network. A password must always be entered for an encrypted wireless network.
- 4. Confirm with the **OK** function button.
- 5. Enter the password for the desired wireless network. The device connects to the desired wireless network.
- 6. In the event of problems setting up the wireless network: Contact system administrator.
- *Result* A wireless network has been automatically configured for the device and specified as the default wireless network.

From now on, the device will always automatically establish a connection to the default wireless network that has been set up when a WiFi connection is required. Following successful data upload, the WiFi connection is disconnected.

4.8.2 Adding a wireless network manually

- Prerequisite
 The WiFi interface of the device has been activated (see "12.3.1 Activating the WiFi interface", page 303).
 - The **Allow wireless network configuration** menu item in the operator menu has been activated by the operator (see "13.11 Communication settings", page 353).
 - The network configuration of the wireless network to be added manually is known.
 - The user menu is activated (see "10.1 Navigating the user menu", page 260).
 - 1. Select Communication settings | Configure wireless network.

14:27		\triangle	Menu	
(;; k k k k k k k k k k k k k k k k k k				
Configure wireless network				
Add wireless network automatically				
Add wireless network manually				
Edit wireless network	k			
Back				
	Monitor	Back	ОК	

2. Select the Add wireless network manually menu item.

14:30		\triangle	Menu
\$ № *			
Add wireless network manually			
Network name (SSID))	WM	
Password		****	
DHCP			R
Add wireless networ	k		
Back			
		Death	01/
	Monitor	Back	OK

- 3. Make the settings for the wireless network.
- 4. Select the Add wireless network menu item.
- Confirm with the **OK** function button. The wireless network is added as an available wireless network.
- *Result* A wireless network has been manually configured for the device.

4.8.3 Specifying a wireless network as the default wireless network

- Prerequisite
 The WiFi interface of the device has been activated by the operator (see "12.3.1 Activating the WiFi interface", page 303).
 - The **Allow wireless network configuration** menu item in the operator menu has been activated by the operator (see "13.11 Communication settings", page 353).
 - Several wireless networks have been configured.
 - The user menu is activated (see "10.1 Navigating the user menu", page 260).
 - 1. Select Communication settings | Configure wireless network.
 - Select the Edit wireless network menu item. All configured wireless networks are displayed.
 - 3. Select a wireless network as the default wireless network to be configured.
 - 4. Select the **OK** function button.
 - Select the Set wireless network as default menu item. The desired wireless network is specified as the default network.
 - 6. Restart device.
 - *Result* The selected wireless network has been specified as the default wireless network.

5 Function check

A WARNING

Disrupted or failed therapy due to defective device, defective components or defective accessories!

Using defective devices, defective components or defective accessories may result in malfunctions of the device, the components, and the accessories. This may cause the patient and the user serious or life-threatening injury.

- ⇒ Carry out a complete function check prior to every use (see "5.2 Carrying out a function check", page 109).
- ⇒ Only operate the device, components, and accessories if they have no external damage.
- \Rightarrow Replace illegible or damaged labels.
- \Rightarrow Only use devices, components, and accessories which have passed the function check.
- \Rightarrow Have defective devices repaired.
- ⇒ Have defective components and defective accessories repaired, or replace them.
- \Rightarrow Observe the applicable intervals for function check and service.

After being switched on, the device carries out an automatic function check which checks that all the key functions are working.

The device also offers a step-by-step guide to carrying out a function check (see "5.2 Carrying out a function check", page 109).

When the device is switched on, permanent test routines are active in the background which alert the user to malfunctions via an alarm.

The correct functioning of some components cannot be checked with the support of software. In such cases, users must assess functionality themselves (e.g. external damage to the device, functionality of the ECG cable or the NIBP cuff).

5.1 Intervals

Part concerned	Interval
Device including accessories	 Before every use After every hygienic reprocessing After every repair After every software update

5.2 Carrying out a function check

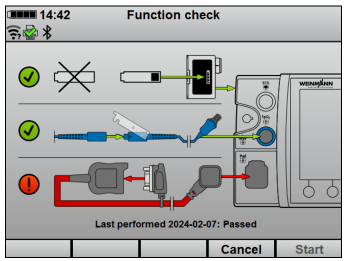
Requirement • The device is disconnected from the patient.

• A battery with a battery status of at least 1 LED is inserted in the device.

Alternatively:

- The SD card is in the SD card slot.
- The master cable is connected.
- The pulse oximetry sensor connecting cable with pulse oximetry sensor is connected.
- Only when WEINMANN Connect is used: A default wireless network has been configured (see "12.3 Configuring default network", page 302).
- Only when WEINMANN Connect is used: The device has been registered with WEINMANN Connect (see "12.4 Registering the device with WEINMANN Connect", page 308).
- 1. Check the following parts for external damage:
 - Device
 - Defibrillation electrode packaging
 - Master cable
 - Paddles
 - ECG cable
 - Pulse oximetry sensor connecting cable

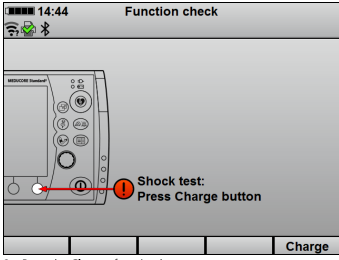
- Pulse oximetry sensor
- NIBP connecting tube
- NIBP cuff
- Printer
- 2. If necessary: Replace parts.
- Check the expiry date on the packaging for the ECG electrodes and the defibrillation electrodes.
 If necessary: Replace ECG electrodes and/or defibrillation electrodes.
- Switch on the device (see "6.1 Switching on the device", page 125). The start menu appears.
- 5. Select the **Function check** menu item in the start menu.



The automatic function check starts. The date and event of the last function check carried out are displayed.

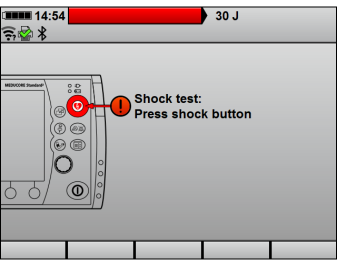
- 6. Connect the function test resistor to the master cable.
- Once all components are marked with a green check mark: Press the **Start** function button. The function check starts and can now no longer be canceled.

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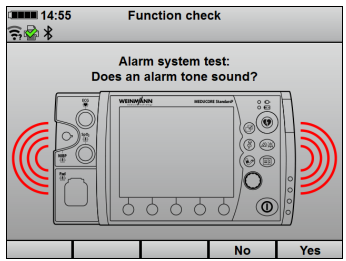


8. Press the **Charge** function button.

The shock capacitor is charged to 30 J and the shock energy is maintained for 30 s.

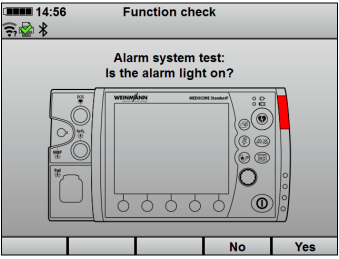


9. When the shock capacitor is fully charged and the shock button flashes: Press shock button (.

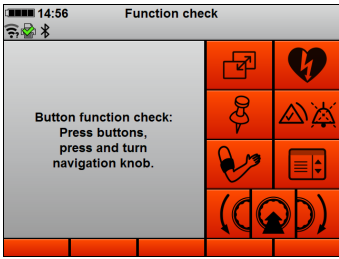


10. If an alarm tone is emitted: Press the **Yes** function button.

11. If no alarm tone is emitted: Press the **No** function button.

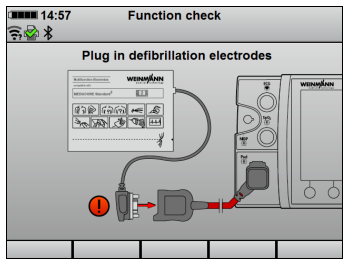


12. When the alarm light is red: Press the **Yes** function button.



13. If the alarm light is not red: Press the **No** function button.

- 14. In the button function check, press all of the controls one after the other except for the On/Off button (①).
- 15. To cancel the button function test: Press menu button () twice.



16. If defibrillation electrodes are used:

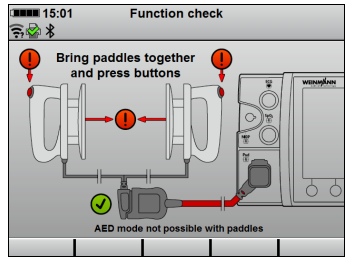
- Disconnect the function test resistor from the master cable.
- Connect the defibrillation electrodes to the master cable.

or

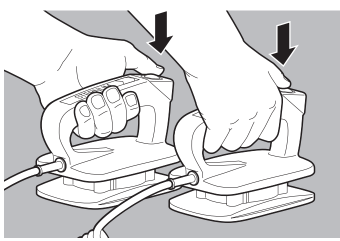
If the adapter cable for connection of corPatch easy defibrillation electrodes to MEDUCORE Standard² and corPatch easy defibrillation electrodes are used:

- Disconnect the master cable with function test resistor from the device.
- Connect the adapter cable for connection of corPatch easy defibrillation electrodes to MEDUCORE Standard² with corPatch easy defibrillation electrodes to the device.
- Confirm the connection with the **OK** function button.

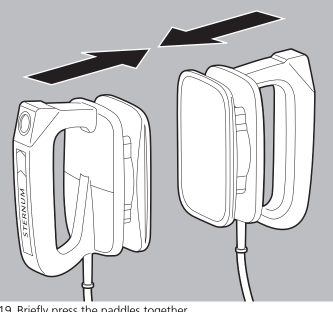
Only when WEINMANN Connect is used: The network test begins. The device tests the connection to WEINMANN Connect. If it can establish the connection successfully, the device automatically uploads the result of the function check to WEINMANN Connect.



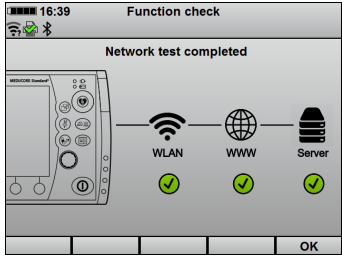
17. If paddles are being used: Disconnect the function test resistor from the master cable and connect paddles to the master cable.



18. Press both buttons on the paddles.

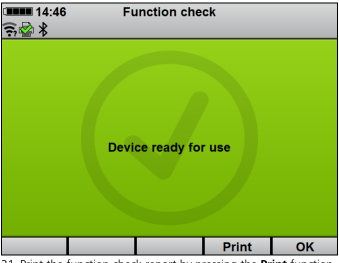


19. Briefly press the paddles together. Only when WEINMANN Connect is used: The network test begins. The device tests the connection to WEINMANN Connect. If it can establish the connection successfully, the device automatically uploads the result of the function check to WEINMANN Connect.



20. Once the network test is completed: Press the **OK** function button.

The status report appears (example: Function check passed):



 Print the function check report by pressing the **Print** function button (only with Printing option) (see "6.16.5 Printing a function check report", page 208).

22. Only if WEINMANN Connect is used and the network test has failed: Upload the function check to WEINMANN Connect manually using the **Upload** function button (see "5.3 Uploading function check to WEINMANN Connect manually", page 120).



Risk of injury due to device not ready for use!

If you operate the device after a failed function check, the patient may be injured.

 \Rightarrow Only operate the device after it passes a function check.

23. Proceed with the device according to the following table:

Display	Meaning	Action
Device ready for use	Function check passed	Use device without restriction.
Device ready for use The Upload function button is displayed (only when WEINMANN Connect is used).	 Function check passed Network test failed: Function check not uploaded to WEINMANN Connect automatically 	 Use device without restriction. Upload function check to WEINMANN Connect manually using the Upload function button (see 5.3, p. 120). Eliminate the cause of the failed network test.
Device ready for use The service symbol flashes in the start menu.	 Function check passed Information about the scheduled service 	Use device without restriction.Have device checked.
Device ready for use The service symbol flashes in the start menu and the Upload function button is displayed (only when WEINMANN Connect is used).	 Function check passed Information about the scheduled service Network test failed: Function check not uploaded to WEINMANN Connect automatically 	 Use device without restriction. Have device checked. Upload function check to WEINMANN Connect manually using the Upload function button (see 5.3, p. 120).
Device not ready for use	Function check failed or Function check canceled	 Check the parts listed in the display and replace them if necessary. Repeat the function check. If the function check is still failed: Contact the manufacturer or a technician expressly authorized by WEINMANN Emergency.

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Display	Meaning	Action
Device not ready for use The Upload function button and the cause of the failed network test are displayed (only when WEINMANN Connect is used).	 Function check failed or function check interrupted Network test failed: Function check not uploaded to WEINMANN Connect automatically 	 Check the parts listed in the display and replace them if necessary. Eliminate the cause of the failed network test. Repeat the function check. If the function check is still failed: Contact the manufacturer or a technician expressly authorized by WEINMANN Emergency. If the Upload function button and the cause of the failed network test continue to be displayed once the function check has been repeated: Upload function check to WEINMANN Connect manually using the Upload function button (see 5.3, p. 120). Contact system administrator or WEINMANN Emergency Technical Service.

- 24. Finish function check with the **OK** function button. The start menu appears.
- 25. Connect the ECG cable to the ECG connection for ECG cable.
- 26. Connect the NIBP cuff to the NIBP connection for the NIBP connecting tube using the NIBP connecting tube.
- *Result* The function check is complete. The device is ready for use.

5.3 Uploading function check to WEINMANN Connect manually

A default network has been configured (see "12.3 Configuring default network", page 302).

- The device has been registered with WEINMANN Connect (see "12.4 Registering the device with WEINMANN Connect", page 308).
- A function check has been carried out.
- The network test failed.
- The status report of the function check is displayed.



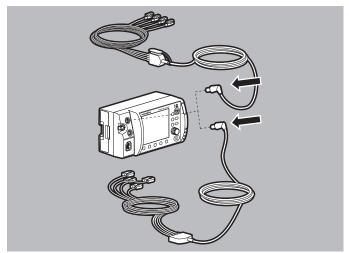
- Select the **Upload** function button. All configured networks are displayed.
- 2. Select a network within range.
- Confirm with the **OK** function button. The device connects to the selected network and uploads the result of the function check to WEINMANN Connect.
- 4. Confirm with the **OK** function button.

Result The function check has been uploaded to WEINMANN Connect.

5.4 Checking ECG cables

In addition to visual inspection of the ECG cables (see "5.2 Carrying out a function check", page 109),

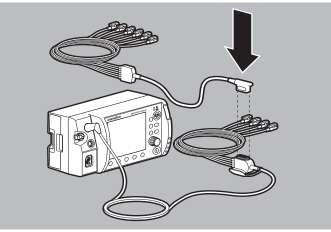
WEINMANN Emergency recommends checking the function of ECG cables at regular intervals.



1. Connect the ECG cable to the ECG connection for ECG cable.

Alternatively:

Connect the ECG cable to the ECG connection for ECG cable using the connection for 12-lead ECG extension cable.



- To check 12-lead ECG recording and assessment: Connect the 12-lead ECG extension cable to the ECG cable with connection for 12-lead ECG extension cable.
- 3. Connect the terminals of the ECG cables used to an ECG simulator:
 - ECG simulator, 6-lead ECG, shockable WM 45444
 - ECG simulator, 12-lead ECG, shockable WM 45445



In principle, any ECG simulator can be used. Alternatively, the ECG cables can also be tested on a volunteer test subject.

- 4. Switch on the ECG simulator and set a sinus rhythm.
- Switch on the device (see "6.1 Switching on the device", page 125). The start menu appears.
- 6. Select the **Adult** patient group (see "6.4 Selecting patient group", page 127).
- 7. If the ECG curve is shown in the center curve field: Press the **Lead** button to display all ECG leads.
- 8. Assess the ECG leads:
 - All ECG leads must be displayed.
 - All ECG leads must display the set sinus rhythm.

- Shaking the cable must not result in one or more ECG leads either not being displayed or being displayed with faults.
- 9. If ECG leads are not shown or are shown with faults: Replace ECG cable.
- 10. Switch off the device (see "6.2 Switching the device off", page 126).
- 11. Disconnect the ECG simulator from the ECG cable used.
- *Result* The ECG cables have been checked.

5.5 Checking NIBP cuff and NIBP connecting tube

In addition to visual inspection of the NIBP cuff and NIBP connecting tube (see "5.2 Carrying out a function check", page 109), WEINMANN Emergency recommends checking the function of parts at regular intervals:

- Carry out non-invasive blood pressure measurement on a volunteer test subject (see "6.12 Non-invasive blood pressure measurement (NIBP measurement)", page 189).
- 2. Watch for escaping air during the measurement to ensure that the system is free from leaks.
- 3. If the NIBP connecting tube leaks: Replace the NIBP connecting tube.
- 4. If the NIBP cuff leaks: Replace the NIBP cuff.
- If non-invasive blood pressure measurement takes too long or produces implausible results: Check whether the patient group and NIBP cuff are suitable for the test subject and that the NIBP cuff has been attached correctly (see "6.12.1 Preparing noninvasive blood pressure measurement (NIBP measurement)", page 189).
- 6. Repeat non-invasive blood pressure measurement.

- If the non-invasive blood pressure measurement again takes too long or produces incorrect results: Contact WEINMANN Emergency or a technician expressly authorized by WEINMANN Emergency.
- *Result* The NIBP cuff and NIBP connecting tube have been checked.

5.6 Carrying out a function check on the printer (only with Printing option)

At the end of the device function check you can print out a function check report. By assessing this report, you can check that the printer is working properly. This printer function check is not part of the device function check.

- A printer has been connected to the device (see "6.16.1 Connecting the printer to the device", page 203).
 - The printer is switched on.
 - 1. Carry out a function check of the device (see "5.2 Carrying out a function check", page 109).
 - When the status report appears: Print the function check report using the **Print** function button (see "6.16.5 Printing a function check report", page 208).
 - 3. Assess whether the printer is working properly:
 - The function check report has been printed.
 - The function check report is legible in its entirety and not interrupted.
 - No other printer defect is apparent.
 - *Result* The printer function check has been completed.

6 **Operation**

6.1 Switching on the device

Requirement • The ECG cable is not connected to the patient.

- The defibrillation electrodes or paddles are not connected to the patient.
- A fully charged battery is inserted in the device.
- 1. Briefly press the On/Off button 0.

An automatic self-test starts, which runs through the following items:

- Alarm light flashes and test tone sounds
- The start screen appears
- Shock standby indicator comes on

The self-test is successful when all of the steps mentioned have been completed.

When doing so, please note:

The start menu appears.The device starts with the presets from the operator menu.
 The device starts in the mode previously active and with the preset start view. The settings in the user menu from the last session are retained. The device assigns the session data to the last session.
The device skips the test of the ECG module and the defibrillation module.

2. If one or more conditions are not met: Do not operate the device.

- 3. Carry out a function check (see "5.2 Carrying out a function check", page 109).
- *Result* The device is switched on.

6.2 Switching the device off

Requirement The device is switched on (see "6.1 Switching on the device", page 125).

- 1. Press and hold the On/Off button 0 for at least 2 seconds.
- *Result* The device is completely switched off.

6.3 Navigating in the device

	Result			
Action	In a menu	Within a menu item	In the start menu	In a mode
Press the function button	The function is shown in the display, directly above the function button (e.g. AED or Back).			
O Turn the navigation knob counterclockwise	Navigate upwards	Decrease value	Navigate upwards	-
O Turn the navigation knob clockwise	Navigate downwards	Increase value	Navigate downwards	-
O Press the navigation knob	Select menu item	Confirm the set value	Select menu item	Activate the application menu
Press the menu button	Close the menu	Close the menu	Activate the operator menu	Activate the user menu

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	Result			
Action	In a menu	Within a menu item	In the start menu	In a mode
Press the view button	-	-	-	Switch view: Parameter view Curve view In 12-lead ECG function mode (only with 12-lead ECG option): Change the number of leads displayed
(8) Press the event button	-	-	-	Opens the events list.
Press the NIBP button	-	-	-	 Activate NIBP function mode (press for < 2 s) Start NIBP measurement (press for > 2 s)

6.4 Selecting patient group

When you select a patient group, the presets which the operator specified for this patient group are loaded. If the operator has not specified presets, the factory settings are loaded.

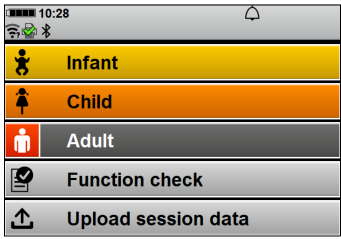
Requirement The device is switched on (see "6.1 Switching on the device", page 125).

A WARNING

Risk of injury due to incorrectly selected patient group!

If the wrong patient group is selected, the shock energy may be insufficient or too high for the selected patient group and may injure the patient.

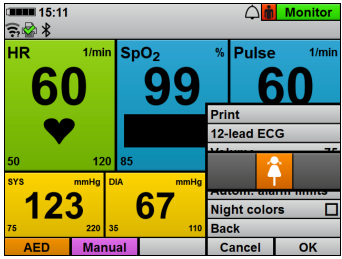
- \Rightarrow Adapt the patient group to the patient.
- \Rightarrow If the patient group is incorrect: Change the patient group in the application menu.



1. When the start menu is active: Select the patient group with the navigation knob.

Alternatively:

If the timer has expired: The device automatically selects the **Adult** patient group.

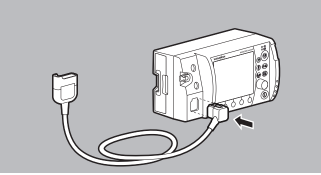


- During ongoing operation: Open the application menu with the navigation knob and change the patient group. When doing so, please note: The Infant patient group is not available in AED mode.
- *Result* The selected patient group is shown in the top right-hand corner next to mode display.

6.5 Preparing for shock delivery

6.5.1 Preparing for shock delivery using defibrillation electrodes

The following section describes how to connect the defibrillation electrodes to the device and attach them to the patient's torso. The specifications in the instructions for use provided by the manufacturer of the defibrillation electrodes and the information on the packaging of the defibrillation electrodes are key for use of the defibrillation electrodes. Observe these instructions for use and the packaging information.

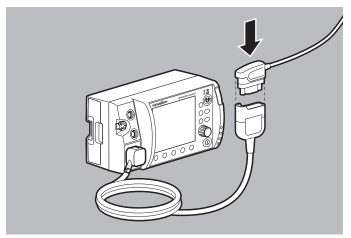


1. Connect the master cable connector to the Pad connection for master cable on the device.

Risk of injury due to incorrectly selected size of defibrillation electrodes!

If the wrong size of defibrillation electrodes is selected, this may result in sub-optimal defibrillation results or in burns.

- ⇒ Select the correct size of defibrillation electrodes in line with currently applicable guidelines and not based on the weight specifications given on the packaging.
- 2. Select defibrillation electrodes suitable for adults (Adult) or children (Pediatric).
- 3. In the case of defibrillation electrodes for children (Pediatric): Tear open the defibrillation electrode packaging and take out the defibrillation electrodes.



 Attach the Pad connector of the defibrillation electrodes to the master cable.
 When define a place are placed at the Pad energy of the placed at
When doing so, please note: The Pad connector must be plugged in firmly.

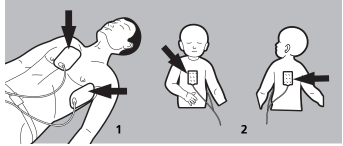
5. Bare the patient's torso.

A WARNING

Risk of injury from incorrect positioning of the defibrillation electrodes!

Incorrectly positioned defibrillation electrodes lead to a suboptimal defibrillation/cardioversion result.

- ⇒ Select the correct electrode position in line with currently applicable guidelines.
- \Rightarrow Maintain distance from ECG electrodes.



- 6. Select the desired electrode position on the patient's torso:
 - Position **1**: Sternum-apex
 - Position 2: Anterior-posterior (can also be used for adults)

🛦 WARNING 🛛

Risk of injury due to air/moisture between defibrillation electrodes and the patient's skin!

Air (e.g. in the case of hirsute patients) or moisture between the defibrillation electrodes and the patient's skin prevent correct shock delivery and may result in burns to the skin and unsuccessful defibrillation/cardioversion.

- \Rightarrow Remove hair from hirsute patients.
- \Rightarrow Rub the patient's skin dry.
- \Rightarrow Wipe down oily skin with an alcohol pad.
- \Rightarrow Press the defibrillation electrodes on firmly.
- 7. Remove hair from the torso.
- 8. Rub damp spots on the torso dry.
- 9. Wipe down oily skin with an alcohol pad.
- 10. In the case of defibrillation electrodes for adults (Adult): Tear open the defibrillation electrode packaging and take out the defibrillation electrodes.
- 11. Remove the protective film from the defibrillation electrodes.

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12. Attach defibrillation electrodes and press in place firmly.

A WARNING

Risk of injury from defibrillation electrodes not adhering correctly!

If defibrillation electrodes have been attached incorrectly and are attached again, they no longer adhere correctly and may lead to inadequate shock delivery. This may injure the patient.

- \Rightarrow Always keep spare defibrillation electrodes to hand.
- \Rightarrow If defibrillation electrodes are attached incorrectly: Always use new defibrillation electrodes.
- 13. If defibrillation electrodes are attached incorrectly: Use new defibrillation electrodes instead of the old defibrillation electrodes.
- 14. Stroke out any air trapped under the defibrillation electrodes.
- *Result* Shock delivery via defibrillation electrodes is prepared.

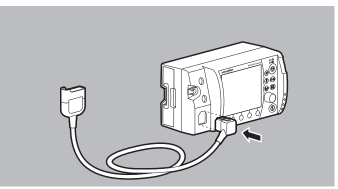
6.5.2 Preparing for shock delivery using paddles

A WARNING

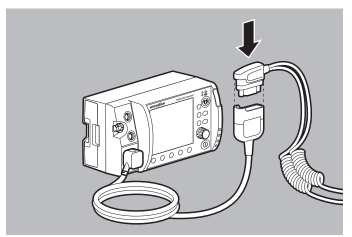
Risk of injury from using paddles with the incorrect software version!

On devices with a software version < 2.1, the device does not detect paddles so they cannot be used correctly. This may injure the patient and the user.

⇒ Only use paddles on devices running software version 2.1 or above.



1. Connect the master cable connector to the Pad connection for master cable on the device.



 Connect the Pad connector of the paddles to the master cable. When doing so, please note: The Pad connector must be plugged in firmly.

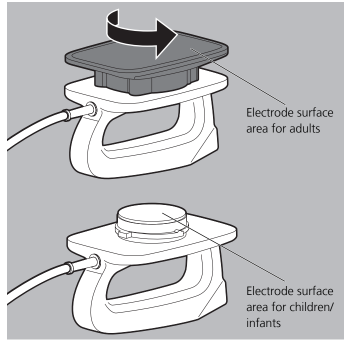
A WARNING

Risk of injury due to air/moisture between the paddles and the patient's skin!

Air (e.g. in the case of hirsute patients) or moisture between the paddles and the patient's skin prevent correct shock delivery and may result in burns to the skin and unsuccessful defibrillation/ cardioversion.

- \Rightarrow Remove hair from hirsute patients.
- \Rightarrow Rub the patient's skin dry.
- \Rightarrow Wipe down oily skin with an alcohol pad.
- \Rightarrow Always use electrode gel with paddles.
- ⇒ Always press paddles on firmly with a contact pressure of about 8 kg.
- 3. Remove hair from the torso.
- 4. Rub damp spots on the torso dry.
- 5. Wipe down oily skin with an alcohol pad.

6 Operation



- 6. Select the electrode surface area of the paddles to suit the patient group:
 - Adults: Use attachments for large electrode surface areas.
 - Children/infants: Twist off and remove attachments for large electrode surface areas.
 The attachments for small electrode surface areas are located under the attachments for large electrode surface areas.

Risk of injury due to incorrect handling of electrode gel! Incorrect handling of electrode gel may lead to electric shock, to ineffective shock delivery, and to burns, injuring the patient, user, and bystanders. \Rightarrow Always use electrode gel with paddles. \Rightarrow Do not allow any electrode gel to get between the surface of the electrode and the handle in order to prevent electric shock. \Rightarrow Do not use too much electrode gel to prevent a gel bridge and thus burns on the chest. \Rightarrow Do not use too little electrode gel to keep the resistance to the patient low for effective shock delivery and to prevent burns on the chest. Risk of injury from the incorrect electrode gel! **A** CAUTION The incorrect electrode gel may trigger intolerance reactions and lead to ineffective shock delivery. This may injure the patient. \Rightarrow Only use electrode gel recommended by

- WEINMANN Emergency.
- 7. Wet the electrode surfaces of the paddles completely with electrode gel.
- *Result* Shock delivery via paddles is prepared.

6.6 Semi-automatic defibrillation in AED mode with defibrillation electrodes

The defibrillation sequence in AED mode described here corresponds to the device settings as delivered. The operator menu enables you to adapt the device to users' qualification level and to provide optimal support to users during resuscitation measures whilst taking regional features into account.

- *Requirement* A charged battery is inserted in the device.
 - The device is switched on (see "6.1 Switching on the device", page 125).
 - A patient group is selected (see "6.4 Selecting patient group", page 127).
 - Shock delivery via defibrillation electrodes is prepared (see "6.5.1 Preparing for shock delivery using defibrillation electrodes", page 129).





Risk of injury due to unsuitable AED analysis algorithm in children below one year of age!

The device's AED analysis algorithm is not designed for children below one year of age and may result in injury to the child. \Rightarrow Do not use AED mode on children below one year of age.

Risk of injury due to use of paddles in AED mode! Shocks cannot be delivered using paddles in AED mode. This may result in injury to the patient.

 \Rightarrow Only use defibrillation electrodes in AED mode.

Risk of injury due to missing battery!

Operation with line power without a battery prevents the device being fully ready for use as the shock capacitor in the device cannot charge. This prevents shock delivery and delays the patient's treatment.

 \Rightarrow Only operate the device with the battery inserted.

A CAUTION

Delay in treatment due to simultaneous voice prompts from defibrillator and ventilator!

If the defibrillator in AED mode is used in conjunction with a ventilator (MEDUMAT Easy CPR) which also guides the user through CPR by means of voice prompts, the simultaneous voice prompts from the defibrillator and ventilator may confuse the user and delay treatment.

- \Rightarrow When using the defibrillator in AED mode and a ventilator at the same time: Switch off the ventilator voice prompts.
- Select AED mode using the **AED** function button. When doing so, please note:
 - Depending on the patient group selected, the AED settings from the operator menu which apply to this group are taken as the basis.
 - When defibrillation electrodes for children are connected, shock energy is limited to 100 J. If a higher shock energy was preset in the device, the device reduces the shock energy to 100 J.
 - The Infant patient group is not available since the AED analysis algorithm is not suitable for children under the age of 1 year.
 - No alarms are displayed or emitted in AED mode.
 - Only defibrillation electrodes can be used in AED mode.
 - During cardiac rhythm analysis, the system determines that "---" is displayed instead of heart rate.

A WARNING

Delayed or failed therapy due to defective defibrillation electrodes!

Defective defibrillation electrodes may delay or prevent analysis and continued therapy in AED mode and injure the patient. \Rightarrow Always keep spare defibrillation electrodes to hand.

2. If the AED instruction **Plug in defibrillation electrodes** appears in spite of correctly connected defibrillation electrodes: Use spare defibrillation electrodes.

WARNING

Risk of injury due to incorrectly selected patient group!

If an incorrect patient group is selected in AED mode, shock energy, energy progression and/or metronome frequency, the pause for ventilation, and the compressions/ventilation ratio may be unsuitable for the patient and injure the patient.

- \Rightarrow Adapt the patient group to the patient.
- \Rightarrow If the patient group is incorrect: Change the patient group in the application menu.
- If the patient group is incorrect: Open the application menu with the navigation knob and change the patient group.
 When doing so, please note: In AED mode, only the adult and child patient groups are available in the application menu.
- 4. Follow the voice prompts and AED instructions.
- i

If you operate the device via line power and the inserted battery is defective or if the battery does not have sufficient capacity to charge the shock capacitor, in AED mode the device guides you through CPR without charging for shocking. If you then insert an undamaged and sufficiently charged battery, the device starts cardiac rhythm analysis immediately and prepares to charge for shocking.

- *Result* The device carries out a cardiac rhythm analysis. The cardiac rhythm analysis has one of two results:
 - Shock required (see " Shock required", page 140)

or

 Shock not recommended (see "Shock not recommended", page 142)

Shock required

The device carries out a cardiac rhythm analysis, charges for shock delivery and outputs the message:

Voice prompt	AED instruction
Stand clear of the patient	Stand clear of the patient
Cardiac rhythm is being analyzed	Analysis

If, based on the cardiac rhythm analysis, the device determines that a shock is required, the device outputs the message:

Voice prompt	AED instruction
Shock required	Shock required
Press shock button	Press shock button

The shock button (v) flashes and an alarm tone sounds.

A WARNING

Risk of injury from electric shock!

The electric shock administered to the patient may injure the user or bystanders.

- \Rightarrow Stand clear of the patient.
- ⇒ Keep patient away from liquids (e.g. blood, gel or saline solution).
- ⇒ Do not touch parts in contact with the patient (e.g. bed frames or stretchers).
- \Rightarrow Keep your distance from liquids in contact with the patient.
- ⇒ Clearly warn bystanders to stand clear of the patient or parts in contact with the patient and to keep their distance from liquids in contact with the patient.

A WARNING

Failure of therapy due to defibrillation electrodes accidentally coming loose during shock delivery!

Defibrillation electrodes accidentally coming loose during shock delivery may lead to damage to the device and thus to the failure of treatment. This may injure the patient.

- \Rightarrow Ensure that the defibrillation electrodes are always connected to the device during shock delivery.
- Deliver a shock with the shock button .
 When doing so, please note:
 - If the shock button () is not pressed, the shock capacitor discharges automatically after 15 s.
 - Only with the Printing option: If the printer is connected to the device and the **Defibrillation report** menu item is activated (see "13.9 Printer settings (only with Printing option)", page 347), the printer automatically prints a defibrillation report after shock delivery (see "6.16.4 Printing a defibrillation report", page 208).
- *Result* The patient has been given an electric shock. The shock energy corresponds to the settings in the operator menu. When the device is supplied, the device settings are as follows:

Patient group	Setting
Child	First shock: 75 J Further shocks: 75 J
Adult	First shock: 150 J Further shocks: 200 J

The device guides you through CPR by means of voice prompts, AED instructions, and the metronome (see " Carrying out CPR", page 142). It warns you again to stand clear of the patient after the preset time has elapsed (120 s when device supplied), to allow it to carry out another cardiac rhythm analysis.

Shock not recommended

The device carries out a cardiac rhythm analysis, charges for shock delivery and outputs the message:

Voice prompt	AED instruction
Stand clear of the patient	Stand clear of the patient
Cardiac rhythm is being analyzed	Analysis

If, based on the cardiac rhythm analysis, the device determines that a shock is **not** required, the device outputs the message:

Voice prompt	AED instruction
Shock not recommended	Shock not recommended

1. Carry out CPR (see " Carrying out CPR", page 142).

ResultThe patient does not have a cardiac rhythm which can be
defibrillated. The device guides you through CPR by means of voice
prompts, AED instructions, and the metronome. It warns you again
to stand clear of the patient after the preset time has elapsed
(120 s when device supplied), to allow it to carry out another
cardiac rhythm analysis.

Carrying out CPR

This section describes CPR in AED mode. When supplied, the device carries out CPR with the following parameters; these, however, can be adapted by the operator:

Sotting	Patient grou	Patient group	
Setting	Adult	Child	
CPR phase			
Duration	120 s	120 s	
Pause for ventilation	5 s	5 s	

Setting	Patient group		
	Adult	Child	
Audio outputs			
CPR voice prompts	Deactivated	Deactivated	
Metronome	15:2/30:2	15:2/30:2	
CV ratio	30:2	15:2	
Metronome frequency	100/min	100/min	
Start analysis automatically			
Start analysis automatically	Activated	Activated	

After cardiac rhythm analysis and shock delivery (if necessary), the device instructs you to carry out CPR. A metronome provides a guide for chest compressions.

Voice prompt (optional)	AED instruction
Carry out cardiopulmonary resuscitation	CPR

- 1. Carry out chest compressions:
 - 30 chest compressions for the adult patient group
 - 15 chest compressions for the child patient group
 - Continuous chest compression on intubated patients

When doing so, please note:

- The metronome sets the ideal frequency.
- The device only issues certain AED instructions and voice prompts after CPR is complete (Operator menu | Adult AED settings/Child AED settings | CPR phase | Duration), as complete and correct carrying out of cardiopulmonary resuscitation takes priority.

After 30/15 metronome beats, there is a pause to allow for ventilation:

Voice prompt (optional)	AED instruction
Ventilate twice	Ventilate twice

2. Ventilate the patient twice.

The device outputs the message:

Voice prompt (optional)	AED instruction
Carry out cardiopulmonary resuscitation	CPR

- 3. Repeat the CPR sequence.
- 4. If the patient is intubated:
 - Open the application menu using the navigation knob.
 - Select the **Select** the **Sele**
- 5. If the patients shows vital signs (breathing, response): Take basic patient care steps.
- 6. Before every cardiac rhythm analysis: Check that the defibrillation electrodes are positioned correctly.
- Result CPR has been carried out.

6.7 Manual defibrillation (only with Manual defibrillation option)

6.7.1 Carrying out manual defibrillation using defibrillation electrodes

This function is only available if manual mode has been enabled and activated by the operator: **Operator menu | System settings | Enable options | Manual defibrillation** (see "13.12 System settings", page 358).

If you are the operator of the device and have access to the operator menu, you can disable manual mode: **Operator menu | System settings | Disable functions | Manual mode** (see "13.12 System settings", page 358).

Requirement	• A charged battery is inserted in the device.
	• The device is switched on (see "6.1 Switching on the device", page 125).
	• A patient group is selected (see "6.4 Selecting patient group", page 127).
	• Shock delivery via defibrillation electrodes is prepared (see "6.5.1 Preparing for shock delivery using defibrillation electrodes", page 129).
WARNING	 Risk of injury due to lack of knowledge and failure to follow guidelines in manual mode! The use of manual mode by users without medical qualifications and training in defibrillation/cardioversion and/or failure to follow guidelines may result in injury to the patient, user or bystanders. ⇒ Only use manual mode if the user has a medical qualification and is familiar with device operation and options. ⇒ Follow currently applicable guidelines on defibrillation/cardioversion. ⇒ Observe national and regional provisions on defibrillation/cardioversion. ⇒ Observe organizational guidelines on defibrillation/cardioversion. ⇒ Include the patient condition when deciding on treatment. Risk of injury due to incorrectly selected patient group! If the wrong patient group is selected, the shock energy may be insufficient or too high for the selected patient group and may injure the patient. ⇒ Adapt the patient group to the patient. ⇒ If the patient group is incorrect: Change the patient group in the application menu. 1. If the patient group is incorrect: Open the application menu with the navigation knob and change the patient group.
	2. Select manual mode with the Manual function button. When doing so, please note:

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- It is not possible to switch to parameter view in manual mode. If you activate manual mode from parameter view, the device automatically switches to curve view, since for manual shock delivery the ECG analysis is required in the display.
- Audio alarm output is deactivated in manual mode.
- 3. To activate audio alarm output: Briefly press the alarm button (公政).
- 4. Evaluate the ECG lead.
- If shock is required: Select shock energy using the Energy function button.
 When doing so, please note: When the defibrillation electrodes for children (Pediatric) are connected, the shock energy is automatically restricted to 100 J. It is not possible to set a higher shock energy in manual mode.
- 6. Press the Charge function button. The charge progress bar appears. A rising charging tone sounds until the device is ready to deliver the shock. When the device is charged, a sequence of tones sounds which signals shock standby and the shock button () flashes.
- 7. To cancel shock charging: Cancel shock charging by pressing the **Cancel** function button or by switching to another mode.
- 8. Check the ECG leads to see whether defibrillation is still indicated.

Risk of injury from electric shock!

The electric shock administered to the patient may injure the user or bystanders.

- \Rightarrow Stand clear of the patient.
- ⇒ Keep patient away from liquids (e.g. blood, gel or saline solution).
- ⇒ Do not touch parts in contact with the patient (e.g. bed frames or stretchers).
- \Rightarrow Keep your distance from liquids in contact with the patient.
- ⇒ Clearly warn bystanders to stand clear of the patient or parts in contact with the patient and to keep their distance from liquids in contact with the patient.

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Failure of therapy due to defibrillation electrodes accidentally coming loose during shock delivery!

Defibrillation electrodes accidentally coming loose during shock delivery may lead to damage to the device and thus to the failure of treatment. This may injure the patient.

- \Rightarrow Ensure that the defibrillation electrodes are always connected to the device during shock delivery.
- Deliver a shock with the shock button (). When doing so, please note:
 - If the shock button () is not pressed, the shock capacitor discharges automatically after 30 seconds.
 - Only with the Printing option: If the printer is connected to the device and the **Defibrillation report** menu item is activated (see "13.9 Printer settings (only with Printing option)", page 347), the printer automatically prints a defibrillation report after shock delivery (see "6.16.4 Printing a defibrillation report", page 208).
- *Result* The patient has been given an electric shock.

6.7.2 Carrying out manual defibrillation using paddles

This function is only available if manual mode has been enabled and activated by the operator: **Operator menu | System settings | Enable options | Manual defibrillation** (see "13.12 System settings", page 358).

If you are the operator of the device and have access to the operator menu, you can disable manual mode: **Operator menu | System settings | Disable functions | Manual mode** (see "13.12 System settings", page 358).

Requirement

- A charged battery is inserted in the device.
 - The device is switched on (see "6.1 Switching on the device", page 125).
 - A patient group is selected (see "6.4 Selecting patient group", page 127).

 Shock delivery via paddles is prepared (see "6.5.2 Preparing for shock delivery using paddles", page 133).

A WARNING

Risk of injury due to lack of knowledge and failure to follow guidelines in manual mode!

The use of manual mode by users without medical qualifications and training in defibrillation/cardioversion and/or failure to follow guidelines may result in injury to the patient, user or bystanders. ⇒ Only use manual mode if the user has a medical qualification and is familiar with device operation and options.

- ⇒ Follow currently applicable guidelines on defibrillation/ cardioversion.
- ⇒ Observe national and regional provisions on defibrillation/ cardioversion.
- ⇒ Observe organizational guidelines on defibrillation/ cardioversion.
- \Rightarrow Include the patient condition when deciding on treatment.

Risk of injury due to incorrectly selected patient group! If the wrong patient group is selected, the shock energy may be insufficient or too high for the selected patient group and may injure the patient.

- \Rightarrow Adapt the patient group to the patient.
- \Rightarrow If the patient group is incorrect: Change the patient group in the application menu.
- 1. If the patient group is incorrect: Open the application menu with the navigation knob and change the patient group.
- 2. Adapt the electrode surface area of the paddles to suit the changed patient group (see "6.5.2 Preparing for shock delivery using paddles", page 133).
- 3. Select manual mode with the **Manual** function button. When doing so, please note:
 - It is not possible to switch to parameter view in manual mode. If you activate manual mode from parameter view, the device automatically switches to curve view, since for manual shock delivery the ECG analysis is required in the display.
 - Audio alarm output is deactivated in manual mode.

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4. To activate audio alarm output: Briefly press the alarm button (公論).

A WARNING

Risk of injury from movement artifacts during rapid derivation!

Movement artifacts may falsify the ECG and lead to misinterpretations. This may injure the patient.

- \Rightarrow In the normal case, use 6-lead ECG monitoring.
- \Rightarrow Only use rapid derivation in an emergency.
- 5. Prepare 6-lead ECG monitoring (see "6.10.1 Preparing 6-lead ECG monitoring", page 171).

Alternatively:

Position paddles on the upper torso in line with currently applicable guidelines to carry out a rapid derivation.

- 6. Evaluate the ECG lead.
- 7. If necessary: Select another ECG lead using the **Lead** function button.

Risk of injury due to incorrectly selected shock energy in children!

If the attachments for small electrode surfaces are used in combination with too high a shock energy on the paddles, this may lead to injuries in children/infants.

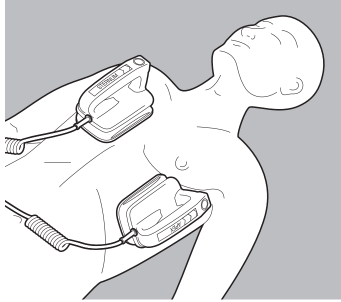
- ⇒ When using the attachments for small electrode surfaces, set a maximum shock energy of 100 J for children/infants.
- 8. If shock is required: Select shock energy using the **Energy** function button.

A WARNING

Risk of injury from incorrect positioning of the paddles!

Incorrect positioning of the paddles may lead to incorrect measurements and unsuccessful defibrillation. This may injure the patient.

- \Rightarrow Select the position of the paddles in accordance with the diagram.
- \Rightarrow Maintain distance from ECG electrodes.



- 9. Position paddles on the upper torso in accordance with currently applicable guidelines.
- 10. Briefly press and release the charge button (CHARGE) on the **APEX** paddle.

When doing so, please note: The shock capacitor can only be charged if the user menu is not activated.

The charge progress bar appears. A rising charging tone sounds until the device is ready to deliver the shock.

- 11. To cancel shock charging: Cancel shock charging by pressing the **Cancel** function button or by switching to another mode.
- 12. Check the ECG leads to see whether defibrillation is still indicated.

A WARNING

Risk of injury due to paddles having an inadequate contact pressure!

Too low a contact pressure of the paddles leads to a high resistance against the patient and may prevent shock delivery (at a resistance > 400 Ω). This may injure the patient.

 \Rightarrow Always press paddles on with a contact pressure of about 8 kg.

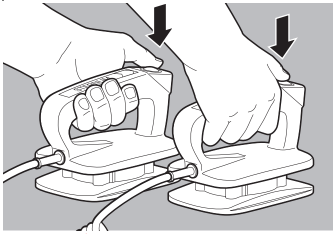
13. Press on paddles with a contact pressure of about 8 kg to keep resistance to the patient low.

A WARNING

Risk of injury from electric shock!

The electric shock administered to the patient may injure the user or bystanders.

- \Rightarrow Stand clear of the patient.
- ⇒ Keep patient away from liquids (e.g. blood, gel or saline solution).
- ⇒ Do not touch parts in contact with the patient (e.g. bed frames or stretchers).
- \Rightarrow Keep your distance from liquids in contact with the patient.
- ⇒ Clearly warn bystanders to stand clear of the patient or parts in contact with the patient and to keep their distance from liquids in contact with the patient.



14. Press both A buttons (SHOCK) on the paddles simultaneously.

When doing so, please note: Only with Printing option: If the printer is connected to the device and the **Defibrillation report** menu item is activated (see "13.9 Printer settings (only with Printing option)", page 347), the printer prints a defibrillation report (see "6.16.4 Printing a defibrillation report", page 208).

Result The patient has been given an electric shock.

6.8 Cardioversion (only with Cardioversion option)

6.8.1 Carrying out cardioversion using defibrillation electrodes

In cardioversion, shock delivery is synchronized with the R wave of the ECG (**SYNC**).

- *Requirement* A charged battery is inserted in the device.
 - The patient is prepared in line with currently applicable guidelines.
 - The device is switched on (see "6.1 Switching on the device", page 125).
 - A patient group is selected (see "6.4 Selecting patient group", page 127).
 - Shock delivery via defibrillation electrodes is prepared (see "6.5.1 Preparing for shock delivery using defibrillation electrodes", page 129).
 - 6-lead ECG monitoring is prepared (see "6.10.1 Preparing 6-lead ECG monitoring", page 171).

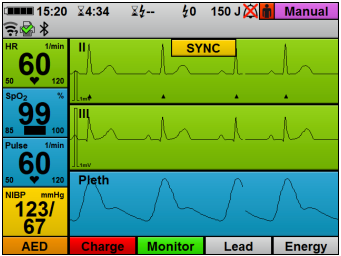
WARNING

Risk of injury from movement artifacts when using defibrillation electrodes for synchronous shock delivery!

Cardioversion cannot be carried out without 6-lead monitoring, as the device may erroneously detect movement artifacts as R waves and deliver a shock at the wrong time. This may injure the patient. ⇒ Always carry out 6-lead ECG monitoring for synchronous shock delivery.

A WARNING	Risk of injury due to lack of knowledge and failure to follow guidelines in manual mode!
	 The use of manual model. The use of manual model by users without medical qualifications and training in defibrillation/cardioversion and/or failure to follow guidelines may result in injury to the patient, user or bystanders. ⇒ Only use manual mode if the user has a medical qualification and is familiar with device operation and options. ⇒ Follow currently applicable guidelines on defibrillation/ cardioversion. ⇒ Observe national and regional provisions on defibrillation/ cardioversion. ⇒ Observe organizational guidelines on defibrillation/
	cardioversion.
	\Rightarrow Include the patient condition when deciding on treatment. Risk of injury due to incorrectly selected patient group!
A WARNING	 If the wrong patient group is selected, the shock energy may be insufficient or too high for the selected patient group and may injure the patient. ⇒ Adapt the patient group to the patient. ⇒ If the patient group is incorrect: Change the patient group in the application menu.
	 If the patient group is incorrect: Open the application menu with the navigation knob and change the patient group.
	2. Select manual mode with the Manual function button. When doing so, please note:
	• It is not possible to switch to parameter view in manual mode. If you activate manual mode from parameter view, the device automatically switches to curve view, since for manual shock delivery the ECG analysis is required in the display.
	• Audio alarm output is deactivated in manual mode.
	3. To activate audio alarm output: Briefly press the alarm button 🔊 .
	4. Evaluate the ECG lead.
	5. If necessary: Select another ECG lead using the Lead function button.

- 6. If cardioversion is required: Select shock energy using the **Energy** function button.
- 7. Open the application menu using the navigation knob.



8. Select and activate the **SYNC** menu item. Manual mode switches to cardioversion:

- The **SYNC** marking indicates that the next shock delivery will be synchronized with the R wave.
- In the ECG, R waves are marked with a triangle **A**.
- 9. Wait until the ECG curve has stabilized.

A WARNING

Delayed therapy due to incorrectly selected feed rate for the ECG!

An incorrectly selected feed rate may make it harder to assess R waves, delaying therapy.

- \Rightarrow Select the feed rate of the ECG so that reliable assessment of R wave detection is possible.
- 10. Select the feed rate of the ECG so that reliable assessment of R wave detection is possible.

A CAUTION

Risk of injury due to shock being delivered at the wrong time during cardioversion!

A cardioversion carried out at the wrong time may lead to cardiac arrhythmias and cause the patient serious or life-threatening injury.

- \Rightarrow Ensure that the ECG is stable.
- \Rightarrow Ensure that the device detects and marks R waves correctly in the ECG.

Risk of injury from pacemaker pulses incorrectly detected as R waves!

Pacemaker pulses may lead to the device interpreting them as R waves and triggering a shock at the wrong time.

- \Rightarrow Ensure that the device detects and marks R waves correctly in the ECG.
- 11. Check the ECG curve, and that the triangles **a** match the R waves.

When doing so, please note:

- The ECG curve must be stable.
- The triangles \blacktriangle must be present.
- The position of the triangles **a** must match the position of the R waves.
- The triangles **A** must not be displaced from heartbeat to heartbeat.
- The triangles 🔺 must mark every R wave reliably.
- 12. Press the **Charge** function button.

The charge progress bar appears. A rising charging tone sounds until the device is ready to deliver the shock. When the device is charged, a sequence of tones sounds which signals shock standby and the shock button () flashes.

- 13. To cancel shock charging: Cancel shock charging by pressing the **Cancel** function button or by switching to another mode.
- 14. Check the ECG leads to see whether cardioversion is still indicated.

	Risk of injury due to movement artifacts!
A WARNING	Movement artifacts erroneously detected as R waves may cause
	the device to deliver a shock at the wrong time and thus injure the
	patient.
	\Rightarrow Prepare the patient for cardioversion in line with currently
	applicable guidelines in order to avoid patient movements.
	\Rightarrow Stand clear of the patient to avoid movement artifacts.
	Risk of injury from electric shock!
	The electric shock administered to the patient may injure the user
	or bystanders.
	\Rightarrow Stand clear of the patient.
	\Rightarrow Keep patient away from liquids (e.g. blood, gel or saline
	solution).
	\Rightarrow Do not touch parts in contact with the patient (e.g. bed frames or stretchers).
	\Rightarrow Keep your distance from liquids in contact with the patient.
	\Rightarrow Clearly warn bystanders to stand clear of the patient or parts in
	contact with the patient and to keep their distance from liquids
	in contact with the patient.
	Failure of therapy due to defibrillation electrodes accidentally
A WARNING	coming loose during shock delivery!
	Defibrillation electrodes accidentally coming loose during shock
	delivery may lead to damage to the device and thus to the failure
	of treatment. This may injure the patient.
	\Rightarrow Ensure that the defibrillation electrodes are always connected
	to the device during shock delivery.
	15 Press and hold the shock button 🝙 to deliver the shock

- When doing so, please note:
 - The device synchronizes shock delivery with the next R wave.
 - If the shock button is not held down, or the device does not detect an R wave after the shock button has been pressed, the shock capacitor discharges automatically after 5 s.

- Only with Printing option: If the printer is connected to the device and the **Defibrillation report** menu item is activated (see "13.9 Printer settings (only with Printing option)", page 347), the printer prints a defibrillation report (see "6.16.4 Printing a defibrillation report", page 208).
- After shock delivery, cardioversion (SYNC marking) remains activated in manual mode. You can set in the operator menu whether further cardioversion or defibrillation is to follow cardioversion (see "13.5 Manual mode settings (only with Manual defibrillation option)", page 336).
- *Result* The patient has undergone cardioversion.

6.8.2 Carrying out cardioversion using paddles

In cardioversion, shock delivery is synchronized with the R wave of the ECG (**SYNC**). The ECG is derived via the ECG cable.

- *Requirement* A charged battery is inserted in the device.
 - The patient is prepared in line with currently applicable guidelines.
 - The device is switched on (see "6.1 Switching on the device", page 125).
 - A patient group is selected (see "6.4 Selecting patient group", page 127).
 - Shock delivery via paddles is prepared (see "6.5.2 Preparing for shock delivery using paddles", page 133).
 - 6-lead ECG monitoring is prepared (see "6.10.1 Preparing 6-lead ECG monitoring", page 171).

Risk of injury from movement artifacts when using paddles for synchronous shock delivery!

Cardioversion cannot be carried out without 6-lead monitoring, as the device may erroneously detect movement artifacts as R waves and deliver a shock at the wrong time. This may injure the patient. \Rightarrow Always carry out 6-lead ECG monitoring for synchronous shock

delivery.

Risk of injury due to lack of knowledge and failure to follow guidelines in manual mode!

The use of manual mode by users without medical qualifications and training in defibrillation/cardioversion and/or failure to follow guidelines may result in injury to the patient, user or bystanders.

- \Rightarrow Only use manual mode if the user has a medical qualification and is familiar with device operation and options.
- ⇒ Follow currently applicable guidelines on defibrillation/ cardioversion.
- ⇒ Observe national and regional provisions on defibrillation/ cardioversion.
- ⇒ Observe organizational guidelines on defibrillation/ cardioversion.

 \Rightarrow Include the patient condition when deciding on treatment.

Risk of injury due to incorrectly selected patient group! If the wrong patient group is selected, the shock energy may be insufficient or too high for the selected patient group and may injure the patient.

- \Rightarrow Adapt the patient group to the patient.
- \Rightarrow If the patient group is incorrect: Change the patient group in the application menu.
- 1. If the patient group is incorrect: Open the application menu with the navigation knob and change the patient group.
- 2. Adapt the electrode surface area of the paddles to suit the changed patient group (see "6.5.2 Preparing for shock delivery using paddles", page 133).
- 3. Select manual mode with the **Manual** function button. When doing so, please note:
 - It is not possible to switch to parameter view in manual mode. If you activate manual mode from parameter view, the device automatically switches to curve view, since for manual shock delivery the ECG analysis is required in the display.
 - Audio alarm output is deactivated in manual mode.
- 5. Evaluate the ECG lead.

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A WARNING

6. If necessary: Select another ECG lead using the **Lead** function button.

A CAUTION

Risk of injury due to incorrectly selected shock energy in children!

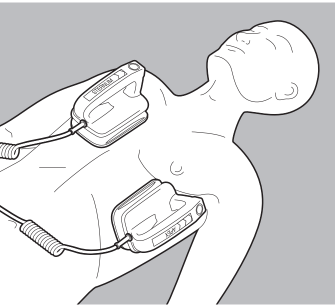
If the attachments for small electrode surfaces are used in combination with too high a shock energy on the paddles, this may lead to injuries in children/infants.

- ⇒ When using the attachments for small electrode surfaces, set a maximum shock energy of 100 J for children/infants.
- 7. If cardioversion is required: Select shock energy using the **Energy** function button.
- 150 J 🔀 💼 ¥4--40 Manual ŝ¢ ∦ 1/min HR SYNC п 120 50 SpO₂ % Ш 100 Pulse 1/min Pleth 50 120 ¥ NIBP mmHq 123/ 67 AED Charge Monitor Lead Energy
- 8. Open the application menu using the navigation knob.

- Select and activate the SYNC menu item. Manual mode switches to cardioversion:
 - The **SYNC** marking indicates that the next shock delivery will be synchronized with the R wave.
 - In the ECG, R waves are marked with a triangle

Risk of injury due to shock delivery with an unstable ECG curve! An unstable ECG curve may lead to shock delivery at the wrong time. This may injure the patient. \Rightarrow Always wait until the ECG curve has stabilized. 10. Wait until the ECG curve has stabilized. Delayed therapy due to incorrectly selected feed rate for the A WARNING ECG! An incorrectly selected feed rate may make it harder to assess R waves, delaying therapy. \Rightarrow Select the feed rate of the ECG so that reliable assessment of R wave detection is possible. 11 Select the feed rate of the ECG so that reliable assessment of R wave detection is possible. Risk of injury due to shock being delivered at the wrong time A WARNING during cardioversion! A cardioversion carried out at the wrong time may lead to cardiac arrhythmias and cause the patient serious or life-threatening injury. \Rightarrow Ensure that the ECG is stable. \Rightarrow Ensure that the device detects and marks R waves correctly in the ECG. Risk of injury from pacemaker pulses incorrectly detected as **A** CAUTION R waves! Pacemaker pulses may lead to the device interpreting them as R waves and triggering a shock at the wrong time. \Rightarrow Ensure that the device detects and marks R waves correctly in the ECG. 12. Check the ECG curve, and that the triangles A match the R waves. When doing so, please note: The ECG curve must be stable. The triangles must be present. the R waves. The triangles A must not be displaced from heartbeat to heartbeat.

The triangles 🔺 must mark every R wave reliably.



- 13. Position paddles on the upper torso in accordance with currently applicable guidelines.
- 14. Briefly press and release the charge button (CHARGE) on the **APEX** paddle.

When doing so, please note: The shock capacitor can only be charged if the user menu is not activated.

The charge progress bar appears. A rising charging tone sounds until the device is ready to deliver the shock.

- 15. To cancel shock charging: Cancel shock charging by pressing the **Cancel** function button or by switching to another mode.
- 16. Check the ECG leads to see whether cardioversion is still indicated.

Risk of injury due to paddles having an inadequate contact pressure!

Too low a contact pressure of the paddles leads to a high resistance against the patient and may prevent shock delivery (at a resistance > 400 Ω). This may injure the patient.

 \Rightarrow Always press paddles on with a contact pressure of about 8 kg.

17. Press on paddles with a contact pressure of about 8 kg to keep resistance to the patient low.

Risk of injury due to movement artifacts!

Movement artifacts erroneously detected as R waves may cause the device to deliver a shock at the wrong time and thus injure the patient.

- \Rightarrow Hold the paddles still after pressing them on.
- \Rightarrow Again, check the ECG curve and that the triangles \blacktriangle match the R waves before shock delivery.
- 18. Again, check the ECG curve and that the triangles a match the R waves before shock delivery.

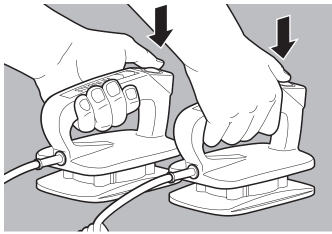
A WARNING

Risk of injury from electric shock!

The electric shock administered to the patient may injure the user or bystanders.

- \Rightarrow Stand clear of the patient.
- ⇒ Keep patient away from liquids (e.g. blood, gel or saline solution).
- \Rightarrow Do not touch parts in contact with the patient (e.g. bed frames or stretchers).
- \Rightarrow Keep your distance from liquids in contact with the patient.
- ⇒ Clearly warn bystanders to stand clear of the patient or parts in contact with the patient and to keep their distance from liquids in contact with the patient.

6 Operation



- 19. Press and hold both A (SHOCK) buttons on the paddles simultaneously.When doing so, please note:
 - The device synchronizes shock delivery with the next R wave.
 - Only with Printing option: If the printer is connected to the device and the **Defibrillation report** menu item is activated (see "13.9 Printer settings (only with Printing option)", page 347), the printer prints a defibrillation report (see "6.16.4 Printing a defibrillation report", page 208).
 - After shock delivery, cardioversion (SYNC marking) remains activated in manual mode. You can set in the operator menu whether further cardioversion or defibrillation is to follow cardioversion (see "13.5 Manual mode settings (only with Manual defibrillation option)", page 336).
- *Result* The patient has undergone cardioversion.

6.9 Pulse oximetry monitoring

6.9.1 Preparing pulse oximetry monitoring

Requirement

- The device is switched on (see "6.1 Switching on the device", page 125).
 - A patient group is selected (see "6.4 Selecting patient group", page 127).

A WARNING

Risk of injury due to incorrect use of the pulse oximetry sensor!

Incorrect use of the pulse oximetry sensor may falsify measurement results and lead to patient injury.

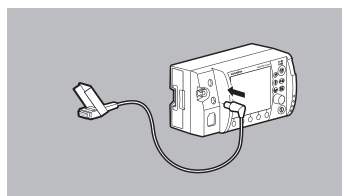
- ⇒ Keep the pulse oximetry sensor away from strong electromagnetic sources (e.g. electrosurgical devices).
- \Rightarrow Do not use the pulse oximetry sensor in areas subject to X-ray (e.g. with MRI devices).
- ⇒ Keep the pulse oximetry sensor away from strong and fluctuating ambient light (including infrared and UV light). If necessary: Cover the pulse oximetry sensor.
- ⇒ Avoid strong movement of the pulse oximetry sensor. If necessary: To relieve strain, loop the pulse oximetry sensor cable and the pulse oximetry sensor connecting cable and fix to the patient with a plaster.
- \Rightarrow Do not attach the pulse oximetry sensor to a limb on which there is already an NIBP cuff or catheter port.
- ⇒ Keep the pulse oximetry sensor away from nail polish and artificial fingernails.
- \Rightarrow Keep the pulse oximetry sensor away from intravascular dyes.
- ⇒ Be aware of deviations from the measurement result with a high proportion of dysfunctional hemoglobins.
- ⇒ Be aware of deviations from the measurement result in the case of severe anemia, venous pulsation and high total bilirubin values.
- ⇒ Be aware of deviations in pulse rate with an intra-aortic balloon pump or certain arrhythmias.

If necessary: Compare the pulse rate with the heart rate determined by ECG monitoring.

 \Rightarrow Be aware of deviations from the measurement result during defibrillation/cardioversion.

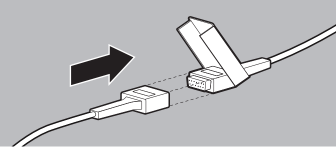
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- \Rightarrow Only use undamaged pulse oximetry sensors.
- ⇒ Only use the pulse oximetry sensors and pulse oximetry sensor connecting cables quoted in the scope of supply and in the accessories.

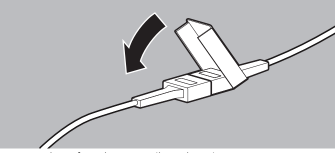


- 1. Connect the SpO₂ connector of the pulse oximetry sensor connecting cable to the SpO₂ connection on the device.
- 2. Select the appropriate pulse oximetry sensor for the patient group:

Pulse oximetry sensor	Patient group	Application site
MCS2-SoftTip pulse oximetry sensor, size S	Ø 7.5 mm - 12.5 mm finger diameter	
MCS2-SoftTip pulse oximetry sensor, size M	Ø 10 mm - 19 mm finger diameter	Finger/big toe
MCS2-SoftTip pulse oximetry sensor, size L	Ø 12.5 mm - 25.5 mm finger diameter	
MCS2-Wrap pulse oximetry sensor	> 10 kg body weight	Finger/hand
MCS2-Earclip pulse oximetry sensor	> 30 kg body weight	Ear
MCS2-Wrap pulse oximetry sensor, adult (Adult), disposable	> 30 kg body weight	
MCS2-Wrap pulse oximetry sensor, child (Pediatric), disposable	10 kg - 50 kg body weight	Finger/big toe
MCS2-Wrap pulse oximetry sensor, infant (Infant), disposable	10 kg - 20 kg body weight	

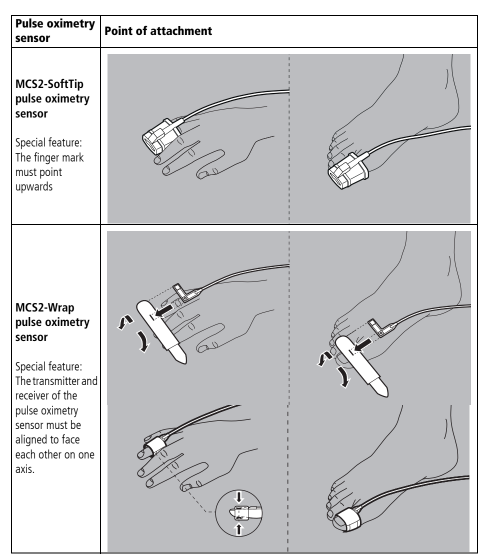


3. Connect the selected pulse oximetry sensor to the pulse oximetry sensor connecting cable.



4. Press the safety closure until you hear it engage.

5. Attach the pulse oximetry sensor:



Pulse oximetry sensor	Point of attachment
MCS2-Earclip pulse oximetry sensor	S
MCS2-Wrap pulse oximetry sensor, disposable Special feature: The transmitter and	
receiver of the disposable pulse oximetry sensor must be aligned to face each other on one axis.	

When doing so, please note:

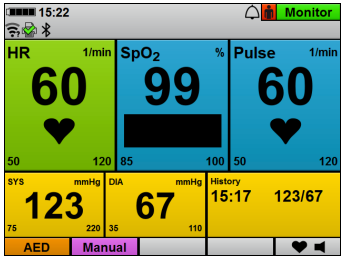
- The site must have a good blood supply.
- When attaching to the finger, use the ring finger or middle finger on the non-dominant hand.
- The pulse oximetry sensor must not be attached too tightly.
- The pulse oximetry sensor must be checked every 4 hours and repositioned if necessary.
- The pulse oximetry sensor must be repositioned if there are any skin changes.

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- 6. Check whether the oxygen saturation values displayed on the device are plausible.
- *Result* A pulse oximetry sensor is connected.

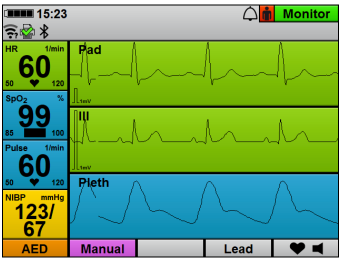
6.9.2 Carrying out pulse oximetry monitoring

- The device is switched on (see "6.1 Switching on the device", page 125).
 - A patient group is selected (see "6.4 Selecting patient group", page 127).
 - A pulse oximetry sensor is connected (see "6.9.1 Preparing pulse oximetry monitoring", page 164).
 - 1. If the patient group is incorrect: Select another patient group (see "6.4 Selecting patient group", page 127).
 - 2. If necessary: Select monitor mode using the **Monitor** function button.
 - 3. If necessary: Switch between parameter view and curve view with the view button (2).



4. In parameter view: Read off the measured values for arterial oxygen saturation (**SpO₂**) and pulse rate (**Pulse**).

Alternatively:



In curve view: Evaluate the SpO₂ curve (**Pleth**) and read off the measured values for arterial oxygen saturation (**SpO₂**) and pulse rate (**Pulse**).

- If necessary: Make the following SpO₂ settings in the user menu (see "10.3.4 SpO₂ settings", page 269):
 - Adapt the feed rate of the plethysmogram.
 - Give audio pulse tone output priority over heart rate tone output so that the tone level is dependent on oxygen saturation even when the ECG cable or defibrillation electrodes are connected to the patient.
- 6. If necessary: Set alarm limits in the user menu (see "10.3.1 Alarm settings", page 263).

Alternatively:

Set automatic alarm limits in the application menu (see "9 Application menu", page 256).

- 7. If necessary: Deactivate the pulse tone with the heart rate tone/ pulse tone function button ♥ ■.
- If there are artifacts in the SpO₂ curve or if signal quality is poor (bar in the SpO₂ parameter field): Reposition the pulse oximetry sensor on the patient's limb.
- *Result* Pulse oximetry monitoring is carried out.

6.10 6-lead ECG monitoring

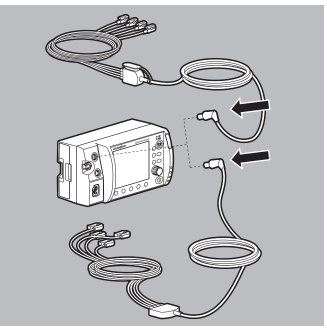
6.10.1 Preparing 6-lead ECG monitoring

- The device is switched on (see "6.1 Switching on the device", page 125).
 - A patient group is selected (see "6.4 Selecting patient group", page 127).

Risk of injury due to live connection for 12-lead ECG extension cable!

The connection for the 12-lead ECG extension cable may be live and lead to an electric shock if contacted. This may injure the user.

 \Rightarrow Always seal the 12-lead ECG extension cable connection with the protective cap if no 12-lead ECG extension cable is connected.



1. Connect the ECG connector of the ECG cable to the ECG connection for ECG cable on the device.

Alternatively:

Connect the ECG connector of the ECG cable to the ECG connection for ECG cable on the device using the connection for the 12-lead ECG extension cable.

2. Bare the patient's torso.

Risk of injury due to air/moisture between ECG electrodes and the patient's skin!

Air (e.g. in the case of hirsute patients) or moisture between the ECG electrodes and the patient's skin impair the quality of the ECG signal and falsify measurement results. This may injure the patient.

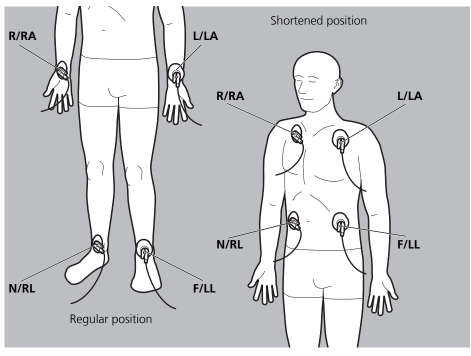
- \Rightarrow Remove hair from hirsute patients.
- \Rightarrow Rub the patient's skin dry.
- \Rightarrow Wipe down oily skin with an alcohol pad.
- 3. Remove hair from the torso.
- 4. Rub damp spots on the torso dry.
- 5. Wipe down oily skin with an alcohol pad.
- 6. Remove the protective film from the ECG electrodes.

A WARNING

Risk of injury from incorrect positioning of the ECG electrodes!

Incorrectly positioned ECG electrodes impair the quality of the ECG signal and falsify measurement results.

- \Rightarrow Select the electrode position according to the illustration.
- \Rightarrow Position ECG electrodes so that defibrillation/cardioversion is possible.
- \Rightarrow Maintain distance from the defibrillation electrodes.
- \Rightarrow Maintain distance from other ECG electrodes.
- \Rightarrow Do not position ECG electrodes on tendons or muscle groups.
- \Rightarrow Do not route individual lines of the ECG cable via ECG electrodes or other lines.



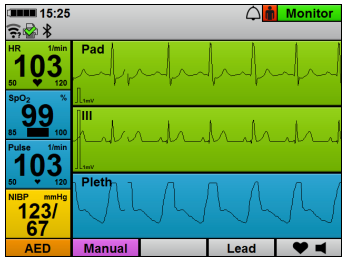
7. Attach and firmly press on the ECG electrodes (limb electrodes) as follows:

Code 1/ERC (Europe)		Code 2/AHA (USA)			
Electrode marking	Color coding	Electrode marking	Color coding	Application site	
Limb electrod	es	•			
R	Red	RA	White	Right arm, shortened: Below right collarbone	
L	Yellow	LA	Black	Left arm, shortened: Below left collarbone	
F	Green	LL	Red	Left leg, shortened: Left groin crease, centrally to leg axis	
N	Black	RL	Green	Right leg, shortened: Right groin crease, centrally to leg axis	

- If ECG electrodes are used at the same time as defibrillation electrodes: Do not allow ECG electrodes and defibrillation electrodes to overlap.
- 9. If necessary: Stroke out any air trapped under the ECG electrodes.
- 10. Clip the ECG cable to the individual ECG electrodes.
- 11. Check whether the ECG curves for ECG recording displayed on the device are plausible.
- *Result* The 6-lead ECG cable and the ECG electrodes are connected. 6-lead ECG monitoring is prepared.

6.10.2 Carrying out 6-lead ECG monitoring

- Requirement
 The device is switched on (see "6.1 Switching on the device", page 125).
 - A patient group is selected (see "6.4 Selecting patient group", page 127).
 - 6-lead ECG monitoring is prepared (see "6.10.1 Preparing 6-lead ECG monitoring", page 171).
 - 1. If the patient group is incorrect: Select another patient group (see "6.4 Selecting patient group", page 127).
 - 2. If necessary: Select monitor mode using the **Monitor** function button.
 - 3. If necessary: Switch between parameter view and curve view with the view button (2).
 - 4. Following a shock delivery: Wait until the ECG has stabilized again.



- 5. Evaluate the ECG leads and heart rate.
- 6. If necessary: Select another ECG lead using the **Lead** function button.
- 7. If necessary: Make the following ECG settings in the user menu (see "10.3.2 ECG settings", page 267):
 - Adapt amplitude scaling in order to adapt the displayed height of the ECG curve to the ECG measuring signal.
 - Set **Autom.** for amplitude scaling to have the displayed height of the ECG curve adapted to the ECG measuring signal automatically.
 - Adapt the feed rate of the ECG curve.
 - Activate the filter to filter interference caused by the power supply network out of the ECG display.
- 8. If necessary: Set alarm limits in the user menu (see "10.3.1 Alarm settings", page 263).

Alternatively:

Set automatic alarm limits in the application menu (see "9 Application menu", page 256).

- 9. If necessary: Switch off heart rate tone/pulse tone with the
 ♥ function button.
 The symbol ♥ ¥ appears.
- 10. If desired: Print a live printout of a 6-lead ECG (see "6.16.2 Printing a live printout of ECG and measured values", page 205).
- *Result* 6-lead ECG monitoring is carried out.

6.11 12-lead ECG recording and assessment (only with 12-lead ECG option)

A WARNING

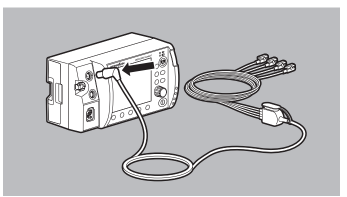
Risk of injury due to incorrect use of 12-lead ECG function mode!

Use of 12-lead ECG function mode by users without medical training and instruction in 12-lead ECG recording and assessment may injure the patient.

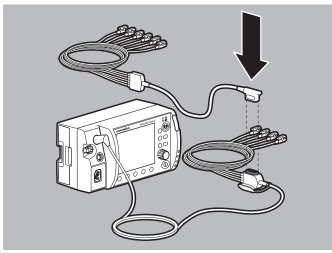
- ⇒ Only use 12-lead ECG function mode if the user is medically trained and has received instruction in 12-lead ECG recording and assessment.
- \Rightarrow Only use 12-lead ECG function mode if the user is familiar with the 12-lead ECG function mode of the device.

6.11.1 Preparing 12-lead ECG recording and assessment

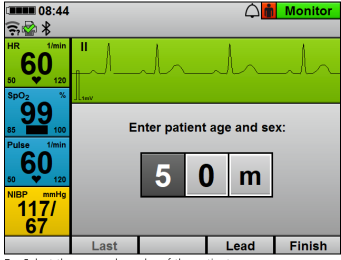
- The device is switched on (see "6.1 Switching on the device", page 125).
 - A patient group is selected (see "6.4 Selecting patient group", page 127).



1. Connect the ECG connector of the ECG cable to the ECG connection for ECG cable on the device using the connection for the 12-lead ECG extension cable.



- Connect the connector of the 12-lead ECG extension cable to the connector of the ECG cable with a connection for the 12lead ECG extension cable.
- 3. Open the application menu using the navigation knob.
- 4. Select the **12-lead ECG** menu item.



- 5. Select the age and gender of the patient.
- 6. Bare the patient's torso.

Risk of injury due to air/moisture between ECG electrodes and the patient's skin!

Air (e.g. in the case of hirsute patients) or moisture between the ECG electrodes and the patient's skin impair the quality of the ECG signal and falsify measurement results. This may injure the patient.

- \Rightarrow Remove hair from hirsute patients.
- \Rightarrow Rub the patient's skin dry.
- \Rightarrow Wipe down oily skin with an alcohol pad.
- 7. Remove hair from the torso.
- 8. Rub damp spots on the torso dry.
- 9. Wipe down oily skin with an alcohol pad.

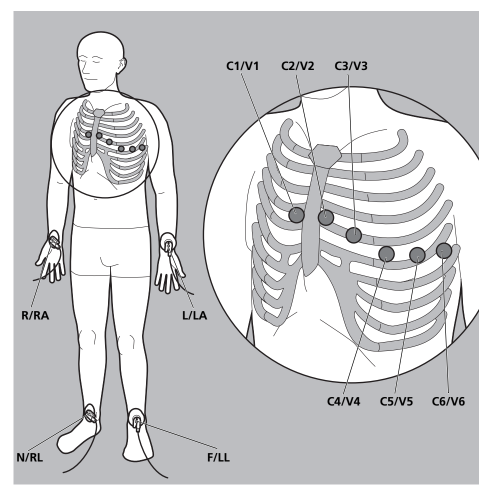
10. Remove the protective film from the ECG electrodes.

|--|--|

Risk of injury from incorrect positioning of the ECG electrodes!

Incorrectly positioned ECG electrodes impair the quality of the ECG signal and falsify measurement results.

- \Rightarrow Select the electrode position according to the illustration.
- \Rightarrow Position ECG electrodes so that defibrillation/cardioversion is possible.
- \Rightarrow Maintain distance from the defibrillation electrodes.
- \Rightarrow Maintain distance from other ECG electrodes.
- \Rightarrow Do not position ECG electrodes on tendons or muscle groups.
- \Rightarrow Do not route individual lines of the ECG cable via ECG electrodes or other lines.
- \Rightarrow For female patients: Always position chest wall electrodes V3/ C3 to V6/C6 underneath the breast.



11. Attach and firmly press on the ECG electrodes as follows:

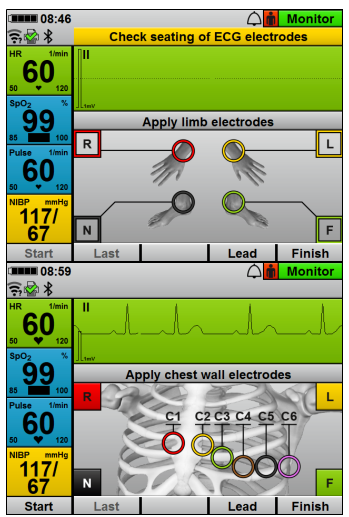
Code 1/ERC (Europe)		Code 2/AHA (USA)				
Electrode marking	Color coding	Electrode marking	Color coding	Application site		
Limb electroc	Limb electrodes					
R	Red	RA	White	Right arm		
L	Yellow	LA	Black	Left arm		
F	Green	LL	Red	Left leg		
Ν	Black	RL	Green	Right leg		

6 Operation

Code 1/ERC (Europe)		Code 2/AHA (USA)		
Electrode marking	Color coding	Electrode marking	Color coding	Application site
Chest wall ele	ectrodes	-		•
C1	Red	V1	Red	Fourth intercostal space on the right-hand edge of the breastbone
C2	Yellow	V2	Yellow	Fourth intercostal space on the left-hand edge of the breastbone
C3	Green	V3	Green	Centrally between V2/ C2 and V4/C4
C4	Brown	V4	Blue	Fifth intercostal space on the medioclavicular line
C5	Black	V5	Orange	Left ventral axillary line at the level of V4/C4
C6	Purple	V6	Purple	Left mid-axillary line at the same level as V4/C4 and V5/C5

12. If ECG electrodes are used at the same time as defibrillation electrodes: Do not allow ECG electrodes and defibrillation electrodes to overlap.

13. If necessary: Stroke out any air trapped under the ECG electrodes.



- 14. Clip the ECG cables to the individual ECG electrodes. Correctly connected ECG electrodes are indicated by a check mark. The limb electrodes are only displayed once all the limb electrodes have been attached. With chest wall electrodes, each chest wall electrode is displayed individually.
- 15. Check whether the ECG curves for 12-lead ECG recording displayed on the device are plausible.

Result The ECG electrodes and ECG cable for 12-lead ECG recording and assessment are connected. 12-lead ECG recording and assessment is prepared.

6.11.2 Carrying out and delivering 12-lead ECG recording and assessment

- Requirement
 The device is switched on (see "6.1 Switching on the device", page 125).
 - A patient group is selected (see "6.4 Selecting patient group", page 127).
 - 12-lead ECG recording and assessment is prepared (see "6.11.1 Preparing 12-lead ECG recording and assessment", page 178).
 - 1. Call up all ECG leads using the **Lead** function button:
 - Check whether all ECG leads are being displayed in a stable manner. If the ECG leads are not stable: Ensure the ECG electrodes are connected to the patient and ECG cable.
 - Check whether the sequence of ECG leads is plausible.
 - Following a shock delivery: Wait until the ECG has stabilized again.
 - 2. If a 12-lead ECG has already been recorded in the current session: Call up the last 12-lead ECG with the **Last** function button.

When doing so, please note:

- You can only call up the last 12-lead ECG of the current session.
- If no 12-lead ECG has yet been recorded in this session, the **Last** function button will not be available.
- To cancel 12-lead ECG function mode: Cancel 12-lead ECG function mode using the **Finish** function button.

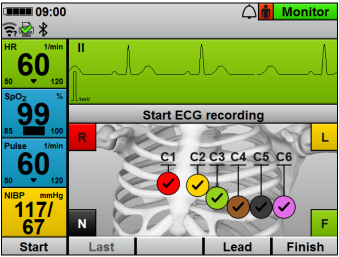
A CAUTION

Risk to therapy from movement artifacts when recording the 12-lead ECG!

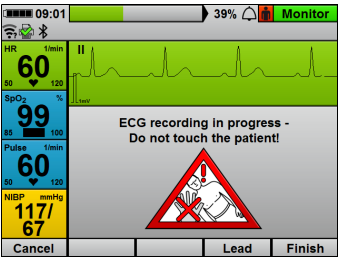
Movement artifacts falsify the 12-lead ECG. They may result in the user or the device interpreting the 12-lead ECG incorrectly, delaying treatment.

During recording of the 12-lead ECG:

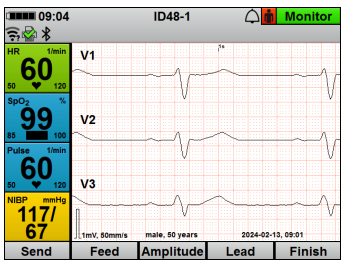
- \Rightarrow Ensure that the patient does not move.
- \Rightarrow Stand clear of the patient.



4. Start 12-lead ECG recording using the **Start** function button. The device starts recording the 12-lead ECG.



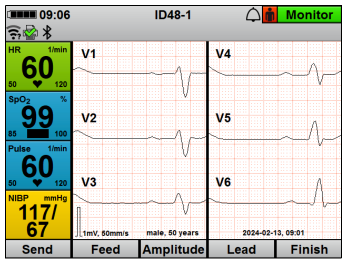
 Wait until the 12-lead ECG recording is finished. When doing so, please note: The patient must not be moved.



6. If the 12-lead ECG has been recorded: Assess the ECG leads.

 If the low-pass filter is incorrect: Set the low-pass filter in the user menu (see "10.3.3 12-lead ECG settings (only with 12lead ECG option)", page 268). WM 68401b 2024-03

 If a different view is required: Set the view in the user menu (see "10.3.3 12-lead ECG settings (only with 12-lead ECG option)", page 268).



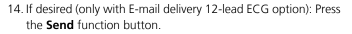
- Switch between the individual views using the view button
 :
 - Display 1 curve
 - Display 3 curves simultaneously
 - Display 6 curves simultaneously
- 10. Select another ECG lead using the **Lead** function button.
- 11. Displace ECG leads using the navigation knob.
- 12. Use the **Feed** function button to change the feed rate of the ECG curve.
- 13. Use the **Amplitude** function button to adapt the amplitude scaling to adapt the displayed height of the ECG curve to the ECG measuring signal.

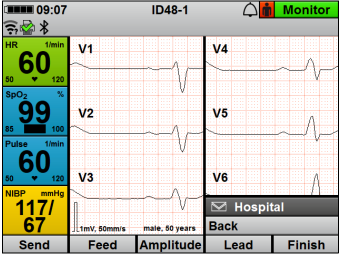
A WARNING

Delayed therapy due to severely delayed reception of the 12-lead ECG!

If the infrastructure of the monitoring station/hospital is not configured by the operator to receive 12-lead ECGs, receipt of the 12-lead ECG may be severely delayed. This may lead to delayed therapy and injure the patient.

- ⇒ Configure e-mail delivery 12-lead ECG correctly and check it (see "12.6 Configuring e-mail delivery 12-lead ECG (only with E-mail delivery 12-lead ECG option)", page 311).
- ⇒ If transmission time is unexpectedly high: Contact the system administrator of the e-mail recipient.





15. Select the recipient's name and send the 12-lead ECG to a predefined recipient by e-mail (see "12.6 Configuring e-mail delivery 12-lead ECG (only with E-mail delivery 12-lead ECG option)", page 311).

Following successful delivery, the symbol 🖄 appears in the display. If delivery is unsuccessful, the symbol 递 appears in the display and an alarm is displayed (see "11.2 Alarm messages", page 280).

- If desired (only with Printing option): Print 12-lead ECG (see "6.16.3 Printing 12-lead ECG (only with 12-lead ECG option)", page 206).
- 17. Exit 12-lead ECG function mode using the **Finish** function button.
- *Result* A 12-lead ECG has been recorded and assessed.

6.12 Non-invasive blood pressure measurement (NIBP measurement)

NIBP measurement technology has been optimized for measuring blood pressure with a normal sinus rhythm. Certain conditions may impair the ability of the non-invasive blood pressure measurement module to record correct measured values.

Risk of injury from falsified measured values during noninvasive blood pressure measurement!

Cardiac rhythm disorders, arteriosclerosis, reduced perfusion, diabetes, pregnancy, pre-eclampsia, arrhythmias (in newborns up to 28 days), kidney problems, shaking, shivering or the use of a cardiac pacemaker may impair the ability of the non-invasive blood pressure measuring module to record correct measured values. Evidence of the safety and efficacy of non-invasive blood pressure measurement has not been obtained for patient groups with these characteristics. This may injure the patient.

- ⇒ Include the status of these patient groups when evaluating the measured values of non-invasive blood pressure measurement.
- ⇒ Only use non-invasive blood pressure measurement on patient groups for whom evidence of the safety and efficacy of non-invasive blood pressure measurement has been provided.

6.12.1 Preparing non-invasive blood pressure measurement (NIBP measurement)

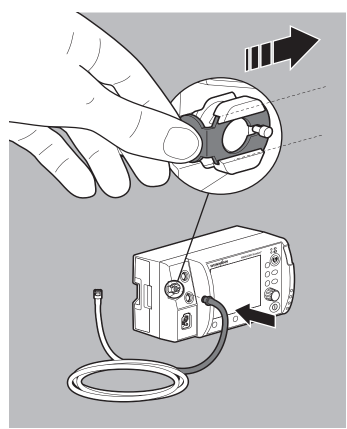
The following section describes how to attach the NIBP cuff. The instructions for use provided by the NIBP cuff manufacturer are binding for attachment. Follow these instructions for use.

A CAUTION

Risk of injury if NIBP cuff selected or put on incorrectly!

An NIBP cuff selected or put on incorrectly may interrupt the blood supply or falsify measurement results due to excessive contact pressure. This may result in injury to the patient.

- ⇒ Always use the NIBP cuff best suited to the patient's limb. Selecting the right NIBP cuff is the key to ensuring a goodquality measured value.
- \Rightarrow Attach the NIBP cuff level with the heart.
- \Rightarrow Attach the NIBP cuff so that the blood supply is not stopped.
- \Rightarrow Avoid moving the NIBP cuff during NIBP measurement.
- \Rightarrow When NIBP measurement is for an extended period: Check the position of the NIBP cuff regularly and, if necessary, reposition the NIBP cuff.
- ⇒ Repeat the NIBP measurement if measurement results are implausible. If the repeat measurement is still implausible, select an alternative method.
- \Rightarrow Do not bend or crush the NIBP cuff tube or the NIBP connecting tube.
- \Rightarrow Do not attach the NIBP cuff to a limb with poor circulation.
- \Rightarrow Do not attach the NIBP cuff to a limb with an intravenous infusion.
- \Rightarrow Do not attach the NIBP cuff to a limb on which there is already a pulse oximetry sensor or another monitoring device.
- \Rightarrow Do not attach the NIBP cuff to a limb with a shunt.
- \Rightarrow Do not attach the NIBP cuff to a limb with open wounds or burns.
- ⇒ In the case of patients who have undergone a mastectomy, do not attach the NIBP cuff to the affected side. In the case of patients who have undergone double mastectomies, attach the NIBP cuff to the non-dominant arm.
- \Rightarrow Only use an undamaged NIBP cuff.
- \Rightarrow Only use the NIBP cuffs and NIBP connecting tubes quoted in the scope of supply and in the accessories.
- \Rightarrow Follow the instructions for use of the NIBP cuff.

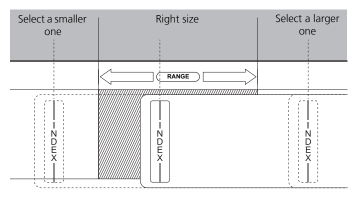


- 1. If the NIBP connection for NIBP connecting tube is latched: Push the latch of the NIBP connection to the right to release the NIBP connecting tube from the NIBP connection.
- 2. Connect NIBP connecting tube to the NIBP connection for NIBP connecting tube.

3. Select the NIBP cuff which is suitable for the patient's limb as per the following table:

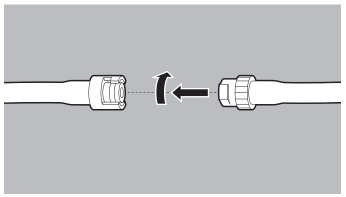
Designation	Color	Limb circumference
Thigh		·
Thigh	Brown	38 cm-50 cm
Upper arm		
Large Adult plus	Dark red	40 cm-55 cm
Adult plus	Dark blue	28 cm-40 cm
Adult	Dark blue	23 cm-33 cm
Small Adult	Turquoise	17 cm-25 cm
Child	Green	12 cm-19 cm
Infant	Orange	8 cm-13 cm

When doing so, please note:

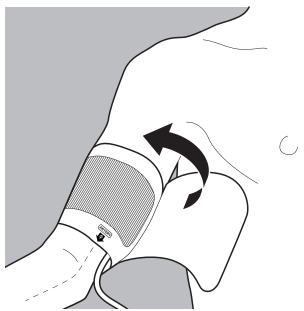


- The index printed on the NIBP cuff must be within the printed range (Range).
- If the index marking does not extend into the printed range (Range): Select a larger NIBP cuff.
- If the index marking extends beyond the printed range (Range): Select a smaller NIBP cuff.
- For disposable NIBP cuffs for newborns: Use adapter tube (WM 45467).

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- 4. Connect the NIBP connecting tube to the tube of the NIBP cuff.
- 5. Turn the two tubes against each other until they lock into place.



6. Attach the empty NIBP cuff to fit snugly around the patient's limb.

When doing so, please note:

• The skin below the NIBP cuff must be undamaged.

- The NIBP cuff must fit snugly around the limb.
- When applying to the arm: The artery marking (ARTERY) of the NIBP cuff must be positioned over the brachial artery and point toward the hand.
- When applying to the leg: The artery marking (ARTERY) of the NIBP cuff must be positioned over the femoral artery and point toward the foot.
- If the NIBP cuff is attached to the arm: The NIBP cuff must be positioned level with the heart.
- *Result* An NIBP cuff suitable for the patient is attached.

6.12.2 Carrying out non-invasive blood pressure measurement (NIBP measurement)

With an individual NIBP measurement, the device inflates the NIBP cuff to the set pressure (initial NIBP cuff pressure). The user can adapt the initial NIBP cuff pressure (**Initial** function button). To determine the patient's systolic and diastolic blood pressure, the air is slowly released from the NIBP cuff whilst measuring the pressure of the pulse wave. The values for diastolic and systolic blood pressure are determined from this and shown in the display. At the end of the NIBP measurement, the device releases the remaining air from the NIBP cuff.

The NIBP measurement can be influenced by various factors:

- Application site of the NIBP cuff
- Patient position (ideal position: Sitting comfortably, legs not crossed, feet flat on the floor, back and arm supported, center of the NIBP cuff level with the right heart atrium)
- Exertion (recommendation: Patient should rest for 5 minutes before the measurement, keep still during the measurement and not speak)
- Physiological condition

The device is switched on (see "6.1 Switching on the device", page 125).

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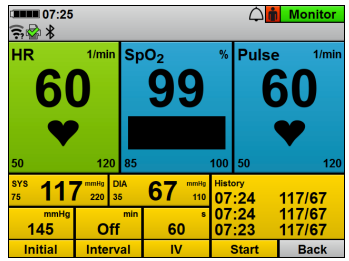
- A patient group is selected (see "6.4 Selecting patient group", page 127).
- A non-invasive blood pressure measurement (NIBP measurement) is prepared (see "6.12.1 Preparing non-invasive blood pressure measurement (NIBP measurement)", page 189).
- Press the NIBP button for < 2 s. The device switches to NIBP function mode.



Risk of injury due to incorrectly selected patient group!

The device only delivers correct measured values if the appropriate patient group is selected. An incorrect patient group may lead to incorrect measurements and injure the patient.

- \Rightarrow Adapt the patient group to the patient.
- \Rightarrow If the patient group is incorrect: Change the patient group in the application menu.
- If the patient group is incorrect: Select another patient group (see "6.4 Selecting patient group", page 127). The NIBP module is configured accordingly in the device with the selected patient group.



- Adapt the initial NIBP cuff pressure to the patient using the Initial function button and navigation knob.
 When doing so, please note: Following successful NIBP measurement, the initial NIBP cuff pressure adapts to the patient (approximately 30 mmHg above the systolic measured value of the previous NIBP measurement).
- 4. Press the **Start** function button.

Alternatively:

Press the NIBP button 😥 for > 2 s. An NIBP measurement starts. After the end of the measurement, the device displays systolic and diastolic arterial pressure.

- 5. Evaluate the NIBP measurement result.
- 6. If implausible measured values are shown:
 - Check whether the NIBP cuff is correctly selected and attached.
 - If necessary: Repeat the NIBP measurement.
- 7. If necessary: Cancel the NIBP measurement with the **Stop** function button.

The device releases the pressure from the NIBP cuff.

8. If necessary: Set alarm limits in the user menu (see "10.3.1 Alarm settings", page 263).

Alternatively:

Set automatic alarm limits in the application menu (see "9 Application menu", page 256).

9. If necessary: Press the **Back** function button.

Alternatively:

Press the NIBP button for < 2 s. The device exits NIBP function mode and switches to the set mode.

- 10. Once the measurement is finished: Remove the NIBP cuff.
- *Result* A non-invasive blood pressure measurement (NIBP measurement) has been carried out.

6.12.3 Carrying out interval measurement

During an interval measurement (**Interval** function button), the device carries out several successive NIBP measurements. The interval duration indicates the time between two successive NIBP measurements.

- Requirement
 The device is switched on (see "6.1 Switching on the device", page 125).
 - A patient group is selected (see "6.4 Selecting patient group", page 127).
 - A non-invasive blood pressure measurement (NIBP measurement) is prepared (see "6.12.1 Preparing non-invasive blood pressure measurement (NIBP measurement)", page 189).
 - Press the NIBP button for < 2 s. The device switches to NIBP function mode.

A CAUTION

Risk of injury due to incorrectly selected patient group!

The device only delivers correct measured values if the appropriate patient group is selected. An incorrect patient group may lead to incorrect measurements and injure the patient.

 \Rightarrow Adapt the patient group to the patient.

- \Rightarrow If the patient group is incorrect: Change the patient group in the application menu.
- If the patient group is incorrect: Select another patient group (see "6.4 Selecting patient group", page 127). The NIBP module is configured accordingly in the device with the selected patient group.
- 3. Press the Interval function button.
- 4. Set the interval duration using the navigation knob.
- 5. Press the **Start** function button.

Alternatively:

Press the NIBP button \bigotimes for > 2 s. An NIBP measurement starts. After the end of the measurement, the device displays the systolic and diastolic arterial blood pressure, the timer counts down, and when it has finished the next measurement starts automatically.

- 6. Evaluate the NIBP measurement result.
- If necessary: Cancel the NIBP measurement with the Stop function button.
 The device releases the pressure from the NIBP cuff.
- 8. If necessary: Set alarm limits in the user menu (see "10.3.1 Alarm settings", page 263).

Alternatively:

Set automatic alarm limits in the application menu (see "9 Application menu", page 256).

9. If necessary: Press the **Back** function button.

Alternatively:

Press the NIBP button for < 2 s. The device exits NIBP function mode and switches to the set mode.

- 10. Once the measurement is finished: Remove the NIBP cuff.
- *Result* An interval measurement has been carried out.

6.12.4 Carrying out venous stasis

With venous stasis (**IV** function button), the device inflates the NIBP cuff and maintains this pressure for the time preset in the operator menu (**Operator menu | NIBP settings | Venous stasis duration**). The venous blood return flow is impeded and the user can puncture one of the patient's veins. Venous stasis can only be carried out on the **Adult** patient group.

If you are the operator of the device and have access to the operator menu, you can disable the **Venous stasis** function: **Operator menu | System settings | Disable functions | Venous stasis** (see "13.12 System settings", page 358).

- The device is switched on (see "6.1 Switching on the device", page 125).
 - The **Adult** patient group has been selected.
 - A mode is set.
 - A non-invasive blood pressure measurement (NIBP measurement) is prepared (see "6.12.1 Preparing non-invasive blood pressure measurement (NIBP measurement)", page 189).
 - Press the NIBP button for < 2 s. The device switches to NIBP function mode.
 - Press the IV function button. The NIBP cuff is inflated to the pressure set in the operator menu. The timer displaying the duration of venous stasis counts down. Throughout venous stasis duration, pressure in the NIBP cuff is maintained.

- 3. Create the intravenous access.
- 4. Once the access has been created: Release the pressure from the NIBP cuff using the **Stop** function button.
- 5. Once venous stasis is finished: Remove the NIBP cuff.
- *Result* Venous stasis has been carried out.

6.13 Using audio alarm output

6.13.1 Canceling audio alarm output

Requirement An alarm is active and is audible.

- 1. Briefly (< 2 s) press the alarm button (
- *Result* Audio alarm output is canceled for this alarm. The symbol appears in the display and no audio signal is outputted for this alarm.

6.13.2 Pausing/muting audio alarm output

- 1. Press and hold the alarm button \bigotimes (> 2 s).
- Result Audio alarm output pauses for the time set in the operator menu (**Operator menu | Alarm settings | Pause audio**). The symbol \bigotimes appears in the display. If you set the time in the operator menu to ∞ (infinite), audio alarm output is permanently paused (audio alarm output is muted). The symbol \bigotimes appears in the display.

A reminder signal can remind you at certain intervals that audio alarm output is paused or muted. You can set the reminder signal in the operator menu (**Operator menu | Alarm settings | Reminder signal**).

6.13.3 Canceling muting or pausing of audio alarm output

- *Requirement* An alarm is active and is muted or paused.
 - 1. Briefly (< 2 s) press the alarm button 🔊

Alternatively:

Switch mode.

Result Muting or pausing of audio alarm output is canceled.

6.14 Changing the volume of the device

- The device is switched on (see "6.1 Switching on the device", page 125).
 - A patient group is selected (see "6.4 Selecting patient group", page 127).
 - 1. Open the application menu using the navigation knob.
 - 2. Select the **Volume** menu item.
 - 3. Select volume and confirm with the navigation knob.
 - *Result* The volume of the device has been changed.

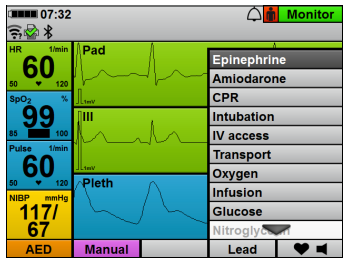
6.15 Saving event manually in the session data set

Measured values and user actions carried out on the device are saved in the internal memory and on the SD card.

With the event button (s), events which cannot be recorded by the device automatically (e.g. intubation, administration of medication, etc.) can be saved in the data set in order retrospectively to assign them chronologically during evaluation.

6 Operation

- The device is switched on (see "6.1 Switching on the device", page 125).
 - A patient group is selected (see "6.4 Selecting patient group", page 127).
 - Press the event button (\$).
 The events list opens.



2. Select and confirm an event from the events list using the navigation knob.



If you are the operator of the device and have access to the operator menu, you can adapt the events list to suit your requirements:

Operator menu | System settings | Events list (see "13.12 System settings", page 358)

Result The device saves the event selected in the events list in the session data set.

6.16 Printing ECGs and reports (only with Printing option)

Delayed therapy due to incorrect storage of the printer!

Storage outside the specified ambient conditions may lead to printer malfunctions or damage the printer. This may delay the patient's therapy.

- \Rightarrow Always store the printer within the specified ambient conditions
- \Rightarrow In the event of problems with the printer: Carry out a function check of the printer (see "5.6 Carrying out a function check on the printer (only with Printing option)", page 124).

6.16.1 Connecting the printer to the device

- Requirement The printer battery is inserted (see "4.7.2 Inserting the printer battery", page 97).
 - The printer paper is inserted (see "4.7.3 Inserting printer paper", page 98).
 - The printer is stowed in the printer case (see "4.7.4 Stowing") the printer in the printer case", page 100).
 - The printer has been paired with the device (see "12.7 Pair ٠ printer with device (only with Printing option)", page 314).

A WARNING

Risk of injury from defective printer!

A defective printer may lead to electric shock if touched. This may injure the patient or the user.

 \Rightarrow Always operate the printer in the printer case.

Delayed therapy due to printer not being ready for use!

The device cannot be connected to the printer if e-mail delivery 12-lead ECG or network data transmission are taking place in parallel.

- \Rightarrow Wait until the e-mail delivery 12-lead ECG or network data transmission is finished.
- \Rightarrow If necessary: Switch the printer on again.

EN

A CAUTION

Printer function restricted due to excessive distance between printer and device!

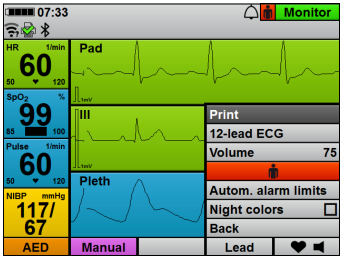
An excessive distance between the printer and the device may lead to special characters in the printout and confuse the user. \Rightarrow Reduce the distance between the printer and the device.

- Position the printer within range of the device. When doing so, please note: The printer must be positioned at least 20 cm away from the patient's body (excluding limbs).
- 2. Press and hold the On/Off button 0 of the printer for 3 s. The status indicator of the printer is green and an audible signal sounds.
- 3. Wait until the printer has connected to the device. When doing so, please note:
 - When connection is successful, the symbol 🕋 appears in the display.
 - The device outputs an audio signal when connected successfully.
 - The connection indicator appears in the printer display when connection is successful.
- 4. Leave the printer switched on. When doing so, please note:
 - The printer battery gets hot during operation.
 - Maximum print time may be reduced at low temperatures.
 - If the device is switched off, the connection to the printer is disconnected. The printer switches off automatically after 1 minute.
- 5. If the paper roll has finished: Insert new printer paper.
- 6. If the printer battery is empty: Insert a charged printer battery.
- 7. To disconnect the connection between the printer and the device: Press and hold the On/Off button of the printer for 3 s.
 The printer switches off.

Result The printer has been connected to the device.

6.16.2 Printing a live printout of ECG and measured values

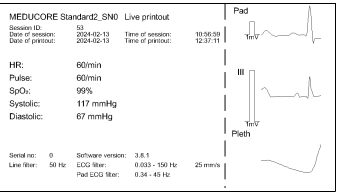
- *Requirement* A printer has been connected to the device (see "6.16.1 Connecting the printer to the device", page 203).
 - 1. If necessary: In the user menu, set the feed rate for the ECG to be printed out (see "10.3.5 Printer settings (only with Printing option)", page 270).



2. Open the application menu using the navigation knob.

3. Select the **Print** menu item.

The printer prints current measured values and the curves currently shown in the display on printer paper for 10 s (shown in abstract form here):



 To stop printing manually: Select the Stop print xx s menu item.



If you are the operator of the device and have access to the operator menu, you can adapt automatic stopping of printing to suit your requirements:

Operator menu | Printer settings | Automatic stop print (see "13.9 Printer settings (only with Printing option)", page 347)

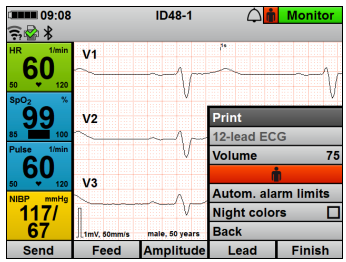
Result The measured values and curves currently shown in the display have been printed.

6.16.3 Printing 12-lead ECG (only with 12-lead ECG option)

- A printer has been connected to the device (see "6.16.1 Connecting the printer to the device", page 203).
 - A 12-lead ECG recording and assessment has been carried out (see "6.11.2 Carrying out and delivering 12-lead ECG recording and assessment", page 184).
 - 12-lead ECG function mode is activated.
 - 1. If necessary: Use the **Feed** function button to change the feed rate of the ECG curve.

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2. If necessary: Use the **Amplitude** function button to change the amplitude scaling in order to adapt the displayed height of the ECG curve to the ECG measuring signal.



3. Open the application menu using the navigation knob.

4. Select the **Print** menu item.

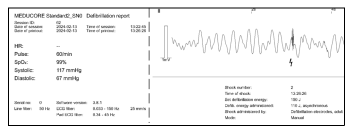
The printer prints the previously recorded measured values and all 12 ECG leads (shown in abstract form here):

MEDUCO Session ID: Date of session Date of recomposed of printed o	on: ding:	ndard2_SN0 48 2024-02-13 2024-02-13 2024-02-13	12-lead ECG ECG ID: Time of session: Time of recording: Time of printout:	ID48-1 08:44:10 09:01:26 09:57:26	1	~
HR: Pulse: SpO2: Systolic: Diastolic:		60/min 60/min 99% 117 mmHg 67 mmHg			II	~
				I	111	
Serial no: Line filter:	0 50 Hz	Software versior ECG filter:	n: 3.8.1 0.033 - 150 Hz			



6.16.4 Printing a defibrillation report

- A printer has been connected to the device (see "6.16.1 Connecting the printer to the device", page 203).
 - The Defibrillation report menu item is activated in the operator menu (see "13.9 Printer settings (only with Printing option)", page 347).
 - Carry out defibrillation or cardioversion. On every shock delivery, the printer prints a defibrillation report (3 s before and 5 s after the shock delivery, shown in abstract form here):

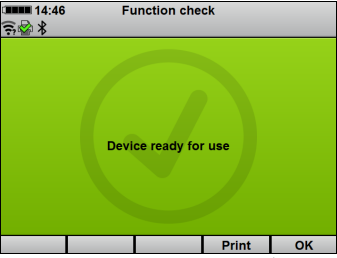


Result A defibrillation report has been printed.

6.16.5 Printing a function check report

Requirement A printer has been connected to the device (see "6.16.1 Connecting the printer to the device", page 203).

1. Carry out a function check (see "5 Function check", page 108).



2. When the status report appears: Select the **Print** function button.

The printer prints a report with the results of the function check (shown in abstract form here):

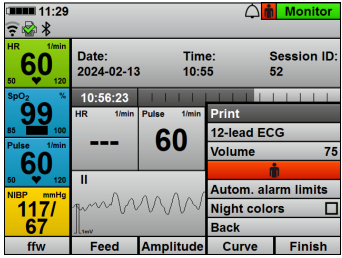
MEDUCORE Standard2 Date of printout: 2024-02-13	Function check report Time of printout: 14:52:01	
Alarm system test: Button test: ECG module:	OK OK OK	Last name:
NIBP module: Defibrillation module: Master cable: Defibrillation electrodes:	OK OK OK	Signature:
SpO₂ module: SpO₂ sensor: SD card:	OK OK OK	
Result: Device ready for use		
Serial no: 0 Software versi	on: 3.8.1	

Result

A function check report has been printed.

6.16.6 Printing replay view (only with Printing and Replay view options)

- A printer has been connected to the device (see "6.16.1 Connecting the printer to the device", page 203).
 - The Printing option is enabled and activated (see "13.12 System settings", page 358).
 - The Replay view option is enabled and activated (see "13.12 System settings", page 358).
 - Replay view shows one event in the session (see "6.17.2 Analyzing the session in replay view on the device (only with Replay view option)", page 214).



1. Open the application menu using the navigation knob.

2. Select the **Print** menu item.

The printer prints the replay view of the event on printer paper (10 s from the time of the event, shown in abstract form here):

MEDUCORE St Session ID: Date of session: Date of printout:	andard2_SN0 52 2024-02-13 2024-02-13	Replay printout Time of session: Time of printout:	10:56:23 11:51:15	
HR: Pulse: SpO ₂ : Systolic: Diastolic:	 60/min 99% 		 	
Serial no: 0 Line filter: 50 Hz	Software version ECG filter: Pad ECG filter:	n: 3.8.1 0.033 - 150 Hz 0.34 - 45 Hz	25 mm/s	

To stop printing manually: Select the Stop print xx s menu item.



If you are the operator of the device and have access to the operator menu, you can adapt automatic stopping of printing to suit your requirements:

Operator menu | Printer settings | Automatic stop print (see "13.9 Printer settings (only with Printing option)", page 347)

Result Replay view of an event (replay printout) has been printed.

6.16.7 Print a session report (only with Printing option)

- A printer has been connected to the device (see "6.16.1 Connecting the printer to the device", page 203).
 - The Printing option is enabled and activated (see "13.12 System settings", page 358).
 - The details included in the session report have been specified in the operator menu (see "13.9 Printer settings (only with Printing option)", page 347).
 - A session has been selected in the session archive (see "6.17.1 Selecting a session in the session archive", page 212).

 Select the **Print session report** menu item. The device prints the session report with all the details specified in the operator menu (abstract example shown here):



When doing so, please note:

- The device only prints a resuscitation report in the session report if a shock was delivered or a resuscitation mode (AED mode or manual mode) was activated for > 2 minutes in the selected session.
- In the resuscitation report, the first two events from the events list are recorded and counted (see "13.12.2 Possible values", page 360).
- 2. To cancel printing of the session report: Select the **Cancel printing** menu item.
- *Result* A session report has been printed.

6.17 Analyzing sessions

6.17.1 Selecting a session in the session archive

- The device is switched on (see "6.1 Switching on the device", page 125).
 - A patient group is selected (see "6.4 Selecting patient group", page 127).
 - A mode is set.
 - At least one session has been carried out with the device.
 - The user menu is activated (see "10.1 Navigating the user menu", page 260).

1. Select the **Session archive** menu item. The session archive appears:

· I1:21				Menu	
	Ses	sion archiv	е		
Date	Time	Duration	CPR	12-lead	
2024-02-13	10:56	25:04	No	No	
2024-02-13	10:55	01:06	Yes	Yes	
2024-02-13	10:50	05:19	No	No	
2024-02-13	10:48	01:19	Yes	Yes	
2024-02-13	10:39	09:45	No	No	
2024-02-13	08:44	104:43	No	Yes	
2024-02-13	08:43	00:40	No	Yes	
Load all sessions					
		Monitor	Back	ОК	
	· .				

2. Select the session by session time and the following criteria:

- Resuscitation (defibrillation or cardioversion) carried out during the session: **Yes** in the **CPR** column
- One or more 12-lead ECGs recorded during the session: **Yes** in the **12-lead** column
- 3. Confirm with the **OK** function button.

11:23		\triangle	Menu
Date: 2024	4-02-13 T	ime: 10:55	
Replay view			
Display 12-lead ECG			
Print session report			
Back			
	Monitor	Back	ОК

4. Analyze the session in replay view on the device (see "6.17.2 Analyzing the session in replay view on the device (only with Replay view option)", page 214).

Alternatively:

Analyze the 12-lead ECG of the session (see "6.17.3 Analyzing and delivering a 12-lead ECG of a session (only with 12-lead ECG option)", page 216).

Alternatively:

Print session report (see "6.16.7 Print a session report (only with Printing option)", page 211).

- 5. Confirm with the **OK** function button.
- *Result* A session has been selected for further analysis in the session archive.

6.17.2 Analyzing the session in replay view on the device (only with Replay view option)

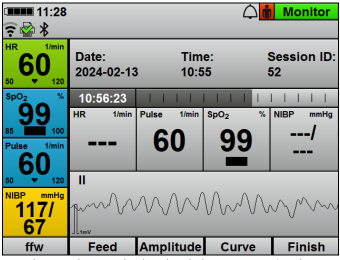
- The Replay view option is enabled and activated (see "13.12 System settings", page 358).
 - A session has been selected in the session archive (see "6.17.1 Selecting a session in the session archive", page 212).

 Select the **Replay view** menu item. The device displays a selection of events during a session.

IIII 11:24			Menu
??? 🖓 🖇			
	Date: 2024-02-13 Ti	me: 10:55	
10:55:45	Monitor mode activated		
10:55:57	12-lead ECG recording	started	
10:56:15			
10:56:23	△ VF/VT		
10:56:30	Manual mode activated		
10:56:33	Function charge button	pressed	
10:56:39	Shock administered (15	0 J)	
10:56:39	Defibrillation report prin	nted	
10:56:46	Monitor mode astated		
	Monitor	Back	ОК

2. Select event.

 Confirm with the **OK** function button. The device displays the event in replay view.



4. Select another ECG lead or the plethysmogram using the **Curve** function button.

- 5. Use the **Amplitude** function button to adapt the amplitude scaling to adapt the displayed height of the ECG curve to the ECG measuring signal.
- 6. Use the **Feed** function button to adapt the feed rate of the ECG curve.
- 7. Use the **ffw** function button to adapt the scrolling resolution.
- 8. Shift the session as from the time of the event using the navigation knob.
- 9. Analyze the event in the session.
- 10. If desired (only with Printing option): Print the replay view of the event (see "6.16.6 Printing replay view (only with Printing and Replay view options)", page 210).
- *Result* A session has been analyzed in replay view on the device.

6.17.3 Analyzing and delivering a 12-lead ECG of a session (only with 12-lead ECG option)

• The 12-lead ECG option is enabled and activated (see "13.12 System settings", page 358).

- If e-mail delivery 12-lead ECG is desired: A default network has been configured (see "12.3 Configuring default network", page 302).
- Select a session with at least one 12-lead ECG (Yes in the 12-lead column) in the session archive (see "6.17.1 Selecting a session in the session archive", page 212).
- 2. Confirm with the **OK** function button.
- Select the Display 12-lead ECG menu item. The device displays all 12-lead ECGs recorded during the selected session.

12:58			\triangle	Menu
(?•				
D	ate: 2024-0	2-13 T	ime: 12:54	
Date	Time	1	2-lead ECG	ID
2024-02-13	12:55		ID57-1	
2024-02-13	12:57		ID57-2	
		Monitor	Back	OK

 Select the desired 12-lead ECG. The device displays the desired 12-lead ECG.

- If desired (only with E-mail delivery 12-lead ECG option): Use the Send function button to deliver the 12-lead ECG to a predefined recipient (see "12.6 Configuring e-mail delivery 12-lead ECG (only with E-mail delivery 12-lead ECG option)", page 311).
- If desired (only with Printing option): Print 12-lead ECG (see "6.16.3 Printing 12-lead ECG (only with 12-lead ECG option)", page 206).
- *Result* A 12-lead ECG from a session has been analyzed on the device.

6.17.4 Analyzing a session using DEFIview

1. Export session data from the internal memory to SD card (see "6.21.1 Exporting session data from the internal memory to SD card", page 221).

Alternatively:

Upload session data to WEINMANN Connect (see "6.21.2 Uploading session data to WEINMANN Connect (only with Session data upload option)", page 221).

- 2. Analyze session data with the DEFlview PC software in accordance with the DEFlview instructions for use.
- *Result* The session data have been analyzed using the DEFlview PC software.

6.18 Reprocessing the device after use

- 1. Remove disposable articles from the patient and dispose of them:
 - Defibrillation electrodes
 - ECG electrodes
 - Disposable pulse oximetry sensor
 - Disposable NIBP cuff for newborns
- 2. Put new disposables with the device.
- Hygienically reprocess the device, components, and accessories (see "8 Hygienic reprocessing", page 238).
 When doing so, please note: It is recommended that hygienic reprocessing is carried out immediately after use, as dried-on residues are harder to remove.
- 4. Ensure that the device and printer batteries have a sufficiently good battery status.
- 5. Insert new printer paper and have spare printer paper to hand (see "4.7.3 Inserting printer paper", page 98).
- Stow components and accessories in the protective transport bag (see "4.2.1 Stowing components and accessories in the protective transport bag", page 76).

Alternatively:

Stow components and accessories on the portable unit (see "4.2.2 Stowing components and accessories on the portable unit", page 83).

- If required: Store the device, components, and accessories in accordance with the conditions for storage (see "15 Storage", page 372).
- *Result* The device is reprocessed following use.

6.19 Pairing the device with an external device via the Bluetooth[®] interface (only with Bluetooth[®] data transmission option)

- Requirement
 The Bluetooth[®] data transmission option is enabled and activated (see "13.12 System settings", page 358).
 - An external device with a Bluetooth[®] interface (Example: System for digital patient data recording) is within range.
 - 1. Activate the user menu (see "10.1 Navigating the user menu", page 260).
 - 2. Select **Communication settings | Pair Bluetooth[®] devices** . The device waits to pair with an external device.
 - 3. Activate the Bluetooth[®] interface of the external device and search for available devices.
 - 4. Select MEDUCORE Standard² on the external device.
 - 5. Check whether the PIN displayed on the device is shown on the external device.
 - If the PINs displayed on the device and on the external device are identical: Confirm pairing on device and external device. Device and external device are paired.
 - *Result* The device and the external device are paired via the Bluetooth[®] interface. The external device can connect to the device and interrogate device data via the Bluetooth[®] interface.

6.20 Uploading data to an external device via Bluetooth[®] interface (only with Bluetooth[®] data transmission option)

- The Bluetooth[®] data transmission option is enabled and activated (see "13.12 System settings", page 358).
 - The device is paired with the external device via the Bluetooth[®] interface (see "6.19 Pairing the device with an external device via the Bluetooth[®] interface (only with Bluetooth[®] data transmission option)", page 219).
 - Start Bluetooth[®] data transmission to an external device. When the Bluetooth[®] connection is set up, the symbol is displayed on the device. If the Bluetooth[®] connection is terminated by the external device, the symbol is displayed.

A device can only be connected to one external device via the Bluetooth[®] interface at a time.

- 2. To update data, restart Bluetooth[®] data transmission on the external device.
- *Result* The Bluetooth[®] connection between the device and the external device has been set up. Device data have been uploaded to the external device.

6.21 Archiving session data

From 60 seconds after the start of a session, the device permanently saves session data to its internal memory and on the SD card if the SD card is in the SD card slot at the beginning of the session. After a session has started, session data are saved only temporarily in the internal memory and not saved on the SD card and are deleted if you switch off the device in < 60 s.

If you switch off the device and then switch it on again after < 30 s, the device saves session data in the session data set already started.

If you switch off the device and then switch it on again after > 30 s, the device creates a new session data set.

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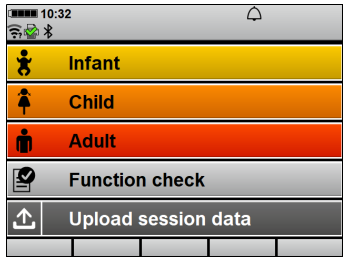
6.21.1 Exporting session data from the internal memory to SD card

Requirement There is an SD card in the SD card slot.

- 1. Activate the operator menu (see "13.1 Activating the operator menu", page 322).
- 2. Select System settings | Import/export configurations | Export internal memory to SD card.
- 3. Confirm with the **OK** function button. The export process starts.
- 4. Remove the SD card (see "4.6.2 Removing the SD card", page 93).
- *Result* Session data and service data have been exported from the internal memory to the SD card.

6.21.2 Uploading session data to WEINMANN Connect (only with Session data upload option)

- The Session data upload option is enabled and activated (see "13.12 System settings", page 358).
 - A default network has been configured (see "12.3 Configuring default network", page 302).
 - The device has been registered with WEINMANN Connect (see "12.4 Registering the device with WEINMANN Connect", page 308).
 - The device has been switched off for > 30 s.
 - 1. Switch on the device. The start menu appears.



- 2. Select the **Upload session data** menu item. All configured networks are displayed.
- 3. Select a network within range.
- Confirm with the **OK** function button. The device connects to the selected network and the session archive appears.

10:36			\triangle		
🛜 🗟 🖇					
	Manual s	ession data	upload		
Date	Time	Duration	CPR	12-lead	
2024-02-13	08:44	104:43	No	Yes	R
2024-02-13	08:43	00:40	No	Yes	$\mathbf{\nabla}$
2024-02-13	08:41	00:08	No	No	$\mathbf{\nabla}$
2024-02-13	07:21	11:58	No	No	区
2024-02-12	15:35	37:57	No	Yes	区
2024-02-12	15:31	01:36	No	No	区
2024-02-12	15:16	13:32	Yes	No	$\mathbf{\nabla}$
2024-02-12	15:09		No	No	
Unload	A11	None	Canaal		
Upload	All	None	Cancel		

- 5. Select individual sessions in the session archive using the navigation knob.
- 6. If desired: Deselect all sessions using the **None** function button.

Alternatively:

Select all sessions using the **All** function button.

- Upload the selected session data to WEINMANN Connect using the **Upload** function button.
 When upload is successful, the symbol support appears in the display. If upload is unsuccessful, the symbol appears in the display and an alarm is displayed (see "11.2 Alarm messages", page 280).
- 8. Confirm with the **OK** function button.
- To cancel the upload: Select the Cancel function button. When doing so, please note: If the upload is canceled, the Upload session data menu item is briefly grayed out in the start menu.
- *Result* The session data have been uploaded to WEINMANN Connect.

6.22 Using service data

Service data are device data which WEINMANN Emergency can use to analyze the device in the event of a fault. Service data do not contain any patient data. There are two options for making service data available to WEINMANN Emergency Technical Service in the event of a fault:

- Upload service data straight to WEINMANN Emergency (see "6.22.1 Uploading service data to WEINMANN Emergency", page 224)
- Export service data to SD card and e-mail them to Technical Service (see "6.22.2 Exporting service data to SD card", page 224)

6.22.1 Uploading service data to WEINMANN Emergency

- A default network has been configured (see "12.3 Configuring default network", page 302).
 - The device has been registered with WEINMANN Connect (see "12.4 Registering the device with WEINMANN Connect", page 308).
 - The user menu is activated (see "10.1 Navigating the user menu", page 260).

Alternatively:

The operator menu is activated (see "13.1 Activating the operator menu", page 322).

1. Select System settings | Service | Upload service data to WEINMANN.

- 2. Confirm with the **OK** function button.
- *Result* The service data have been uploaded to WEINMANN Emergency.

6.22.2 Exporting service data to SD card

- *Requirement* There is an SD card in the SD card slot.
 - The user menu is activated (see "10.1 Navigating the user menu", page 260).

Alternatively:

The operator menu is activated (see "13.1 Activating the operator menu", page 322).

- Select System settings | Service | Export service data to SD card.
- 2. Confirm with the **OK** function button. The export process starts.

- 3. If export is successful: Remove the SD card (see "4.6.2 Removing the SD card", page 93).
- 4. Insert the SD card in the SD card slot of a PC.
- 5. E-mail service data to WEINMANN Emergency Technical Service for further analysis.
- *Result* Service data have been exported to the SD card and made available to WEINMANN Emergency Technical Service.

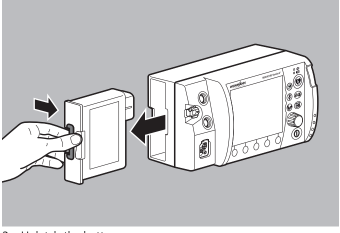
7 Disassembling

7.1 Disassembling the power supply

7.1.1 Removing battery

Requirement The device is switched off.

- 1. When used in a protective transport bag:
 - Open the left-hand side compartment of the protective transport bag.
 - Take components and accessories out of the side compartment.
 - Remove dividers from the side compartment.



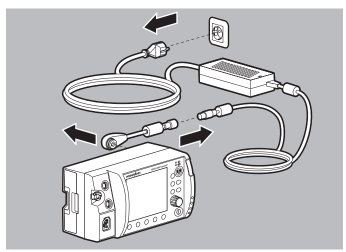
- 2. Unlatch the battery.
- 3. Remove the battery.

Result The battery has been removed.

7.1.2 Disconnecting the device from line power

Requirement The device is switched off.

1. When used in a protective transport bag: Open the rear of the device compartment of the protective transport bag.



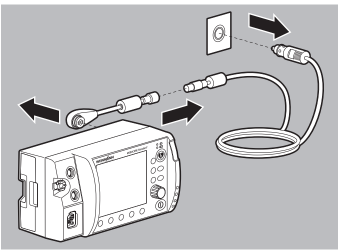
- 2. Disconnect the charging adapter from the power supply connection.
- 3. Disconnect the charging adapter from the power supply unit and charger.
- 4. Disconnect the power supply unit and charger from line power.

Result The device is disconnected from line power.

7.1.3 Disconnecting the device from the 12 V on-board power supply

Requirement The device is switched off.

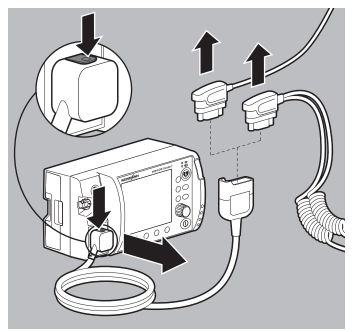
1. When used in a protective transport bag: Open the rear of the device compartment of the protective transport bag.



- 2. Disconnect the charging adapter from the power supply connection.
- 3. Disconnect the charging adapter from the adapter cable for 12 V on-board power supply/circular connector.
- 4. Disconnect the adapter cable for 12 V on-board power supply/ circular connector from the 12 V on-board power supply.
- *Result* The device is disconnected from the 12 V on-board power supply.

7.2 Disassembling defibrillation electrodes/paddles and master cable

- The device is switched off (see "6.2 Switching the device off", page 126).
 - The device is disconnected from the patient.
 - 1. When used in a protective transport bag: Open the viewing window and side compartments of the protective transport bag.



- 2. Unlatch the master cable and disconnect it from the Pad connection.
- 3. Disconnect the Pad connector of the defibrillation electrodes from the master cable.

Alternatively:

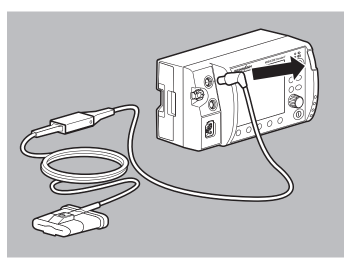
Disconnect the Pad connector of the paddles from the master cable.

- 4. Dispose of used defibrillation electrodes.
- 5. When used in a protective transport bag: Guide the master cable through the opening between the side compartment and the device compartment of the protective transport bag.
- *Result* The defibrillation electrodes/paddles and the master cable have been disassembled.

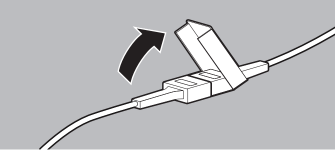
7.3 Disassembling pulse oximetry sensor and pulse oximetry sensor connecting cable

• The device is switched off (see "6.2 Switching the device off", page 126).

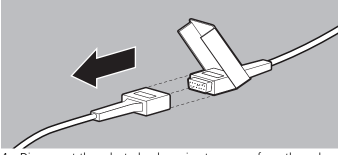
- The device is disconnected from the patient.
- 1. When used in a protective transport bag: Open the viewing window and side compartments of the protective transport bag.



 Disconnect the pulse oximetry sensor connecting cable with the selected pulse oximetry sensor from the SpO₂ connection. WM 68401b 2024-03



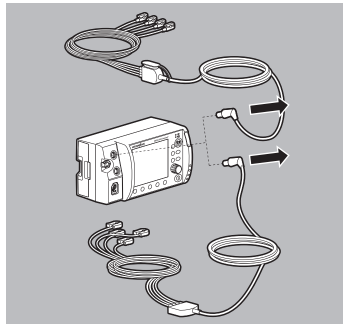
3. Open the safety closure.



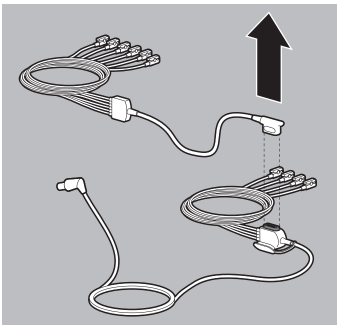
- 4. Disconnect the selected pulse oximetry sensor from the pulse oximetry sensor connecting cable.
- *Result* A pulse oximetry sensor and the pulse oximetry sensor connecting cable have been disassembled.

7.4 Disassembling ECG cable

- The device is switched off (see "6.2 Switching the device off", page 126).
 - The device is disconnected from the patient.



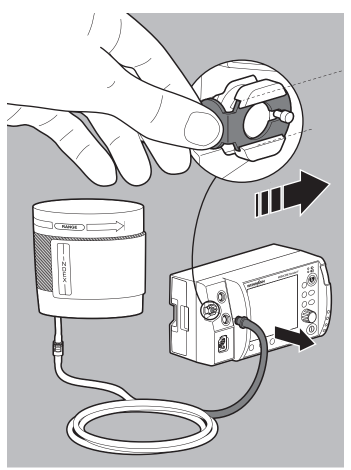
1. Disconnect the selected ECG cable from the ECG connection on the device.



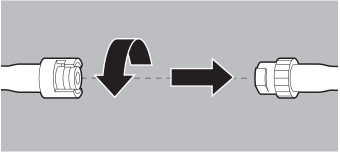
- 2. If the 12-lead ECG extension cable is connected: Disconnect the 12-lead ECG extension cable from the ECG cable with connection for 12-lead ECG extension cable.
- 3. Disconnect the ECG electrodes from the ECG cable clips and dispose of them.
- *Result* An ECG cable has been disassembled.

7.5 Disassembling the NIBP cuff and NIBP connecting tube

- The device is switched off (see "6.2 Switching the device off", page 126).
 - The device is disconnected from the patient.
 - 1. When used in a protective transport bag: Open the viewing window and side compartments of the protective transport bag.



 Disconnect the NIBP connecting tube with the selected NIBP cuff from the NIBP connection on the device. When doing so, please note: The latch of the NIBP connection must be pushed to the right to release the NIBP connecting tube from the NIBP connection.

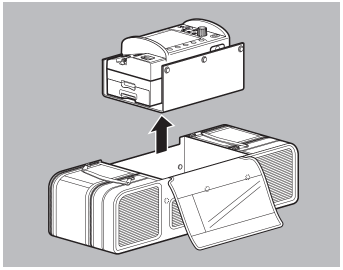


- 3. Open the safety closure.
- 4. Disconnect the NIBP cuff from the NIBP connecting tube.
- 5. If necessary: Pass all tubes through the respective openings in the protective transport bag.
- *Result* An NIBP cuff and the NIBP connecting tube have been disassembled.

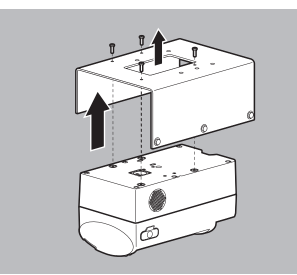
7.6 Disassembling the device from the protective transport bag and the holding plate

Required auxiliary equipment Phillips screwdriver, size PH1

- 1. Place the protective transport bag with its back on a level, firm surface.
- 2. Release the magnetic catch of the protective transport bag and open the viewing window.
- 3. Disconnect all cables and tubes from the device.
- 4. Open the snap fasteners between the holding plate and the protective transport bag.



5. Take the device on the holding plate up out of the protective transport bag.



- 6. Put the device on the holding plate on a smooth, firm surface with the control panel membrane facing down.
- 7. Undo the screws from the holding plate.
- 8. Remove the holding plate.

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Result The device has been disassembled from the protective transport bag and the holding plate.

7.7 Disassembling the device from the portable unit

Follow the instructions for use of the portable unit.

8 Hygienic reprocessing

The following sections set out the procedures necessary for hygienic reprocessing. Read this chapter in full before starting hygienic reprocessing. If you have any questions regarding hygienic reprocessing, contact the manufacturer WEINMANN Emergency, or a technician expressly authorized by it.

A WARNING

Risk of infection resulting from poor hygienic reprocessing! If the device, components or accessories are subjected to incorrect hygienic reprocessing, or to none at all, then in the case of skin contact infections may occur which may seriously or fatally injure the patient or the user. What is more, the device, components, and accessories may be damaged.

- \Rightarrow Subject the device, components, and accessories to hygienic reprocessing after every use.
- ⇒ Only reuse the device, components, and accessories if they have been subjected to hygienic reprocessing in accordance with the cleaning and disinfection plan.
- ⇒ Carry out hygienic reprocessing in accordance with the cleaning and disinfection plan (see "8.7 Cleaning and disinfection plan", page 248).
- ⇒ When reprocessing the device, components, and accessories, use only the recommended cleaning agents and disinfectants.
- ⇒ Follow the instructions for use of the cleaning agent and disinfectant being used.
- \Rightarrow Follow the instructions for use of the components and the accessories.
- \Rightarrow Wear suitable personal protective equipment.
- ⇒ Only have the device, components, and accessories reprocessed by those who have medical training and training in the care of emergency patients.
- ⇒ Only have the device, components, and accessories reprocessed by those who have been trained in hygienic reprocessing methods.

Disrupted and failed therapy due to incorrect use of A WARNING disposables! Reusing and reprocessing disposables might induce unpredictable reactions as a result of aging, embrittlement, wear, thermal stress, and chemical action. This may put the functionality and safety of the device at risk, and cause the patient and user serious or lifethreatening injury. \Rightarrow Do not reuse disposables. \Rightarrow Do not subject disposables to hygienic reprocessing. Loss of mechanical or electrical safety resulting from A WARNING reprocessing of the device and accessories with unsuitable cleaning agents and disinfectants! Using incorrect cleaning agents and disinfectants may cause damage to the surface of the device and accessories, as well as impairing electrical and insulating properties. This may cause the user and the patient serious or life-threatening injury. \Rightarrow Never clean the device, components, and accessories with bleach, bleach solution or compounds containing phenols. \Rightarrow Only use the cleaning agents and disinfectants recommended in these instructions for use and in the instructions for use of the components and accessories (see "8.7 Cleaning and disinfection plan", page 248). Risk of injury and material damage from residues of disinfectants or cleaning agents in the device, components, and accessories! Residues of disinfectants or cleaning agents or moisture may lead to short circuits in the device connections and thus impair device function. This may injure the user and the patient and lead to material damage. \Rightarrow After hygienic reprocessing, check the device, components, and accessories visually for any residues of cleaning agent or disinfectant and remove any residues as necessary. \Rightarrow Carry out a complete function check after every hygienic reprocessing operation. \Rightarrow Do not immerse the device, components, and accessories in

liquids.

MEDUCORE Standard² EN **239**

A WARNING

Failure of therapy due to disinfection by immersion, mechanical reprocessing and sterilization!

If the device, or components or accessories not designed for these processes are subjected to disinfection by immersion, mechanical reprocessing or sterilization, they may be damaged.

⇒ Never subject the device, or components or accessories not designed for these processes to disinfection by immersion, mechanical reprocessing or sterilization.

8.1 Intervals

Part	After every use*	At least 1x weekly
All parts	Х	Х

* In the event of visible dirt or suspected contamination.

8.2 Preparing hygienic reprocessing

• The device is switched off (see "6.2 Switching the device off", page 126).

- The device is disconnected from the patient.
- Battery and SD card are in the device.
- SD card cover is closed.
- If the device is connected to line power or to a 12 V on-board power supply: Disconnect the device from line power (see "7.1.2 Disconnecting the device from line power", page 227).

Alternatively:

Disconnect the device from the 12 V on-board power supply (see "7.1.2 Disconnecting the device from line power", page 227). When doing so, please note: The battery remains in the device for hygienic reprocessing.

2. Disassemble the defibrillation electrodes/paddles and master cable (see "7.2 Disassembling defibrillation electrodes/paddles and master cable", page 229).

- Disassemble the pulse oximetry sensor and pulse oximetry sensor connecting cable (see "7.3 Disassembling pulse oximetry sensor and pulse oximetry sensor connecting cable", page 230).
- 4. Disassemble the ECG cable (see "7.4 Disassembling ECG cable", page 232).
- Disassemble the NIBP cuff and NIBP connecting tube (see "7.5 Disassembling the NIBP cuff and NIBP connecting tube", page 233).
- 6. Remove the remaining accessories from the device.
- 7. In the event of contamination or suspected contamination of inaccessible areas of the device, protective transport bag or portable unit: Disassemble the device from the protective transport bag and holding plate (see "7.6 Disassembling the device from the protective transport bag and the holding plate", page 235).

Alternatively:

Disassemble the device from the portable unit (see "7.7 Disassembling the device from the portable unit", page 237). When doing so, please note: Disassembly is described in the instructions for use of the portable unit.

- Dispose of all disposables properly (see "16 Disposal", page 375).
- *Result* All parts have been prepared for hygienic reprocessing.

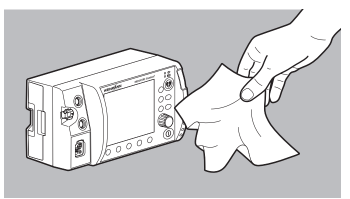
8.3 Cleaning parts manually

- *Requirement* The parts exhibit visible dirt.
 - Hygienic reprocessing is prepared (see "8.2 Preparing hygienic reprocessing", page 240).
 - For parts approved for cleaning, refer to the cleaning and disinfection plan (see "8.7 Cleaning and disinfection plan", page 248).
 - 2. For the agents, dose, and exposure time for the individual parts, refer to the cleaning and disinfection plan.

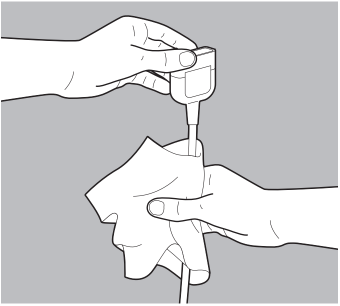
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- 3. Prepare the cleaning solution as specified by the cleaning agent manufacturer.
- 4. To remove all visible dirt: Brush parts thoroughly using a commercially available soft brush suitable for plastic and wetted with the cleaning agent.

When doing so, please note: Uneven surfaces and grooves (e.g. navigation knob) must be brushed especially thoroughly.



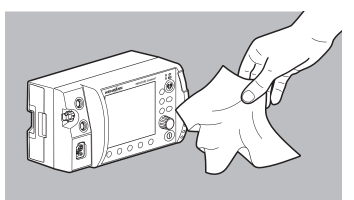
- Wipe over parts with a clean, lint-free cloth moistened with cleaning solution until they look clean. When doing so, please note:
 - Use a new cloth for every cleaning process.
 - All surfaces must be wiped carefully.
 - All surfaces must be wetted with cleaning solution.
 - The exposure time specified in the cleaning and disinfection plan must be observed.
 - Uneven surfaces and grooves need wiping over especially carefully.
 - The inner surfaces of MCS2-SoftTip pulse oximetry sensors must be turned inside out for cleaning.
 - The attachments for large electrode surfaces (adults) must be left on the paddles.
 - The attachments for large electrode surfaces must be removed to clean the attachments for small electrode surfaces (children/infants).



- Surround cables/tubes firmly with the cloth and draw them through the cloth so that they are completely wetted. When doing so, please note: Stretch out coiled cables.
- 7. If visible dirt is still present: Repeat manual cleaning.
- 8. Wipe down parts with a damp cloth to remove residues of the cleaning agent.
- 9. Allow parts to dry completely at room temperature.
- 10. If required: Dry parts manually with a lint-free cloth.
- *Result* Parts have been cleaned manually.

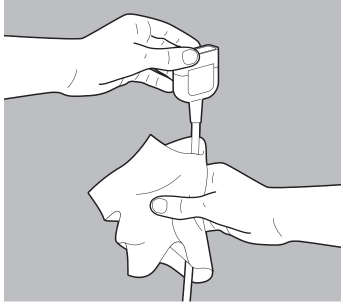
8.4 Disinfecting parts by wiping

- Hygienic reprocessing is prepared (see "8.2 Preparing hygienic reprocessing", page 240).
 - The parts have been cleaned manually and look clean (see "8.3 Cleaning parts manually", page 241).
 - 1. For parts approved for disinfection by wiping, refer to the cleaning and disinfection plan (see "8.7 Cleaning and disinfection plan", page 248).
 - 2. For the agents, dose, and exposure time for the individual parts, refer to the cleaning and disinfection plan.
 - 3. Prepare the solution for disinfection as specified by the disinfectant manufacturer.



- 4. Disinfect parts by wiping with one of the products listed. When doing so, please note:
 - Uneven surfaces and grooves (e.g. navigation knob) must be adequately wetted with disinfectant.
 - The exposure time specified by the manufacturer of the disinfectant must be observed.
 - The inner surfaces of MCS2-SoftTip pulse oximetry sensors must be turned inside out for cleaning.
 - The attachments for large electrode surfaces (adults) must be left on the paddles.

• The attachments for large electrode surfaces must be removed to clean the attachments for small electrode surfaces (children/infants).



- Surround cables/tubes firmly with the cloth and draw them through the cloth so that they are completely wetted. When doing so, please note: Stretch out coiled cables.
- 6. Wait for the exposure time and allow parts to dry completely at room temperature.
- *Result* The parts have been disinfected.

8.5 Subjecting bags to hygienic reprocessing

Requirement Hygienic reprocessing is prepared (see "8.2 Preparing hygienic reprocessing", page 240).

- 1. For the agents, dose, and exposure time for the bags, refer to the cleaning and disinfection plan.
- 2. Soak the bag in mains water for at least 10 minutes.
- 3. Brush contaminated areas under running mains water for at least 2 minutes using a cleaning brush until such areas look clean.
- 4. Rinse off cleaned areas under running mains water for 2 minutes.
- 5. Wash the open bag in a washing machine with cleaning agent and disinfectant but without a spin cycle:
 - Prewash: 5 min at 20 °C
 - Main wash: 30 min at 30 °C
- *Result* A bag has been hygienically reprocessed.

8.6 Preparing parts for reuse

Requirement The parts have been subjected to hygienic reprocessing according to the cleaning and disinfection plan.

- 1. Check all parts for damage resulting from use (e.g. tension cracks or cable breaks).
- 2. Replace damaged parts.
- 3. When the device has been disassembled from the protective transport bag and the holding plate: Mount the device in the protective transport bag (see "4.1.1 Mounting the device in the protective transport bag", page 74).

Alternatively:

When the device has been disassembled from the portable unit: Mount the device on the portable unit (see "4.1.2 Mounting the device on the portable unit", page 76).

4. Stow components and accessories in the protective transport bag (see "4.2.1 Stowing components and accessories in the protective transport bag", page 76).

Alternatively:

Stow components and accessories on the portable unit (see "4.2.2 Stowing components and accessories on the portable unit", page 83).

- Connect the power supply (see "4.3 Connecting a power supply", page 83).
- 6. Carry out a function check (see "5 Function check", page 108).
- If required: Store the device, components, and accessories in accordance with the conditions for storage (see "15 Storage", page 372).
- *Result* The parts are ready for use again.

8.7 Cleaning and disinfection plan

Carry out hygienic reprocessing according to the table below after **every** use:

Part	Manual cleaning (only necessary in the event of visible dirt)	Disinfection by wiping	Disinfection by immersion	Mechanical reprocessing	Sterilization
Device					
Device	Wipe down with neodisher [®] MediClean forte (Dr. Weigert). Dose: 10 ml/l Wipe down all surfaces at least 2x until they look clean.	surfaces with Incidin™ OxyWipe S (Ecolab). Exposure time:	Not permitted	Not permitted	Not permitted

Part	Manual cleaning (only necessary in the event of visible dirt)	Disinfection by wiping	Disinfection by immersion	Mechanical reprocessing	Sterilization
Power supply					
Accu-Pack battery Adapter cable for 12 V on- board power supply/circular connector	Wipe down with neodisher [®] MediClean forte (Dr. Weigert). Dose: 10 ml/l Wipe down all surfaces at least 2x until they look clean.	Wipe down all surfaces with Incidin™ OxyWipe S (Ecolab). Exposure time: 5 min	Not permitted	Not permitted	Not permitted
Power supply unit and charger Charging adapter Charging station for battery	MediClean forte (Dr. Weigert). Dose: 10 ml/l	Not necessary	Not permitted	Not permitted	Not permitted

Part	Manual cleaning (only necessary in the event of visible dirt)	Disinfection by wiping	Disinfection by immersion	Mechanical reprocessing	Sterilization
Defibrillation	cardioversion		1		
MCS2-Connect master cable	Wipe down with neodisher [®]				
MCS2-Hardpads paddles	(Dr. Weigert).	surfaces with			
Function test resistor	Dose: 10 ml/l Wipe down all surfaces at least 2x until they look clean.	Incidin™ OxyWipe S (Ecolab). Exposure time: 5 min	Not permitted	Not permitted	Not permitted
MCS2-Softpads defibrillation electrodes for children/adults (disposable)	Not permitted, a:	s disposables			
Electrode gel (disposable)					

Part	Manual cleaning (only necessary in the event of visible dirt)	Disinfection by wiping	Disinfection by immersion	Mechanical reprocessing	Sterilization
Pulse oximetr				•	
MCS2-Adapt pulse oximetry sensor connecting cable	Wipe down with neodisher [®] MediClean forte (Dr. Weigert). Dose: 10 ml/l Wipe down all surfaces at least 2x until they look clean.	Wipe down all surfaces with Incidin™ OxyWipe S (Ecolab). Exposure time: 5 min	Not permitted	Not permitted	Not permitted
MCS2-SoftTip pulse oximetry sensor	Wipe down with neodisher®				
MCS2-Wrap pulse oximetry sensor	MediClean forte (Dr. Weigert). Dose:	Wipe down all surfaces with Incidin™			
MCS2-Earclip pulse oximetry sensor	10 ml/l Wipe down all	OxyWipe S (Ecolab). Exposure time:	Not permitted	Not permitted	Not permitted
Ear clip for MCS2-Earclip pulse oximetry sensor	surfaces at least 2x until they look clean.	5 min			
MCS2-Wrap pulse oximetry sensors (disposable)			·		·
Strap for attaching MCS2-Wrap pulse oximetry sensor (disposable)	Not permitted, a	s disposables			

Part	Manual cleaning (only necessary in the event of visible dirt)	Disinfection by wiping	Disinfection by immersion	Mechanical reprocessing	Sterilization
6-lead ECG m	onitoring/12-le	ad ECG record	ing and assess	ment	
MCS2-Line ECG cables (all versions) ECG cable separator	neodisher [®] MediClean forte (Dr. Weigert). Dose: 10 ml/l Wipe down all surfaces at least 2x until they look clean.	surfaces with Incidin™ OxyWipe S (Ecolab). Exposure time: 5 min	Not permitted	Not permitted	Not permitted
ECG electrodes		ctions for use of t			
	blood pressure	measurement	: (NIBP monito	ring)	1
NIBP connecting tube Adapter tube for connecting NIBP disposable cuffs for neonates	Wipe down with neodisher [®] MediClean forte (Dr. Weigert). Dose: 10 ml/l Wipe down all surfaces at least 2x until they look clean.	surfaces with Incidin™ OxyWipe S (Ecolab). Exposure time:	Not permitted	Not permitted	Not permitted
NIBP cuffs (reusable)	Follow the instructions for use of the NIBP cuffs				
NIBP cuffs (disposable)	Not permitted, as disposables				

Part Printing	Manual cleaning (only necessary in the event of visible dirt)	Disinfection by wiping	Disinfection by immersion	Mechanical reprocessing	Sterilization
Printer Printer battery Charging station for printer battery with power supply unit and charger	MediClean forte (Dr. Weigert). Dose:	Wipe down all surfaces with Incidin™ OxyWipe S (Ecolab).	Not permitted	Not permitted	Not permitted
Quadruple charging station for printer battery including power supply unit and charger	look clean.	Exposure time:			

Part	Manual cleaning (only necessary in the event of visible dirt)	Disinfection by wiping	Disinfection by immersion	Mechanical reprocessing	Sterilization
Bags and por	table unit				_
Printer case MCS2-Bag protective transport bag	Wipe down smooth surfaces with neodisher [®] MediClean forte (Dr. Weigert). Dose: 10 ml/l Wipe down all surfaces at least 2x until they look clean.	Not permitted	Not permitted	Wash in a washing machine with no spin cycle. Prewash: 5 min at 20 °C Main wash: 30 min at 30 °C Cleaning agent: Turbo Usona (Ecolab) Disinfectant: Turbo Oxysan (Ecolab)	Not permitted
Holding plate	Wipe down with neodisher [®] MediClean forte (Dr. Weigert). Dose: 10 ml/l Wipe down all surfaces at least 2x until they look clean.	surfaces with Incidin™ OxyWipe S (Ecolab). Exposure time: 5 min	Not permitted	Not permitted	Not permitted
Portable unit	Follow the instru	ctions for use of t	he portable unit		

Part Accessories fo	Manual cleaning (only necessary in the event of visible dirt) or training	Disinfection by wiping	Disinfection by immersion	Mechanical reprocessing	Sterilization
ECG simulator, 6-lead ECG,	Wipe down with				
shockable ECG simulator, 12-lead ECG, shockable	neodisher [®] MediClean forte (Dr. Weigert). Dose:				
Adapter cable for connecting to Ambu/Laerdal practice manikin	· · ·	Not necessary	Not permitted	Not permitted	Not permitted
Adapter cable for connecting ShockLink [®]	2x until they look clean.				



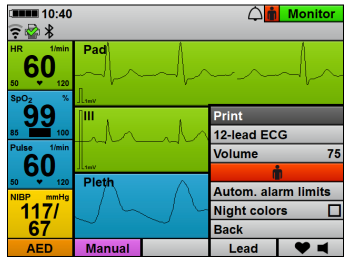
The applicable instructions are those in the instructions for use from the manufacturers of the individual components or parts. Follow these instructions for use.

9 Application menu

The application menu contains functions and settings which can be accessed quickly and easily during the session.

9.1 Navigating the application menu

- The device is switched on (see "6.1 Switching on the device", page 125).
 - The patient group is set (see "6.4 Selecting patient group", page 127).
 - A mode is set.
 - 1. To call up the application menu: Press the navigation knob. The application menu appears (example):



- 2. Select setting using the navigation knob.
- 3. Change setting using the navigation knob.

Alternatively:

Press the navigation knob to activate/deactivate a setting.

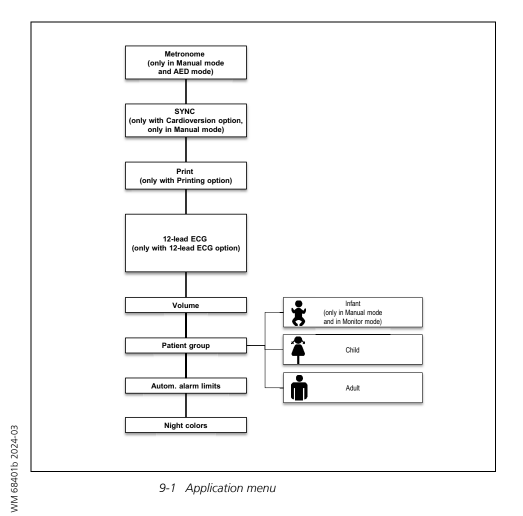
4. To exit the menu without making any entries: Select the **Back** menu item.

Alternatively:

Wait 3 seconds.

Result Functions are carried out or settings are made.

9.2 Menu structure



9.3 Settings

Parameter	Possible values	Description	Factory setting
Metronome (only in manual and AED modes)	Off 15:2 (only in manual and AED modes, child) 30:2 (only in manual and AED modes, adult)	Here you can set the metronome rhythm.	 AED mode adult: 30:2 AED mode child: 15:2 Manual mode: off
SYNC (only with Cardioversion option, only in manual mode)	Activated M Deactivated	Here you can set whether shocks are to be delivered synchronized. Once a shock has been delivered, then depending on the settings in the operator menu (Operator menu Manual mode settings Defibrillation after cardioversion) another cardioversion or defibrillation can be carried out.	Deactivated
12-lead ECG (only with 12-lead ECG option)	-	Here you activate the 12-lead ECG function mode. If you start 12-lead ECG function mode from AED or manual mode, the device automatically switches to monitor mode in the background.	-
Print (only with Printing option)	-	 Here you can print the following printouts: Live printout 12-lead ECG printout Replay printout The Printing menu item is grayed out if the printer is not connected. When the printer is printing, the menu item is called Stop print xx s. A printing process can be finished prematurely by pressing this menu item. A counter indicates the time remaining until the end of printing. 	-

Parameter	Possible values	Description	Factory setting
Volume	25 % 50 % 75 % 100 %	Here you can set the volume of the device for the current session. The operator specifies minimum volume (see "13.12 System settings", page 358).	75 %
Patient group	Adult Child Adult Child Adult Child Adult Child Adult Child Adult Child Adult Child Adult Adult Child Adult Ad	Here you can select the patient group.	Adult
Autom. alarm limits	-	The device automatically specifies the alarm limits for the physiological alarms. The alarm limit corridor is 10 %, 20 % or 30 % above or below the physiological values measured at the time of activation. The alarm limit corridor can be set in the user menu (see "10 User menu", page 260).	20 %
Night colors	Activated Deactivated	Here you can set whether the device is to display night colors.	Deactivated

10 User menu

The user menu contains functions and settings which affect the current session and which are not saved permanently as device presets (except: date and time settings).

If the device was switched off for < 30 s and if, while switched on previously, patient measured values were determined or a manual event was saved, the settings made previously in the user menu are retained.

10.1 Navigating the user menu

Requirement	٠	The device is switched on (see "6.1 Switching on the device",
		page 125).

- The patient group is set (see "6.4 Selecting patient group", page 127).
- A mode is set.
- Press the menu button (I). The user menu appears:

13:49		\triangle	Menu
(î;			
	User menu		
Alarm settings			
ECG settings			
12-lead ECG settings			
SpO ₂ settings			
Printer settings	_	_	
Communication setti	ngs		
System settings			
Session archive			
Back			
	Monitor	Back	OK

2. Select the setting with the navigation knob and confirm.

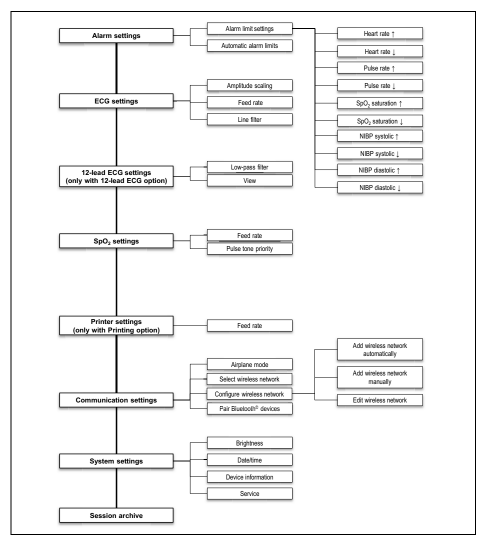
- 3. Change the setting with the navigation knob and confirm.
- 4. To exit the menu: Select the **Back** function button.

Alternatively:

Press the menu button 📳.

Result Settings are made and apply to the current session.

10.2 Menu structure



10-1 User menu

WM 68401b 2024-03

10.3 Settings

10.3.1 Alarm settings

WARNING

Risk of injury due to alarm limits which are too high or too low!

Alarm limits which are either too high or too low may prevent the device triggering an alarm, thereby putting the patient at risk.

⇒ Always set alarm limits which have been adapted to the patient.

13:50	\triangle	Menu
(;; 🖓 ¥		
Alarm settings		
Alarm limit settings		
Automatic alarm limits		20 %
Back		
Monitor	Back	OK
Wonitor	Back	OK

10-2 Alarm settings submenu

10 User menu

Parameter		Possible values	Description	Factory setting
	Heart rate †	35/min - 250/min, in increments of 5	Here you can set the upper (†) and lower (↓) limit values as of which the device is to trigger an alarm. According to the values set, the setting ranges of the upper and lower limit values depend on each other as follows: The setting range for the upper alarm limit ends at least one setting value above the set value for the lower alarm limit. The setting range for the lower alarm limit ends at least one setting value below the set value for the upper alarm limit.	120/min
	Heart rate 🖡	30/min - 245/min, in increments of 5		50/min
	Pulse rate 1	35/min - 250/min, in increments of 5		120/min
	Pulse rate ↓	30/min - 245/min, in increments of 5		50/min
	SpO_2 saturation \uparrow	66 % - 100 %		100 %
	SpO_2 saturation \downarrow	65 % - 99 %		85 %
Alarm limit settings Adult	NIBP systolic †	45 mmHg - 260 mmHg, in increments of 5		220 mmHg
	NIBP systolic ↓	40 mmHg - 255 mmHg, in increments of 5		75 mmHg
	NIBP diastolic †	25 mmHg - 200 mmHg, in increments of 5		110 mmHg
	NIBP diastolic ↓	20 mmHg - 195 mmHg, in increments of 5		35 mmHg

Parameter		Possible values	Description	Factory setting
	Heart rate †	35/min - 250/min, in increments of 5	Here you can set the upper (1) and lower (1) limit values as of which the device is to trigger an alarm. According to the values set, the setting ranges of the upper and lower limit values depend on each other as follows: The setting range for the upper alarm limit ends at least one setting value above the set value for the lower alarm limit. The setting range for the lower alarm limit ends at least one setting value below the set value for the upper alarm limit.	150/min
	Heart rate ↓	30/min - 245/min, in increments of 5		50/min
	Pulse rate †	35/min - 250/min, in increments of 5		150/min
	Pulse rate ↓	30/min - 245/min, in increments of 5		50/min
	SpO ₂ saturation †	66 % - 100 %		100 %
	SpO_2 saturation \downarrow	65 % - 99 %		85 %
Alarm limit settings Child	NIBP systolic †	45 mmHg - 230 mmHg, in increments of 5		145 mmHg
	NIBP systolic ↓	40 mmHg - 225 mmHg, in increments of 5		75 mmHg
	NIBP diastolic †	25 mmHg - 160 mmHg, in increments of 5		100 mmHg
	NIBP diastolic ↓	20 mmHg - 155 mmHg, in increments of 5		35 mmHg

Parameter		Possible values	Description	Factory setting
	Heart rate 1	35/min - 250/min, in increments of 5		200/min
	Heart rate ↓	30/min - 245/min, in increments of 5	Here you can set the upper (1) and lower (1) limit values as of which the device	100/min
	Pulse rate †	35/min - 250/min, in increments of 5		200/min
	Pulse rate ↓	30/min - 245/min, in increments of 5	According to the values set, the setting ranges of the	100/min
	SpO_2 saturation \uparrow	66 % - 100 %	upper and lower limit values	95 %
	SpO_2 saturation \downarrow	65 % - 99 %	follows: The setting range for the upper alarm limit ends at least one setting value above the set value for the lower alarm limit. The setting range for the lower alarm limit ends at least one setting value below the set value for the upper alarm limit.	85 %
Alarm limit settings Infant	SpO ₂ saturation 1	45 mmHg - 130 mmHg, in increments of 5		100 mmHg
	SpO ₂ saturation ↓	40 mmHg - 125 mmHg, in increments of 5		50 mmHg
	NIBP systolic †	25 mmHg - 100 mmHg, in increments of 5		70 mmHg
	NIBP systolic ↓	20 mmHg - 95 mmHg, in increments of 5		30 mmHg
Automatic alarm limits	-	10 % 20 % 30 %	Here you can set the automatic alarm limits. The device automatically specifies the alarm limits for the physiological alarms. The deviation is 10 %, 20 % or 30 % from the measured values at the time of activation.	20 %

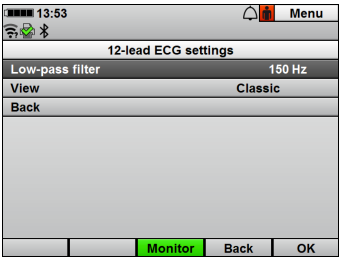
10.3.2 ECG settings

I3:52			Menu
Ŝ №			
E	ECG setting	S	
Amplitude scaling		Auto	om.
Feed rate			25 mm/s
Line filter			区
Back			
	Monitor	Back	ок
	wonitor	Back	UK

10-3 ECG settings submenu

Parameter	Possible values	Description	Factory setting
Amplitude scaling	Autom. 2 mm/mV 5 mm/mV 10 mm/mV 20 mm/mV	Here you can set the amplification of the ECG signal and thus the height of the ECG curve. If the Autom . setting has been selected, the amplification of the ECG signal and thus the height of the ECG curve are adapted automatically so that a maximum height for lead II is displayed.	Autom.
Feed rate	12.5 mm/s 25 mm/s 50 mm/s	Here you can set the feed rate of the ECG curve display and so change the time resolution.	25 mm/s
Line filter	Activated Deactivated	By activating the line filter, you can reduce ECG interference caused by the power supply network.	Activated

10.3.3 12-lead ECG settings (only with 12-lead ECG option)



10-4 12-lead ECG settings submenu

Parameter	Possible values	Description	Factory setting
Low-pass filter	50 Hz Here you can set a low-pass filter to filter out artifacts.		150 Hz
View	Classic Cabrera	 Here you can set the sequence in which the leads are displayed in 12-lead ECG function mode: Classic: I, II, III, aVR, aVL, aVF, V1, V2, V3, V4, V5, V6 Cabrera: aVL, I, -aVR, II, aVF, III, V1, V2, V3, V4, V5, V6 	Classic

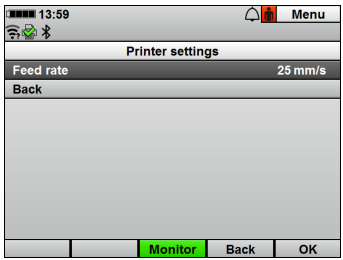
10.3.4 SpO₂ settings

13:55		\triangle	Menu
€ 🗟 🖇			
S	pO ₂ setting	S	
Feed rate			25 mm/s
Pulse tone priority			区
Back			
	Manifan	Deek	01/
	Monitor	Back	ОК

10-5 SpO₂ settings submenu

Parameter	Possible values	Description	Factory setting
Feed rate	12.5 mm/s 25 mm/s 50 mm/s	Here you can set the feed rate of the SpO_2 curve display and so change the time resolution.	25 mm/s
Pulse tone priority	Activated C Deactivated	Here you can set whether pulse tone is to take priority over heart rate tone. The tone level of the pulse tone is adapted to measured oxygen saturation.	Activated

10.3.5 Printer settings (only with Printing option)



10-6 Printer settings submenu

Parameter	Possible values	Description	Factory setting
Feed rate	12.5 mm/s 25 mm/s 50 mm/s	Here you can set the feed rate at which the printer prints a live printout and a replay printout.	25 mm/s

10.3.6 Communication settings

14:01			\triangle	Menu	
??? 🖓 🖇					
	Comm	unication s	ettings		
Airplane n	node				
Select wir	eless netwo	ork			
Configure	wireless n	etwork			
Pair Bluet	ooth® devi	ces			
Back					
		Monitor	Back	ОК	

10-7 **Communication settings** submenu

Parameter	Possible values	Description	Factory setting
Airplane mode	Activated Deactivated	Here you can activate or deactivate all the wireless interfaces of the device (example: WiFi interface).	Deactivated
Select wireless network	-	Here you can select a network configured in the operator menu as the default network. If you cancel the process, the menu item is grayed out briefly.	-

Parameter		Possible values	Description	Factory setting
Configure wireless network	Add wireless network automatically	-	Here you can search for an available wireless network and automatically add it as the default network.	-
	Add wireless network manually	-	Here you can configure a wireless network manually. Contact your IT system administrator for required information about the settings.	-
	Edit wireless network	-	Here you can edit the settings for a wireless network that has already been configured and specify a wireless network as the default network.	-
Pair Bluetooth [®] devices		-	Here you can pair your device with a compatible system for digital patient data recording. If you cancel the process, the menu item is grayed out briefly.	-

10.3.7 System settings

14:02				Menu
?; 🗟 🖇				
	Sy	stem settin	gs	
Brightnes	S			70 %
Date/time				
Device inf	ormation			
Service				
Back				
		Manifan	Deek	01/
		Monitor	Back	ОК

10-8 System settings submenu

Parameter		Possible values	Description	Factory setting
Brightness		10 %-100 %, in 10 % increments	Here you can set the display brightness.	70 %
	Year		Here you can set the date and	-
	Month	-	time. The device adopts a new date or a new time only if it remains switched on for at least 1 min after the date	-
	Day			-
Date/time	Hour			-
	Minute		or time have been changed. The change to date or time only takes effect once the device is switched on again.	-

Parameter		Possible values	Description	Factory setting
	Telephone contact	-	Here you can find out the telephone number the operator has assigned to the device (example: The number for a cellphone on the emergency vehicle).	-
	Serial number	-	Here you can find out the device serial number. This is also located on the device information label.	-
	Device ID	-	Here you can find out the device ID. This is required to purchase options.	-
Device information	MAC Bluetooth [®] module (only with Printing and Blue- tooth [®] data transmission options)	-	Here you can find out the MAC address of the Bluetooth [®] module.	-
	MAC WLAN module	-	Here you can find out the MAC address of the WLAN module.	-
	Last function check	-	Here you can find out when a function check was last carried out.	-
	Function check result	-	Here you can find out whether the last function check carried out was passed.	-
	Days until next service	-	Here you can find out how many days to go until the next service is due.	-
	Next service	-	Here you can find out when the next service is due.	-

Parameter		Possible values	Description	Factory setting
Device information	Device software	-	Here you can find out which software version is currently installed on the device.	-
Service	Upload service data to WEINMANN	-	Here you can upload the device service data to WEINMANN Emergency. If you cancel the process, the menu item is grayed out briefly.	-
	Export service data to SD card	-	Here you can export the device service data to an SD card.	-

10.3.8 Session archive

14:03				Menu
	0			
	Ses	sion archiv	e	
Date	Time	Duration	CPR	12-lead
2024-02-13	13:37	26:37	No	No
2024-02-13	13:22	14:27	Yes	No
2024-02-13	13:18	02:21	No	No
2024-02-13	13:04	14:06	Yes	Yes
2024-02-13	13:03	00:10	No	No
2024-02-13	12:58	04:30	No	Yes
2024-02-13	12:54	04:23	No	Yes
Load all sessions				
		Monitor	Back	ОК

10-9 Session archive submenu

Parameter	Possible values	Description	Factory setting
Session archive	-	 Here you can have all the sessions saved in the internal memory of the device displayed (approx. the last 9 h). The following actions are possible for any session: Analyze the session in replay view on the device (only with Replay view option) (see 6.17.2, p. 214) Analyze the 12-lead ECG of a session (only with 12-lead ECG option) (see 6.17.3, p. 216) Print a session report (only with Printing option) (see 6.16.7, p. 211) The session archive is only available if at least one of the options mentioned is enabled. 	-

11 Alarms and faults

11.1 General instructions

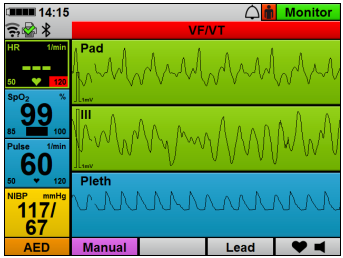
The device emits an alarm as long as the cause continues to exist. Once the cause of the alarm no longer exists, the device no longer emits the alarm. Some alarms are excepted from this. The duration of these alarms is marked in the tables.

The device emits physiological and technical alarms. Every alarm has a certain priority:

Priority	Color in alarm line	Meaning
High priority	Red	High-priority alarms warn of imminent fatal or irreversible patient injuries or of device faults.
Medium priority	Yellow	Medium-priority alarms warn of immediate reversible patient injuries or of minor device faults.
Low priority	Turquoise	Low-priority alarms warn of delayed minor injuries or inconvenience to the patient or minor restrictions on the device.

If more than one alarm is active, the device handles this as follows:

- Multiple alarms of different priorities: The device displays the alarm with the highest priority. Alarms with a lower priority do not appear until the higher-priority alarms are no longer active.
- Multiple alarms of identical priorities: The device displays the alarms in turn.



11-1 Alarm line with VF/VT alarm (example)

The device displays alarms as follows:

- As text in the alarm line in the display
- As audio alarm tones (via the loudspeaker)
- With the alarm light (in the top right-hand corner on the front of the device)

Alarms are outputted according to their priority:

Type of alarm signal	High priority	Medium priority	Low priority
Alarm line			
Flashing frequency	2 Hz	0.5 Hz	Not flashing (constantly on)
Size	90 mm x 5 mm		
Color	Red	Yellow	Turquoise
Signal ratio	1:1	1:1	100 % (on)

Type of alarm signal	High priority	Medium priority	Low priority
Audible alarm			
Number of pulses	5	3	1
Duration of pulses	120 ms	200 ms	200 ms
Pulse interval	240 ms	200 ms	Not applicable
Pulse frequency	480 Hz	840 Hz	1200 Hz
Alarm light			
Color	Red	Off	Off
Frequency	2 Hz	Not applicable	Not applicable
Signal ratio	33 % on	Not applicable	Not applicable
Size	10 mm x 35 mm	•	•

The device additionally displays physiological alarms by means of the respective parameter field flashing.

The device's alarm behavior depends on the mode selected:

- In AED mode, the device does not emit alarms. The device provides information by means of AED instructions and voice prompts about technical conditions which lead to AED mode being impaired.
- Audio alarm output is deactivated in manual mode. It can be reactivated at any time by pressing the alarm button. In the operator menu, the operator can specify whether audio alarm output is to be activated when manual mode is entered.

11.2 Alarm messages

11.2.1 High-priority alarm (red)

Alarm	Cause	Remedy
Asystole	Asystole detected	Act in accordance with currently applicable guidelines.
		If medically indicated, carry out suitable treatment.
Battery defective	Battery defective	Let the device run on battery power without line power until it switches off. Fully recharge battery (see 4.4.2, p. 86). If the device continues to display the alarm: Change battery (see 4.4.4, p. 88).
Battery empty	Very low battery status	Change battery (see 4.4.4, p. 88). Connect device to line power (see 4.3, p. 83) and charge battery (see 4.4.2, p. 86).
Check defibrillation electrodes (Manual mode)	Defibrillation electrodes no longer attached/not attached correctly to the patient	Attach defibrillation electrodes to the patient correctly (see 6.5.1, p. 129).
Check patient contact	Paddles are not in contact/not correctly in contact with the patient	Check contact between paddles and patient. If the alarm persists: Cancel shock delivery.
Defibrillation module defective	Internal defibrillation module defective	Have the device repaired.
Insert battery (Manual mode)	Battery not inserted/not inserted correctly	Insert battery correctly (see 4.3, p. 83).
	Defibrillation electrodes not connected/not connected correctly to the master cable	Connect defibrillation electrodes to the master cable correctly (see 6.5.1, p. 129).
Plug in Pad connector (Manual mode)	Paddles not connected/not connected correctly to the master cable	Connect paddles to the master cable correctly (see 6.5.2, p. 133).
	Master cable not connected/not connected correctly to the device	Connect the master cable to the device correctly (see 6.5.1, p. 129).
Shock delivery unsuccessful (Manual mode)	No patient contact during shock delivery	Wait 10 seconds. Charge the shock capacitor again using the Charge function button.

Alarm	Cause	Remedy
		Act in accordance with currently applicable guidelines.
VF/VT*	Ventricular fibrillation or ventricular tachycardia detected	Switch to AED mode or manual mode and carry out defibrillation, if medically indicated (see 6.5, p. 129).

* In the operator menu, the device can be preset so that the VF/VT alarm can be deactivated and activated by the user. It is also possible in the operator menu to preset whether the device is to start with the VF/VT alarm activated or deactivated.

11.2.2 Medium-priority alarm (yellow)

Alarm	Cause	Remedy
Battery weak	Low battery status	 Change battery (see 4.4.4, p. 88). Connect the device to line power (see 4.3, p. 83).
Check defibrillation electrodes (Monitor mode)	Defibrillation electrodes no longer attached/not attached correctly to the patient	Attach defibrillation electrodes to the patient correctly (see 6.5.1, p. 129).
Check seating of ECG electrodes	ECG electrodes not attached/not attached correctly to the patient	Attach ECG electrodes to the patient correctly (see 6.10.1, p. 171).
	ECG cable not connected correctly to the ECG electrodes	Connect ECG cable to ECG electrodes correctly (see 6.10.1, p. 171).
Close paper compartment	Paper compartment cover of the printer open	Close the paper compartment cover.
	Device outside the range of the network selected in the operator menu	Bring the device within range of the selected network. If necessary: Select a different network in the operator menu (see 12.3, p. 302).
Connection failed (Alarm deactivated after 60 s)	Device unable to connect to the network selected in the operator menu	Check login data (see 12.3, p. 302).
		Check the signal strength of the selected network. If necessary: Select a different network in the operator menu (see 12.3, p. 302).

Alarm	Cause	Remedy
ECG module defective	Internal ECG module defective	Have the device repaired.
	Server not accessible	Check connection to network (see 12.3, p. 302).
E-mail delivery error		Check preset e-mail recipient (see 12.6, p. 311).
(Alarm deactivated after 60 s)	Connection between device and network interrupted	Check connection to network (see 12.3, p. 302). If necessary: Select a different network.
		Check date/time (see 10.3.7, p. 273).
Excessive NIBP cuff pressure (Alarm deactivated after 10 s)	Pressure in the NIBP cuff increased to such a degree as a result of external influences that the safety valve opened	Check which influences resulted in the pressure increase and eliminate them before the next measurement.
Heart rate †	Measured heart rate is above the set upper alarm limit	If medically indicated, carry out suitable treatment.
		Adapt alarm limits (see 10.3.1, p. 263).
Heart rate > 250/min	Measured heart rate is above 250/ min (device shows as heart rate)	If medically indicated, carry out suitable treatment.
Heart rate ↓	Measured heart rate is below the set lower alarm limit	If medically indicated, carry out suitable treatment.
neart rate +		Adapt alarm limits (see 10.3.1, p. 263).
Heart rate < 30/min	Measured heart rate is below 30/ min (device shows as heart rate)	If medically indicated, carry out suitable treatment.
Incost printor papar	Printer paper completely used up	Change printer paper (see 4.7.3, p. 98).
Insert printer paper	No printer paper inserted	Change printer paper (see 4.7.3, p. 98).
Memory error	Internal memory module defective	Have the device repaired.
NIBP diastolic †	Measured diastolic blood pressure	If medically indicated, carry out suitable treatment.
(Alarm deactivated after 10 s)	is above the set upper alarm limit	Adapt alarm limits (see 10.3.1, p. 263).

Alarm	Cause	Remedy
NIBP diastolic ↓ (Alarm deactivated after 10 s)	Measured diastolic blood pressure is below the set lower alarm limit	If medically indicated, carry out suitable treatment. Adapt alarm limits (see 10.3.1, p. 263).
NIBP measurement error (Alarm deactivated after 10 s)	NIBP measurement cannot be carried out	 Make sure that the patient lies still during the measurement. Rule out the risk of patient vibration. Use a suitable size of NIBP cuff. Ensure that the NIBP cuff is properly attached. Ensure that there is no clothing between the NIBP cuff and the patient.
NIBP module defective (Alarm active as long as the condition prevails)	Internal NIBP module defective	Have the device repaired.
NIBP outside measuring range (Alarm deactivated after 10 s)	Blood pressure cannot be measured and displayed as it is outside the device's measuring range	Use an alternative blood pressure measurement device with a larger measuring range.
No NIBP signal (Alarm deactivated after 10 s)	NIBP module unable to detect a pulse wave signal	 Make sure that the patient lies still during the measurement. Rule out the risk of patient vibration. Use a suitable size of NIBP cuff. Ensure that the NIBP cuff is properly attached. Ensure that there is no clothing between the NIBP cuff and the patient.
NIBP safety shutdown	Pressure in the NIBP cuff is too	Remove the NIBP cuff from the
(Alarm deactivated after 10 s)	high for too long	patient. Have the device repaired.

Alarm	Cause	Remedy
NIBP signal disrupted (Alarm deactivated after 10 s)	Unable to detect pulse wave signal for NIBP measurement reliably	 Make sure that the patient lies still during the measurement. Rule out the risk of patient vibration. Use a suitable size of NIBP cuff. Ensure that the NIBP cuff is properly attached. Ensure that there is no clothing between the NIBP cuff and the patient.
NIBP systolic † (Alarm deactivated after 10 s)	Measured systolic blood pressure is above the set alarm limit	If medically indicated, carry out suitable treatment. Adapt alarm limits (see 10.3.1, p. 263).
NIBP systolic ↓ (Alarm deactivated after 10 s)	Measured systolic blood pressure is below the set alarm limit	If medically indicated, carry out suitable treatment. Adapt alarm limits (see 10.3.1, p. 263).
NIBP tube blocked (Alarm deactivated after 10 s)	NIBP connecting tube kinked or blocked	Check NIBP connecting tube and replace if necessary.
NIBP tube leaking (Alarm deactivated after 10 s)	NIBP connecting tube or NIBP cuff are leaking or incorrectly connected. Sufficient pressure cannot be built up.	Check NIBP connecting tube and NIBP cuff and replace if necessary.
	·	Wait 10 seconds.
No printer connected	Connection between device and	Switch printer on again.
(Alarm deactivated after 10 s)	printer interrupted	Position the printer within range of the device.
Plug in ECG connector	ECG connector of the ECG cable not connected/not connected correctly to the device	Connect ECG connector of the ECG cable to the device correctly (see 6.10.1, p. 171).
	Defibrillation electrodes not connected/not connected correctly to the master cable	Connect defibrillation electrodes to the master cable correctly (see 6.5.1, p. 129).
Plug in Pad connector (Monitor mode)	Paddles not connected/not connected correctly to the master cable	Connect paddles to the master cable correctly (see 6.5.2, p. 133).
	Master cable not connected/not connected correctly to the device	Connect the master cable to the device correctly (see 6.5.1, p. 129).

Alarm	Cause	Remedy
Pulse rate †	Measured pulse rate is above the set alarm limit	If medically indicated, carry out suitable treatment. Adapt alarm limits (see 10.3.1, p. 263).
Pulse rate ↓	Measured pulse rate is below the set alarm limit	If medically indicated, carry out suitable treatment. Adapt alarm limits (see 10.3.1, p. 263).
Server not accessible (Alarm deactivated after 60 s)	Server not accessible as there is no Internet connection	Check using an alternative network whether the configured network is the cause.
Settings lost	Settings had to be reset to factory settings.	Re-enter presets in the operator menu. If this recurs: Have the device repaired.
SpO ₂ †	Measured oxygen saturation is above the set alarm limit and SpO_2 signal quality is $\geq 40 \%$	If medically indicated, carry out suitable treatment. Adapt alarm limits (see 10.3.1, p. 263).
SpO₂↓	Measured oxygen saturation is below the set alarm limit and SpO_2 signal quality is $\geq 40 \%$	If medically indicated, carry out suitable treatment. Adapt alarm limits (see 10.3.1, p. 263).
SpO ₂ module defective	Internal SpO ₂ module defective	Have the device repaired.
Unknown defibrillation	Unknown defibrillation electrodes connected to the master cable	Only use defibrillation electrodes from WEINMANN Emergency.
electrodes	Unknown paddles connected to the master cable	Only use paddles from WEINMANN Emergency.
	Data transmission canceled by user	• Restart data transmission (see 12.6, p. 311).
Upload canceled (Alarm deactivated after 60 s)	Data transmission interrupted during transmission	 Check date/time (see 10.3.7, p. 273). Check connection settings (see 12.6, p. 311). Restart data transmission (see 12.6, p. 311).
Upload failed	Connection to network interrupted	Check connection to network (see 12.3, p. 302). If necessary: Select a different network.

11.2.3 Low-priority alarm (turquoise)

Alarm	Cause	Remedy
Battery operation (Alarm deactivated after 10 s)	Line power too low Line power disconnected by removing from the wall mounting Power outage	Restore line power.
Check seating of SpO ₂ sensor	Pulse oximetry sensor not attached/not attached correctly to the patient	Attach pulse oximetry sensor to the patient correctly (see 6.9.1, p. 164).
Erroneous date	Date read-out error	Insert battery (see 4.3, p. 83). Reset date (see 10.3.7, p. 273).
	Internal module defective	If this recurs: Have the device repaired.
	Real Time Clock battery defective or empty	Have the device repaired.
Insert battery (Monitor mode)	Battery not inserted/not inserted correctly	Insert battery correctly (see 4.3, p. 83).
Insert SD card	No SD card present/detected	Insert new SD card (see 4.6.1, p. 92), switch off device (see 6.2, p. 126), and switch device back on after at least 30 s (see 6.1, p. 125).
Internal memory full	No SD card inserted and session duration > 4 h-10 h	To continue to record the session: Insert SD card (see 4.6.1, p. 92), copy data to SD card, and resume session with SD card.
		Resume session and do not record any more data.
Plug in SpO ₂ sensor	SpO ₂ connector of the pulse oximetry sensor connecting cable not connected/not connected correctly to the device	Connect SpO ₂ connector of the pulse oximetry sensor connecting cable to the device correctly.
Printer battery empty	Status of printer battery very low	Change printer battery (see 4.7.2, p. 97).
		Charge printer battery (see 4.7.1, p. 94).

Alarm	Cause	Remedy
SD card defective	SD card write/read error	Insert new SD card (see 4.6.1, p. 92), switch off device (see 6.2, p. 126), and switch device back on after at least 30 s (see 6.1, p. 125).
		Resume session and do not record any more data.
SD card full	SD card full	Insert new SD card (see 4.6.1, p. 92), switch off device (see 6.2, p. 126), and switch device back on after at least 30 s (see 6.1, p. 125).
		Erase data on current SD card.
SpO ₂ sensor or cable defective	Pulse oximetry sensor connecting cable defective	Replace pulse oximetry sensor connecting cable.
	Pulse oximetry sensor defective	Replace pulse oximetry sensor.
SpO ₂ signal quality ↓ *	SpO ₂ signal quality < 40 %	Attach pulse oximetry sensor to the patient correctly (see 6.9.1, p. 164).
		If medically indicated, carry out suitable treatment.
Undata anavatar manu accase	Access code for the operator menu is 000000 (status when device supplied)	Wait 10 seconds.
Update operator menu access code (Alarm deactivated after 10 s)		Change operator menu access code (see "13.12 System settings", page 358).

* If the device outputs this alarm, it will no longer output a simultaneously occurring SpO_2 measured value alarm. If signal quality drops to < 20 %, the device will also stop displaying the SpO_2 measured value and the plethysmogram.

11.3 Faults

If you are unable to remedy faults with the aid of the table, you should contact the manufacturer WEINMANN Emergency, or technicians expressly authorized by WEINMANN Emergency, to have the device repaired. To avoid more serious damage, do not continue using the device.

11.3.1 Device

Fault	Cause	Remedy
Device cannot be switched on	Battery not correctly inserted in device, or battery empty	Check battery.
	Battery empty and device not connected to line power	Check power supply.
	Device defective	Have the device repaired.
Device cannot be switched off	Operating error	Press and hold On/Off button \textcircled{O} for at least 2 s.
	Device defective	 Disconnect the device from line power. Remove the battery. Have the device repaired.
Yellow display with ERROR and error code	Temporary device fault	 Switch device off (see 6.2, p. 126) and back on again (see 6.1, p. 125). Carry out a function check (see 5.2, p. 109).
	Device defective	Have the device repaired.
Function check does not start	Battery weak or empty	 Insert a battery with a battery status of at least 2 LEDs. Restart the function check.
	Master cable not detected	Connect master cable correctly.
	Master cable defective	Replace the master cable.
	Function test resistor not detected	Connect function test resistor correctly.
	Function test resistor defective	Replace the function test resistor.
	Pulse oximetry sensor connecting cable not detected	Connect pulse oximetry sensor connecting cable correctly.
	Pulse oximetry sensor connecting cable defective	Replace pulse oximetry sensor connecting cable.
	Pulse oximetry sensor not detected	Connect pulse oximetry sensor correctly.
	Pulse oximetry sensor defective	Replace pulse oximetry sensor.
Brightness of the display too low	Brightness of display set too low	Increase brightness of the display.
	Night colors activated	Deactivate night colors.

Fault	Cause	Remedy
	Volume set to 25 %	Increase volume in application menu (see 9, p. 256).
Alarm output too quiet		Increase volume preset in operator menu (see 13.12, p. 358).
	Battery empty and device not connected to line power	Check power supply.
Energy failure/black screen	Temporary device fault	 Switch device off (see 6.2, p. 126) and back on again (see 6.1, p. 125). Carry out a function check (see 5.2, p. 109).
	Device defective	Have the device repaired.
	Loose contact on navigation knob	
Settings cannot be made with the navigation knob	No gap between navigation knob and control panel membrane	Have the device repaired.
	Device defective	
Device failure Alarm light flashing Audio alarm output 	Device defective	Have the device repaired.

11.3.2 Battery and charging station

Fault	Cause	Remedy
Red fault indicator on when status	Battery defective	Replace battery.
button on battery is pressed, or red battery status indicator on device on	Battery temperature outside the permitted range (> 70 °C)	Use battery within permitted temperature range (see 18.5, p. 389).
Battery does not respond when status button is pressed	Battery has run down completely and has shut down to prevent deep discharge	 Charge battery in the device for 24 h (see 4.4.2, p. 86). After 24 h: Green LED is on: Battery fully charged and ready for use. Red LED or no LED on: Battery defective. Replace battery.
		Charge the battery in the charging station (see 4.4.3, p. 87): Charging process takes longer than usual.
Device runtime with battery operation too short	Battery has reached end of its lifetime	Replace battery.

11 Alarms and faults

Fault	Cause	Remedy
Battery not charging although it is not full	Battery temperature < 0 °C or > 45 °C	Charge battery within permitted temperature range (see 18.5, p. 389).
	Battery defective	Replace battery.
	Battery not inserted correctly in	Check battery.
	charging station	Insert battery correctly.
Status LEDs are not flashing and battery is not charging	Charging station not connected to	Connect the charging station to line power (see 4.5.3, p. 90).
	Charging station not connected to the power supply	Connect the charging station to a 12 V on-board power supply (see 4.5.4, p. 90).
	Charging station defective	Have charging station repaired.
	Battery capacity at $> 90 \%$	Press the status button on the battery to check battery status.

11.3.3 Defibrillation/cardioversion

Fault	Cause	Remedy
	Defibrillation electrodes not attached correctly to the patient	Attach defibrillation electrodes to the patient correctly (see 6.5.1, p. 129).
The Check defibrillation	Defibrillation electrodes defective	Replace defibrillation electrodes.
electrodes alarm occurs although the defibrillation electrodes are attached to both the patient and the device via the master cable	Pad connector not connected correctly to the master cable	Connect Pad connector of the defibrillation electrodes to the master cable correctly.
	Master cable not connected correctly to the device	Connect the master cable to the device correctly.
	Master cable defective	Replace the master cable.
	Defibrillation module defective	Have the device repaired.

11.3.4 6-lead ECG monitoring/12-lead ECG recording and assessment

Fault	Cause	Remedy
The Plug in ECG connector		Check plug connection.
alarm occurs although the ECG	Device not detecting connected	Change ECG cable.
cable is connected to the device		Have the device repaired.
The ECG cable is not connected to	Device incorrectly detecting ECG cable as connected	Have the device repaired.
the device but the Plug in ECG connector alarm does not occur	The alarm only occurs if an ECG has already been successfully derived since switching on	-
The Check seating of ECG electrodes alarm occurs	ECG electrodes not attached correctly to the patient	Attach ECG electrodes to the patient correctly (see 6.10.1, p. 171).
although the ECG electrodes are attached to the patient		Change ECG cable.
		Have the device repaired.
ECG cable not attached/not	Device incorrectly detecting ECG cable as attached to the patient	Have the device repaired.
attached correctly to the patient, but the Check seating of ECG electrodes alarm does not occur	The alarm only occurs if an ECG has already been successfully derived since switching on	-

11.3.5 Pulse oximetry monitoring

Fault	Cause	Remedy
The Plug in SpO₂ sensor alarm occurs although the pulse oximetry	Device does not detect connected	Check plug connection.
		Change pulse oximetry sensor connecting cable.
sensor is connected to the device	pulse oximetry sensor	Change pulse oximetry sensor.
		Have the device repaired.
The pulse oximetry sensor is not	Device incorrectly detecting pulse oximetry sensor as connected	Have the device repaired.
connected to the device but the Plug in SpO₂ sensor alarm does not occur	The alarm only occurs if an SpO ₂ signal has already been successfully detected since switching on	-
The Check seating of SpO₂ sensor alarm occurs although the pulse oximetry sensor is attached	Pulse oximetry sensor not attached correctly to the patient	Attach pulse oximetry sensor to the patient correctly (see 6.9.1, p. 164).
		Change pulse oximetry sensor connecting cable.
to the patient		Change pulse oximetry sensor.
		Have the device repaired.
The pulse oximetry sensor not attached/not attached correctly to the patient, but the Check seating of SpO2 sensor alarm does not occur	Device incorrectly detecting pulse oximetry sensor as attached to the patient	Have the device repaired.
	The alarm only occurs if an SpO ₂ signal has already been successfully detected since switching on	-

Fault	Cause	Remedy
		Remove or reduce light source.
		Protect pulse oximetry sensor from light incidence.
	Strong ambient light or direct light, UV light or infrared light	Attach pulse oximetry sensor to another site which is better protected from light.
		Remove the patient and pulse oximetry sensor from the light.
		Cover the pulse oximetry sensor.
Implausible measured values	Intravascular dyes (e.g. methylene blue)	Impairment of the measurement result cannot be remedied. Measures to treat patient based on medical indication.
	Nail varnish, artificial fingernails	 Rotate the pulse oximetry sensor by 90°. Clean finger nail. Select a different suitable measuring point.
	Significant patient movement	Fix pulse oximetry sensor connecting cable in a strain relief loop on patient using adhesive tape.

11.3.6 Non-invasive blood pressure measurement (NIBP monitoring)

Fault	Cause	Remedy
	NIBP cuff attached incorrectly	Reattach NIBP cuff (see 6.12.1, p. 189).
	NIBP cuff leaking	Replace the NIBP cuff.
	Adapter tube for connecting disposable NIBP cuffs not connected correctly	Connect adapter tube for connecting disposable NIBP cuffs correctly.
Implausible measured values	Adapter tube for connecting disposable NIBP cuffs leaking	Replace the adapter tube for connecting disposable NIBP cuffs.
	Unsuitable patient position	Reposition the patient.
	NIBP module defective	Have the device repaired.
	NIBP cuff size not suitable (too large or too small)	Use a suitable NIBP cuff size.
	Device defective	Have the device repaired.
NIBP measurement cannot be started	NIBP cuff and/or NIBP connecting tube not connected correctly	Connect NIBP cuff and/or NIBP connecting tube correctly.
NIBP measurement keeps failing	NIBP cuff is constricted and cannot be inflated	Remedy NIBP cuff constriction.
	Movement results in artifacts	Reduce movements.
	NIBP cuff or NIBP connecting tube damaged	Replace NIBP cuff and/or NIBP connecting tube.
NIBP cuff cannot be fully inflated	Connection of NIBP cuff and/or NIBP connecting tube interrupted	Connect NIBP cuff and/or NIBP connecting tube.
	Adapter tube for connecting disposable NIBP cuffs not connected correctly	Connect adapter tube for connecting disposable NIBP cuffs correctly.
	Device defective	Have the device repaired.

11.3.7 Printing

Fault	Cause	Remedy
Printer not connecting to the	E-mail delivery 12-lead ECG or network data transmission occurring in parallel	Wait until e-mail delivery 12-lead ECG or network data transmission is finished. If necessary: Switch the printer on again.
device	Poor reception quality	Put printer within range of the device.
	Feed button (FEED) pressed for longer than 10 s	Do not press feed button (FEED) for longer than 10 s.
Connection between device and printer fails	Description	Put printer within range of the
Connection between printer and device is extremely slow	Poor reception quality	device.
Special characters in printout or	Electromagnetic radiation in airplane mode	Do not use the printer in airplane mode.
printer fails in airplane mode	Printer is at the limit of device range	Reduce the distance between device and printer.

11.3.8 Data transmission

Fault	Cause	Remedy
	WiFi is deactivated	Activate network (see 12.3.1, p. 303).
	Airplane mode is activated	Deactivate airplane mode (see 10.3.6, p. 271).
		Bring device within range of the network. If necessary, activate a mobile network.
	Unable to connect network	Configure default network (see 12.3, p. 302)
E mail with 12 load ECC door not		or configure default network (see 4.8, p. 103).
reach recipient		Check whether network is connected to the Internet.
		Check whether port 443 is enabled.
		Request connection data for the network from the system administrator.
	Device not registered with WEINMANN Connect	Register the device with WEINMANN Connect (see 12.4, p. 308).
	E-mail has ended up in the recipient's spam folder	Contact system administrator.

Fault	Cause	Remedy
	WiFi is deactivated	Activate network (see 12.3.1, p. 303).
	Airplane mode is activated	Deactivate airplane mode (see 10.3.6, p. 271).
	Unable to connect network	Bring device within range of the network. If necessary, activate a mobile network.
		Configure default network (see 12.3, p. 302)
		or
Function check is not uploaded to WEINMANN Connect		configure default network (see 4.8, p. 103).
		Check whether network is connected to the Internet.
		Check whether port 443 is enabled.
		Request connection data for the network from the system administrator.
	Device not registered with WEINMANN Connect	Register the device with WEINMANN Connect (see 12.4, p. 308).

11 Alarms and faults

Fault	Cause	Remedy
	WiFi is deactivated	Activate network (see 12.3.1, p. 303).
	Airplane mode is activated	Deactivate airplane mode (see 10.3.6, p. 271).
		Bring device within range of the network. If necessary, activate a mobile network.
		Configure default network (see 12.3, p. 302)
Session data are not uploaded to WEINMANN Connect	Unable to connect network	or configure default network (see 4.8, p. 103).
		Check whether network is connected to the Internet.
		Check whether port 443 is enabled.
		Request connection data for the network from the system administrator.
	Device not registered with WEINMANN Connect	Register the device with WEINMANN Connect (see 12.4, p. 308).
	WiFi is deactivated	Activate network (see 12.3.1, p. 303).
	Airplane mode is activated	Deactivate airplane mode (see 10.3.6, p. 271).
		Bring device within range of the network. If necessary, activate a mobile network.
	Unable to connect network	Configure default network.
Service data are not uploaded to WEINMANN Emergency		Check whether network is connected to the Internet.
		Check whether port 443 is enabled.
		Request connection data for the network from the system administrator.
	Device not registered with WEINMANN Connect	Register the device with WEINMANN Connect (see 12.4, p. 308).

Fault	Cause	Remedy	
E-mail with 12-lead ECG does not reach the recipient, or does so with a severe delay	E-mail with 12-lead ECG blocked/	Add sender's address to list of safe senders. If necessary: Contact system administrator.	
	delayed by recipient's server		
E-mail does not contain a 12-lead ECG in the attachment	E-mail with 12-lead ECG blocked/ delayed by recipient's server	Add sender's address to list of safe senders. If necessary: Contact system administrator.	

12 Configuration by the operator

12.1 Configuring the device

WARNING

Risk of injury due to incorrectly set parameters or too few/too many enabled functions in the operator menu!

Incorrectly set parameters or too few/too many enabled functions in the operator menu may result in incorrect settings in the user menu or too limited/too extensive device functions. This may cause critical operating situations and injure the patient.

- ⇒ The operator menu should only be used by an operator familiar with the settings in the operator menu and their impact on the user menu and device functions.
- \Rightarrow Adapt the device functions to the user's know-how.
- \Rightarrow Protect the operator menu with a secure access code (at least one letter and one number).
- 1. Activate operator menu (see "13.1 Activating the operator menu", page 322).
- 2. Check whether all the required options have been enabled (see "12.2 Enabling options", page 301).
- 3. Configure default wireless network (see "12.3 Configuring default network", page 302).
- Register the device with WEINMANN Connect (see "12.4 Registering the device with WEINMANN Connect", page 308).
- Set date and time (see "12.5 Setting the date and time", page 309).
- Configure e-mail delivery 12-lead ECG (only with E-mail delivery 12-lead ECG option) (see "12.6 Configuring e-mail delivery 12-lead ECG (only with E-mail delivery 12-lead ECG option)", page 311).
- Pair printer with device (only with Printing option) (see "12.7 Pair printer with device (only with Printing option)", page 314).

- Pair device with external device via the Bluetooth[®] interface (only with Bluetooth[®] data transmission option) (see "12.8 Pairing the device with an external device via the Bluetooth[®] interface (only with Bluetooth[®] data transmission option)", page 316).
- 9. Make further settings in the operator menu.
- 10. Change operator menu access code (see "13.12 System settings", page 358).
- *Result* The device is ready for use.

12.2 Enabling options

- *Requirement* There is an enable code for a new option.
 - 1. Activate the operator menu (see "13.1 Activating the operator menu", page 322).
 - 2. Select System settings | Enable options | Enter enable code.
 - Enter the enable code for the new option. The new option appears as a selection under System settings | Enable options.
 - 4. Activate/deactivate a new option with a check mark.
 - *Result* A new option is enabled for use and activated/deactivated.

12.3 Configuring default network

A WARNING

Risk of injury due to delayed delivery!

The user can deliver a 12-lead ECG to a teleconsultation service for further diagnosis. Delivering a 12-lead ECG is purely to obtain a second opinion, and does not replace adequately trained specialists at the session location. Specialist staff at the session location are responsible for treatment within the scope of their training and in accordance with the specifications of the operator. If the 12-lead ECG reaches the teleconsultation service very late or not at all for technical reasons, delayed treatment may result in a potentially life-threatening situation. The operator is responsible for ensuring that the necessary configuration for 12-lead ECG delivery has been carried out correctly.

- ⇒ When configuring e-mail delivery 12-lead ECG: Test e-mail delivery.
- \Rightarrow If the IT infrastructure permits it: Configure several networks for various devices or session locations so that if one network is unavailable, it is possible to select another network.
- \Rightarrow Check the configuration of the networks at regular intervals and update them if necessary.
- \Rightarrow Ensure that the e-mail addresses are correct and complete.
- \Rightarrow Check e-mail addresses at regular intervals and update them if necessary.

12.3.1 Activating the WiFi interface

- 1. Activate the operator menu (see "13.1 Activating the operator menu", page 322).
- 2. Select System settings | Regional settings | WLAN region.

WLAN region		
AD - Andorra		0
AE - United Arab Emirates (the)		0
AF - Afghanistan		0
AI - Anguilla	_	0
AL - Albania	_	0
AM - Armenia		0
AN - Netherlands Antilles		0
AR - Argentina	_	0
AS - American Samoa		0
AT - Austria		0
AU - Australia		0
	Back	ОК

- 3. Select network region.
- 4. Select the **Back** function button 2x.
- 5. Select Communication settings | Configure wireless network.
- 6. Select the **Activate wireless network** menu item.
- *Result* The WiFi interface of the device has been activated.

12.3.2 Add network automatically

Requirement The WiFi interface of the device has been activated (see "12.3.1 Activating the WiFi interface", page 303).

Configure wireless network		
Activate wireless network		冈
Add wireless network automatic	ally	
Add wireless network manually		
Edit wireless network	_	
Allow wireless network configur	ration	冈
Back		
	Back	ок
	Back	ок

 Select the Add wireless network automatically menu item. When doing so, please note: Some smartphones automatically switch off the WiFi hotspot. In order to ensure that the device can use the smartphone's hotspot, the dialog for switching on the WiFi hotspot must be open in the smartphone whilst the device sets up the WiFi connection.

All the networks in range of the device are displayed.

New risks from integrating the device in an IT network! A WARNING When integrating the device in an IT network, new risks may arise for patients, users or third parties which need to be determined, analyzed, and managed. \Rightarrow As the operator of the IT network, reassess changes to the IT network. When doing so, please note: Modified IT network configuration Connection of additional elements to the IT network Removal of additional elements from the IT network Updates of devices connected to the IT network Upgrades of devices connected to the IT network Delayed therapy due to integrating the device in a network with a login page! If the device is integrated in a network with a login page, it is not possible to upload a 12-lead ECG by e-mail.

 \Rightarrow Do not use a network with a login page.

- Select the desired network. When doing so, please note: The device must only be connected to an encrypted network. A password always has to be entered on an encrypted network.
- 3. Confirm with the **OK** function button.
- 4. Enter the password for the desired network. The device connects to the desired network.
- 5. In the event of problems setting up the network: Contact system administrator.
- *Result* A network has been configured for the device automatically and specified as the default network.

From now on, the device always automatically sets up a connection to the default network set up when a WiFi connection is required. Following successful data transmission, the WiFi connection is disconnected.

12.3.3 Adding network manually

- The WiFi interface of the device has been activated (see "12.3.1 Activating the WiFi interface", page 303).
 - The network configuration of the network to be added manually is known.

Configure wireless	network	
Activate wireless network		R
Add wireless network automatic	ally	
Add wireless network manually		
Edit wireless network		
Allow wireless network configur	ation	$\mathbf{\nabla}$
Back		
	Back	ОК

1. Select the Add wireless network manually menu item.

Add wireless network manually		
Network name (SSID)	WM	_
Password	****	
DHCP	_	
IP address		
Subnet mask		
Gateway		
Primary DNS server		
Secondary DNS server		
Add wireless network		
Back		
	Back	ОК

- 2. Make the settings for the network.
- 3. Select the Add wireless network menu item.
- Confirm with the **OK** function button. The network is added as an available network.
- *Result* A network has been manually configured for the device.

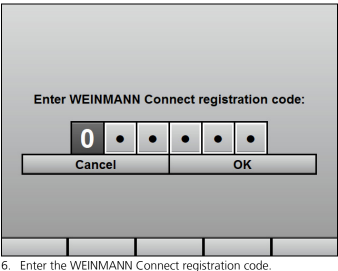
12.3.4 Specifying a network as the default network

- The WiFi interface of the device has been activated (see "12.3.1 Activating the WiFi interface", page 303).
 - Several networks have been configured.
 - 1. Select the **Edit wireless network** menu item. All configured networks are displayed.
 - 2. Select a network as the default network to be configured.
 - 3. Select the **OK** function button.
 - Select the Set wireless network as default menu item. The desired network is specified as the default network.
 - 5. Restart device.
 - *Result* The selected network has been specified as the default network.

12.4 Registering the device with WEINMANN Connect

Requirement A default network has been configured (see "12.3 Configuring default network", page 302).

- 1. Apply to WEINMANN Emergency Customer Service for a user account for WEINMANN Connect by telephone or by e-mail.
- 2. Log in to WEINMANN Connect at www.weinmannconnect.com using the user account data (in accordance with the WEINMANN Connect user manual).
- Add a new device to WEINMANN Connect in accordance with the WEINMANN Connect user manual. When doing so, please note: A WEINMANN Connect registration code is generated for the device.
- 4. Activate the operator menu (see "13.1 Activating the operator menu", page 322).
- 5. Select Communication settings | Register device with WEINMANN Connect.



Enter the WEINMANN Connect registration code. The device is registered with WEINMANN Connect. WM 68401b 2024-03

- Upon successful registration: Upload service data to WEINMANN Connect with the OK function button.
- *Result* The device has been registered with WEINMANN Connect and the service data have been uploaded to WEINMANN Connect. The device can upload data to WEINMANN Connect with immediate effect.

12.5 Setting the date and time

12.5.1 Setting time zone

- 1. Activate operator menu (see "13.1 Activating the operator menu", page 322).
- 2. Select System settings | Regional settings.
- Activate the Automatic clock change menu item. The Time zone preselection and Time zone menu items are displayed. The device can now automatically switch from winter time to summer time and vice versa.
- 4. Select the **Time zone preselection** menu item. A list of possible time zone areas is displayed.
- 5. Select the time zone area for the device.
- 6. Confirm with the **OK** function button.
- 7. Select the **Back** function button.
- Select the **Time zone** menu item. A list of all the time zones in the selected time zone area is displayed.
- 9. Select the time zone for the device.
- 10. Confirm with the **OK** function button.
- *Result* A time zone has been set for the device. The device automatically switches from winter time to summer time and vice versa.

12.5.2 Updating the date and time automatically

- Prerequisite
 A time zone has been set for the device (see "12.5.1 Setting time zone", page 309).
 - The device has been registered with WEINMANN Connect (see "12.4 Registering the device with WEINMANN Connect", page 308).
 - The operator menu is activated (see "13.1 Activating the operator menu", page 322).
 - 1. Select Communication settings | Date and time update.
 - 2. Select the **Update date/time automatically** menu item.
 - *Result* During each function check, the device establishes a connection to WEINMANN Connect and automatically synchronizes the date and time with WEINMANN Connect.

12.5.3 Updating the date and time manually

- A time zone has been set for the device (see "12.5.1 Setting time zone", page 309).
- The device has been registered with WEINMANN Connect (see "12.4 Registering the device with WEINMANN Connect", page 308).
- The operator menu is activated (see "13.1 Activating the operator menu", page 322).
- 1. Select Communication settings | Date and time update.
- Select the Update date/time manually menu item.
 All the wireless networks in range of the device are displayed.
- 3. Select the desired wireless network.
- Press the **OK** function button. The device connects to the desired wireless network. Date and time are updated.
- *Result* Date and time have been updated manually.

12.6 Configuring e-mail delivery 12-lead ECG (only with E-mail delivery 12-lead ECG option)

A WARNING

Risk of injury due to delayed delivery!

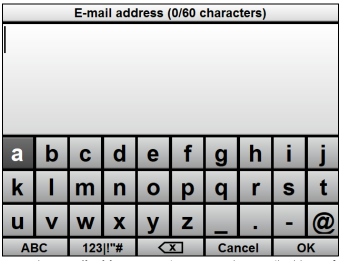
The user can deliver a 12-lead ECG to a teleconsultation service for further diagnosis. Delivering a 12-lead ECG is purely to obtain a second opinion, and does not replace adequately trained specialists at the session location. Specialist staff at the session location are responsible for treatment within the scope of their training and in accordance with the specifications of the operator. If the 12-lead ECG reaches the teleconsultation service very late or not at all for technical reasons, delayed treatment may result in a potentially life-threatening situation. The operator is responsible for ensuring that the necessary configuration for 12-lead ECG delivery has been carried out correctly.

- ⇒ When configuring e-mail delivery 12-lead ECG: Test e-mail delivery.
- \Rightarrow If the IT infrastructure permits it: Configure several networks for various devices or session locations so that if one network is unavailable, it is possible to select another network.
- ⇒ Check the configuration of the networks at regular intervals and update them if necessary.
- \Rightarrow Ensure that the e-mail addresses are correct and complete.
- ⇒ Check e-mail addresses at regular intervals and update them if necessary.

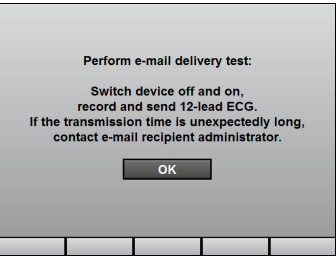
The 12-lead ECG option is enabled and activated (see "13.12 System settings", page 358).

- The E-mail delivery 12-lead ECG option is enabled and activated (see "13.12 System settings", page 358).
- A default network has been configured (see "12.3 Configuring default network", page 302).

- The device has been registered with WEINMANN Connect (see "12.4 Registering the device with WEINMANN Connect", page 308).
- 1. Activate the operator menu (see "13.1 Activating the operator menu", page 322).
- 2. Select Communication settings | E-mail delivery 12-lead ECG.
- 3. Select Contacts | New entry.
- 4. In the **Recipient name** menu item, enter the name of the recipient.
- 5. Confirm with the **OK** function button.



- 6. In the **E-mail address** menu item, enter the e-mail address of the recipient using the navigation knob.
- 7. Confirm with the **OK** function button.



8. Test e-mail delivery in accordance with the instructions in the display.

When doing so, please note: The duration for setting up the connection and delivering the e-mail may vary depending on reception quality.

- 9. Confirm with OK.
- 10. If the e-mail delivery was not successful: Check the setup of the network and e-mail delivery 12-lead ECG.
- 11. If the e-mail is delivered successfully but transmission time is unexpectedly long: Contact the system administrator of the email recipient.
- 12. Select the **Back** function button 2x.
- 13. Make the following settings for the 12-lead ECG to be delivered:
 - Feed rate and duration
 - Amplitude scaling
 - View
- *Result* E-mail delivery 12-lead ECG has been configured.

12.7 Pair printer with device (only with Printing option)

A charged printer battery is inserted (see "4.7.2 Inserting the printer battery", page 97).

- The printer paper is inserted (see "4.7.3 Inserting printer paper", page 98).
- The Printing option is enabled and activated (see "13.12 System settings", page 358).
- Press the On/Off button U of the printer for > 5 s. The status indicator of the printer is green and an audible signal sounds.
- Switch on the device (see "6.1 Switching on the device", page 125). The start menu appears.
- 3. Activate the operator menu (see "13.1 Activating the operator menu", page 322).
- 4. Select Printer settings | Pair printer.
- 5. Position the printer next to the device.
- Press the **OK** function button. The device displays printers available in the vicinity.
- 7. If necessary: Cancel printer search and connection setup by pressing the **Cancel** function button.
- 8. If several printers are available: Briefly press the On/Off button
 O of the printer.
 The MAC address of the printer is shown in the printer display.
- 9. To identify the printer to be paired: Compare the MAC address in the printer display with the MAC addresses shown in the device.

S	Select printe	r	
00:15:AA:BB:CC:DD ("WOOSIM")			
		Cancel	OK

- 10. Select the printer marked "WOOSIM" and the correct MAC address.
- 11. Check whether the PIN shown in the device display is shown in the printer display.
- 12. If the PIN is shown in the printer display: Confirm PIN with the printer's feed button (FEED).
- 13. Confirm the PIN in the device display with the OK function button.The connection to the printer is set up. The printer automatically prints a printer connection log.
- 14. Check the printer connection log and confirm successful printout with the **OK** function button.
- 15. Exit operator menu.
- *Result* A printer has been paired with the device. From now on, the device will always set up a connection to this printer automatically when the device and printer are switched on and are within range.

12.8 Pairing the device with an external device via the Bluetooth[®] interface (only with Bluetooth[®] data transmission option)

- Requirement
 The Bluetooth[®] data transmission option is enabled and activated (see "13.12 System settings", page 358).
 - An external device with a Bluetooth[®] interface (Example: System for digital patient data recording) is within range.
 - 1. Activate the operator menu (see "13.1 Activating the operator menu", page 322).
 - Select Communication settings | Pair Bluetooth[®] devices. The device waits to pair with an external device.
 - 3. Activate the Bluetooth[®] interface of the external device and search for available devices.
 - 4. Select MEDUCORE Standard² on the external device.
 - 5. Check whether the PIN displayed on the device is shown on the external device.
 - If the PINs displayed on the device and on the external device are identical: Confirm pairing on device and external device. Device and external device are paired.
 - *Result* The device and the external device are paired via the Bluetooth[®] interface. The external device can connect to the device and interrogate device data via the Bluetooth[®] interface.

12.9 Importing/exporting configurations

There are two options for exporting and importing a configuration (all settings in the operator menu):

• With the aid of an SD card (see "12.9.1 Exporting a configuration to an SD card/importing a configuration from an SD card", page 317)

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• With the aid of WEINMANN Connect (see "12.9.2 Exporting a configuration to WEINMANN Connect/importing a configuration from WEINMANN Connect", page 318)

12.9.1 Exporting a configuration to an SD card/ importing a configuration from an SD card

A configuration exported to an SD card can be imported into another device. If the transfer is made with the aid of an SD card, only the configuration is transferred. Contacts must be transferred separately (only with E-mail delivery 12-lead ECG option).

A WARNING

Risk of injury from different alarm presets in the same or similar devices!

Different alarm presets in the same or similar devices in different application areas may confuse the user and result in injury to the patient.

 \Rightarrow Select the same alarm presets in the same or similar devices.

Exporting configuration to an SD card

- *Prerequisite* An SD card is inserted in the device (see "4.6.1 Inserting an SD card", page 92).
 - 1. Activate operator menu (see "13.1 Activating the operator menu", page 322).
 - 2. Select System settings | Import/export configurations | Export configuration to SD card.
 - 3. Confirm with the **OK** function button. The export process starts.
 - When the export process is finished: Confirm with the **OK** function button.
 - 5. Remove the SD card (see "4.6.2 Removing the SD card", page 93).
 - *Result* The configuration has been exported to an SD card.

Importing configuration from an SD card

Prerequisite An SD card with the configuration of the same or another device is inserted in the device (see "4.6.1 Inserting an SD card", page 92).

- 1. Activate operator menu (see "13.1 Activating the operator menu", page 322).
- 2. Select System settings | Import/export configurations | Import configuration from SD card.
- 3. Confirm with the **OK** function button. The import process starts.
- When the import process is finished: Confirm with the **OK** function button.
- *Result* The configuration of the same or another device has been imported from an SD card.

12.9.2 Exporting a configuration to WEINMANN Connect/importing a configuration from WEINMANN Connect

It is only possible to transfer a configuration between the device and WEINMANN Connect from WEINMANN Connect version 3.0 upward. During a transfer between the device and WEINMANN Connect, contacts are also transferred simultaneously (only with E-mail delivery 12-lead ECG option).

Exporting configuration to WEINMANN Connect

- 1. Activate operator menu (see "13.1 Activating the operator menu", page 322).
- Select System settings | Import/export configurations | Export configuration to WEINMANN Connect.
 All the wireless networks in range of the device are displayed.
- 3. Select the desired wireless network.
- 4. Confirm with the **OK** function button. The export process starts.

- 5. When the export process is finished: Confirm with the **OK** function button.
- *Result* The configuration and the contacts (only with E-mail delivery 12-lead ECG option) have been exported to WEINMANN Connect.

Importing configuration from WEINMANN Connect

- 1. Activate operator menu (see "13.1 Activating the operator menu", page 322).
- Select System settings | Import/export configurations | Import configuration from WEINMANN Connect.
 All the wireless networks in range of the device are displayed.
- 3. Select the desired wireless network.
- 4. Confirm with the **OK** function button. The import process starts.
- 5. When the import process is finished: Confirm with the **OK** function button.
- *Result* The configuration and the contacts (only with E-mail delivery 12-lead ECG) have been imported from WEINMANN Connect.

12.10 Carrying out a software update

Requirement

- A charged battery is inserted in the device.
- The device is connected to line power.
- The device is switched off and an SD card with new software is in the SD card slot.



Risk of injury due to lack of function check after software update!

If no function check is carried out following a software update, errors occurring during the software update may go undetected. This may cause the patient or the user serious or life-threatening injury.

 \Rightarrow Carry out a function check after every software update.

A WARNING

Failure of therapy due to failure of the battery and lack of line power!

If the power supply to the device is cut during the software update due to a battery failure and lack of line power, the device may be damaged to such an extent that it needs to be repaired.

- ⇒ Always connect the device to line power with a charged battery for software updates.
- 1. If the software is provided as a ZIP file: Unzip the software.
- Place the file in the SD card's root directory. When doing so, please note: The file must not be in a sub-folder.
- 3. Activate the operator menu (see "13.1 Activating the operator menu", page 322).
- 4. Select System settings | Software update.
- 5. Select new software with the navigation knob.
- 6. Run the software update with the **Start** function button.
- Wait until the software update has finished. After the end of the software update, the device displays the message SUCCESS.
- 8. Restart the device with the **Restart** function button. The device restarts.
- 9. Activate the operator menu (see "13.1 Activating the operator menu", page 322).
- 10. Select **Device information | Device**. The device displays the installed software version.
- 11. Press and hold the On/Off button (1) for at least 2 s to switch off the device and save the settings.
- 12. Carry out a function check (see "5.2 Carrying out a function check", page 109).
- 13. If necessary: Set the date and time (see "10.3.7 System settings", page 273).
- *Result* A software update has been carried out.

13 Operator menu

A WARNING

Risk of injury due to incorrectly set parameters or too few/too many enabled functions in the operator menu!

Incorrectly set parameters or too few/too many enabled functions in the operator menu may result in incorrect settings in the user menu or too limited/too extensive device functions. This may cause critical operating situations and injure the patient.

- ⇒ The operator menu should only be used by an operator familiar with the settings in the operator menu and their impact on the user menu and device functions.
- \Rightarrow Adapt the device functions to the user's know-how.
- \Rightarrow Protect the operator menu with a secure access code (at least one letter and one number).

WARNING

Risk of injury from incorrectly secured access to functions!

Too simple an access code gives the user access to functions with which he or she is not familiar or in which he or she is not trained. This may cause the patient or the user serious or life-threatening injury.

 \Rightarrow Protect functions with a secure access code (at least one letter and one number).

The operator menu contains device presets which are permanently stored. If the device was switched off for longer than 30 seconds and is switched on again, the operator menu device presets are loaded. If the device was switched off for less than 30 seconds and a patient was previously connected, the user menu settings are restored.

The operator menu also contains functions relevant solely to the operator and not to the user.

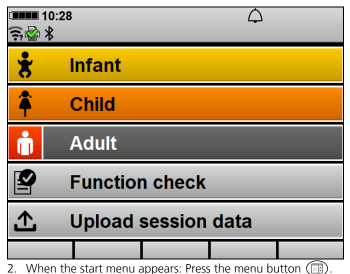
Examples:

- Enable options (see "12.2 Enabling options", page 301)
- Configure default network (see "12.3 Configuring default network", page 302)
- Configure e-mail delivery 12-lead ECG (see "12.6 Configuring e-mail delivery 12-lead ECG (only with E-mail delivery 12-lead ECG option)", page 311)
- Carry out software updates (see "12.10 Carrying out a software update", page 319)
- Disable functions

13.1 Activating the operator menu

Requirement A battery with a battery status of at least 2 LEDs is inserted in the device.

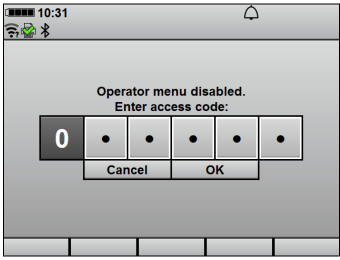
1. Switch on the device. The start menu appears:



3. If the start menu disappears before the menu button is pressed and the device switches to normal mode: Restart device.



The operator menu is protected by an access code. When the device is supplied, this code is 000000. WEINMANN Emergency urgently recommends changing this access code when the device is put into operation. As long as the access code has not been changed, the **Update operator menu access code** alarm is outputted for 10 seconds.



 Select the access code for the operator menu using the navigation knob.
 The operator menu appears:

The operator menu appears:

Operator mer	าน	
AED settings		
Alarm settings		
Manual mode settings	_	_
ECG settings		
SpO ₂ settings		
NIBP settings		
Printer settings		
12-lead ECG settings		
Communication settings		
System settings		
Device information		
	Back	ОК

5. If any digit is entered incorrectly: Wait 5 s.

Alternatively:

Fill all fields with values and select Cancel.

Alternatively:

When input of all numerals is complete, select an individual digit and correct it.

Result The operator menu is activated.

13.2 Navigating the operator menu

Requirement The operator menu is activated (see "13.1 Activating the operator menu", page 322).

- 1. Select setting using the navigation knob.
- 2. Change setting using the navigation knob.
- To exit the menu: Select the **Back** function button.

Alternatively:

Press the menu button 🗐.

- 4. Restart device.
- *Result* Settings have been made and apply to all sessions.

13.3 AED settings

You can make presets for AED mode in the **AED settings** submenu.

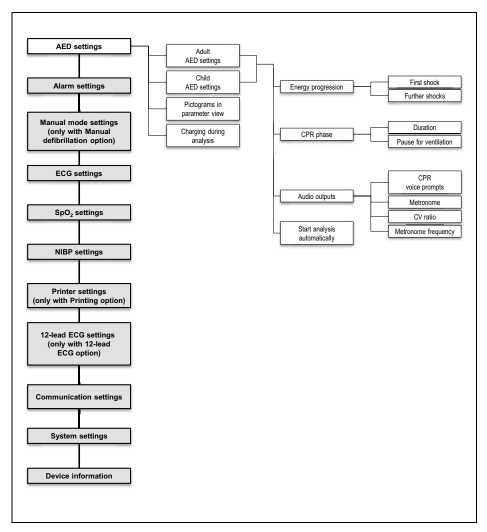
The factory settings are selected to meet currently applicable ERC guidelines. Adjustments to AED settings may result in the user no longer being supported in compliance with currently applicable ERC guidelines.

The AED settings enable you to adapt the device to the users' qualification level and to provide optimal support to users during resuscitation measures whilst taking regional features into account.

AED setting	s	
Adult AED settings		
Child AED settings		
Pictograms in parameter view		
Charging during analysis		
Back		
	Back	ОК

13-1 AED settings submenu





13-2 Operator menu: AED settings submenu

Parameter		Possible values	Description	Factory setting		
Adult AED se	Adult AED settings/Child AED settings					
Energy	First shock	- 1 J to 200 J	Here you can set the shock energy of the first shock in AED mode.	 AED mode adult: 150 J AED mode child: 75 J 		
progression	Further shocks	13 10 200 3	Here you can set the shock energy of further shocks in AED mode.	 AED mode adult: 200 J AED mode child: 75 J 		
	Duration	120 s to 300 s	Here you can set the time interval between cardiac rhythm analyses. The device only outputs certain AED instructions and voice prompts after CPR is complete, as complete and correct carrying out of CPR takes priority.	120 s		
CPR phase	Pause for ventilation	3 s to 8 s	 Here you can set the time interval for ventilation between chest compressions. When choosing a setting, consider whether voice prompts are deactivated: When voice prompts are activated, the pause begins after the last metronome sound and ends when the voice prompts begin. When voice prompts are deactivated, the pause begins after the last metronome sound and ends when the last metronome sound and ends when the first metronome sound begins. 	5 s		

13.3.2 Possible values

Parameter		Possible values	Description	Factory setting
Adult AED set	tings/Child AED	settings		
	CPR voice prompts	Activated C Deactivated	Here you can set whether the voice prompts for guiding chest compressions and for ventilation are to be given. The voice prompt <i>Ventilate</i> <i>twice</i> is only given if the metronome is set to the rhythm 15:2/30:2 .	Deactivated
	Metronome	off 15:2/30:2	Here you can set the metronome rhythm.	15:2/30:2
Audio outputs	CV ratio	30:2 15:2	 Here you can set the compression/ventilation ratio. 15:2: After 15 metronome beats, there is a pause to allow for ventilation. 30:2: After 30 metronome beats there is a pause to allow for ventilation. 	 AED mode adult: 30:2 AED mode child: 15:2
	Metronome frequency	100/min to 120/min	Here you can set the metronome frequency.	100/min

Parameter		Possible values	Description	Factory setting
Adult AED sett	ings/Child AED	settings		
Start analysis automatically	-	Activated Deactivated	Here you can set whether cardiac rhythm analysis starts automatically, or manually at the touch of a button.	Activated
Pictograms in parameter view	-	Activated Deactivated	Here you can set whether pictograms instead of parameters are displayed in parameter view in AED mode.	Deactivated
Charging during analysis	-	Activated M Deactivated	Here you can set whether the shock capacitor is to be charged at the same time as cardiac rhythm analysis, and so independently of the analysis result. If this setting is deactivated, the lifetime of the shock capacitor is extended.	Activated

13.4 Alarm settings

A WARNING

A WARNING

Risk of injury due to alarm limits which are too high or too low!

Alarm limits which are either too high or too low may prevent the device triggering an alarm, thereby putting the patient at risk.

⇒ Always set alarm limits which have been adapted to the patient.

Risk of injury from different alarm presets in the same or similar devices!

Different alarm presets in the same or similar devices in different application areas may confuse the user and result in injury to the patient.

 \Rightarrow Select the same alarm presets in the same or similar devices.

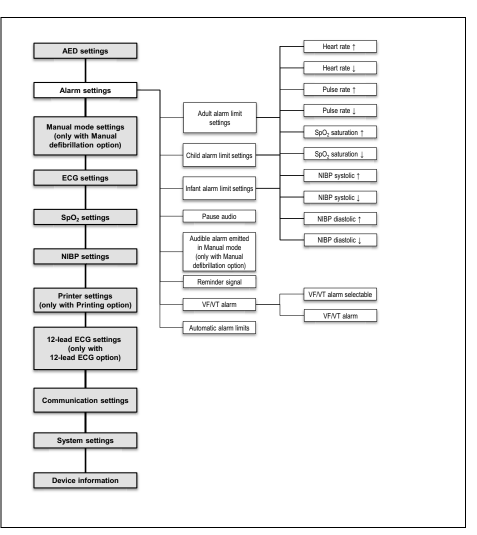
You can make presets for the alarm system in the **Alarm settings** submenu.

The factory settings are selected to meet the guidelines required by the standards and achieve a high level of safety. Changes to the alarm settings may lead to an increased risk for the user and the patient, for example if the alarm limit range is expanded or if **Pause audio** is extended.

You can adapt the device to the patient group and the application area using the relevant alarm settings.

Alarm settings				
Adult alarm limit settings				
Child alarm limit settings				
Infant alarm limit settings				
Pause audio		2 min		
Audible alarm emitted in Manual	mode			
Reminder signal		2 min		
VF/VT alarm				
Automatic alarm limits	_	20 %		
Back				
	Back	ОК		

13-3 Alarm settings submenu



13.4.1 Menu structure

13-4 Operator menu: Alarm settings submenu

13.4.2 Possible values

Parameter		Possible values	Description	Factory setting
	Heart rate †	35/min - 250/min, in increments of 5	Here you can set the upper (†) and lower (↓) limit	120/min
	Heart rate ↓	30/min - 245/min, in increments of 5		50/min
	Pulse rate 1	35/min - 250/min, in increments of 5	values for the patient group as of which the device is to emit an alarm.	120/min
	Pulse rate ↓	30/min - 245/min, in increments of 5	According to the values set, the setting ranges of the upper and lower limit values	50/min
	SpO_2 saturation f	66 % - 100 %		100 %
	SpO ₂ saturation \downarrow 65 % - 99 % depend on each other as		85 %	
Adult alarm limit settings	NIBP systolic 1	45 mmHg - 260 mmHg, in increments of 5	 least one setting value above the set value for the lower alarm limit. The setting range for the lower alarm limit ends at least one setting value below the set value for the upper alarm limit. 	220 mmHg
-	NIBP systolic ↓	40 mmHg - 255 mmHg, in increments of 5		75 mmHg
	NIBP diastolic 1	25 mmHg - 200 mmHg, in increments of 5		110 mmHg
	NIBP diastolic ↓	20 mmHg - 195 mmHg, in increments of 5		35 mmHg

Parameter		Possible values	Description	Factory setting
-	Heart rate †	35/min - 250/min, in increments of 5	 Here you can set the upper (↑) and lower (↓) limit values for the patient group as of which the device is to emit an alarm. According to the values set, the setting ranges of the upper and lower limit values depend on each other as follows: The setting range for the upper alarm limit ends at least one setting value above the set value for the lower alarm limit. The setting range for the lower alarm limit ends at least one setting value above the set value for the lower alarm limit. 	150/min
	Heart rate ↓	30/min - 245/min, in increments of 5		50/min
	Pulse rate 1	35/min - 250/min, in increments of 5		150/min
	Pulse rate ↓	30/min - 245/min, in increments of 5		50/min
	SpO_2 saturation †	66 % - 100 %		100 %
	SpO_2 saturation \downarrow	65 % - 99 %		85 %
Child alarm limit settings	NIBP systolic 1	45 mmHg - 230 mmHg, in increments of 5		145 mmHg
	NIBP systolic ↓	40 mmHg - 225 mmHg, in increments of 5		75 mmHg
	NIBP diastolic †	25 mmHg - 160 mmHg, in increments of 5		100 mmHg
	NIBP diastolic 4	20 mmHg - 155 mmHg, in increments of 5		35 mmHg

13 Operator menu

Parameter		Possible values	Description	Factory setting
	Heart rate 1	35/min - 250/min, in increments of 5		200/min
	Heart rate ↓	30/min - 245/min, in increments of 5	Here you can set the upper (↑) and lower (↓) limit values for the patient group	100/min
	Pulse rate †	35/min - 250/min, in increments of 5	as of which the device is to emit an alarm.	200/min
	Pulse rate ↓	30/min - 245/min, in increments of 5	According to the values set, the setting ranges of the	100/min
	SpO ₂ saturation †	66 % - 100 %	upper and lower limit values	95 %
	SpO_2 saturation \downarrow	65 % - 99 %	depend on each other as	85 %
Infant alarm limit settings	NIBP systolic †	45 mmHg - 130 mmHg, in increments of 5	 follows: The setting range for the upper alarm limit ends at least one setting value above the set value for the lower alarm limit. The setting range for the lower alarm limit ends at least one setting value below the set value for the upper alarm limit. 	100 mmHg
	NIBP systolic ↓	40 mmHg - 125 mmHg, in increments of 5		50 mmHg
	NIBP diastolic 1	25 mmHg - 100 mmHg, in increments of 5		70 mmHg
	NIBP diastolic ↓	20 mmHg - 95 mmHg, in increments of 5		30 mmHg
Pause audio	-	1 min 2 min 5 min 10 min ∞ (infinite)	Here you can set the length of time for which audio alarm output is paused. If you select ∞ (infinite), audio alarm output is paused permanently (audio alarm output is muted).	2 min
Audible alarm emitted in Manual mode (only with Manual defibrillation option)	-	Activated Markov Deactivated	Here you can set whether audio alarm output is active or inactive when the user calls up manual mode.	Deactivated

Parameter		Possible values	Description	Factory setting
Reminder signal	-	Off 1 min 2 min 5 min	Here you can set the time after which a reminder signal is to remind you that audio alarm output is paused or muted.	2 min
VF/VT alarm	VF/VT alarm selectable	Activated Deactivated	Here you can set whether the VF/VT alarm can be activated/deactivated by the user in the user menu.	Deactivated
	VF/VT alarm	Activated Deactivated	Here you can set whether the VF/VT alarm is to be activated when the device starts.	Activated
Automatic alarm limits	-	10 % 20 % 30 %	Here you can set the automatic alarm limits. The device automatically specifies the alarm limits for the physiological alarms. The deviation is 10 %, 20 % or 30 % from the measured values at the time of activation.	20 %

13.5 Manual mode settings (only with Manual defibrillation option)

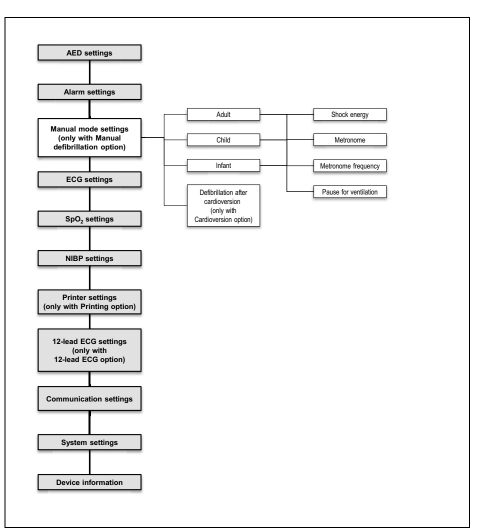
You can make presets for shock energies in the **Manual mode** settings submenu.

The factory settings are selected to meet currently applicable ERC guidelines. Adjustments to manual mode settings may result in the user no longer being supported in compliance with currently applicable ERC guidelines.

You can adapt the device to the patient group using the respective settings for manual mode.

Manual mode settings				
Adult				
Child				
Infant				
Defibrillation after cardioversion	1			
Back				
	Back	ОК		

13-5 Manual mode settings submenu



13.5.1 Menu structure

13-6 Operator menu: Manual mode settings submenu

13.5.2 Possible values

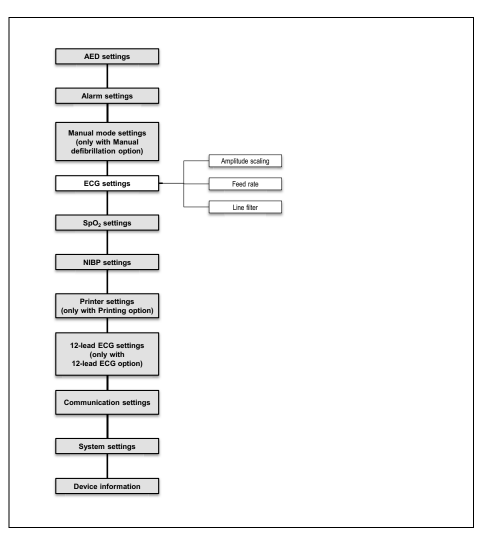
Parameter		Possible values	Description	Factory setting
	Shock energy	1 J to 200 J	Here you can set what defibrillation energy is to be preset in manual mode.	150 J
Adult	Metronome	off 15:2 30:2	Here you can set the metronome rhythm.	off
	Metronome frequency	100/min to 120/min	Here you can set the metronome frequency.	100/min
	Pause for ventilation	3 s to 8 s	Here you can set the time interval for ventilation between chest compressions.	5 s
	Shock energy	1 J to 200 J	Here you can set what defibrillation energy is to be preset in manual mode.	75 J
Child Infant	Metronome	off 15:2 30:2	Here you can set the metronome rhythm.	off
	Metronome frequency	100/min to 120/min	Here you can set the metronome frequency.	100/min
	Pause for ventilation	3 s to 8 s	Here you can set the time interval for ventilation between chest compressions.	5 s
Defibrillation after cardioversion (only with Cardioversion option)	-	Activated C Deactivated	Here you can set whether defibrillation is to follow cardioversion (box checked) or cardioversion is to follow cardioversion (box unchecked).	Deactivated

13.6 ECG settings

In the **ECG settings** submenu, you can make the presets for showing ECG curves in the display.

ECG settings				
Amplitude scaling	Auto	om.		
Feed rate		25 mm/s		
Line filter		$\mathbf{\nabla}$		
Back				
	-	-		
	Back	OK		

13-7 ECG settings submenu



13.6.1 Menu structure



13.6.2 Possible values

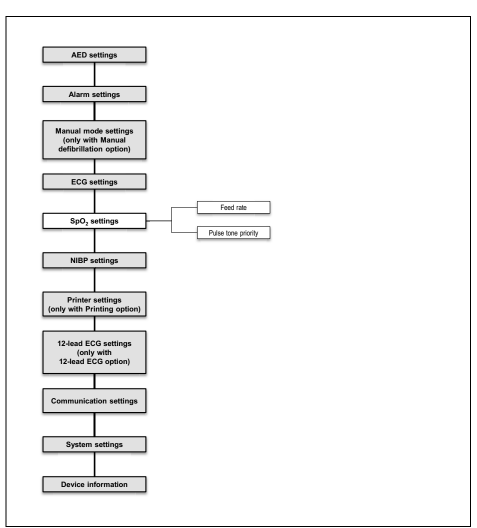
Parameter	Possible values	Description	Factory setting
Amplitude scaling	Autom. 2 mm/mV 5 mm/mV 10 mm/mV 20 mm/mV	Here you can set the amplification of the ECG signal and thus the height of the ECG curve.	Autom.
Feed rate	12.5 mm/s 25 mm/s 50 mm/s	Here you can set the feed rate of the ECG curve display and so change the time resolution.	25 mm/s
Line filter	Activated Deactivated	By activating the line filter, you can reduce ECG interference caused by the power supply network.	Activated

13.7 SpO₂ settings

In the **SpO₂ settings** submenu, you can make the presets for display of the plethysmogram and output of the pulse tone.

SpO ₂ setting	s	
Feed rate		25 mm/s
Pulse tone priority		$\mathbf{\nabla}$
Back		
	Back	ОК

13-9 SpO₂ settings submenu



13.7.1 Menu structure

13-10 Operator menu: SpO₂ settings submenu

13.7.2 Possible values

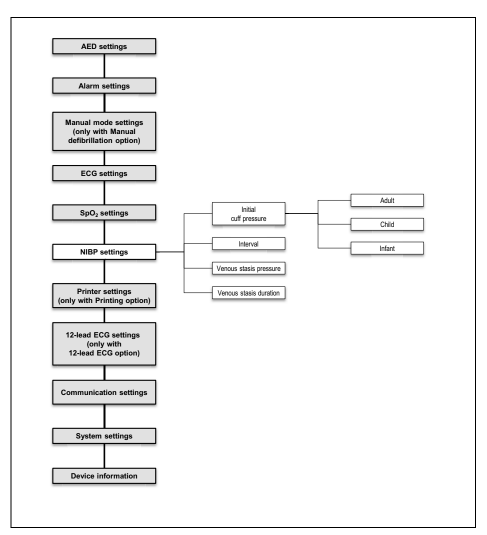
Parameter	Possible values	ossible values Description		
Feed rate 12.5 mm/s 25 mm/s 50 mm/s		Here you can set the feed rate of the SpO_2 curve display and so change the time resolution.	25 mm/s	
Pulse tone priority Deactivated		Here you can set whether pulse tone is to take priority over heart rate tone. The tone level of the pulse tone is adapted to measured oxygen saturation.	Activated	

13.8 NIBP settings

In the **NIBP settings** submenu, you can make the presets for blood pressure measurement and venous stasis. You can adapt the device to the patient group using the specific NIBP settings.

NIBP setting	S	
Initial cuff pressure		
Interval		Off
Venous stasis pressure		80 mmHg
Venous stasis duration		60 s
Back		
	Back	ОК

13-11 NIBP settings submenu



13.8.1 Menu structure

13-12 Operator menu: NIBP settings submenu

13.8.2 Possible values

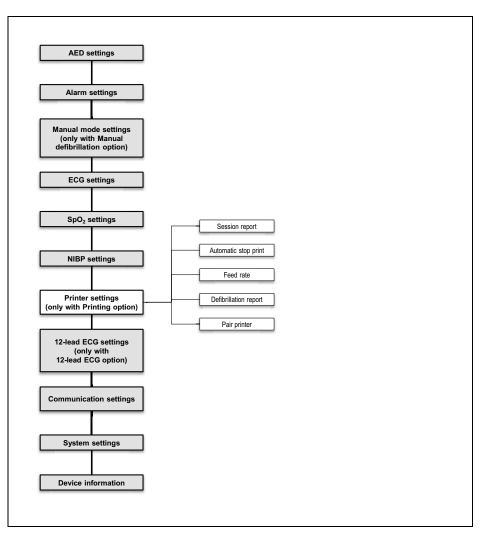
Parameter		Possible values	Description	Factory setting
	Adult	120 mmHg - 280 mmHg, in increments of 5	Here you can set the NIBP cuff pressure to which the device must inflate the NIBP cuff for a particular patient	180 mmHg
Initial cuff pressure	Child	80 mmHg - 170 mmHg, in increments of 5		120 mmHg
	Infant	60 mmHg - 140 mmHg, in increments of 5	group.	90 mmHg
Interval	-	Off 0:30 min 0:45 min 1:00 min 1:15 min 1:30 min 1:45 min 2:00 min 3:00 min 5:00 min 10:00 min 15:00 min 30:00 min 60:00 min	Here you can set after how long another NIBP measurement is to be started automatically.	Off
Venous stasis pressure	-	60 mmHg - 140 mmHg, in increments of 5	Here you can set the pressure to which the NIBP cuff is inflated for venous stasis (IV function button).	80 mmHg
Venous stasis duration	-	10 s - 120 s	Here you can set for venous stasis (IV function button) the time for which pressure is to be maintained in the NIBP cuff.	60 s

13.9 Printer settings (only with Printing option)

In the **Printer settings** submenu, you can make the presets for the printer and connect the device to a printer.

Printer setting	gs	
Session report		
Automatic stop print		10 s
Feed rate		25 mm/s
Defibrillation report	_	$\mathbf{\nabla}$
Pair printer	_	_
Back		
	Back	ОК

13-13 **Printer settings** submenu



13.9.1 Menu structure



13.9.2	Possible values
--------	-----------------

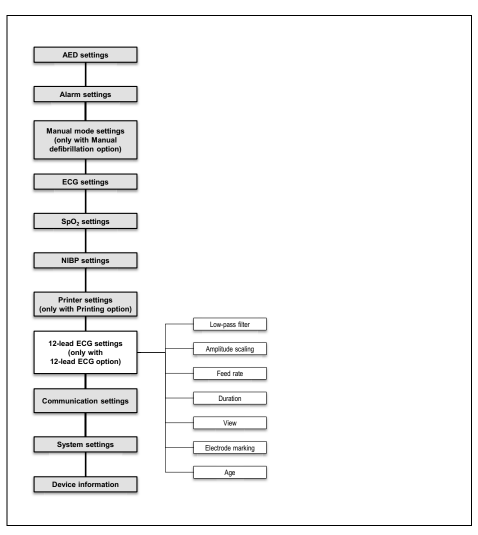
Parameter		Possible values	Description	Factory setting
Session report	-	Resuscitation report Vital signs Events Events and vital signs	Here you can set the details the session report contains.	Resuscitation report Vital signs Events
Automatic stop print	-	5 s to 120 s	Here you can set the time after which printing of a live printout or replay printout finishes automatically.	10 s
Feed rate	-	12.5 mm/s 25 mm/s 50 mm/s	Here you can preset the feed rate at which the printer prints a live printout and a replay printout.	25 mm/s
Defibrillation report	-	Activated Deactivated	Here you can set whether a defibrillation report is printed in the event of defibrillation or cardioversion (only with the Cardioversion option) if the printer is connected to the device and is ready for use.	Activated
Pair printer	-	-	Here you can pair the printer with a device (see "12.7 Pair printer with device (only with Printing option)", page 314). The connection is always set up automatically whenever the device and printer are switched on and are within range.	-

13.10 12-lead ECG settings (only with 12lead ECG option)

You can make presets for the 12-lead ECG in the **12-lead ECG** settings submenu.

12-lead ECG	settings	
Low-pass filter	150 Hz	
Amplitude scaling	10 mm/mV	
Feed rate	50 mm/s	
Duration	5 s	
View Classic		
Electrode marking	ERC	
Age	50	
Back		
	Back OK	

13-15 12-lead ECG settings submenu



13.10.1 Menu structure

13-16 Operator menu: 12-lead ECG settings submenu

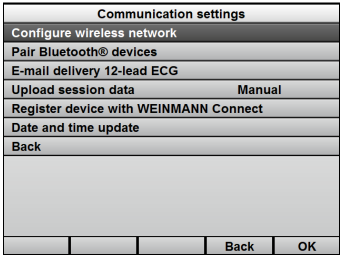
13.10.2 Possible values

Parameter		Possible values	Description	Factory setting
Low-pass filter	-	50 Hz 150 Hz	Here you can set a low-pass filter to filter out artifacts.	150 Hz
Amplitude scaling	-	2 mm/mV 5 mm/mV 10 mm/mV 20 mm/mV	Here you can set the amplification of the ECG signal and thus the height of the ECG curve.	10 mm/mV
Feed rate	-	12.5 mm/s 25 mm/s 50 mm/s	Here you can set the feed rate of the ECG curve display and so change the time resolution.	50 mm/s
Duration	-	5 s 10 s	Here you can set the recording duration of the 12-lead ECG.	5 s
View	-	Classic Cabrera	Here you can set the sequence in which the leads are displayed in the 12-lead ECG.	Classic
Electrode marking	-	ERC AHA	Here you can set whether the positions of the ECG electrodes in 12-lead ECG function mode are named in accordance with European (ERC) or American (AHA) coding and the colors assigned accordingly.	ERC
Age	-	0 to 99 years	Here you can set the patient's age.	50

13.11 Communication settings

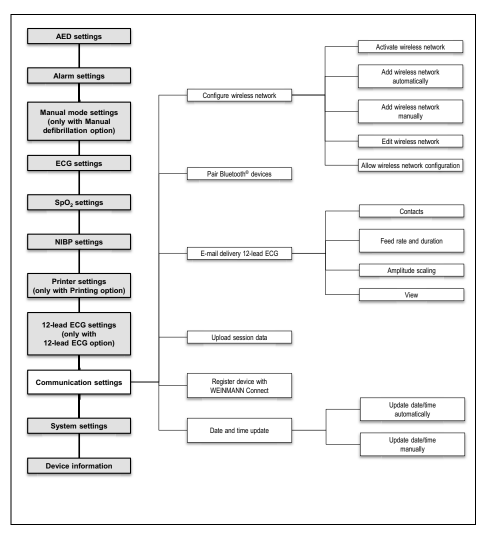
You can make presets for the following functions in the **Communication settings** submenu:

- Configure default network (see "12.3 Configuring default network", page 302)
- Pair the device with an external device via the Bluetooth[®] interface (see "12.8 Pairing the device with an external device via the Bluetooth[®] interface (only with Bluetooth[®] data transmission option)", page 316)
- Deliver 12-lead ECGs by e-mail (see "12.6 Configuring e-mail delivery 12-lead ECG (only with E-mail delivery 12-lead ECG option)", page 311)
- Upload session data to WEINMANN Connect (see "6.21.2 Uploading session data to WEINMANN Connect (only with Session data upload option)", page 221)
- Register the device with WEINMANN Connect (see "12.4 Registering the device with WEINMANN Connect", page 308)
- Set date and time (see "12.5 Setting the date and time", page 309)



13-17 Communication settings submenu





13-18 Operator menu: Communication settings submenu

13.11.2 Possible values

Parameter		Possible values	Description	Factory setting
	Activate wireless network	Activated Deactivated	Here you can activate or deactivate the WiFi interface of your device.	Deactivated
	Add wireless network automatically	-	Here you can search for an available network and automatically add it as the default network.	-
Configure wireless network	Add wireless network manually	-	Here you can configure a network manually. Contact your IT system administrator for required information about the settings.	-
	Edit wireless network	-	Here you can edit the settings for a network already configured and specify a network as the default network.	-
	Allow wireless network configuration	Activated Deactivated	Here you can specify that the user is permitted to configure a wireless network independently.	Activated
Pair Bluetooth [®] devices	-	-	Here you can pair your device with a compatible system for digital patient data recording. If you cancel the process, the menu item is grayed out briefly.	-

13 Operator menu

Parameter		Possible values	Description	Factory setting
E-mail delivery 12-lead ECG	Contacts	-	Here you can enter the names and e-mail addresses for e-mail delivery 12-lead ECG.	-
	Feed rate and duration	Feed: 50 mm/s Duration: 5 s Feed: 25 mm/s Duration: 5 s Feed: 25 mm/s Duration: 10 s	Here you can set whether a single-sided or double-sided PDF of the 12-lead ECG is generated. Single-sided PDF: Feed: 25 mm/s Duration: 5 s Double-sided PDF: • Feed: 50 mm/s Duration: 5 s • Feed: 25 mm/s Duration: 10 s	Single-sided PDF with feed: Feed: 50 mm/s Duration: 5 s
	Amplitude scaling	2 mm/mV 5 mm/mV 10 mm/mV 20 mm/mV	Here you can set the amplification of the ECG signal and thus the height of the ECG curve for e-mail delivery 12-lead ECG.	10 mm/mV
	View	Classic Cabrera	Here you can set the sequence of ECG leads for e- mail delivery 12-lead ECG.	Classic
Upload session data	-	Manual Automatic	Here you can set whether all session data or only session data you select are automatically uploaded to WEINMANN Connect.	Manual

Parameter		Possible values	Description	Factory setting
Register device with WEINMANN Connect	-	-	Here you can enter the registration code generated by WEINMANN Connect in order to register your device with WEINMANN Connect.	-
Date and time update (displayed only if the Automatic clock change menu item is activated)	Update date/time automatically	Activated Deactivated	Here you can set the date and time to synchronize with WEINMANN Connect automatically.	Deactivated
	Update date/time manually	-	Here you can synchronize the date and time with WEINMANN Connect manually.	-

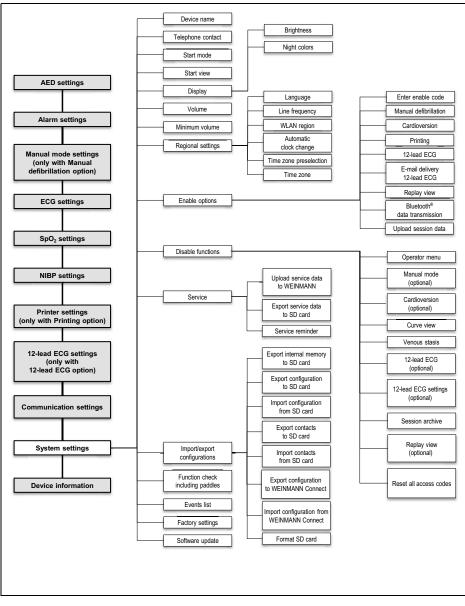
13.12 System settings

You can make presets for the system in the **System settings** submenu.

System settings						
Device name	MEDUCORE Standard2					
Telephone contact						
Start mode	Monitor					
Start view	Curve					
Display						
Volume	75 %					
Minimum volume	25 %					
Regional settings						
Enable options						
Disable functions						
Service						
	Back OK					

13-19 System settings submenu

13.12.1 Menu structure



13-20 Operator menu: System settings submenu

13.12.2 Possible values

Parameter		Possible values	Description	Factory setting
Device name	-	-	Here you can enter the device name, the location of the device or some other identification. The device name appears on the printout and in the PDF of the 12-lead ECG. It assigns the 12-lead ECG to the device with which the 12-lead ECG was created.	MEDUCORE Standard2
Telephone contact	-	-	Here you can enter a telephone number for queries. The telephone contact appears in the Device information menu item and in the PDF of the 12-lead ECG.	-
Start mode	-	Monitor AED Manual	Here you can set in which mode the device is to start. Manual mode is not available if parameter view has been selected as the start view.	Monitor
Start view	-	Curve Parameter	Here you can set in which view the device is to start. Parameter view is not available if manual mode has been selected as the start mode.	Curve
Display	Brightness	10 %-100 %, in increments of 10	Here you can set the display brightness.	70 %
	Night colors	Activated Deactivated	Here you can set whether the device is to start with night colors.	Deactivated

Parameter		Possible values	Description	Factory setting
Volume	-	25 % - 100 %, in increments of 25	Here you can set the volume of the device.	75 %
Minimum volume	-	25 % - 100 %, in increments of 25	Here you can specify the minimum volume which can be set by the user.	25 %
	Language	Languages available in the device (depends on software version)	Here you can set the language of display texts and voice prompts. The device shows the available languages in their own respective language and with their international language abbreviation (e.g. for English: English (en US)).	Customer-specific
	Line frequency	50 Hz 60 Hz	Here you can select the ECG filter in order to suppress interference caused by your regional power supply network.	50 Hz
Regional settings	WLAN region	All available network regions	Here you can specify your network region.	WORLD (default)
	Automatic clock change	Activated C Deactivated	Here you can set the device to switch from winter time to summer time and vice versa automatically. When the Automatic clock change menu item is activated, the Date and time update menu item in the Communication settings submenu is activated.	Deactivated
	Time zone preselection	All available time zone areas	Here you can limit the number of time zones. The menu item is only available if the Automatic clock change menu item is activated.	Europe

13 Operator menu

Parameter		Possible values	Description	Factory setting
Regional settings	Time zone	All time zones within the selected time zone area	You can select your time zone here. The menu item is only available if the Automatic clock change menu item is activated.	Amsterdam
Enable options	-	-	 Here you can enable options and activate and deactivate them after enabling them (see "3.10 Options", page 72). The following options are possible: Manual defibrillation Cardioversion Printing 12-lead ECG E-mail delivery 12-lead ECG Replay view Bluetooth[®] data transmission Session data upload 	-

Parameter		Possible values	Description	Factory setting
Parameter Disable functions	Operator menu Manual mode (only with Manual defibrillation option) Cardioversion (only with Cardioversion option) Curve view Venous stasis 12-lead ECG (only with 12- lead ECG option) 12-lead ECG settings (only with 12-lead	- -	 Description Here you can disable functions for the user and protect them from unauthorized access using an individual access code. When doing so, please note: Manual mode cannot be disabled if it has been selected as the start mode. Curve view cannot be disabled if it has been selected as the start view. If manual mode and curve view are disabled at the same time, then when manual mode is enabled in the session, curve view is automatically enabled at the same time. The access code request 	-
	option) Reset all access codes	-	in the operator menu cannot be removed. Here you can reset all access codes to the factory setting 000000.	-

Parameter		Possible values	Description	Factory setting
	Upload service data to WEINMANN	-	Here you can upload your device service data to WEINMANN Emergency. If you cancel the process, the menu item is grayed out briefly.	-
Service	Export service data to SD card	-	Here you can export your device service data to an SD card.	-
	Service reminder	Activated Deactivated	Here you can set whether the device is to remind you of the next service \leq 30 days before the next service.	Activated
	Export internalHere you can export thememory to SD-cardthe SD card.	internal device memory to	-	
Import/export configurations	Export device configuration to SD card	-	Here you can export the current settings of the operator menu to the SD card so that they can be imported by another device.	-
	Import device configuration from SD card	-	Here you can import the settings exported to an SD card from the operator menu of another device into your device.	-
	Export contacts to SD card (only with E-mail delivery 12-lead ECG option)		Here you can export the contacts you have configured for e-mail delivery 12-lead ECG from your device to an SD card in order to have them imported by another device.	-
	Import contacts from SD card (only with E-mail delivery 12-lead ECG option)	-	Here you can import the contacts you have configured for e-mail delivery 12-lead ECG from an SD card to your device.	-

Parameter		Possible values	Description	Factory setting
	Export configuration to WEINMANN Connect	-	Here you can export the current settings of the operator menu to WEINMANN Connect.	-
Import/export configurations	Import configuration from WEINMANN Connect	-	Here you can import the settings exported to WEINMANN Connect into your device.	-
	Format SD card	-	Here you can format the inserted SD card. This deletes all data from the SD card.	-
Function check including paddles	-	Activated C Deactivated	Here you can set whether the paddles are included in the function check. If you activate paddles on the device, the function check including paddles will be activated automatically.	Deactivated

13 Operator menu

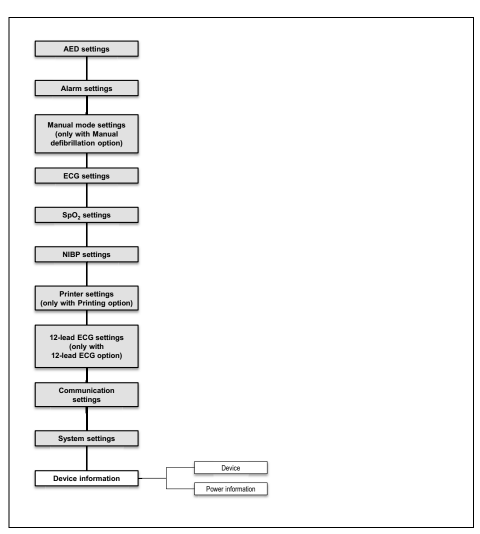
Parameter		Possible values	Description	Factory setting
	Number of events	1 to 50	Here you can set the number of events which can be selected.	28 events preset
Events list	Event 1 to 50 (28 preset)	-	Here you can adapt the separate events individually (maximum 15 characters). The two top events are recorded and counted in the session report.	Furosemide Midazolam Albuterol Atrovent Prednisolone Clemastine Morphine Fentanyl Ketamine Ondansetron Butylscopolamine Metamizole
Factory settings	-	-	Here you can reset the device to its factory settings.	-
Software update	-	-	Here you can carry out a software update.	-

13.13 Device information

You will find information about the device and the battery in the **Device information** submenu.

Device information			
Device			
Power information			
Back			
	Back	ОК	

13-21 Device information submenu



13.13.1 Menu structure

13-22 Operator menu: Device information submenu

Information displayed	Description
Device	
Telephone contact	Here you can find out the telephone number which the operator has assigned to the device for teleconsulting queries.
Serial number	Here you can find out the device serial number. This is also located on the device information label.
Device ID	Here you can find out the device ID. This is required to purchase options.
MAC Bluetooth [®] module (only with Printing and Bluetooth [®] data transmission options)	Here you can find out the MAC address of the Bluetooth $^{\ensuremath{\mathbb{B}}}$ module.
MAC WLAN module	Here you can find out the MAC address of the WLAN module.
Last function check	Here you can find out when a function check was last carried out.
Function check result	Here you can find out whether the last function check carried out was passed.
Days until next service	Here you can find out how many days to go until the next service is due.
Next service	Here you can find out when the next service is due.
Device software	Here you can find out which software version is currently installed on the device.
Power information	
Here you can find out inf for remote diagnoses.	ormation about the battery and the line voltage. This information may be of use

13.13.2 Information displayed

14 Maintenance

DANGER

Risk of injury due to electric shock when the device is opened! The device contains a shock capacitor. Opening the device leads to

serious injuries or death from electric shock.

- \Rightarrow Do not open the device.
- ⇒ The device should only be opened by WEINMANN Emergency or by technicians authorized by WEINMANN Emergency.
- ⇒ Measures such as repairs and maintenance should only be carried out by the manufacturer or by a technician expressly authorized by it.

Risk of injury from impaired device functions or defects, lack of testing and excessive lifetime!

If the device is not correctly tested, or is operated for too long, impaired device functions, deviations or defects may occur. The device cannot be used correctly, and may injure the patient and the user.

⇒ Observe the intervals for maintenance, for Technical Safety Check ("Sicherheitstechnische Kontrolle" in accordance with § 11 of the German regulation MPBetreibV), and for metrological check.

 \Rightarrow Do not operate the device for longer than 10 years.

14.1 General instructions

Maintenance, Technical Safety Check (in Germany only), and measures such as inspections and repairs must only be carried out by the manufacturer or by a technician expressly authorized by it.

14.2 Intervals

Part concerned	Interval	Maintenance by	
	Maintenance-free		
Device	Technical Safety Check every year	Manufacturer or a technician	
	Metrological check every two years	expressly authorized by manufacturer	

Part concerned	Interval	Maintenance by
Battery	Maintenance-free When stored in the device: Charge every 3 months. When not stored in the device: Charge batteries with serial numbers $< 20,000$ every 5 months and batteries with serial numbers $\ge 20,000$ every 9 months. Recommendation: Replace battery after 2 years.	
Charging station	Maintenance-free Recommendation: Replace charging station after 12 years	
Accessories*	Please follow the instructions for use supplied with the accessories. Should the accessories not come with their own instructions for use, the same intervals as for the device apply.	

*Additional information for Germany:

In accordance with the Technical Safety Check ("Sicherheitstechnische Kontrolle" in accordance with § 11 of the German regulation MPBetreibV), we as the manufacturer recommend that all accessories for use of the MEDUCORE Standard² that are connected to it likewise be subject to a safety check (STK) at the same interval as the device.

14.3 Sending in the device

A WARNING

Risk of infection from contaminated parts!

The device, accessories, and other parts may be contaminated, and infect technicians with bacteria and viruses. Parts sent in which are clearly contaminated will be disposed of at the cost of the sender by WEINMANN Emergency or by technicians authorized by WEINMANN Emergency.

- \Rightarrow Clean and disinfect parts before sending them in.
- \Rightarrow Do not send in parts which are potentially contaminated.
- 1. Disassemble parts.
- Clean and disinfect parts (see "8 Hygienic reprocessing", page 238).
- 3. Send parts to WEINMANN Emergency or to technicians expressly authorized by WEINMANN Emergency.

15 Storage

A WARNING

Disrupted or failed therapy due to device being defective or not ready for use following incorrect storage!

Incorrect storage may damage the device, components, and accessories, and lead to disruption or failure of therapy. This may cause serious or life-threatening injury to the patient.

- ⇒ Observe storage conditions and storage times (see "18 Technical data", page 385).
- \Rightarrow Store the device and accessories in a dry location.
- ⇒ Following storage at extreme ambient conditions outside ambient operating conditions: Store the device, components, and accessories at room temperature for at least 12 hours before starting to use them.
- ⇒ Protect the device, components, and accessories from UV light and direct sunlight.

15.1 Storing the device

- 1. Switch off the device (see "6.2 Switching the device off", page 126).
- 2. Disassemble the power supply (see "7.1 Disassembling the power supply", page 226).
- Clean and disinfect the device (see "8 Hygienic reprocessing", page 238).
- 4. Store the device in a dry place.
- *Result* The device is stored in a dry place.

15.2 Storing the battery

- The device and the battery have been cleaned and disinfected (see "8 Hygienic reprocessing", page 238).
 - The battery is fully charged.
 - If available: The replacement battery is fully charged.
 - 1. Insert the battery in the battery compartment and store the device in a dry place.

Alternatively:

Store the battery in a dry place outside the device.

No therapy due to extended storage of the battery without recharging!

Storing the battery for an extended period of time without recharging may result in the rapid shutdown of and irreparable damage to the battery.

- ⇒ When the battery is stored in the device without a power supply: Charge battery every 3 months.
- ⇒ If the battery is not stored in the device: Charge batteries with serial numbers < 20,000 every 5 months and batteries with serial numbers \ge 20,000 every 9 months.
- 2. Charge battery at regular intervals:

Type of storage	Charging interval	
In device without a power supply	Every 3 months	
Outside the device	 Batteries with serial numbers < 20,000: Charge every <p>5 months. </p> Batteries with serial numbers ≥ 20,000: Charge every 9 months. 	

Result The battery is stored in a dry place and is ready for use.

A WARNING

15.3 Storing printer

A CAUTION

Delayed therapy due to incorrect storage of the printer!

Storage outside the specified ambient conditions may lead to printer malfunctions or damage the printer. This may delay the patient's therapy.

- \Rightarrow Always store the printer within the specified ambient conditions.
- \Rightarrow In the event of problems with the printer: Carry out a function check of the printer (see "5.6 Carrying out a function check on the printer (only with Printing option)", page 124).

15.4 Storing printer battery

- If the printer is not being used for an extended period: Charge printer battery (see "4.7.1 Charging the printer battery", page 94).
- 2. Store the printer battery **separately** from the printer.
- 3. Charge the printer battery after no more than 12 months.
- *Result* The printer battery is stored in a dry place and ready for use.

16 Disposal

16.1 Electronic waste





Environmental hazard from electronic waste!

Electronic waste poses an environmental hazard, and must be subjected to proper disposal.

- \Rightarrow Do not dispose of electronic waste in domestic waste.
- ⇒ Contact WEINMANN Emergency or a licensed, certified electronic waste dealer for proper disposal.

The following products are categorized as electronic waste:

- Device
- Master cable
- Defibrillation electrodes
- Paddles
- Pulse oximetry sensor connecting cable
- Pulse oximetry sensor
- ECG cable
- ECG electrodes
- Power supply unit and charger
- Charging station
- Function test resistor
- Printer
- Charging station for printer battery
- Power supply unit and charger for charging stations for printer battery

16.2 Battery/printer battery





Environmental hazard from used batteries!

Used batteries are a risk to the environment and must be subjected to proper disposal.

- \Rightarrow Do not dispose of used batteries in domestic waste.
- ⇒ Contact WEINMANN Emergency or a licensed, certified electronic waste dealer for proper disposal.

16.3 Plastics

Dispose of plastics in a proper manner applicable to plastics at the end of their useful lives.

16.4 Contaminated parts

Do not dispose of contaminated parts in domestic waste. Use a licensed, certified specialist waste management contractor to dispose of contaminated parts properly.

17.1 Version supplied (example)

This sub-section describes just one version supplied. Functions, accessories, and other parts depend on the version purchased, and are not available in every case.

Medical devices and accessories are marked with a UDI-DI. Other parts do not have a UDI-DI. For parts made by other manufacturers (third-party products) you can request the UDI-DI from the manufacturer.

Designation	Supplementary information	UDI-DI	Article no.
MEDUCORE Standard ² , basic device	_	04054685271510	WM 45310
Accu-Pack rechargeable battery	_	04054685157616	WM 45045
MCS2-Connect master cable	_	04054685267162	WM 45397
MCS2-Softpads defibrillation electrodes for adults	Third-party product	_	WM 45418
MCS2-SoftTip pulse oximetry sensor, size M	Reusable	04054685265977	WM 45432
MCS2-Adapt pulse oximetry sensor connecting cable	_	04054685265953	WM 45430
MCS2-Line ECG cable, ERC, 2 m	_	04054685265205	WM 45451
ECG cable separator	-	-	WM 45450
NIBP connecting tube, 2 m	Third-party product	-	WM 45481
NIBP cuff, adult plus, 28- 40 cm arm circumference		-	WM 45464
Kit, mounting elements	-	-	WM 17806
SD card, 32 GB	-	-	WM 39510
Function test resistor	-	-	WM 45428

Designation	Supplementary information	UDI-DI	Article no.
MEDUCORE Standard ² instructions for use	_	_	WM 68401

17.2 Accessories and other parts

This sub-section describes accessories and other parts in accordance with the Medical Device Regulation (MDR). Accessories are marked with a UDI-DI. Other parts do not have a UDI-DI. For parts made by other manufacturers (third-party products) you can request the UDI-DI from the manufacturer.

Supplementary information	UDI-DI	Article no.
•		
-	-	WM 45499
Requirement: Manual defibrillation option is enabled	-	WM 45620
-	-	WM 45621
-	-	WM 45622
Requirement: 12-lead ECG option is enabled	-	WM 45626
-	_	WM 45628
-	-	WM 45624
-	-	WM 45627
	information Requirement: Manual defibrillation option is enabled - Requirement: 12-lead	information UDI-DI - - Requirement: Manual defibrillation option is enabled - - - - - - - - - - - - - - - - - - - - - - - - -

Designation	Supplementary information	UDI-DI	Article no.
Power supply			
Accu-Pack battery	-	04054685157616	WM 45045
Charging station for battery WM 45045		-	WM 45190
Power supply unit and charger 100 W	Third-party product	-	WM 28937
Charging adapter for charging with power supply unit and charger or 12 V adapter cable	-	-	WM 28979
Adapter cable for 12 V on-board power supply/ circular connector	_	_	WM 28356
Defibrillation/cardio	version (WEINMANN	l Emergency)	
MCS2-Connect master cable	-	04054685267162	WM 45397
MCS2-Hardpads paddles	-	04054685281618	WM 45498
Defibrillation/cardio	version (other man	ufacturers)	
MCS2-Softpads defibrillation electrodes for adults	_	_	WM 45418
MCS2-Softpads defibrillation electrodes for children	_	_	WM 45419
Set of 12, electrode gel	-	-	WM 14292
Pulse oximetry moni	toring (WEINMANN	Emergency)	
MCS2-Adapt pulse oximetry sensor connecting cable	_	04054685265953	WM 45430
MCS2-SoftTip pulse oximetry sensor, size S	Reusable	04054685265960	WM 45431
MCS2-SoftTip pulse oximetry sensor, size M	Reusable	04054685265977	WM 45432
MCS2-SoftTip pulse oximetry sensor, size L	Reusable	04054685265984	WM 45433

Designation	Supplementary information	UDI-DI	Article no.
Set of 24, MCS2-Wrap pulse oximetry sensor, adult	Disposable	04054685266776	WM 45436
Set of 24, MCS2-Wrap pulse oximetry sensor, infant	Disposable	04054685266783	WM 45437
Set of 24, MCS2-Wrap pulse oximetry sensor, child	Disposable	04054685266790	WM 45439
MCS2-Wrap pulse oximetry sensor	Reusable	04054685265991	WM 45434
Set of 10, strap for fastening MCS2-Wrap pulse oximetry sensor	Disposable	04054685267230	WM 45442
MCS2-Earclip pulse oximetry sensor	Reusable	04054685266004	WM 45435
Set of 5, ear clip for MCS2-Earclip pulse oximetry sensor	_	-	WM 45443
ECG cable separator	ng/12-lead ECG record	ling and assessment	WM 45450
MCS2-Line ECG cable, ERC, 2 m	_	04054685265205	WM 45451
MCS2-Line ECG cable, ERC, 3 m	_	04054685265212	WM 45452
MCS2-Line ECG cable with connection for 12- lead ECG extension cable, ERC, 2.4 m	_	04054685275099	WM 45455
MCS2-Line ECG cable with connection for 12- lead ECG extension cable, ERC, 3.4 m	-	04054685275105	WM 45456
MCS2-Line 12-lead ECG extension cable, ERC	-	04054685275075	WM 45447
MCS2-Line ECG cable, AHA, 2 m	_	04054685265229	WM 45453

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Designation	Supplementary information	UDI-DI	Article no.
MCS2-Line ECG cable, AHA, 3 m	-	04054685265236	WM 45454
MCS2-Line ECG cable with connection for 12- lead ECG extension cable, AHA, 2.4 m	-	04054685275112	WM 45457
MCS2-Line ECG cable with connection for 12- lead ECG extension cable, AHA, 3.4 m	-	04054685275129	WM 45458
MCS2-Line 12-lead ECG extension cable, AHA	_	04054685275082	WM 45448
ECG simulator, 6-lead ECG, shockable*	_	-	WM 45444
ECG simulator, 12-lead ECG, shockable*	-	-	WM 45445
Non-invasive blood g	pressure measuremen	t (NIBP monitoring) (o	ther manufacturers)
NIBP connecting tube, 2 m	-	-	WM 45481
NIBP connecting tube, 3 m	_	-	WM 45482
NIBP cuff, infant, 8- 13 cm arm circumference	Reusable	-	WM 45460
NIBP cuff, child, 12- 19 cm arm circumference	Reusable	-	WM 45461
NIBP cuff, small adult, 7- 25 cm arm circumference	Reusable	-	WM 45462
NIBP cuff, adult, 23- 33 cm arm circumference	Reusable	_	WM 45463
NIBP cuff, adult plus, 28- 40 cm arm circumference	Reusable	_	WM 45464
NIBP cuff, large adult plus, 40-55 cm arm circumference	Reusable	_	WM 45465
NIBP cuff, adult, 38- 50 cm thigh circumference	Reusable	_	WM 45466

Designation	Supplementary information	UDI-DI	Article no.
Adapter tube for connecting NIBP disposable cuffs for neonates	_	_	WM 45467
Set of 20, NIBP cuff, neonate, size 1, 3-6 cm arm circumference	Disposable	-	WM 45468
Set of 20, NIBP cuff, neonate, size 2, 4-8 cm arm circumference	Disposable	_	WM 45469
Set of 20, NIBP cuff, neonate, size 3, 6-11 cm arm circumference	Disposable	_	WM 45470
Set of 20, NIBP cuff, neonate, size 4, 7-13 cm arm circumference	Disposable	-	WM 45471
Set of 20, NIBP cuff, neonate, size 5, 8-15 cm arm circumference	Disposable	-	WM 45472

Designation	Supplementary information	UDI-DI	Article no.
Printing			
Set, printer*	-	-	
Comprising:		·	
Printer	Third-party product	-	
Printer battery	Third-party product	-	
Charging station for printer battery with power supply unit and charger	-	-	WM 45640
Printer case	-	-	
	1	I	
Printer battery*	Third-party product	-	WM 45616
Charging station for printer battery with power supply unit and charger*	_	-	WM 45617
Charging station for printer battery with power supply unit and charger*, quadruple	_	-	WM 45618
Printer case	_	-	WM 45619
Set of 10, printer paper	-	-	WM 14698
Portable units/prote	ctive transport bag	5	
MCS2-Bag protective transport bag	-	04054685268473	WM 45490
MCS2-Bag protective transport bag, large	-	04054685281623	WM 45590
Insert for accommodating MCS2- Hardpads paddles in MCS2-Bag protective transport bag, large	_	_	WM 45202
LIFE-BASE portable unit	-	UDI-DI on request	Article no. on request
			·
Miscellaneous			
Function test resistor	-	-	WM 45428
Holding plate	-	-	WM 9714

Designation	Supplementary information	UDI-DI	Article no.
Raised countersunk head screw ISO 7047 M4 x 8	-	-	WM 51398
DEFIview PC software	-	-	WM 45120
Kit, mounting elements	-	-	WM 17806
SD card, 32 GB	-	-	WM 39510
Adapter cable for connection to Ambu or Laerdal practice manikin*	Third-party product	_	WM 45424
Adapter cable for connecting ShockLink [®]	_	-	WM 45369
Adapter cable for connection of corPatch easy defibrillation electrodes to MEDUCORE Standard ²	-	-	WM 45429

* Not suitable for use in the vicinity of the patient. Printer and printer battery must only be used in the vicinity of the patient if they are in the printer case.

18 Technical data

18.1 Device

A WARNING

Risk of injury due to ignoring the technical data of components and accessories!

The technical data of components and accessories may differ from device technical data. Ignoring the technical data of components and accessories may injure the patient, the user, and bystanders. \Rightarrow Observe the technical data of components and accessories.

 \Rightarrow Follow the instructions for use of the components and

accessories.

Specification	Device
Product class according to Directive 93/42/EEC	llb
Dimensions (W x H x D)	242 mm x 137 mm x 130 mm
Weight:	
Without battery With battery	2.25 kg 2.75 kg
Continuous operation: • Temperature range • Humidity • Air pressure • Altitude above mean sea level Unless stated otherwise, these conditions also apply to all components and all accessories.	0 °C to +40 °C 15 % rh to 95 % rh, no condensation 540 hPa to 1100 hPa -500 m to 5000 m
Transient operation (temporary operation at the session location): • Temperature range • Humidity • Air pressure • Altitude above mean sea level Unless stated otherwise, these conditions also apply to all components and all accessories.	-20 °C to +55 °C 15 % rh to 95 % rh, no condensation 540 hPa to 1100 hPa -500 m to 5000 m

Specification	Device
Storage/transport (see "15 Storage",	
page 372):	
Temperature range	-40 °C to +70 °C
Humidity	15 % rh to 95 % rh, no condensation
Air pressure	540 hPa to 1100 hPa
Altitude above mean sea level	-500 m to 5000 m
Unless stated otherwise, these	
conditions also apply to all components and all accessories.	
	12 V to 15.1 V
Electrical connection (rated voltage)	
Max. power consumption	30 W
Disconnection from line power	Pulling out the power plug disconnects the device from line power on all poles.
Operation with on-board power supply: • Rated voltage	12 V
 Max. internal resistance of on- 	
board power supply	500 mΩ
Maximum current consumption	< 3 A
Operating mode	Continuous duty
Classification acc. to EN 60601-1:	
 Type of protection against elec. shock 	Protection class II
• Degree of protection against elec. shock (pulse oximetry sensor)	Degree of protection BF - defibrillation-proof
 Degree of protection against elec. shock (ECG electrodes) 	Degree of protection CF - defibrillation-proof
 Degree of protection against elec. shock (defibrillation electrodes/ 	Degree of protection BF - defibrillation-proof
paddles)	
 Degree of protection against elec. shock (NIBP cuff) 	Degree of protection BF - defibrillation-proof
Degree of protection against	
Ingress of solid objects	
Ingress of dust	IP55: Protected against dust and water jets
 Ingress of water with harmful effect 	
Mechanical resistance to falls	1 m
Electromagnetic compatibility (EMC)	Test parameters and limit values can be requested from the
acc. to EN 60601-1-2:	manufacturer WEINMANN Emergency if required.
Radio interference suppression	EN 55011
Radio interference immunity	EN 61000-4 (parts 2 to 6, 8, and 11)

Specification	Device
Electromagnetic compatibility (EMC) in accordance with EN 60601-12 and EUROCAE ED-14G (RTCA DO 160 G)	EUROCAE ED-14G (RTCA DO 160 G), Section 21, Category M EUROCAE ED-14G (RTCA DO 160 G), Section 20, Category T The device is not intended for connection to the power supply network in aircraft in accordance with RTCA DO 160 G.
Frequency band	WLAN: 2.4 GHz to 2.5 GHz Bluetooth [®] interface: 2.4 GHz to 2.4835 GHz
Transmission power	WLAN: maximum 18 dBm \pm 1 dB Bluetooth [®] interface: maximum 7 dBm
Types of modulation of Bluetooth $^{\textcircled{B}}$ interface	1 Mbps: GFSK (BDR) 2 Mbps: π/4-DQPSK (EDR) 3 Mbps: 8-DPSK (EDR)
Types of modulation of WLAN interface	Modulation 802.11b CCK and DSSS 802.11a/g/n OFDM
Display	5.7" TFT color display Resolution VGA 640 pixels x 480 pixels
Sound pressure level range for alarms	39 dBA to 83 dBA
Sound pressure level range for QRS beeps	35 dBA to 77 dBA
Sound pressure level range for pulse tones	43 dBA to 83 dBA
Mechanical resistance to shock and vibration	 EN 1789 EN 60601-1-12 (Categories: Secured in a rescue vehicle, secured in an airplane, secured in a helicopter, portable at the site of the emergency) EUROCAE ED-14G (RTCA DO 160 G) (Section 7 - Shock and Crash Cat. A; Section 8 - Vibration Fixed-Wing Airplanes Cat. S and Helicopters Cat. U2)
Type of emergency vehicle	Secured in rescue vehicle, airplane and helicopter as well as portable at the site of the emergency

Specification	Device
	EN 60601-1
	EN 60601-1-2
	EN 60601-1-6
	EN 60601-1-8
	EN 60601-1-12
	EN 60601-2-4
Standards used	EN 60601-2-25
Standards used	EN 60601-2-27
	EN 60601-2-49
	EN 80601-2-30
	EN 80601-2-61
	EN 1789
	EUROCAE ED-14G (RTCA DO 160 G)
	MIL-STD 810 G CAT 12/CAT 13/CAT 14/CAT 20
	EN 60601-2-4
	EN 60601-2-25
Essential performance	EN 60601-2-27
Essential performance	EN 60601-2-49
	EN 80601-2-30
	EN 80601-2-61
Resuscitation report	ERC 2015
Expected lifetime	10 years
CE marking	C E 0197
The right to make design modific	ations is reserved

18.2 Defibrillation electrodes

Specification	Defibrillation electrodes
Conductive surface of the defibrillation electrodes	Per defibrillation electrode for adults: 86 cm ² Per defibrillation electrode for children: 54 cm ²
Temperature range for storage	0 °C to 50 °C
Temperature range for operation	0 °C to 50 °C

18.3 Paddles

Specification	Paddles
	Per paddle for adults: 75 cm ² Per paddle for children: 19.6 cm ²
Temperature range for storage	-40 °C to +55 °C

18.4 Master cable

Specification	Master cable
Lifetime	5000 connector/connection cycles

18.5 Battery

Specification	Battery
Туре	Li-ion
Dimensions (W x H x D)	97 mm x 127 mm x 33 mm
Weight	450 g
Shock capacity (at 20 °C with new and fully charged battery)	350 shocks at 200 J
Monitoring capacity	Approx. 5 h
Battery operating time for resuscitation	4 h with shocks of 200 J
Rated capacity	4.2 Ah (45.6 Wh/46.4 Wh/46.8 Wh)
Rated voltage	10.8 V
Charging time (0 % to 90 %)	Approx. 3.5 h
Charging temperature	0 °C to +45 °C
Lifetime	At least 300 cycles or 2 years
Storage	Storage above 60 °C maximum 1 week

18.6 Power supply unit and charger

Specification	Power supply unit and charger
Manufacturer	PROTEK POWER
Туре	PMP120F-13-K24
Operation:	
Temperature range	0 °C to +40 °C
Humidity	5 % rh to 95 % rh, no condensation
Air pressure	700 hPa to 1100 hPa
Altitude above mean sea level	-500 m to 3000 m
Input current	1.4 A to 0.6 A
Input voltage	100 V-240 V~/50 Hz-60 Hz
Rated voltage output	15 V
Disconnection from line power	Pulling out the power plug disconnects the device from line power on all poles.

18.7 Charging station

Specification	Charging station
Dimensions (W x H x D)	175 mm x 46 mm x 135 mm
Supply voltage	12 V to 15.1 V direct voltage
Weight	380 g (tare weight)
Operation:	
Temperature range	0 °C to +45 °C
Humidity	5 % rh to 95 % rh, no condensation
Storage:	
Temperature range	-30 °C to +70 °C
Humidity	5 % rh to 95 % rh, no condensation
Short circuit resistance	Yes
	WEINMANN Emergency-specific plug connection
Connection to power supply network	 Hard-wired power connection on the rear of the
	charging station for on-board power supply
Electromagnetic compatibility (EMC) acc. to	RTCA DO 160 F
EN 60601-1-2:	EN 55011
Radio interference suppression	EN 61000-4 (Parts 2 and 3)
Radio interference immunity	EN 1789
Degree of protection against ingress of drips	IPX1
Power consumption	Maximum 30 W, on standby maximum 0.4 W

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18.8 Printer and printer paper

Type WSP-i Size 120 m Weight 424 g Print method Direct Resolution 203 d Bluetooth [®] interface • Bl Bluetooth [®] interface • Cl Fr • Cl Range 1 m Display • LC Printer • Cl Degree of protection against • LC Ingress of solid objects IP54: I Ingress of dust IP54: I Ingress of water with harmful effect IP54: I Operation: -10 °C • Temperature range -10 °C • Humidity 30 % Storage: -10 °C • Temperature range -10 °C • Humidity 10 % Printer battery 7.4 V 1800 13.32	er and printer paper
Type WSP-i Size 120 m Weight 424 g Print method Direct Resolution 203 d Bluetooth [®] interface • Bl Size 1 m Display • L0 Printer • L0 Degree of protection against • L0 Ingress of solid objects IP54: I Ingress of dust IP54: I Ingress of dust 1.5 m Operation: - 10 °C Temperature range -10 °C Humidity 30 % Storage: - 10 °C Printer battery 7.4 V 1800 13.32	
Size 120 m Size 120 m Weight 424 g Print method Direct Resolution 203 d Bluetooth [®] interface • Bl Bluetooth [®] interface • Cl Fr • Cl Range 1 m Display • LC Printer • Cl Degree of protection against • LC Ingress of solid objects IP54: I Ingress of dust IP54: I Ingress of dust 1.5 m Operation: -10 °C • Temperature range -10 °C • Humidity 30 % Storage: -10 °C • Temperature range -10 °C • Humidity 10 % Printer battery 7.4 V 1800 13.32	m Systems Inc.
Weight 424 g Print method Direct Resolution 203 d Bluetooth [®] interface • Bl Bluetooth [®] interface • Cl Fr • Cl Range 1 m Display • LC Printer • Cl Degree of protection against • LC Ingress of solid objects IP54: I Ingress of dust IP54: I Ingress of dust 1.5 m Operation: -10 °C Temperature range -10 °C Humidity 30 % Storage: -10 °C Printer battery 7.4 V 1800 13.32	350
Print method Direct Resolution 203 d Bluetooth [®] interface 0 Bluetooth [®] interface 0 Cl Fr Range 1 m Display 0 Printer Degree of protection against 1 Ingress of solid objects 1 Ingress of dust 1 Ingress of water with harmful effect 1 Mechanical resistance to falls 1.5 m Operation: -10 °C Humidity 30 % Storage: -10 °C Humidity 10 % Printer battery 7.4 V 1800 13.32	m x 130.5 mm x 58.4 mm
Resolution 203 d Bluetooth [®] interface • Bl Bluetooth [®] interface • Cl Fr • Cl Range 1 m Display • LC Printer • Cl Degree of protection against • LC Ingress of solid objects IP54: I Ingress of dust IP54: I Ingress of vater with harmful effect 1.5 m Operation: -10 °C Humidity 30 % Storage: -10 °C Temperature range -10 °C Humidity 10 % Printer battery 7.4 V 1800 13.32	
Bluetooth [®] interface • Bl Bluetooth [®] interface • Cl • Fr • Ingress Display • LC Printer • Ra Degree of protection against • Ingress of solid objects • Ingress of dust IP54: I • Ingress of dust IP54: I • Ingress of water with harmful effect Ingress of water with harmful effect Mechanical resistance to falls 1.5 m Operation: -10 °C • Temperature range -10 °C • Humidity 30 % Storage: -10 °C • Temperature range -10 °C • Humidity 10 % Printer battery 7.4 V 1800 13.32	thermal printing
Bluetooth [®] interface - C Fr Range 1 m Display - LC Printer Degree of protection against Ingress of solid objects Ingress of dust Ingress of dust Ingress of water with harmful effect Mechanical resistance to falls 1.5 m Depration: -10 °C Humidity 30 % Storage: -10 °C Humidity 10 % Printer battery 7.4 V 1800 13.32	pi: 8 dots/mm
Display Output Display Output Display Printer Degree of protection against Ingress of solid objects Ingress of dust Ingress of dust Ingress of water with harmful effect Mechanical resistance to falls Ingreation: Temperature range Humidity Storage: Temperature range Humidity I0 % Printer battery 7.4 V 1800 13.32	uetooth [®] interface: Version 3.0 ass: 2 equency band: 2402 MHz to 2480 MHz
Printer Degree of protection against Ingress of solid objects Ingress of dust Ingress of water with harmful effect Mechanical resistance to falls Degreation: Temperature range -10 °C Humidity Storage: Temperature range -10 °C Humidity 10 % Printer battery 7.4 V 1800 13.32	
Degree of protection against Ingress of solid objects Ingress of dust Ingress of water with harmful effect Mechanical resistance to falls Deperation: Temperature range Humidity Storage: Temperature range Temperature range Temperat	D display with blue background illumination solution 128 pixels x 32 pixels
 Ingress of solid objects Ingress of dust Ingress of water with harmful effect Mechanical resistance to falls Temperature range Humidity Storage: Temperature range <li< td=""><td></td></li<>	
Operation: -10 °C • Temperature range -10 °C • Humidity 30 % Storage: -10 °C • Temperature range -10 °C • Humidity 10 % Printer battery 7.4 V 1800 13.32	Protected against dust and splashes
Temperature range -10 °C Humidity 30 % Storage: -10 °C Temperature range -10 °C Humidity 10 % Li-ion Recha Printer battery 7.4 V 1800 13.32	
Temperature range -10 °C Humidity 10 % Li-ion Recha Printer battery 7.4 V 1800 13.32	to +50 °C h to 80 % rh, no condensation
Printer battery 7.4 V 1800 13.32	to +70 °C h to 90 % rh, no condensation
14 cos	geable OC nAh
Maximum printing time with printer battery (1 ses: 2 12-1	sions ion: 60 min standby, 2 live printouts and ead ECG printouts)
	out: 100 V-250 V~/50 Hz-60 Hz
Charging time, printer battery 4 h (fu	utput: 8.4 V DC/0.8 A

Specification	Printer and printer paper
Printer paper	
Printer paper	 Type of paper: Thermal paper Paper width: 80 mm Paper thickness: 60 ± 5 μm Roll diameter: up to Ø 50 mm
Operation: • Temperature range • Humidity	-10 °C to +50 °C 30 % rh to 80 % rh, no condensation
Transport < 48 h: • Temperature range • Humidity	-10 °C to +50 °C 30 % rh to 80 % rh, no condensation
Storage > 48 h: • Temperature range • Humidity	18 °C to 28 °C 40 % rh to 60 % rh, no condensation

18.9 CARDIObiphasic defibrillation system

Specification	Defibrillation system
Operating mode	Semi-automatic (AED mode)
Operating mode	Manual (manual mode)
Shock form	Biphasic, current-limited, impedance-compensated
Patient impedance:	
Maximum	200 Ω
Minimum	5 Ω
	Adjustable:
Shock sequence	Energy progression:
Shock Sequence	First shock: 1 J to 200 J
	Further shocks: 1 J to 200 J
Analysis time	8 s under typical conditions
Duration of analysis and shock charging in	
AED mode:	
With fully charged source of energy	8 s to 15 s
After 15 discharges	8 s to 15 s
After 6 discharges	8 s to 15 s
Duration of switching on device, analysis, and shock	
charging in AED mode:	
With fully charged source of energy	< 25 s
After 15 discharges	< 25 s
After 6 discharges	< 25 s
CPR phase adjustable	120 s to 300 s

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Specification	Defibrillation system
Energy level adjustable	1 J to 200 J (see "18.18 The CARDIObiphasic shock pulse", page 404)
Duration of shock charging in manual mode	2 s to 9 s (depending on the selected shock energy)
Impedance compensation	Yes
Defibrillator voltage	2 kV
Recovery time for derivation of defibrillation electrodes after a defibrillation shock	5 s

18.10 6-lead ECG monitoring/12-lead ECG recording and assessment

Specification	6-lead ECG monitoring/12-lead ECG recording and assessment
Maximum patient impedance	500 kΩ
Detected heart rate (with ECG derivation via ECG cable or derivation via defibrillation electrodes)	30 bpm to 250 bpm
Suitability for direct use on the heart	Type CF
Displayed heart rates with pacemaker pulses (with ECG derivation via ECG cable)	If pacemaker pulses are detected, "-" is displayed instead of the heart rate.
Pacemaker pulses detected by the device: Amplitude Pulse width Overshoots	2 mV to 700 mV 0.5 ms to 2 ms Detectable at the quoted amplitude and pulse width in lead I. Pacemaker pulse marking displayed in all ECG leads.
Device operating mode in the event of a power supply interruption < 30 s	The device saves all user settings and restores them when the device is restarted.
Detection of detached electrodes	Yes
Noise suppression	Yes
Respiration detection	No
Max. T wave suppression	0.9 mV
Delay time between R wave detection and shock delivery	< 60 ms (measured with Fluke Impulse 7000DP)
Detection range for R wave detection	30 bpm to 180 bpm 0.5 mV to 4.5 mV
Heart rate averaging	The mean heart rate is obtained by averaging up to 7 heartbeat intervals, updated with every heartbeat or at least every second.

Specification	6-lead ECG monitoring/12-lead ECG recording and assessment
Heart rate accuracy (handling irregular cardiac rhythms)	All complexes are detected. The heart rate display is between the shortest and the longest RR interval detected.
Inputs and input impedance	> 2.5 MΩ
Alarm time for tachycardia (acc. to 60601-2-27)	 Signal B1: 1 mV, 206/min: 11 s Signal B1: 0.5 mV, 206/min: 11 s Signal B1: 2 mV, 206/min: 10 s Signal B2: 2 mV, 195/min: 9 s Signal B2: 1 mV, 195/min: 9 s Signal B2: 4 mV, 195/min: 8 s
Response time of heart rate display: Steep rise Steep drop	80/min to 120/min: 4.5 s 80/min to 40/min: 8 s
Displayed heart rate for varying ECG complexes according to 60601-2-27, Section 201.7.9.2.9.101	 Signal A1: 40/min Signal A2: 30/min Signal A3: 60/min Signal A4: 70/min to 95/min
Recovery time after a defibrillation shock	3 s
Band width: ECG in monitor mode	0.03 Hz to 150 Hz (ECG leads I, II, III, aVR, aVL, aVF, V1, V2, V3, V4, V5, V6) 0.34 Hz to 45 Hz (Pad ECG lead)
ECG in 12-lead ECG function mode	0.03 Hz to 50 Hz or 0.03 Hz to 150 Hz for all ECG leads displayed (depending on setting in user menu)
Protection against malfunction caused by electrosurgery	Yes
Leakage currents	Leakage currents may add up.
Feed rate of ECG curve (display and printout)	6-lead ECG: 12.5 mm/s, 25 mm/s, 50 mm/s 12-lead ECG: 25 mm/s, 50 mm/s

18.11 CARDIOlogic ECG analysis system

Specification	ECG analysis system
Analysis time (VF/VT)	 Duration of analysis in AED mode: 8 s under typical conditions Duration of analysis for VF/VT alarm: 8 s with continuous measurement
ECG derivation used for analysis	Lead II of the defibrillation electrodes is primarily used for analysis. If the defibrillation electrodes are not connected, lead II of the ECG electrodes is used.
Impedance measurement	Checks electrode contact
Pulses from implanted pacemakers may affect or prevent correct dete tesponse to implanted cardiac of arrhythmia. As a result, not all rhythms which can be defibrillated be detected, and the device may not recommend shock delivery und certain circumstances.	
Asystole threshold	0.2 mV

The performance of the rhythm detector was validated with a representative cross-section of ECG data from the following databases:

- Creighton University Ventricular Tachyarrhythmia Database
- American Heart Association ECG Database
- MIT-BIH Malignant Ventricular Arrhythmia Database

Effectiveness results are shown in the tables below:

Rhythms	Scope of sample	Objectives for effectiveness	Sensitivity in %	Specificity in %			
Shockable							
Coarse VF (Amplitude > 0.35 mV)	944	> 90 %	98.31 %	-			
Rapid VT (f > 180/min)	252	> 75 %	94.05 %	-			
Non-shockable							
Non-shockable rhythms	3070	> 99 %	-	99.64 %			

These parameters were determined on the basis of the following data:

	Shockable signals	Non-shockable rhythms (normal sinus rhythm (NSR), supraventricular tachycardia (SVT), atrial fibrillation/flutter (AF), ventricular extrasystoles (PVC), pacemaker ECG)	
Shock	1165 (A)	11 (B)	
No shock	31 (C)	3059 (D)	

- A = Number of correct positive decisions
- B = Number of false positive decisions
- C = Number of false negative decisions
- D = Number of correct negative decisions
- This results in the following values:

	Formula for the calculation	Calculation	Result
Sensitivity	$\frac{A}{A+C}$	<u>1165</u> 1165 + 31	97.41 %
Specificity	$\frac{D}{B+D}$	<u> </u>	99.64 %
Precision	$\frac{A + D}{A + B + C + D}$	<u>1165 + 3059</u> 1165 + 11 + 31 + 3059	99.02 %
False positive rate	$\frac{B}{B+D}$	<u>11</u> 11 + 3059	0.36 %
True forecast value	$\frac{A}{A+B}$	<u>1165</u> 1165 + 11	99.06 %

18.12 Pulse oximetry monitoring

The statistical distribution of the measured values of pulse oximetry sensors means that an average of two-thirds of all measurements are within the $\pm A_{rms}$ value of CO-oximeters as a reference method (see table). To determine bpm accuracy, comparative measurements were carried out with an original sensor and original monitor.

Specification	Pulse oximetry		
SpO ₂ display range	45 % to 100 %		
Accuracy (for all the pulse oximetry sensors quoted in	70% to 100 %: $\leq 2 \%$		
these instructions for use)	45 % to 70 %: Not specified		
Signal quality display	Alarm output if SpO ₂ signal quality is $<$ 40 %		
Pulse rate	30 bpm to 300 bpm \pm 3 bpm at SpO ₂ > 45 %		
Reference methods for determining accuracy of pulse rate	Oxitest simulator testing		
Wavelengths at maximum intensity	660 nm/890 nm (2.5 mW to 4.5 mW)		
Curve form	Normalized		
	12.5 mm/s		
Feed rate for plethysmogram (display and printout)	25 mm/s		
	50 mm/s		
Update rate of SpO ₂ mean	8 s (It may take up to 16 s for the correct SpO_2 value		
	to be displayed.)		
Averaging	12 s		
Delay in data updating	500 ms		
Delay in alarm condition	20 s		
Delay in alarm generation	< 1 s		
	The pulse oximetry sensor is calibrated to display		
Functional oxygen saturation	functional oxygen saturation, and must not be		
	calibrated with a function tester.		
	• The pulse oximetry sensor is latex-free. No		
	material used in its production contains latex		
Dis some settle iliter	protein. The materials with which the patient		
Biocompatibility	comes into contact have undergone extensive		
	biocompatibility tests. Further information is available on request.		
	Acc. to EN ISO 10993-10		
	 ACC. TO FIN ISO 1033-10 		

18.13 Non-invasive blood pressure measurement (NIBP monitoring)

Specification	Non-invasive blood pressure measurement
Manufacturer	SunTech Medical
Measuring method	Oscillometric, diastolic values correspond to phase 5 Korotkoff sounds
Measurement range (systolic):	
Adults	40 mmHg to 260 mmHg
Children	40 mmHg to 230 mmHg
Infants	40 mmHg to 130 mmHg
Measurement range (diastolic):	
Adults	20 mmHg to 200 mmHg
Children	20 mmHg to 160 mmHg
Infants	20 mmHg to 100 mmHg
Accuracy (during operation between 0 °C and 50 °C)	±3 mmHg

18.14 Operation/data management

Specification	Operation/data management
Display	Illuminated symbolsDevice status indicators
Audio output	Voice promptsAlarm tonesAudible signals
Session documentation	Automatic recording of measured values for ECG, SpO ₂ and NIBP and of event data
Data transmission via SD card	SD card with 32 GB
Data transmission via WLAN	 Data transfer types: 12-lead ECG, function check results, session data, service data WLAN: 802.11a/b/g/n Security: WPA2-PSK Protocol: https Port: 443
Data transmission via Bluetooth $^{f R}$ interface	Data transfer type: Session data
Data evaluation	Via DEFlview PC software

18.15 Alarm delay times

Alarm	Delay time
Battery defective	10.0 s
Insert battery	10.0 s
Asystole	
 Patient is connected with this condition 	9.6 s
 Patient already connected 	8.5 s
ECG module defective	10.0 s
Heart rate 1 (with derivation via defibrillation	
electrodes)	
Patient is connected with this condition	3.0 s
 Patient already connected 	1.7 s
Heart rate ↓ (with derivation via defibrillation electrodes)	
Patient is connected with this condition	9.7 s
 Patient already connected 	6.0 s
Heart rate † (with derivation via ECG electrodes)	
 Patient is connected with this condition 	8.0 s
 Patient already connected 	2.0 s
Heart rate ↓ (with derivation via ECG electrodes)	
 Patient is connected with this condition 	14.9 s
 Patient already connected 	5.0 s
Pulse rate †	10.0 s
Pulse rate 🖡	10.0 s
SpO ₂ saturation †	10.0 s
SpO ₂ saturation ↓	10.0 s
VF/VT	
 Patient is connected with this condition 	11.3 s
 Patient already connected 	5.3 s

18.16 Saving of session data

Storage medium	Memory hours assuming typical use		
Internal device memory (100 MB)*	09 h 00 min		
SD card WM 39510 (32 GB)*	1675 h		

 Since different curve forms can be compressed to differing degrees, the actual number of storage hours may deviate. The data are typical values.

18.17 Electromagnetic compatibility (EMC)

Medical electrical equipment is subject to special precautions in relation to electromagnetic compatibility (EMC). It must be installed and put into operation in accordance with the EMC information contained in the accompanying documentation.

In addition to the performance defined in the standard, this device has the following essential performance:

- Delivery of a current pulse for defibrillation and cardioversion
- Distinction between cardiac rhythms to be defibrillated and not to be defibrillated
- Display of electrocardiogram (ECG)
- Display of plethysmogram and hemoglobin oxygen saturation (SpO₂)
- Display of systolic and diastolic arterial blood pressure (NIBP)

A WARNING

Delay in treatment due to power supply network disruption!

Disruptions to the power supply network may cause visible disruptions to the ECG. This may lead to incorrect measurement results and alarms and in AED mode, lead to shock delivery at the wrong time. This may delay therapy and injure the patient.

⇒ If there is major disruption to the power supply network, only operate the device with a battery.

A WARNING

Risk of injury from overly powerful high-frequency interference signals or electromagnetic fields!

Overly strong high-frequency interference signals from portable high-frequency communications equipment (e.g. radios) or electromagnetic fields may lead to incorrect analysis, incorrect measurement results, and incorrect alarms, and thus impair the functioning of the device and injure the patient.

- \Rightarrow Maintain separation distances.
- ⇒ With portable high-frequency communications devices, maintain a minimum distance of 30 cm from the device, components, and accessories.

Recommended separation distances between portable and mobile HF telecommunications devices and MEDUCORE Standard²

MEDUCORE Standard² is intended for use in an electromagnetic environment in which HF interference variables are controlled. The customer or user of the MEDUCORE Standard² can avoid electromagnetic interference by maintaining a minimum distance between portable and mobile HF telecommunications equipment (transmitters) and MEDUCORE Standard² (depending on the output power of the HF telecommunications equipment, see below).

	Separation distance depending on transmission frequency in m				
Rated power of	150 kHz -	150 kHz -	When used as a monitor		
HF device in W	80 MHz	80 MHz in the ISM bands	80 MHz - 800 MHz	800 MHz - 2.5 GHz	
0.01	0.12	0.12	0.4	0.77	
0.1	0.38	0.38	1.3	2.4	
1	1.2	1.2	4	7.7	
10	3.8	3.8	13	24	
100	12	12	40	77	

Recommended separation distances between portable and mobile HF telecommunications devices and MEDUCORE Standard²

	Separation distance depending on transmission frequency in m			
Rated power of	When used as a defibrillator		Defibrillator: No unintended energy delivery	
HF device in W	150 kHz - 80 MHz	150 kHz - 80 MHz in the ISM bands	80 MHz - 800 MHz	800 MHz - 2.5 GHz
0.01	0.12	0.27	0.06	0.12
0.1	0.38	0.66	0.15	0.38
1	1.2	2.7	0.6	1.2
10	3.8	6.6	1.5	3.8
100	12	27	6	12

Electromagnetic emission

Electromagnetic emission

MEDUCORE Standard² is designed for operation in the electromagnetic environment described below. The operator or the user must ensure that he or she operates MEDUCORE Standard² in an environment of this kind.

Emission measurements	Compliance	Electromagnetic environment guidelines		
	Group 1	MEDUCORE Standard ² uses high-frequency energy exclusively for internal functions. As the high-frequency emission is very low, interference with electronic devices in the vicinity from MEDUCORE Standard ² is unlikely.		
HF emissions according to CISPR 11	Class B	 MEDUCORE Standard² is suitable for use in the following areas: In all facilities, including those in residential settings which are connected directly to a public power supply network In automobiles, airplanes and ships 		

Electromagnetic immunity

Electromagnetic immunity
MEDUCORE Standard ² is designed for operation in the electromagnetic environment described below. The operator or the user must ensure that he or she operates MEDUCORE Standard ² in an environment of this kind.

Interference immunity tests	IEC 60601 test level	Compliance level
Electrostatic discharge (ESD) acc. to IEC 61000-4-2	\pm 8 kV contact \pm 2 kV, \pm 4 kV, \pm 8 kV, \pm 15 kV air	\pm 8 kV contact \pm 2 kV, \pm 4 kV, \pm 8 kV, \pm 15 kV air
Radiated HF interference acc. to IEC 61000-4-3	20 V/m 80 MHz to 2.7 GHz	20 V/m
Rapid transients/bursts acc. to IEC 61000-4-4	± 2 kV 100 kHz spike frequency	± 2 kV 100 kHz spike frequency
Surges acc. to IEC 61000-4-5	\pm 0.5 kV, \pm 1 kV conductor/ conductor \pm 0.5 kV, \pm 1 kV, \pm 2 kV conductor/ground	\pm 0.5 kV, \pm 1 kV conductor/ conductor \pm 0.5 kV, \pm 1 kV, \pm 2 kV conductor/ground

Electromagnetic immunity				
	3 V	3 V		
Conducted interference induced	0.15 MHz to 80 MHz	0.15 MHz to 80 MHz		
by high-frequency fields acc. to	6 V in the ISM bands between	6 V in the ISM bands between		
IEC 61000-4-6	0.15 MHz and 80 MHz	0.15 MHz and 80 MHz		
	80 % AM at 1 kHz	80 % AM at 1 kHz		
	0 % U _T ; 0.5 cycle	0 % U _T ; 0.5 cycle		
	At 0°, 45°, 90°, 135°, 180°, 225°,	At 0°, 45°, 90°, 135°, 180°, 225°,		
Voltage ding chart interruptions	270° and 315°	270° and 315°		
Voltage dips, short interruptions and voltage fluctuations acc. to IEC 61000-4-11	0 % U _T ; 1 cycle	0 % U _T ; 1 cycle		
	and	and		
	70 % U _T ; 25/30 cycles	70 % U _T ; 25/30 cycles		
	Single phase at 0°	Single phase at 0°		
	0 % U _T ; 250/300 cycles	0 % U _T ; 250/300 cycles		

Mobile HF communications equipment

Mobile HF o	Mobile HF communications equipment					
Test frequency in MHz	Frequency band in MHz ^{a)}	Radio service ^{a)}	Modulation ^{a)}	Maximum power output in W	Distance in m	Interfer- ence immunity test level in V/m
385	380 to 390	TETRA 400	Pulse modulation ^{b)} 18 Hz	1.8	0.3	27
450	430 to 470	GMRS 460 FRS460	FM ^{c)} ± 5 kHz deviation 1 kHz sine	2	0.3	28
810		GSM 800/900				
870	800 to 960	TETRA 800 iDEN 820	Pulse modulation ^{b)}	2	0.3	28
930	800 10 960	CDMA 850 18 Hz LTE Band 5	2	0.5	20	
1720		GSM 1800				
1845		CDMA 1900				
1970	1700 to 1900	GSM 1900 DECT LTE Band 1, 3, 4, 25 UMTS	Pulse modulation ^{b)} 217 Hz	2	0.3	28

Test frequency in MHz	Frequency band in MHz ^{a)}	Radio service ^{a)}	Modulation ^{a)}	Maximum power output in W	Distance in m	Interfer- ence immunity test level in V/m
2450	2400 to 2570	Bluetooth [®] WLAN 802.11 b/g/n RFID 2450 LTE Band 7	Pulse modulation ^{b)} 217 Hz	2	0.3	28
5240 5500 5785	5100 to 5800	WLAN 802.11 a/n	Pulse modulation ^{b)} 217 Hz	0.2	0.3	9
Note: If necessary, the distance between the transmitting antenna and MEDUCORE Standard ² can be reduced to 1 meter to achieve the interference immunity test level. The 1 meter test distance is permitted according to IEC 67000-4-3. ^{a)} For some radio services, only the frequencies for the uplink have been included in the table. ^{b)} The HF carrier must be modulated with a square wave with a 50 % signal ratio.						

^{c)} As an alternative to frequency modulation (FM), pulse-width modulation with a 50 % signal ratio at 18 Hz can be used, as this would represent the worst-case scenario even if it is not the actual modulation.

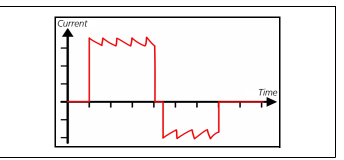
Further technical data can be obtained from the manufacturer WEINMANN Emergency on request.

18.18 The CARDIObiphasic shock pulse

A characteristic of the CARDIObiphasic shock pulse is that it limits the maximum current. This greatly reduces the risk of myocardial damage, which is mainly caused by excessively high electric currents, especially where patient impedance is low.

18.18.1 Functional principle

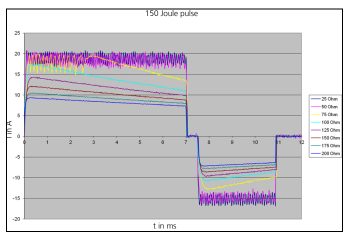
A setpoint controls the delivered current, producing a sawtoothshaped pulse.



18-1 Basic shape of the CARDIObiphasic shock pulse

The mean ratio of electric charge delivered between the second (negative) and the first (positive) phase is 0.38.

For safety reasons, voltages of no higher than 2000 V are used. The resulting currents as a function of patient impedance are shown in the following graph by way of example.



18-2 Currents as a function of patient impedance

18.18.2 Accuracy of the delivered energies

The shock pulse is generated on a current-controlled basis (I = const.) and for a fixed duration (t = const.). Patient impedance is the only variable. The shock pulse energy is the result of current, pulse duration and patient impedance. Since current and pulse duration are specified, the shock pulse can be individually adapted for each patient.

The current control of the shock pulse ensures that patients with low impedance receive the same current as patients with high impedance since it is not energy but current which is key for defibrillation. Furthermore, current control prevents patients with a low patient impedance from being shocked with an overly high current.

Due to current control and the dependency on patient impedance of the energy delivered, it is possible that the energy delivered will deviate from the selected energy (see table below):

	Energy delivered as a function of patient impedance*							Accuracy of energy	
Selected energy in J	Patient impedance in Ω								
energy in J	25	50	75	100	125	150	175	200	delivered
1	0.4	1	2.5	2.4	2	1.8	1.7	1.6	±3 J
5	2.6	5.4	7.8	6.8	6	5.4	4.9	4.4	±3 J
10	5.2	10.8	13.2	11.6	10.2	9.3	8.4	7.6	±3 J
15	7.8	15.7	20.4	18.4	16.5	15.3	13.6	12.4	±3 J
20	10.2	19.9	25.6	23.4	21	19.2	17.5	16	±15 %
30	15.4	30.2	42.1	40.2	36.7	33.9	31.1	28.8	±15 %
40	19.8	39	51.4	49.8	46	42.3	38.8	36.4	±15 %
50	26	50.2	72.1	72	65.7	60.3	55.6	51.6	±15 %
75	38.4	73.8	106.6	107.7	100.7	92.9	86	79.9	±15 %
100	53	102	135.6	134.1	123.2	113.3	104.9	96.3	±15 %
120	63.6	122.3	153.7	148.3	137.2	127.1	116.8	107.9	±15 %
150	79.6	152	181.7	170.9	157.7	146.3	134.7	126.7	±15 %
200	128.9	201.2	211.6	208.3	205.6	196	187.5	179.5	±15 %

*Selected energy differs from delivered energy due to the laws of physics. Delivered energy depends on patient impedance (see also IEC 60601-2-4:2010 Chapter 201.12.1 and Annex AA).

19 Labeling

Symbol	Description
REF	Article no.
SN	Serial number
SN	
LOT	Production batch number
	Unique Device Identifier (UDI): Allows individual products to be uniquely identified in the market
MD	Medical device
	Manufacturer
M	Date of manufacture
<u>[]</u>	Consult instructions for use
	Refer to instructions for use
\triangle	Attention
€ €€ 0197	CE mark (confirms that the product complies with the applicable European directives)
F©	FCC mark (confirms that the product has a market license in the USA for devices with wireless technologies)
No.	KC mark (confirms that the product complies with applicable South Korean directives)

Symbol	Description
(5) (11)	China RoHS label (confirms that the product does not emit toxic substances for the number of years indicated)
(1) (1) (1) (1) (1) (1) (1) (1) (1) (1)	Safety check label (STK, only applies to Germany): Indicates when the next Technical Safety Check ("Sicherheitstechnische Kontrolle" in accordance with § 11 of the German regulation MPBetreibV) is required.
1112/1 9 MTK 8 2000 4 6 5 2000 4	Metrological check label (only applies to Germany): Indicates when the next metrological check in accordance with § 14 of the MPBetreibV is required.
2	Disposable item, do not reuse
	Storage temperature range limits
	Storage humidity range limits
	Air pressure range limits
Ť	Keep dry
Ţ	Fragile
\Box	Expiry date
X	Latex-free
NON	Non-sterile
X	Do not dispose of in household waste
- † -	Defibrillation-proof Type BF applied part
-	Defibrillation-proof Type CF applied part

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Symbol	Description			
	Direct voltage			
	Type of protection against electric shock: Protection class II device			
IP55	Degree of protection against			
	Ingress of solid objects			
IPX1	 Ingress of dust Ingress of water with harmful effect 			
IPX7	Degree of protection against temporary immersion in water			
	Degree of protection against temporary immersion in water Degree of protection against water dripping at an angle, 15° relative to the normal			
IPX2	operating position			
	Input voltage			
	Output voltage			
\rightarrow	Input (12 V - 15.1 V, 30 W)			
12-15,1 V-	Input voltage (12 V - 15.1 V)			
	Line power indicator: The LED comes on when the charging station is connected to line power.			
	Battery fault, if red fault indicator on			
0000	Battery status			
CLICK	Audibly engage battery			
Pad	Connection for master cable			
SpO2	Connection for pulse oximetry sensor connecting cable			
ECG	Connection for ECG cable			
NIBP	Connection for NIBP connecting tube			
E _{max} = 200 J	Maximum energy delivered			

19 Labeling

Symbol	Description
CHARGE	Marks the button which can be used to charge the shock capacitor.
Яноск	Marks the two buttons which have to be pressed simultaneously for shock delivery.
APEX	Marks the paddle which has to be held at the apex position.
STERNUM	Marks the paddle which has to be held at the sternum position.
(\mathbf{b})	On/Off button
	ng, see the following instructions for use: on electrodes odes

20 Warranty

Starting from the date of purchase, WEINMANN Emergency offers the customer a limited manufacturer's warranty on a new original WEINMANN Emergency product or spare part installed by WEINMANN Emergency in accordance with applicable warranty terms and conditions for the particular product and the warranty periods listed below. The warranty terms and conditions are available on the Internet at www.weinmann-emergency.com. On request, we will also send you the warranty terms and conditions.

If you wish to make a warranty claim, consult your authorized dealer.

Product	Warranty periods
WEINMANN Emergency devices, including accessories (excluding masks) for oxygen therapy and emergency medicine	2 years
Masks including accessories, batteries (unless otherwise stated in the technical documentation), sensors, breathing circuits	6 months
Disposable products	None

21 EC Declaration of Conformity on Medical Devices

WEINMANN Emergency Medical Technology GmbH + Co. KG hereby declares that the product complies fully with the respective regulations of the Medical Device Directive 93/42/EEC. The unabridged text of the EC Declaration of Conformity on Medical Devices can be found on our website at www.weinmannemergency.com.



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Ø Made in Germany