Instructions for use

Babytherm 8004/8010

WARNING
To properly use this medical device, read and comply with these instructions for use.

Open intensive care, resuscitation
Typographical conventions

1 Consecutive numbers indicate steps of action, with the numbering restarting with "1" for each new sequence of actions.

- Bullet points indicate individual actions or different options for action.
  - Dashes indicate the listing of data, options, or objects.

A Letters in illustrations denote elements referred to in the text.

Trademarks

<table>
<thead>
<tr>
<th>Trademark</th>
<th>Trademark owner</th>
</tr>
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<tr>
<td>Babytherm®</td>
<td>Dräger</td>
</tr>
<tr>
<td>ThermoPad® IM</td>
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<tr>
<td>BillLux®</td>
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</tr>
<tr>
<td>Scale-Tronix®</td>
<td>Welch Allyn</td>
</tr>
<tr>
<td>Dismozon®</td>
<td>Bode Chemie</td>
</tr>
<tr>
<td>Aclichlor®</td>
<td>Ecolab</td>
</tr>
<tr>
<td>BruTab 6S®</td>
<td>Brulin</td>
</tr>
<tr>
<td>Klorsept®</td>
<td>Medentech</td>
</tr>
<tr>
<td>Descogen®</td>
<td>Antiseptica</td>
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<td>Dismozon®</td>
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<td>Ecolab USA</td>
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<tr>
<td>Perform®</td>
<td>Schülke &amp; Mayr</td>
</tr>
<tr>
<td>Mikrozid®</td>
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<td>Buraton®</td>
<td></td>
</tr>
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</table>

Duties of the operating organization

The tasks described in this document specify the requirements that have to be met by each respective target group.

The operating organization of this product must ensure the following:

- The target group has the required qualifications (e.g., has undergone specialist training or acquired specialist knowledge through experience).
- The target group has been trained to perform the task.
- The target group has read and understood the chapters required to perform the task.
Description of the target groups

The target groups may only perform the following tasks if they meet the corresponding requirements.

Users

<table>
<thead>
<tr>
<th>Task</th>
<th>Requirement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Use of the product in accordance with the</td>
<td>Specialist medical knowledge in the use of the</td>
</tr>
<tr>
<td>intended use</td>
<td>product</td>
</tr>
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</table>

Service personnel

<table>
<thead>
<tr>
<th>Task</th>
<th>Requirement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Installation</td>
<td>Specialist knowledge in electrical engineering</td>
</tr>
<tr>
<td>Basic service work</td>
<td>mechanics</td>
</tr>
<tr>
<td>(inspection, maintenance according to the</td>
<td>Experience in the servicing of medical devices</td>
</tr>
<tr>
<td>&quot;Maintenance&quot; chapter)</td>
<td>Experience in complex service work on this</td>
</tr>
<tr>
<td></td>
<td>product</td>
</tr>
</tbody>
</table>

Dräger recommends arranging a service contract with DrägerService.

Specialized service personnel

<table>
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<tr>
<th>Task</th>
<th>Requirement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Installation</td>
<td>Specialist knowledge in electrical engineering</td>
</tr>
<tr>
<td>Basic and complex service work (inspection,</td>
<td>mechanics</td>
</tr>
<tr>
<td>maintenance, repair)</td>
<td>Experience in the servicing of medical devices</td>
</tr>
<tr>
<td></td>
<td>Experience in complex service work on this</td>
</tr>
<tr>
<td></td>
<td>product</td>
</tr>
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</table>

Safety information definitions

**WARNING**
A WARNING statement provides important information about a potentially hazardous situation which, if not avoided, could result in death or serious injury.

**CAUTION**
A CAUTION statement provides important information about a potentially hazardous situation which, if not avoided, may result in minor or moderate injury to the user or patient or in damage to the medical device or other property.

**NOTE**
A NOTE provides additional information intended to avoid inconvenience during operation.

Instructions for use Babytherm 8004/8010
Abbreviations and symbols

For explanations refer to the “Overview” chapter, "Abbreviations" and "Symbols" sections.
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For your safety and that of your patients

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For your safety and that of your patients

General safety information

The following WARNING and CAUTION statements apply to general operation of the medical device.

WARNING and CAUTION statements specific to subsystems or particular features of the medical device appear in the respective sections of these Instructions for Use or in the Instructions for Use of another product being used with this device.

Strictly follow the instructions for use

WARNING
Risk of incorrect operation and of incorrect use
Any use of the medical device requires full understanding and strict observation of all sections of these instructions for use. The medical device may only be used for the purpose specified under "Intended use" on page 15 and in conjunction with appropriate patient monitoring (see page 10). Strictly observe all WARNING and CAUTION statements throughout these instructions for use and all statements on medical device labels. Failure to observe these safety information statements constitutes a use of the medical device that is inconsistent with its intended use.

Service

WARNING
Risk of medical device failure and of patient injury
The medical device must be inspected and serviced regularly by service personnel. Repair and complex maintenance carried out on the medical device must be performed by experts. If the above is not complied with, medical device failure and patient injury may occur. Observe chapter "Service".

Safety checks

The medical device must be subject to regular safety checks. See chapter "Service".

Metrological checks*

The device must be subjected to regular metrological checks as per the medical device operator ordinance (MPBetreibV), see chapter "Service".

* Applies only to the Federal Republic of Germany
For your safety and that of your patients

Accessories

**WARNING**
Risk due to incompatible accessories

Dräger has tested only the compatibility of accessories listed in the current list of accessories. If other, incompatible accessories are used, there is a risk of patient injury due to medical device failure.

Only use the medical device with accessories listed in the current list of accessories, see "Order list" on page 123.

Not for use in areas of explosion hazard

**WARNING**
Risk of fire

The medical device is not approved for use in areas where combustible or explosive gas mixtures are likely to occur.

Safe connection with other electrical equipment

**WARNING**
Risk of patient injury

Electrical connections to equipment not listed in these instructions for use or the assembly instructions may only be made if approved by the respective manufacturer.

Connection to other devices

Device combinations (Dräger devices + Dräger devices or Dräger devices + third-party devices) approved by Dräger (see instructions for use of individual devices) meet the requirements of the following standards:

- IEC 60601-1 (3rd edition)
  Medical electrical equipment
  Part 1-1: General requirements for safety and essential performance

- IEC 60601-1-2
  Medical electrical equipment
  Part 1-2: General requirements for safety and essential performance
  Collateral standard: Electromagnetic compatibility; Requirements and tests

- IEC 60601-1-8
  Medical electrical equipment
  Part 1-8: General requirements for safety
  Collateral standard: General requirements, tests, and guidance for alarm systems in medical equipment and medical electrical systems

If a device combination is not approved by Dräger, proper operation of the devices can be compromised.

The operator must ensure that the device combination meets the applicable standards.

Strictly observe instructions for use and assembly instructions of all connected devices.
Information on electromagnetic compatibility

General information on electromagnetic compatibility (EMC) according to international EMC standard IEC 60601-1-2:

Medical electrical equipment is subject to special precautionary measures concerning electromagnetic compatibility (EMC) and must be installed and put into operation in accordance with the EMC information provided (see page 113).

Portable and mobile high-frequency communication equipment can affect medical electrical equipment.

Patient monitoring

The user of the medical device is responsible for choosing a suitable patient monitoring system that provides appropriate information on medical device performance and patient condition.

Patient safety can be achieved by a wide variety of means ranging from electronic surveillance of medical device performance and patient condition to direct observation of clinical signs.

The responsibility for selecting the best level of patient monitoring lies solely with the user of the medical device.

Functional safety

The essential performance features are as follows:

For the radiant heater

– Skin temperature mode
  – The heating levels are controlled based on the difference between the target value set for the skin temperature and the measured skin temperature.
  – If a maximum temperature difference of 0.5 °C is reached, a visual and audible alarm is triggered.

For the mattress heater

– The mattress temperature is controlled based on the difference between the set target value and the measured mattress temperature.
  – If the temperature difference is more the 1 °C or the temperature exceeds 41 °C, a visual and audible alarm is triggered.

The medical device is equipped with basic safety features to reduce the possibility of patient injury while the cause of an alarm is remedied.

CAUTION

Risk of patient injury

Do not make therapeutic decisions based solely on individual measured values and monitoring parameters.
**Product-specific safety information**

<table>
<thead>
<tr>
<th>WARNING</th>
<th>WARNING</th>
</tr>
</thead>
<tbody>
<tr>
<td>Risk of injury</td>
<td>Risk of injury</td>
</tr>
<tr>
<td>Blockage, damage, and foreign objects may lead to injuries.</td>
<td>Simultaneous use of a skin temperature sensor and a defibrillator may result in device malfunction and unsuccessful use of the defibrillator.</td>
</tr>
<tr>
<td>Prior to installing, check all system components for blockage, damage, and foreign objects and replace if necessary.</td>
<td>Remove the skin temperature sensor from the patient before using the defibrillator.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>WARNING</th>
<th>CAUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Do not use the medical device near high-frequency surgical equipment or tomographs (e.g., MRI, NMT)!</td>
<td>It is the responsibility of the doctor to draw conclusions from the skin temperature measurements taken.</td>
</tr>
<tr>
<td>The functional integrity of the medical device may be impaired and the patient put at risk.</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>WARNING</th>
<th>WARNING</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baby control mode must not be used with children who are in shock or who have fever.</td>
<td>Risk of overheating of the radiant warmer</td>
</tr>
<tr>
<td></td>
<td>Risk of overheating of the radiant warmer</td>
</tr>
<tr>
<td></td>
<td>If the radiant warmer is continuously operated at high output, e.g., in skin temperature mode with a temperature sensor and without a patient, the life span of the infrared radiator may be reduced.</td>
</tr>
<tr>
<td></td>
<td>Do not operate the device in skin temperature mode with a temperature sensor connected and without a patient.</td>
</tr>
<tr>
<td></td>
<td>Only use the integrated scale to determine the weight of the patient.</td>
</tr>
<tr>
<td></td>
<td>Failure to follow the instructions for use can lead to severe inaccuracies in measuring the patient's weight.</td>
</tr>
<tr>
<td></td>
<td>To safeguard critical therapeutic decisions based on the patient's weight, the weighing result should be checked against a reference measurement on an external scale.</td>
</tr>
<tr>
<td></td>
<td>Risk of malfunction</td>
</tr>
<tr>
<td></td>
<td>Unallowed modifications to the medical device lead to malfunctions.</td>
</tr>
<tr>
<td></td>
<td>This medical device may not be changed without permission from Dräger.</td>
</tr>
</tbody>
</table>
For your safety and that of your patients

**WARNING**
Risk of crushing
Movable device parts or attached components may cause crushing due to clamping.
Pay special attention to edges, movable parts, and corners when working with the following components:
- Column cover
- Drawers
- Swivel arms for mounted devices
- Radiator housing
- Height adjustment between trolley and drawer

**WARNING**
Risk of patient hyperthermia or hypothermia
Set the patient temperature according to the patient’s needs.
Check that the temperature sensor is correctly positioned and secured.

**WARNING**
Risk of tipping over during transport
The medical device may tip over if handled incorrectly.
- The medical device may only be moved by people who have the physical ability to do so.
- Lower the trolley to its minimum height.
- Adjust the bed to a horizontal position.
- Do not let the accessories protrude.

**WARNING**
Risk of incorrect medication
The heating of transdermal medication (patches) can result in the administration of higher doses of active ingredient.
When using transdermal medication, the patient must be monitored at close intervals.

**CAUTION**
Damage to materials due to disinfectants
Disinfectants can damage disposable products.
This may result in incorrect measurements.
Do not disinfect disposable products.

**CAUTION**
Risk of patient injury
The mattress may shift when inclined.
Avoid inclining the mattress excessively and do not wrap the sheet around the mattress.

**WARNING**
Risk of damage to the device and patient injury
Reuse, reprocessing, or resterilization of disposable products can lead to a failure of the medical device and cause injury to the patient.
Do not reuse, reprocess, or resterilize disposable products. Disposable products are designed, tested, and manufactured exclusively for single use.

**CAUTION**
Risk of patient injury
All data transferred through the MEDIBUS interface are for information only and must not be used as the sole basis for diagnostic or therapeutic decisions. The data accessible via this interface are not intended for distributed alarm systems conforming to IEC 60601-1-8 (in the context of remote monitoring systems).
For your safety and that of your patients

**WARNING**
Risk due to liquid penetrating into device and radiant warmer

Penetrating liquid may cause the following:
- Damage to the device
- Electric shock
- Device malfunctions

Ensure that no liquid penetrates the device.

**CAUTION**
To avoid damage to the device electronics, comply with the following:
- Only switch the device on and off with the On/Off switch.
- Do not operate the device outside the specified voltage range.
- Do not operate the device outside the specified ambient temperature range.
- Do not obstruct the ventilation slots on the rear of the control unit.
- Do not place objects that may impede cooling on the base plate above the voltage input module.
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**Intended use**

Babytherm 8004/8010 is an open care unit for warming premature babies, neonates and infants with a body weight of up to 8 kg. The unit may be used in operating theaters, neonatal wards, children’s wards, obstetric units and paediatric intensive care units. The unit may be operated by clinical staff or by nursing staff under the supervision of a doctor.

**Therapy and nursing uses**

- In intensive care and obstetric units for resuscitation
- Normal and intensive care
- For infant warming and compensation of heat loss
- Thermal stability with mattress heating
- Gentle heat therapy with a combination of radiant warmer and mattress heating
- Cooling patients with fever (temperature of mattress below core temperature)
- Adjustable bed angle for head-up or head-down position
- Weaning infants from incubators
- O2 and nebuliser therapy (with appropriate accessories)
- Lowering the bilirubin levels when using the phototherapy option.

**Contraindications**

- In patients suspected of malignant hyperthermia.

**WARNING**

Risk of explosion and fire

This medical device is neither approved nor certified for use in areas where oxygen concentrations greater than 25 Vol%, combustible or explosive gas mixtures are likely to occur.

Risk of device malfunctions and/or patient injury and user injury

Magnetic fields can negatively influence the correct functioning of the medical device and therefore endanger the patient or user.

Do not use the medical device near nuclear magnetic resonance imaging (MRI, NMR, NMI).
Application

**Babytherm 8004 heating features**

- Radiant warmer operation with a preset heat output:
  
  the radiant warmer heat output is controlled in manual mode and therapy time at higher heat output levels is monitored.

- Radiant warmer operation with baby control mode:
  
  temperature is automatically monitored and controlled.

**Babytherm 8010 heating features**

- Radiant warmer operation with a preset heat output:
  
  The radiant warmer heat output is controlled in manual mode and therapy time at higher heat output levels is monitored.

- Radiant warmer operation with baby control mode:
  
  temperature is automatically monitored and controlled.

- Heated gel mattress, operating independently of the radiant warmer:
  
  the temperature of the heating pads is automatically monitored and adjusted to attain and maintain a preselected temperature at the contact surface between the infant and the mattress.
Options

- Height adjustment
- One or two swivel cabinets
- Side panels (150 mm or 230 mm)
- Infusion holder
- Phototherapy
- Bed canopy
- Swivel tray
- RS 232 interface

Scale

It is designed to determine the weight of the patient with the bed in a horizontal or inclined position.

MEDIBUS

Software protocol for the transfer of data between Babytherm 8004/8010 and an external medical or non-medical device (e.g., patient monitors or computers for data management systems) via an RS 232 port, see "MEDIBUS for Dräger Pediatric Devices" (9029205).

WARNING

All transferred data are for information only and should not be used as basis for diagnostic or therapeutic decisions.

WARNING

Risk of injury for patients and users

All systems consisting of medical devices and other electrical equipment and which are not computers, printers, etc., must be assembled by service personnel.

The system must meet the requirements of the IEC/EN 60601-1-1 and IEC/EN 60601-1-2 standards.
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Instructions for use Babytherm 8004/8010

1 Radiant warmer
2 End panel
3 Side panel
4 Inner panel
5 Handle with inner release bar for tilting the bed
6 Swivel cabinet, optional (left and/or right)
7 Chassis with 4 castors, 2 of which are lockable
8 Height-adjustable column, optional
9 Foot controls for height-adjustable column (8)
10 Connection for skin temperature sensor:
   top socket, yellow – for measuring/controlling
   skin temperature; lower socket, white – for
   measuring peripheral skin temperature
11 Connection for equipotential bonding
12 Identification and rating plate
13 Accessory mounting rail
14 Bed assembly
15 Gel mattress for use with mattress heater; foam
   mattress for use without mattress heater
16 Hose grommets
17 Stand, with shelf and 2 side rails
18 Control unit with operating panel
19 Swivel joint for radiant warmer, ±90°
20 X-ray tray at rear – Babytherm 8004 only
21 Bed canopy
Overview

Radiant warmer

1 Handle
2 Red central alarm lamp*
3 Heater elements
4 Procedure light/night light
5 Phototherapy lights (optional), set of 6

* Country-specific function (e.g., not available in China)
Overview

Control unit

1. **ON/OFF indicator:**
The green LED is lit when the unit is ON. The red LED is lit following a power failure.

2. **OK button for confirming messages and settings.** If the OK button is not pressed within 10 seconds of entering a new setting, the previous settings remain active.

3. **Red LED Inop.:** indicates device malfunction.

4. **Button for checking LEDs, displays, horn and audible alarm.**

5. **Button for muting the audible alarm.**

6. **Yellow alarm LED:** lit for Caution alarm level.

7. **Red alarm LED:** flashes for Warning alarm level.

8. **Text display:** provides advisory messages for the user and prompts the user to confirm/acknowledge.

9. **Display and control panel for mattress heater, Babytherm 8010**

10. **Display and keypad for operating the radiant warmer.**

11. **On/Off buttons for procedure light; yellow LED in button is lit when procedure light is on.**

12. **On/Off button for night light; yellow LED in button is lit when night light is on.**

13. **On/Off button for phototherapy; yellow LED in button is lit when phototherapy lights are on (blue background).**

14. **On/Off button for radiant warmer; yellow LED in button is lit when radiant warmer is on.**
Overview

Display and control panel for mattress heater, Babytherm 8010

1 Display of measured (actual) mattress temperature
2 Indicator for upper extended range >38 °C
3 Indicator for lower extended range <36 °C
4 Keys (buttons) for setting the desired mattress temperature
5 Displayed of set (desired) mattress temperature

Display and control panel for the radiant warmer

1 Display of measured temperature for skin temperature
2 Foot symbol for peripheral temperature; lit when the peripheral skin temperature sensor is connected.
3 Display of the measured value for peripheral skin temperature; only possible when the peripheral skin temperature sensor is connected.
4 Display of radiant warmer heating levels; levels 1 to 3: green, levels 4 to 10: yellow.
5 Keys (buttons) for setting the radiant warmer output level
6 Button for toggling between man. (manual mode) and Skin (skin temperature mode); the yellow LED lights up according to the selected mode.
7 Keys (buttons) for setting the desired skin temperature
8 Display for the skin temperature setting
Overview

Controller, rear view

1 Inputs for connecting the Nurse call from up to three equipment units at the workstation (connections for central alarm)

2 Output for connection to internal paging systems, Nurse call (connection for central alarm)

3 RS 232 interface (MEDIBUS), optional

4 RS 232 interface (modem), optional

5 Controller ON/OFF switch; □ = OFF, ○ = ON.

6 Cover
Overview

Scale

The scale in the Babytherm consists of a weighing unit located under the bed and a display unit.

**Scale display unit**

1. Screen for displaying the weight
2. Measuring unit indicator for the weight display
3. Indicator for lifting the patient
4. Indicator for lowering the patient
5. *Weigh* button for starting the weighing operation with zero calibration
6. *Reweight* button for starting the weighing operation without zero calibration
7. *kg* button for displaying the weight in kilograms
8. *Lb* button for displaying the weight in pounds*
9. *Start/Standby* button for selecting standby or operating mode

---

* The verifiable scale can only display kg.
## Symbols

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Explanation</th>
<th>Symbol</th>
<th>Explanation</th>
</tr>
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<tbody>
<tr>
<td>🚨</td>
<td>Warning! Strictly follow these instructions for use</td>
<td>🌱</td>
<td>Do not attach oxygen sources here (minimum distance: 20 cm)</td>
</tr>
<tr>
<td>⚡</td>
<td>ESD warning label</td>
<td>⛈</td>
<td>Protect from moisture</td>
</tr>
<tr>
<td>🛠️</td>
<td>Disposal instructions</td>
<td>📋</td>
<td>Fragile, handle with care</td>
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<tr>
<td>🖐️</td>
<td>Height adjustment for intermittent operation</td>
<td>🤼</td>
<td>Quantity</td>
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<tr>
<td>🤹‍♂️</td>
<td>Temperature limitation</td>
<td>🛋️</td>
<td>MR unsafe</td>
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<tr>
<td>🌡️</td>
<td>Atmospheric pressure</td>
<td>🍃</td>
<td>Do not use this equipment in the vicinity of magnetic resonance scanners.</td>
</tr>
<tr>
<td>💧</td>
<td>Relative humidity</td>
<td>🌈</td>
<td>Weight: main devicet</td>
</tr>
<tr>
<td>🔔</td>
<td>Button for suppressing the acoustic alarm signal and for confirming alarms</td>
<td>🔔</td>
<td>Weight: load</td>
</tr>
<tr>
<td>🔥</td>
<td>Attention! (safety sign)</td>
<td>🔴</td>
<td>Polyethylene with low density, class 4</td>
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<tr>
<td>💥</td>
<td>Risk of crushing (hand)</td>
<td>⚠️</td>
<td>Potential equalization connector</td>
</tr>
<tr>
<td>💥</td>
<td>Risk of crushing (foot)</td>
<td>🚨</td>
<td></td>
</tr>
<tr>
<td>🚨</td>
<td>Note the conditions for tipping stability, see &quot;Use of accessories&quot; on page 27</td>
<td>📚</td>
<td></td>
</tr>
<tr>
<td>🕵️‍♀️</td>
<td>Use light-proof eye cover</td>
<td>🕵️‍♀️</td>
<td></td>
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<tr>
<td>🕵️‍♂️</td>
<td>Follow instructions for use</td>
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<td>🏛️</td>
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<tr>
<td>📅</td>
<td>Date of manufacture</td>
<td>🕒</td>
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</tr>
<tr>
<td>📞</td>
<td>Revision index</td>
<td>📞</td>
<td></td>
</tr>
<tr>
<td>🏆</td>
<td>Part number</td>
<td>🏆</td>
<td></td>
</tr>
<tr>
<td>🏆</td>
<td>Serial number</td>
<td>🏆</td>
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Instructions for use Babytherm 8004/8010

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*Overview*
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Use of accessories

WARNING
Risk of tipping over
Stability may be impaired by the attachment of additional accessories/loads. In order to ensure stability (IEC 60601-1), a one-sided accessory load of 25 Nm1) must not be exceeded when tilted at 10°.

1) corresponds to approx. 5 kg at a distance of 20 cm from the stand column, at a height of 2.00 m when tilted at 10°.

Ensuring stability
Stability is ensured when the unit is placed on a surface with a slope of not more than 10°.

The following restrictions apply when adding accessories:

- Do not exceed the maximum load-bearing capacity of the stand = 30 kg.
- If accessories are positioned on a column of the stand, e.g., on two swivel trays, the maximum load for this column is 5 kg.

Recommended measures for increasing stability
Attach accessory loads as low and as close to the stand columns as possible and avoid one-sided loading.

1 Stand, consisting of 2 stand columns
2 Shelf
3 Standard rail on stand column
4 Compact rail
5 Standard rail, rear
6 Base plate
7 Swivel cabinet
To ensure stability*, observe the following restrictions when using accessories:

- Do not exceed a maximum patient weight of 8 kg
- Observe a maximum weight limit of 20 kg for the shelf and a maximum mounting height of 40 cm above the base plate
- Maximum loading of 2.4 kg for a swivel cabinet, corresponding to a maximum total weight of 9 kg
- Mount swivel cabinets on opposite sides, if present.

When attaching one-sided stand column accessories without swivel cabinets

Maximum permissible one-sided accessory load on the stand column:
- 5 kg at a height of 200 cm and at a distance of 20 cm

When attaching swivel cabinets and stand column accessories

1 For one-sided mounting of the swivel cabinet

Maximum permissible one-sided accessory load on the stand column:
- 1 kg at a height of 200 cm and at a distance of 20 cm
  or
- 2 kg at a height of 130 cm and at a distance of 10 cm

Example:
1.5 kg (approx. 1 L of infusion liquid including bracket) at a distance of 20 cm from the stand column, at a height of 180 cm corresponds to a tilting moment of approx. 7 Nm (at an inclination of 10°).

2 For opposite/diagonal mounting of swivel cabinets

Maximum permissible one-sided accessory load on the stand column:
- 2.5 kg at a height of 200 cm and at a distance of 20 cm
  or
- 4.5 kg at a height of 130 cm and at a distance of 10 cm

3 For same-sided mounting of swivel cabinets

Maximum loading of 2.4 kg = maximum total weight of 9 kg per cabinet
- No one-sided accessory load on the stand column
- Maximum loading of the shelf restricted to 2.5 kg

Maximum load

- Individual wheel load < 100 kg
- Tilting moment < 25 Nm
- Lifting columns/standard housing < 50 kg

of which

- Weight of patient < 8.0 kg
- Swivel cabinet < 2.4 kg
- Shelf < 20 kg
- Stand column < 5.0 kg

Attach accessory loads as low and as close to the stand columns as possible and avoid one-sided loading

Example:
1.5 kg (approx. 1 L of infusion liquid including bracket) at a distance of 20 cm from the stand column, at a height of 180 cm corresponds to a tilting moment of approx. 7 Nm (at an inclination of 10°).

* IEC 60601-1 tilt stability ensured up to an inclination of 10°
**Shelf**

The unit is fitted with one shelf.

Make sure that the shelf is firmly in position.

1. Screw rail to the left and/or right of the shelf.
   - Place desired auxiliary equipment on the rails and shelf. Max. 20 kg per shelf.
2. Fasten infusion holder to stand column at the required height.

**Compact rail/rail brackets**

This rail is designed to hold auxiliary equipment, e.g. O2 flowmeter, O2 monitor, infusion pumps.

3. Fasten the compact rail(s) to one column of the stand at the required height.
4. Fasten rail brackets and
   - small equipment bar to one column of the stand at the required height.

**Swivel tray**

For small articles and parts. Max. load 1 kg.

- Position the swivel tray on one of the columns of the stand and tighten the hand screw.
  - Recommended height: height of the mattress.
- Make sure that the swivel path is kept clear of obstructions!

**X-ray tray**

Babytherm 8004 only

**NOTE**

Do not use the X-ray tray in the extended position to rest on while writing!
Do not place any objects on the tray!
Do not lean on the tray!
Risk of damage!

**Preparation**

Preparation

2. Lower X-ray tank into the bed – the pegs engage in the holes in the X-ray tank.

- Prepare the warming bed, see page 30.

Warming bed

The bed is enclosed by two side panels and two end panels that can be opened independently of one another.

The side panels are provided with holes for secretion and drainage hoses.

The end panels have flexible grommets for hoses and cables.

Fitting the panels

1. Insert the lower mounting pins of the panel hinge-pieces into the guide slots on either side.

With the panels on the long side, ensure that the handle points outwards.

- Position the panel semi-upright,

2. press the panel down firmly until the pins lock into position at the bottom of the slots.

- Fold the panel up into the vertical position and allow it to drop into the locking position.

- Fit all four panels in this way.

Make sure the panels fit correctly!

To open:

- Lift the panel all the way up until the upper pins come out of their slots, then fold the panel down.

With lively infants or older children:

- Use 230 mm high panels.

Fitting the inner panels

- Insert the pins into the tapered holes in the corners of the housing and press down lightly until the inner panel is firmly in place.

Make sure the panels fit correctly!

WARNING

Inner panels should be used with all patients, regardless of whether the side panels are up or down!

Otherwise the patient may fall out!

009
Preparation

**Tilting the bed**

1. Pull the handle of the locking mechanism out towards the front.
2. Press handle down = head-up position. Pull handle up = head-down position.
   - Release the handle: the bed automatically locks into the selected position.

The bed can be tilted in finely graduated steps.

**Bed canopy**

Optional

Higher side panels (230 mm) are recommended when using the canopy.

- Check that the bed canopy is closed.

When the bed canopy is not required:

- Attach bracket to standard rail.
- Hang bed canopy from bracket.

Maximum tilt angle:
- Head-up position: 20°
- Head-down position: 15°

Preferred positions:
- Horizontal,
- 10° tilt head-up and
- 10° tilt head-down.

The mattress may slip at maximum tilt.

*In this case do not place a sheet over the mattress.*
Preparation

Bronchial aspirator

Attaching the bronchial aspirator to the compact rail

1. Attach the compact rail at the desired position on the stand, see page 29.
2. Clamp the bronchial aspirator to the compact rail.
3. Mount bracket with projecting support to one of the positions provided on the column.
   For units with height adjustment, place the bracket so that the bottle holder is outside the path of the swivel cabinet.
4. Place the bottle holder on the projecting support.
5. Connect hoses.
6. Fit the hose clip (optional) to the compact rail.
7. Clip aspiration hose into the hose clip.
8. Screw on the connecting hose and plug the connector into the socket of the central supply system (park position).

Weight and maximum load of accessories

The total weight is the sum of the dead weight of the accessory and the maximum load.

The information in the following table refers to the dead weight of the accessories and the maximum total weight. The maximum total weight is the sum of the dead weight of the accessory and the corresponding maximum load without being attached to the device. To ensure stability, comply with the loading restrictions on Page 28 when attaching accessories to the device.

<table>
<thead>
<tr>
<th>Accessories</th>
<th>Dead weight</th>
<th>Maximum total weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>Swivel cabinet</td>
<td>6.5 kg (14.3 lbs)</td>
<td>9 kg (19.8 lbs)</td>
</tr>
<tr>
<td>Cabinet fixing</td>
<td>0.5 kg (1.1 lbs)</td>
<td>3 kg (6.6 lbs)</td>
</tr>
<tr>
<td>Swivel tray, compl.</td>
<td>1 kg (2.2 lbs)</td>
<td>3 kg (6.6 lbs)</td>
</tr>
<tr>
<td>Shelf</td>
<td>1 kg (2.2 lbs)</td>
<td>5 kg (11 lbs)</td>
</tr>
<tr>
<td>Infusion holder, compl.</td>
<td>0.5 kg (1.1 lbs)</td>
<td>3 kg (6.6 lbs)</td>
</tr>
<tr>
<td>Drainage canister hook</td>
<td>0.04 kg (0.08 lbs)</td>
<td>2 kg (4.4 lbs)</td>
</tr>
<tr>
<td>Ventilation hose bracket</td>
<td>0.5 kg (1.1 lbs)</td>
<td>1 kg (2.2 lbs)</td>
</tr>
<tr>
<td>Compact rail</td>
<td>0.9 kg (2 lbs)</td>
<td>5 kg (11 lbs)</td>
</tr>
<tr>
<td>Integrated handle, compl. (rear of device)</td>
<td>0.5 kg (1.1 lbs)</td>
<td>5 kg (11 lbs)</td>
</tr>
<tr>
<td>Integrated handle, compl. (for shelf/scale)</td>
<td>0.25 kg (0.55 lbs)</td>
<td>2 kg (4.4 lbs)</td>
</tr>
<tr>
<td>Fairfield rail Babytherm, short</td>
<td>0.25 kg (0.55 lbs)</td>
<td>2 kg (4.4 lbs)</td>
</tr>
<tr>
<td>Fairfield rail Babytherm, long</td>
<td>0.5 kg (1.1 lbs)</td>
<td>5 kg (11 lbs)</td>
</tr>
<tr>
<td>Small equipment pole 500</td>
<td>0.5 kg (1.1 lbs)</td>
<td>13.5 kg (29.8 lbs)</td>
</tr>
</tbody>
</table>
Preparation

### Accessories for O₂ therapy

<table>
<thead>
<tr>
<th>Accessories</th>
<th>Dead weight</th>
<th>Maximum total weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>Small equipment pole 1000</td>
<td>0.9 kg (2 lbs)</td>
<td>13.5 kg (29.8 lbs)</td>
</tr>
<tr>
<td>Rail holder, short</td>
<td>0.3 kg (0.66 lbs)</td>
<td>13.5 kg (29.8 lbs)</td>
</tr>
<tr>
<td>Basket 150</td>
<td>0.3 kg (0.66 lbs)</td>
<td>2 kg (4.4 lbs)</td>
</tr>
<tr>
<td>Basket 300</td>
<td>0.4 kg (0.88 lbs)</td>
<td>2 kg (4.4 lbs)</td>
</tr>
<tr>
<td>Basket 600</td>
<td>0.6 kg (1.32 lbs)</td>
<td>2 kg (4.4 lbs)</td>
</tr>
<tr>
<td>Connecting hose clip (central supply)</td>
<td>0.1 kg (0.22 lbs)</td>
<td>1.5 kg (3.3 lbs)</td>
</tr>
<tr>
<td>Waste bag holder</td>
<td>0.7 kg (1.54 lbs)</td>
<td>2 kg (4.4 lbs)</td>
</tr>
</tbody>
</table>

**WARNING**

Risk of fire

Oxygen reaching the area around the power supply connector will result in an increased risk of fire.

Only attach oxygen cylinders and devices containing oxygen at the points intended for this purpose.

**WARNING**

Risk of patient injury due to operating noise

Administering oxygen may cause additional background noise.

Protect the patient from such background noise.
Preparation

**O2 flowmeter**
- Prepare the flowmeter in accordance with its specific Instructions for Use.

To mount the flowmeter on the compact rail:
1. Press the slider all the way down; hook the flowmeter to the compact rail and release the slider.
2. Screw on the O2 connecting hose.
3. Insert the connector of the connecting hose into the O2 delivery socket and press it all the way in.

**Nebuliser**
- Prepare the nebuliser in conformity with its specific Instructions for Use.
4. Screw the nebuliser securely to the flowmeter.
5. Fit the spiral hose to the nebuliser port.

**O2 monitor**
- Fasten O2 monitor, complete with its holder, to the compact rail.
- Place the O2 sensor on the bed and feed the cable through the hole on the front.
- Plug the connector of the sensor into the measuring unit.

**O2 supply via injector**
When using the injector, the higher side panels (230 mm) and the bed canopy must be used (Page 30 and following pages).

6. Press locking lever on mounting flange and push injector into flange.
- Allow locking lever on injector to engage.
- The injector is now locked.

Connect up hose.
**Always monitor the O2 concentration when using O2 supply.**

**Fitting the hose bracket**
- Insert the hose bracket into the hole in the left side of the head panel and tighten with knurled screw.
Preparation

Drainage canister hook

1 Insert hook horizontally into hole in the Babytherm casing.
2 Pivot hook downwards.

NOTE
The weighing result may be inaccurate if the hook and the scale are used at the same time, see chapter "Using the scale" on page 79.

Storage

Units with height-adjustable column:
– One or two swivel cabinets each with 2 swivel compartments and 1 swivel tray, optional.

Units without height adjustment:
– Two open compartments in the column
– One swivel cabinet with two swivel compartments and one swivel tray, optional.
For clearly organised storage of required equipment and accessories.

Information about the nurse call

With the nurse call, high-priority alarm messages (Warning) are forwarded to a central hospital alarm system. Alarm messages of medium priority (Caution) and low priority (Note) are not forwarded. The nurse call will still be activated if the internal acoustic alarm generator of the medical device is defective.

CAUTION
Risk of failure of nurse call
The transfer of information can be interrupted as a result of a malfunction in a component in the link between the nurse call and the central hospital alarm system.
Check the display on the screen of the medical device regularly.
Preparation

**CAUTION**

Risk of limited patient monitoring

The nurse call provides no meaningful data regarding the function of the medical device or the condition of the patient. Do not use the nurse call as the sole source of alarm information.

- Check the display on the screen of the medical device regularly.
- Check the alarms on the medical device directly.

**In-hospital transport**

**WARNING**

Tipping over of the device

The device may tip over and hurt patients, users, or bystanders due to excessive loading or a change of the center of gravity. The device and accessories may become damaged by tipping.

Do not position the device on inclined surfaces and do not apply weight by pushing, resting or leaning on one side of the device.

Observe the maximum load of all shelves, trays, and fixtures.

Stability may be impaired by attaching additional accessories or loads.

Make sure the device does not tip over during transport due to incorrect positioning of accessories on the standard rail.

- Monitor the patient’s core temperature.
- If optional height adjustment is fitted, lower the unit to its lowest position.
- Swivel the swivel cabinets inwards.
- Swivel the swivel tray inwards.
- Remove X-ray cartridge from X-ray tray (see page 78).

- Accessories projecting beyond either side of the unit must be removed or folded in.
- Switch off heating systems and disconnect the mains plug.
- Fit optional bed canopy to protect patient from draughts.

Immediately after transport:

- Plug in the mains plug and switch on the heating systems.
Testing Readiness for Operation

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Before initial operation

**WARNING**
Risk of electric shock and device malfunction
If the device is connected to a power socket with incorrect mains voltage or without a protective ground, the user can be injured and the device damaged.
Only connect the power cable to a grounded power socket, see “Technical data”.

**WARNING**
Risk of electric shock
The connection of devices to auxiliary power sockets can lead to an increased leakage current. If the protective ground of one of these devices fails, the leakage current may rise above the permissible values. Only connect with the approval of the respective device manufacturer. Have the leakage current checked by service personnel.
If the permissible value is exceeded, use a mains power socket on a wall instead of the auxiliary power socket of the device.

**Mains power supply**
The power supply can be removed from the medical device.

1. Check that the mains voltage matches the values indicated on the rating plate next to the power cable.
2. Plug into the mains.

**Establishing potential equalization**
Even small electrical potential differences between housing surfaces of different devices in the vicinity of the patient can be reduced by potential equalization.

1. Connect one end of the potential equalization cable to a potential equalization pin.
2. Connect the other end of the potential equalization cable to the potential equalization connector on the operating table or the wall. Do not use the connection as a PE connection.
3. Connect additional devices via an additional potential equalization connector with the second potential equalization pin.
Testing Readiness for Operation

Scale

NOTE
To safeguard during transport, the weighing unit inside Babytherm has been protected against vibration and shock. To ensure correct operation, this protection must be removed before using the scale.

Before using the scale:
- Loosen the locking screws (4x) under the bed by about 1 cm.

Before transporting Babytherm:
- Tighten the locking screws (4x) under the bed.

Before each use

- Check that the unit has been disinfected.
- Check that the side panels are locked securely into place. The panel hardware must be visible above the bed surface.
- Check that the side panels are free from cracks and sharp edges.
- Check that the correct mattress is in the cot;
  use with mattress heating: gel mattress (Babytherm 8010),
  use without mattress heating: foam mattress (Babytherm 8004).
- Check that the bed tilts properly and locks securely into position.
- Check that the required accessories and therapy equipment are available and in proper working order.
- Check that the gas supply is available and sufficient for the accessories to be used.
- Check that the cables and hoses are correctly and securely installed. Never route cables or hoses over the panels because they might be pinched or crushed when folding up the panels or fitting the bed canopy.
- Check whether both infrared radiators are providing warmth by placing your hand carefully at a safe distance from the radiant warmer.
Testing Readiness for Operation

Checking height adjustment (optional)

Test height adjustment system if fitted:

1. Briefly press right pedal. The bed is raised.

2. Briefly press left pedal. The bed is lowered.
   - Remember the maximum load for height adjustment is 50 kg.
   - Hoses and cables should be long enough to ensure a secure connection even in the top or bottom height adjustment position.
   - Do not place any objects in the raising/lowering path.
   - Adjust to a comfortable working height.

Switching on and activating the self-test

- Press the On/Off button until it engages = ON. Babytherm now runs a self-test to check important functions.

3. The following message appears on the display:
   *All displays on, horn on*
   All displays are switched on for approx. 2 seconds:
   - All LEDs are lit; the numeric displays show **88.88** and a continuous tone is emitted.

4. The green LED is lit.

5. After about 2 seconds, the unit displays the radiant warmer mode:
   - **man.** (manual mode) – no skin temperature sensor in place
   - **Skin** (baby control mode) – yellow skin temperature sensor for core temperature in place
   - The corresponding LED flashes.

6. The preset desired values are displayed.

7. If the text display reads **Battery discharged** and the yellow LED is lit, the battery is being charged for the power failure alarm. It takes about 30 minutes to charge the battery.
7 The yellow LED is extinguished.
8 If the red Inop. LED = operating fault is lit, see page 89.

Checking LEDs, displays and audible alarm

1 Press the button.
2 The following message appears on the display:
   *All displays on, horn on*
   - All indicators are switched on for approx. 2 seconds:
   - All LEDs are lit; the numeric displays show 88.88 and a continuous tone is emitted.

3 The red LED should light up and a continuous alarm tone should sound.
4 Disconnect the power plug.
5 Reconnect the power. The unit will continue to operate with the values set before the power failure alarm.

Testing the lights

4 Press the button. The bed will be illuminated by the working light.
5 Press the button again. The working light will be switched off.
6 Press the button. The bed will be lit by the night light.
7 Press the button again. The night light will be switched off.
Testing Readiness for Operation

Testing phototherapy (optional)

6 Press the button. The yellow LED in the button will start to flash.

8 The following message appears on the display:

   Photo therapy: Use eye protection
   Press OK button to start XX:XX:XX

7 Press OK button = phototherapy lights switched on.

8 The following message appears on the display for 5 seconds:

   Photo therapy duration XX:XX:XX

6 The yellow LED in the button is now continuously lit.

6 Press button.

8 The following message appears on the display:

   Photo therapy OFF
   Press OK button to confirm XX:XX:XX

7 Press OK button = phototherapy switched off.

6 The yellow LED in the button will go out.
# Operation

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Precautions

Patient care

**WARNING**

Never leave the patient unattended when the side panels are down. Risk of infant falling out of the cot.

When operating the side panels and bed canopy, take care not to pinch any parts of the patient’s body or any hoses or other articles, e.g. bedding.

The side panels must be securely locked in position, and the panel hardware should be visible above the surface of the bed.

Inner panels must be used with all patients, regardless of whether the side panels are open or closed!

With lively infants and/or older children, the 230 mm high side panels should be used.

Observe these infants and/or children particularly closely at all times.

**WARNING**

It is the responsibility of the doctor to draw conclusions from the skin temperature measurements taken.

**CAUTION**

At regular intervals, check whether both infrared radiators are providing warmth by placing your hand carefully at a safe distance from each radiant heater.

**NOTE**

Increased heat is directed to the patient when the heated gel mattress, radiant warmer and phototherapy are operated in combined mode.

This should be taken into account when setting the heating system. Follow the instructions for setting the two heating systems, separately and in combination (Page 59 and following pages). Changes in ambient conditions, e.g. draughts, can affect the patient’s temperature balance.

Heat therapy/phototherapy

**CAUTION**

Risk of damage to the device

Do not bring any heat-sensitive objects or devices into the area heated by the radiant warmer.

The devices might overheat and device malfunctions might occur.

**WARNING**

Monitor the patient’s core temperature regularly using a separate thermometer!

Adjust the temperature settings to the needs of the patient. Consider additional heat sources such as sunlight, examination lights, and phototherapy lights. Do not place any objects between the radiant warmer and the patient.

Mattress heater

**WARNING**

In the Babytherm 8004, use foam mattresses (e.g. SoftBed) only, do not use gel mattresses. Risk of hypothermia.

In the Babytherm 8010 use gel mattresses only, do not use foam mattresses.

Keep clear of sharp objects – risk of damage to the gel mattress.

Do not fold or kink the gel mattress.

To transport the gel mattress, roll it up.

Always adhere to the warm-up time of the gel mattress. Risk of hypothermia.

Wait for about 1 hour before placing a patient in the Babytherm, to allow the gel mattress to warm up sufficiently.
### Operation

**WARNING**
Do not switch off the device while an infant is lying on the Gel mattress. Risk of hypothermia.

**CAUTION**
Check for defects and damage before using SoftBed. Only use an undamaged foam mattress.

**Radiant warmer**

**CAUTION**
The use of the radiant warmer can cause an unnoticed increase in the patient’s water loss.

**WARNING**
Do not place any objects on top of the radiant warmer.
Ventilation would be impeded, and both heater and object may be damaged.

**WARNING**
Do not touch the top of the radiant warmer or the protective screen. Risk of burns!

**CAUTION**
When the bed is tilted, those parts of the patient’s body that are closer to the radiant warmer will receive more heat.
Check the skin temperature of such body parts at regular intervals.

**WARNING**
Do not use flammable cleaning agents or medication when the device is in operation! Risk of fire!
Do not place medication or infusion solutions in the heated area.

**WARNING**
Do not use the baby control mode for infants in a state of shock.
When in shock, skin temperature is much lower than normal.
Infants would be overheated by the baby control mode. For infants in a state of shock, set the heater output to operating mode man. and measure the core temperature every 15 minutes.

**WARNING**
Do not use the baby control mode for infants with fever.
In this case, skin temperature is much higher than normal. Baby control mode in these cases would lead to hypothermia.

**Oxygen therapy**

**WARNING**
The risks to the system are increased when oxygen is in use!
Avoid naked flame and lit cigarettes!
Textiles, plastics and oils are more easily ignited and burn with greater intensity in an oxygen-rich atmosphere.
Keep oxygen fittings and seals free of oil and grease.
Open the valves slowly.
Do not use Babytherm in the presence of flammable anaesthetic gases or disinfectants.
Do not use or store flammable liquids such as alcohol, ether or acetone in the Babytherm.
Do not use electrical devices under the bed canopy. Devices that are approved for operation in potentially explosive areas are excluded.
Operation

Physiological dangers of oxygen

**CAUTION**
Only enrich the O₂ concentration under the instructions of a doctor.
Only do this in accordance with the arterially measured O₂ partial pressure in the patient’s blood. Otherwise there is a danger of hyperoxemia (which may cause damage to the eyes) or hypoxemia (which may cause damage to the brain). During oxygen therapy, the oxygen concentration must be constantly monitored, e.g. with MiniOx 3000.

**NOTE**
Use of the bed canopy in conjunction with operation of the integrated phototherapy system leads to a significant reduction in the bilirubin-effective radiant intensity and a very uneven distribution of the radiation on the bed.

**WARNING**
Risk of excessive exposure to radiation
Do not place a phototherapy light on the bed canopy. The distance to the patient is too small.
Observe the minimum distance of the phototherapy light (see the corresponding instructions for use).

**NOTE**
Higher side panels (230 mm) are recommended when using the canopy.

**WARNING**
When the bed canopy is in place, the side panels can be folded down. Do not leave the Babytherm unattended with the side panels open, as the infant may fall out.

**NOTE**
The bed canopy should not be used as a convenient place to lay objects, clothing, etc.

X-ray tray

**CAUTION**
Do not use the X-ray tray in the extended position to rest on while writing!
Do not place any objects on the tray!
Do not lean on the tray!
Risk of damage!

**NOTE**
Remove X-ray cartridge from the X-ray tray during transport.

Bed canopy

The infant's core temperature should be monitored when the bed canopy is in use.

**CAUTION**
The radiant warmer is less effective when the bed canopy is used. The patient’s core temperature should be monitored in order to prevent adverse effects on the temperature balance.

**NOTE**
Higher side panels (230 mm) are recommended when using the canopy.

**WARNING**
When the bed canopy is in place, the side panels can be folded down. Do not leave the Babytherm unattended with the side panels open, as the infant may fall out.

**NOTE**
The bed canopy should not be used as a convenient place to lay objects, clothing, etc.

Configuration mode

Do not perform care/therapy during configuration of the device, because the normal equipment functions (e.g., temperature measurement, alarms) are switched off.
Switching on

After switching on, the system runs a self-test and then proposes the following default operating mode for the radiant warmer:

- No skin temperature sensor connected = **man.** (manual)
- Yellow skin temperature sensor for near-core temperature connected = **Skin** (skin temperature control mode).

1. The yellow LED for the mode selected will flash in the button.

Manual operation

The heater is controlled manually in the case of short-term treatment or for infants in shock and for whom baby control mode must not be used.

The radiant warmer delivers a preset power regardless of the infant's body temperature!

CAUTION

When in clinical use do not leave this device unattended! Measure the patient's core temperature regularly using a separate thermometer!

Check, whether both infrared radiators are providing warmth by placing your hand carefully at a safe distance from the radiant warmer.

If the system is still in **Skin** mode:

2. Press button until the yellow **man.** LED in the button starts to flash.
3. The following message appears on the display:
   
   **Set radiant heater level**
   
   **Press OK button to confirm**

   4. Babytherm will propose the most recently calculated heater output setting as the default. (After switching on, the default heater level is always 3.)

   Each LED segment represents one heater output level:
   
   1 to 3 = green LEDs, radiant warmer power ≤10 mW/cm²
   
   4 to 10 = yellow LEDs, radiant warmer power >10 mW/cm².

5. Press OK button = confirm selected heating level.

Or

6. Press  or  button until the desired heater output setting is displayed.

   Display of new selected heater output level.

   5 Press OK button = confirm selected heating level. If the OK button is not pressed within 10 seconds, the message disappears and the previous setting will remain in effect.

   4 Display of new selected heater output level.

   2 Yellow LED in the button is lit continuously: **man.** mode is now active.
**Operation**

**15 minutes alarm**

To remind the user to monitor the core temperature constantly at higher heater output levels,

- an audible alarm is emitted every 15 minutes at heater level 4 and above.

1. At the same time: the yellow LED flashes.
- The red central alarm lamp on the radiant warmer starts flashing.

2. The heater level display flashes and

3. The following message appears in the text display:

   **15min. patient temp check required**

   **Press OK button to acknowledge**

4. Press the button

   or

5. Press the OK button.

   **The heating will be switched on again.**

   The audible alarm will stop and the yellow LED and the central alarm lamp will go out. The heating level display will be lit again.

   If a skin temperature sensor is connected, the measured skin temperature will be displayed.

   However, no adjustment based on the skin temperature will be made!

---

![Diagram of Babytherm 8010](image-url)
Using baby control mode

In this operating mode, the skin temperature of the infant is adjusted towards the set value. The sensor attached to the skin measures the skin temperature. The radiant warmer output is adjusted according to the temperature difference between the skin temperature and the desired value.

Therefore:

**WARNING**
Do not use for infants in a state of shock. When in shock, skin temperature is much lower than normal. If automatic baby control mode is used, the infants will be overheated.

Manual operation, see page 47.

**WARNING**
Do not use for infants with fever. In this case, skin temperature is much higher than normal. If automatic baby control mode is used, it could induce hypothermia.

**WARNING**
Risk of patient injury
The core body temperature is not measured with the skin temperature sensor.
The core body temperature must be measured with an appropriate sensor.

Check the set value or set the heating level manually, see page 47.

Connecting the skin temperature sensor

1. Plug the yellow sensor connector fully into the yellow connection socket.
2. Feed the cable through one of the flexible grommets in the cot.
3. Remove the protective film from the ThermoPad and place the skin temperature sensor on the pad.
   - Attach the sensor probe to the appropriate area of the patient’s skin with the adhesive pad.
   - If the infant is lying on its back:
     attach the sensor to the abdomen in the liver region.
   - If the infant is lying on its front:
     Attach the sensor to the back, preferably in the kidney region.

Do not attach the sensor underneath the infant; otherwise the measured value for the skin temperature would be distorted by the mattress.

- Secure the sensor cable with adhesive tape (plaster).
- Regularly check that the skin temperature sensor is in the correct position.
- Measure the patient’s core temperature regularly using a separate thermometer!
Operation

**WARNING**
Do not confuse the position of adhesion of the yellow skin temperature sensor with the white skin temperature sensor!

Skin temperature is regulated via the yellow skin temperature sensor. An incorrectly positioned sensor can put the patient at risk of overheating!

**WARNING**
If the yellow skin temperature sensor becomes detached, it will measure the air temperature and so the infant may be at risk of overheating.

**For single use:**
- Only use Dräger approved skin temperature sensors.

**CAUTION**
Only use Dräger ThermoPads. Using other types of adhesive pads may result in incorrect temperature measurements and therefore can affect the safety and effectiveness of the skin temperature measurement.

**CAUTION**
Never use the skin temperature sensor to measure rectal temperature.

**NOTE**
Allow at least 5 minutes for the skin temperature sensor to adjust to the temperature of the infant.

If a skin temperature sensor is connected when the apparatus is switched on:

1. The device proposes the Skin temperature control operating mode; the yellow Skin LED in the button flashes.
2. The following prompt **Check skin sensor position**
   - *Press OK button to confirm* appears in the display:
3. Press the OK button.
4. The following message appears in the text display:
   - **Set skin temperature**
   - *Press OK button to confirm*
5. Press the OK button to accept the proposed setting.
   - *Press OK button = confirm setting.*
6. The new set value is displayed.
7. The yellow Skin LED lights up, indicating that Skin temperature control mode is active.
8. The measured skin temperature value is displayed.

**WARNING**
Do not confuse the position of adhesion of the yellow skin temperature sensor with the white skin temperature sensor!

Skin temperature is regulated via the yellow skin temperature sensor. An incorrectly positioned sensor can put the patient at risk of overheating!

**WARNING**
If the yellow skin temperature sensor becomes detached, it will measure the air temperature and so the infant may be at risk of overheating.

**CAUTION**
Only use Dräger ThermoPads. Using other types of adhesive pads may result in incorrect temperature measurements and therefore can affect the safety and effectiveness of the skin temperature measurement.

**CAUTION**
Never use the skin temperature sensor to measure rectal temperature.

**NOTE**
Allow at least 5 minutes for the skin temperature sensor to adjust to the temperature of the infant.
7 The heater output level display changes according to the difference between the current measured skin temperature of the infant and the set skin temperature.

If the device is in man. mode:

1 Press button until the yellow Skin LED starts to flash. The device proposes the Skin temperature control mode. The prompts in the display should be acknowledged as above.

Allow time for the system to reach steady state.

Deviations between the set and measured skin temperature are normally corrected within 5 to 15 minutes.

An infant’s skin temperature changes frequently, for instance as a result of food intake or treatment. Deviations of a few tenths of a degree are normal.

NOTE
Therefore: Only change the set value for the skin temperature if the core temperature needs to be corrected.

Check whether both infrared radiators are providing warmth by placing your hand carefully at a safe distance from the radiant warmer.

Outside measuring range

1 Three dashes light up in the middle of the display. After 10 seconds:
   – The audible alarm sounds,
   2 the red LED flashes and
   – the red central alarm lamp on the radiant warmer starts flashing.
1 The three dashes on the display flash.

3 The following message appears on the display:
   Plug in skin temp. sensor
   Press OK button to acknowledge
   Or
   Skin temp. sensor fault
   Press OK button to acknowledge

   Immediately connect the sensor plug or change the skin temperature sensor. The audible alarm can be muted for 15 minutes:
4 Press the button
Operation

or

5 Press the OK button.
   – The audible alarm will be muted,
2 the red LED goes out and
   – the red central alarm lamp on the radiant warmer goes out.
3 The following message appears on the display:
   Plug in skin temp. sensor

Immediately connect the sensor plug or change the skin temperature sensor.

If the error cannot be remedied immediately:

6 Switch to man. mode, see page 47.

For deviations greater than ±0.5 °C between the set and measured skin temperature:

![Diagram]

1 The skin temperature display flashes.
   – The audible alarm sounds,
2 the yellow LED starts flashing and
   – the red central alarm lamp on the radiant warmer starts to flash.
3 The following message appears on the display:
   Skin temp. deviation above 0.5 °C
   Press OK button to acknowledge

The permitted skin temperature deviation can be set in configuration mode, see page 70.

A default skin temperature deviation of ±0.5 °C is set by the manufacturer.

The audible alarm can be muted for 15 minutes:

4 Press button
   or
5 Press the OK button.
   – The audible alarm stops.
2 The yellow LED lights up.
   – The red central alarm lamp on the radiant warmer goes out.
1 The skin temperature display flashes.
3 The following message appears on the display:
   Skin temp. deviation above 0.5 °C

After the measured skin temperature has returned to a value within ±0.5 °C of the set temperature, the yellow LED goes out, the audible alarm stops and the message on the display disappears.

Before temporarily removing the skin temperature sensor from the skin, switch to man. mode, see page 47.

Skin temperature above 39 °C

![Diagram]

1 The skin temperature display flashes.
   – The audible alarm sounds.
Operation

2 The red LED starts flashing.
   – The red central alarm lamp on the radiant warmer starts flashing.
3 The following message appears on the display:
   **Skin temperature above 39°C**
   **Press OK button to acknowledge**
2 The red LED lights.
   The audible alarm can be muted for 2 minutes:
4 Press button
   or
5 Press the OK button.
   – The audible alarm stops.

2 The red LED is lit continuously.
   – The red central alarm lamp on the radiant warmer goes out.
3 The following message appears on the display:
   **Skin temperature above 39°C**
The radiant warmer is switched off in *man.* mode:
1 Skin temperature display is lit.
2 The red LED starts flashing.
The alarm is canceled automatically when the measured skin temperature drops below 38.5 °C again.

ThermoMonitoring

For better assessment of the thermal condition of the child, Dräger recommends measuring both the core and the **peripheral temperature**.

Connecting the peripheral skin temperature sensor

1 Plug the white sensor fully into the white socket.
2 Route the sensor cable through one of the flexible grommets in the bed.
3 Remove the protective film from the adhesive pad and place the skin temperature sensor on the pad.

- Attach the sensor with the ThermoPad to the patient’s extremities, preferably the foot.
- Secure the sensor cable with adhesive tape (plaster).

Displaying the peripheral skin temperature
Operation

4 The peripheral temperature is displayed as soon as the peripheral skin temperature sensor is attached.

5 The symbol for the peripheral skin temperature lights up.

If 3 dashes appear on the display, see “System faults - radiant warmer” on page 92. The measured value of the peripheral temperature has no effect on the radiant warmer control. Both skin temperatures can be displayed when the radiant warmer is operating in man. mode.

Data output via interface, optional

The core and peripheral temperature can be displayed in graphic form.

Prerequisites:
- Interface option
- MediCable connecting lead
- Monitor compatible with the MEDIBUS protocol and complying with the requirements of EN 60601-1 and EN 60601-1-2.
- Following the associated Instructions for Use.

Switching off the radiant warmer

6 Press button.

7 The following message appears on the display:

Radiant heater OFF

Press OK button to confirm

8 Press the OK button.
The radiant warmer will be switched off.

7 The following message appears on the display:

Radiant heater turned OFF

6 Press button again = radiant warmer can be switched on again, see page 47.
Using heated gel mattress

Observe the warm-up time of the gel mattress.

Wait until the desired mattress heating value is reached before placing the infant in the Babytherm 8010.

Do not switch off the mattress heater while an infant is lying on the gel mattress. Risk of hypothermia.

Setting the mattress temperature

The set value can be adjusted in increments of 0.1 °C. To set a temperature within the normal range of 36 °C to 38 °C:

1. The display shows the current set value for the mattress temperature.
2. Press \( \downarrow \) or \( \uparrow \) button until the desired setting is displayed.
3. The following message appears on the display:
   
   **Set mattress temperature**
   
   **Press OK button to confirm**

If the setting is not confirmed within the next 10 seconds, the message will disappear and the previous setting will remain in effect.

4. Press the OK button = confirm new setting.
5. The new set value for the mattress temperature is displayed.

Extending the upper limit of the setting, range from 38 °C to 38.5 °C:

6. Press \( \downarrow \) button until the set value 38 °C is displayed.
7. The following message appears on the display:
   
   **Mattress temperature setting**
   
   **Confirm temp. above 38 °C with OK**

8. Press OK button = confirm extended range.
9. The yellow \( >38 °C \) indicator flashes.

5. Press \( \downarrow \) button until the desired setting appears in the display.
6. The following message appears on the display:
   
   **Set mattress temperature**
   
   **Press OK button to confirm**

8. Press the OK button = confirm new setting.
9. The new set value is displayed.
10. The yellow 38 °C indicator is lit continuously.
Operation

Extending the lower limit of the setting, range from 36 °C to 30 °C:

Only use low temperatures if prescribed by doctor. Monitor patient very closely.

1 Press button until the set value 36 °C is displayed.
2 The following message appears on the display:
   **Mattress temperature setting**
   **Confirm temp. below 36 °C with OK**
3 Press OK button = confirm extended range.
4 The yellow <36 °C indicator flashes.
5 Press button until the desired setting appears in the display.
3 The following message appears on the display:
   **Set mattress temperature**
   **Press OK button to confirm**
4 Press the OK button = confirm new setting.
2 The new set value is displayed.
5 The yellow <36 °C indicator is lit continuously.

Outside measuring range

If the temperature is outside the measuring range from 5 °C to 45 °C:

- 3 dashes at the bottom of the display = temperature below 5 °C.
- The following message appears on the display:
  **Mattress temp. below 5°C**
  - Wait until the mattress temperature exceeds 5 °C.
- 3 dashes at the top of the display = temperature above 45 °C.
- Wait until the mattress temperature falls below 45 °C.

Deviation the from set temperature

If the deviation between the set and measured mattress temperature is greater than ±1 °C:
Operation

– The audible alarm sounds.
1 The yellow LED starts flashing and
– the red central alarm lamp on the radiant warmer starts flashing.
2 Measured mattress temperature value flashes,
3 The following message appears in the text display:
   **Mattress temp. deviation above 1°C**
   **Press OK button to acknowledge**
The alarm can be muted for 15 minutes:
4 Press the button
or
5 Press the OK button.
   – The audible alarm stops.
1 The yellow LED lights up.
   – The red central alarm lamp on the radiant warmer goes out.
3 The following message appears in the text display:
   **Mattress temp. deviation above 1°C**
2 Measured mattress temperature value flashes
When the measured mattress temperature returns to within ±1 °C of the set value:
1 The yellow LED goes out and
   – the audible alarm is switched off.
After initially switching on the system:
– the audible alarm is suppressed during the warm-up phase:
1 The yellow LED is lit.

**Mattress temperature above 40 °C**

![](image)

1 Red LED flashes.
– The red central alarm lamp on the radiant warmer starts flashing.
2 Display flashes.
3 The following message appears in the text display:
   **Mattress temp. above 40°C**
   **Press OK button to acknowledge**
The alarm can be muted for 10 minutes:
4 Press the button
or
5 Press the OK button.
1 The red LED starts flashing.
   – The red central alarm lamp on the radiant warmer goes out.
3 The following message appears in the text display:
   **Mattress temp. above 40°C**
The alarm stops automatically as soon as the mattress heater temperature drops below 39 °C.
Using the bed canopy

The infant's core temperature should be monitored when the bed canopy is in use.

CAUTION
The radiant warmer is less effective when the bed canopy is used. The patient's core temperature should be monitored in order to prevent adverse effects on the temperature balance.

NOTE
Use of the bed canopy in conjunction with operation of the integrated phototherapy system leads to a significant reduction in the bilirubin-effective radiant intensity and a very uneven distribution of the radiation on the bed.

WARNING
When the bed canopy is in place, the side panels can be folded down. Do not leave the Babytherm unattended with the side panels open, as the infant may fall out.

Installing the bed canopy

Check whether the bed canopy is in the closed position; otherwise:

1 Close bed canopy = turn lock until it engages.

2 Hold bed canopy handles with both hands and place over the side panels.

It is advisable to use the higher side panels (230 mm) when using the bed canopy.

Opening/closing the bed canopy

1 Unlock bed canopy = turn lock

2 Open bed canopy = swivel handles upwards until they engage. The infant is now accessible for care/medical treatment.

2 Close bed canopy = swivel handles forwards.
Removing the bed canopy

3. Put bed canopy in closed position = turn lock until it engages.

- Remove bed canopy by holding the handles with both hands (follow instructions on the bed canopy).

- Hang bed canopy from bracket.

Recommended heater settings for heat therapy

Instructions for setting the mattress heater and radiant warmer, alone or in combination.

Using the mattress heater with a gel mattress without using the radiant warmer (Babytherm 8010 only)

Normal setting:
The core temperature of the patient adapts to the mattress temperature over a relatively long period. The mattress heater should therefore be set to the appropriate core temperature for the patient, e.g., between 38.0 °C and 38.3 °C for a premature baby and 37.0 °C for a normal-term neonate. Monitor the core temperature and adjust the mattress temperature setting to the needs of the patient!

- Always cover and/or clothe the patient. Do not place blankets or other insulating material under the patient, since they obstruct heat transfer to and from the bed (warming/cooling).

- Warm up the patient:
  Set the mattress temperature to the desired core temperature. If necessary, slightly higher to reduce the warm-up time.

- Cool the patient:
  Set the mattress temperature to a value lower than the current core temperature, e.g., to 36 °C.
Using radiant warmer only

Monitor the core temperature and adjust the mattress temperature setting to the needs of the patient!

- If the distance between the radiant warmer and the patient on the bed is reduced, the irradiance increases.
- If the distance between the radiant warmer and the patient on the bed is increased, the irradiance is reduced.

- The radiant warmer is less effective when the bed canopy is used.
  (The acrylic glass bed canopy provides only limited permeability to the infrared radiation of the radiant warmer.)

- Do not cover or dress the patient. Otherwise, the effect of the radiant warmer is uncontrollably reduced.

Swiveling the radiant warmer and tilting the bed

**WARNING**
Risk of uneven warming of the patient

If the bed is tilted and the radiant warmer is swiveled, those parts of the patient’s body that are closer to the radiant warmer will receive more heat.

Check the skin temperature of such body parts and the body core temperature manually at regular intervals.

Swiveling the radiant warmer

The radiant warmer is freely adjustable and remains aligned towards the bed when it is swiveled.

- Manually swivel the radiant warmer to the right or left.
- Check the patient’s temperature at regular intervals.
Tilting the bed

- Tilt the bed, see Page 31.
- Check the patient's temperature at regular intervals.

1. Use foam mattress. Do not use a gel mattress when operating without mattress heating.

**WARNING**
An unheated gel mattress will cool the patient down.

Using radiant warmer in combination with mattress heater and gel mattress (Babytherm 8010 only)

**Normal setting:**
Set the mattress temperature to the **core temperature** appropriate for the patient, e.g., between 38.0 °C and 38.3 °C for a premature baby and 37.0 °C for a normal-term neonate.

Set the skin temperature setting to the **skin temperature** appropriate for the patient, e.g., 37.0 °C and 36.5 °C for a normal-term neonate.

- To increase patient temperature:
  Set mattress temperature to the desired core temperature. If necessary, set skin temperature to a slightly higher value. For a very hypothermic patient:
  Set the mattress temperature and the skin temperature setpoint so that an increase in the patient temperature of approximately 1 °C per hour is achieved.

Radiant warmer without mattress heater Babytherm 8004
Operation

- To stabilize or maintain patient temperature:
  Set the mattress temperature and skin temperature to the current core and skin temperature, respectively.

- To decrease patient temperature:
  Set mattress temperature to the desired core temperature. If necessary, set skin control temperature to a slightly lower value.
Using phototherapy (optional)

**WARNING**
Only use the specified phototherapy lights, see "Technical data" on page 102. Using lights of other types can affect the safety and effectiveness of the phototherapy.

**WARNING**
Always use patient eye protection when using phototherapy.
Keep a constant check that the eye protection is correctly positioned.

**WARNING**
The patient's bilirubin levels must be measured regularly.
Beware of possible toxic effects due to bilirubin photoisomers.

**WARNING**
Phototherapy can upset the patient’s water balance.

**WARNING**
Dangerous body temperatures can occur when using reflective foil.

**CAUTION**
Direct eye contact with the radiation source for an extended period when in use should be avoided.

**CAUTION**
Protect patients in the immediate vicinity of the phototherapy device from the radiation emitted, e.g. by using goggles and shields.

**CAUTION**
Users should avoid remaining in the area of irradiation for a prolonged time. Risk of sunburn.

**NOTE**
Use of the bed canopy in conjunction with operation of the integrated phototherapy system leads to a significant reduction in the bilirubin-effective radiant intensity and a very uneven distribution of the radiation on the bed.

The integrated phototherapy light supplies additional heat to the patient.

- In manual mode, reduce the heat output of the radiant warmer.
- Use Baby control mode, see Page 49.

Simultaneous operation of the phototherapy unit and the radiant warmer in manual mode, level 1, 2 or 3, without recurring alarms is not permitted because the irradiance of 10 mW/cm² defined by the relevant standards might be exceeded.

**CAUTION**
Risk of hyperthermia
Monitor the patient’s core temperature regularly using a separate thermometer.

- Swing the radiant warmer over the patient. Position the phototherapy lights vertically above the patient, as otherwise the effect of the phototherapy will be diminished.
Operation

Switching on

1. Press button. The yellow LED in the button will start to flash.

2. The following message appears on the display:
   - Photo therapy: Use eye protection
   - Press OK button to start XX:XX:XX.

3. Press OK button = phototherapy lights switched on.

4. The yellow LED in the button is now continuously lit.

Resetting the duration of therapy for phototherapy

1. Press button for 3 seconds.

2. The following message appears in the text display:
   - Photo therapy duration XX:XX:XX
   - Press OK to reset the counter

3. Press OK button = counter reset.

4. The display now reads:
   - Photo therapy duration 00:00:00.

Switching off

4. Press button.

5. The following message appears on the display:
   - Photo therapy OFF
   - Press OK button to confirm XX:XX:XX

6. Press OK button = phototherapy switched off.

4. The yellow LED in the button will go out.

The counter is reset automatically when the unit is switched off.

When using the radiant warmer in "man." mode:

- Reduce the heating level: Reduce the radiant warmer heating level by about 3 increments compared to operation without phototherapy.
- When using the radiant warmer in the Skin, the radiant warmer output is automatically adjusted to the patient’s heat requirements.
Operation

Switching lighting On/Off

1 Press the button. The bed will be illuminated by the working light.

1 Press the button again. The working light will be switched off.

2 Press the button. The bed will be lit by the night light.

2 Press the button again. The night light will be switched off.
Alarms

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alarms ................................. 69
Safety information

**WARNING**
Risk due to failure to hear alarms
The user must remain within the hearing range of the acoustic alarm signal. This operator position makes it possible to quickly detect an alarm and to respond accordingly. The distance to the medical device must be appropriate for the volume of the alarm signal.

**WARNING**
Risk due to differing basic alarm settings
When operating several devices with differing basic alarm settings in one room, this can result in confusion over the alarm signals. Basic alarm settings can be changed without a password.
Adjust the basic alarm settings of the devices to the patient needs.

Display of alarms

**Optical alarm signals**
The alarm LED always signals the alarm with the highest alarm priority. The alarm message in the display always refers to the active and unacknowledged alarm with the highest alarm priority. Due to this alarm behavior, the display and the alarm LED may indicate different alarm priorities at the same time.
If there are several unacknowledged alarms, a request to acknowledge the alarms with OK is displayed.

**Generating and viewing the alarm list**
The alarm list is generated by acknowledging the active alarms. The alarms in the alarm list are sorted according to priority.

- Acknowledge the active alarms with the button or the OK button.
The alarm list can be viewed at any time.
- View the alarm list by acknowledging with the OK button.
If the cause of an alarm has been resolved, the alarm will be removed from the list.

**Acoustic alarm signals**
It is always the unacknowledged alarm with the highest priority that is acoustically signaled. For information on muting, see chapter “Acknowledging and suppressing alarms”.

Instructions for use Babytherm 8004/8010 67
Alarms

Central alarm*

The red lamp on the radiant warmer flashes when an audible alarm is emitted. As soon as the alarm is acknowledged, the lamp goes out.

Alarm priorities

The color of the alarm LED and the alarm tone sequence indicate the priority of the active alarm.

<table>
<thead>
<tr>
<th>Color</th>
<th>Signal</th>
<th>Alarm tone sequence</th>
<th>Priority of the alarm message</th>
<th>Required action</th>
</tr>
</thead>
<tbody>
<tr>
<td>Red</td>
<td><em>Inop.</em> LED is lit</td>
<td>Continuous</td>
<td>Device failure</td>
<td>High-priority alarm</td>
</tr>
<tr>
<td>Red</td>
<td>Power failure LED is lit</td>
<td>Continuous</td>
<td>Power failure</td>
<td></td>
</tr>
<tr>
<td>Red</td>
<td>Alarm LED flashes</td>
<td>Intermittent</td>
<td>Warning</td>
<td>***Immediate action is necessary in order to avert imminent danger.</td>
</tr>
<tr>
<td>Yellow</td>
<td>Alarm LED flashes (unacknowledged alarm), alarm LED is lit (acknowledged alarm)</td>
<td>Intermittent</td>
<td>Caution</td>
<td><strong>Prompt action is necessary in order to avert a danger.</strong></td>
</tr>
<tr>
<td>---</td>
<td>---</td>
<td>---</td>
<td>Note</td>
<td>Attention and action are necessary.</td>
</tr>
</tbody>
</table>

* Country-specific function (e.g., not available in China)
Acknowledging and suppressing alarms

The alarm tone sequence can be muted for the duration of the alarm suppression time; however, the alarm message will be displayed, see "Message – Cause – Remedy".

Muting the acoustic alarm signal when there is no active alarm present

- Press the button
  The LED in the button lights.
  If an alarm occurs during the muting period, Babytherm 8004/8010 reacts as follows:
  - The alarm message is displayed.
  - The acoustic alarm signal is suppressed.
  - The alarm LED on the display lights up according to the alarm priority.
- To acknowledge the alarm, press the button or the OK button.
  If no further alarm has occurred, the muting can be canceled with the button.
  Babytherm 8004/8010 reacts as follows:
  - Alarms that are already muted remain muted until their suppression time elapses.
  - The LED in the button is lit during the suppression time of the active alarms.
  - New alarms and recurring alarms are signaled visually and acoustically.

Muting the acoustic alarm signal of active alarms

- To acknowledge the alarm, press the button or the OK button.
  Babytherm 8004/8010 reacts as follows:
  - The alarm tone sequence is muted during the suppression time.
  - The alarm message is displayed.
  - The alarm LED lights up according to the alarm priority.
  If a new alarm occurs during the suppression time of an active alarm, Babytherm 8004/8010 reacts as follows:
  - The acoustic alarm signal sounds.
  - The alarm message of the new alarm is displayed.
  - The alarm LED lights up according to the priority of the highest active alarm (acknowledged or unacknowledged).
- To acknowledge the alarm, press the button or the OK button.
Configuration mode

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Configuration mode

Safety information

WARNING
No care/therapy should be carried out during configuration of the system, because the normal equipment functions (e.g. temperature measurement, alarms) are switched off.

Start settings

The alarm settings are automatically stored and are retained after a power failure of any duration.

The alarm configuration set last is effective after every restart of the device. If the power supply is interrupted, there is no time delay to restore the alarm configuration set last. The alarm settings must be adapted to the patient's needs.

The alarm settings can be changed without a further password entry.

Factory settings

The following table shows the factory settings of the parameters and their deviations. The factory settings are set in bold text.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Babytherm 8004/8010</th>
</tr>
</thead>
<tbody>
<tr>
<td>Skin temperature</td>
<td>0.3 °C to 1.0 °C</td>
</tr>
<tr>
<td></td>
<td>0.5 °C</td>
</tr>
</tbody>
</table>

Capabilities available in configuration mode

<table>
<thead>
<tr>
<th>Code</th>
<th>Activity</th>
</tr>
</thead>
<tbody>
<tr>
<td>c01</td>
<td>Show current software version</td>
</tr>
<tr>
<td>c02</td>
<td>Set skin temperature deviation</td>
</tr>
<tr>
<td>c03</td>
<td>Set volume of soft alarm</td>
</tr>
<tr>
<td>c04</td>
<td>Operating hours of phototherapy</td>
</tr>
<tr>
<td>c05</td>
<td>Test control panel buttons</td>
</tr>
<tr>
<td>c06</td>
<td>Test Nurse call</td>
</tr>
<tr>
<td>c07</td>
<td>Set language</td>
</tr>
<tr>
<td>c08</td>
<td>Set display contrast</td>
</tr>
<tr>
<td>Err</td>
<td>Error mode</td>
</tr>
</tbody>
</table>

The significance of the keys and displays on the control panel is different from that applicable when working with the radiant warmer!
Configuration mode

Activate configuration mode

1 Press OK button for 3 seconds. A brief acoustic signal sounds.

2 The following message appears on the display:
   Configuration Mode
   Press OK button to start.

1 Press OK button again. A brief acoustic signal sounds. Configuration mode is now active. The equipment is in mode c01 and shows the current software version.

Show software version
Configuration mode

1 The following message appears on the display:
   Configuration Mode
   Mode c01 : Software version.

2 The current software version is indicated: e.g.,
   1.n.

3 The code for the current mode is displayed: c01

Set permissible skin temperature deviation

1 Briefly press the button.

2 The following message appears on the display:
   Configuration Mode
   Mode c01 : Max. skin temperature fault.

3 The abbreviation for this mode appears: SdE.

4 The code for the current mode appears: c01.

5 Indication of the set permissible skin
temperature deviation at which an alarm is not
yet generated. Factory setting: 0.5 °C.

6 Press the or button until the required
value appears on the display. Values can be set
from 0.3 to 1.0 °C in increments of 0.1.

Set start volume for alarm tone sequence (soft alarm)

1 Briefly press the button.

2 The following message appears on the display:
   Configuration Mode
   Mode c03 : Soft alarm start volume.

3 The abbreviation for this mode appears: SSL.

4 The code for the current mode appears: c03.

5 Indication of the set volume. Factory setting: 2.
   The alarm tone sounds briefly.

6 Press the or button until the required
value appears on the display. Settings from 1 to
8 are possible. The alarm tone sequence
sounds briefly at the set volume.
Configuration mode

Show operating hours for phototherapy

1 Briefly press the button.
2 The following message appears on the display: Configuration Mode Mode c04 : Time counter photo therapy.
3 Indication of the operating hours up to 1000 hours.
4 Press button for 3 seconds. The counter is reset to 0. This is necessary when changing the phototherapy lights.

Test buttons on control panel

1 Briefly press the button.
2 The following message appears on the display: Configuration Mode Mode c05 : Keyboard test (inactive).
3 The abbreviation for this mode appears: but
4 The code for the current mode appears: c05
5 Press the or button to activate the test.

Button numbers

- Press the buttons to be tested,
1 A string of digits appears on the display: 1st digit = number of buttons currently pressed, 2nd and 3rd digit = number of last button pressed.
- Press the button for 2 seconds. The test is ended.

Instructions for use Babytherm 8004/8010
**Test Nurse call**

1. Briefly press the button.
2. The following message appears on the display: *Configuration Mode Mode c06 : Nurse Call Relay.*
3. The abbreviation for this mode appears: **Nuc**
4. The code for the current mode appears: **c06**.
5. Display 0 = nurse call relay open.
7. Press button until the relay switches.

**Select language**

1. Briefly press the button.
2. The following message appears on the display: *Configuration Mode Mode c07 : Display language -> English.*
3. The abbreviation for this mode appears: **LAN**.
4. The code for the current mode appears: **c07**.
5. Indication of the language number. Factory setting 2 = English.
6. Press the or button until the number of the required language appears on the display. Possible settings: 1 to 11.
Configuration mode

Set display contrast

1 Briefly press the button.
2 The following message appears on the display:
   Configuration Mode
   Mode c08 : Display contrast.
3 The abbreviation for this mode appears: CON.
4 The code for the current mode appears: c08.
5 Indication of the set contrast. Factory setting: 128.
6 Press the or button until the required contrast has been set. Possible settings: 1 to 255 in increments of 1.

Read error memory

1 Press button until error mode is active.
2 The following message appears on the display:
   Configuration Mode
   Error memory.
3 The error number appears: FXX.
4 Err is displayed.
5 The frequency of the error is indicated.
6 Press or button and the next error is displayed.
7 Press the button. The equipment runs a self-test.
8 Press button for 3 seconds. The error memory is erased.

Return to radiant warmer mode

- Press the OK button.
O2 therapy

CAUTION
Only enrich the O2 concentration under the instructions of a doctor. Only do this in accordance with the arterially measured O2 partial pressure in the patient’s blood. Otherwise there is a danger of hyperoxemia (which may cause damage to the eyes) or hypoxemia (which may cause damage to the brain).
Monitor the O2 concentration during the O2 therapy.

Bronchial aspiration
Use bronchial aspirator in accordance with its specific Instructions for Use.

Injector and closed bed canopy

- Place canopy in position (see page 58).
- Set O2 concentration on the injector: 30, 40, 50 Vol.% O2.
- Set O2 supply on flowmeter.

<table>
<thead>
<tr>
<th>Injector setting</th>
<th>Vol.% O2</th>
<th>30</th>
<th>40</th>
<th>50</th>
</tr>
</thead>
<tbody>
<tr>
<td>O2 supply</td>
<td>L/min</td>
<td>5</td>
<td>10</td>
<td>15</td>
</tr>
</tbody>
</table>

- Monitor the O2 concentration.

After use
- Open the bed canopy.
- Close the valve on the O2 flowmeter = turn clockwise.

Dismantle and reprocess the therapy equipment, see page 82.
Taking X-rays

Babytherm 8010, Babytherm 8004 without X-ray tray

- Swivel radiant warmer to one side. The infant will continue to be warmed.
- Place the X-ray film cartridge directly underneath the patient.
- Position the X-ray machine above the bed surface.

Babytherm 8004 with X-ray tray

The X-ray tray has no lock and can be completely removed from the Babytherm.

- Pull out X-ray tray and insert X-ray cartridge, placing it according to the position of the infant. Use the scale on the side panels and X-ray tray as a guide.
- Push X-ray tray in.
- Swivel radiant warmer to one side and position X-ray machine.
- After taking X-ray, remove X-ray cartridge from tray and push tray in again.
- After removing the X-ray machine, swivel the radiant warmer back into position.
Using the scale

**WARNING**
Only use the integrated scale to determine the weight of the patient.

Failure to follow the instructions for use can lead to severe inaccuracies in measuring the patient's weight. To safeguard critical therapeutic decisions based on the patient’s weight, the weighing result should be checked against a reference measurement on an external scale.

The scale is located directly under the bed. During the weighing operation, the entire bed and any articles on top of it are weighed. By lifting the patient, all other weights are subtracted, so that the precise weight of the patient can be calculated. The precision of the weight measurement is not limited by the articles on the bed. The weight of these additional articles reduces the maximum weight that the scale can display by exactly this extra weight. For patients with hose attachments, e.g. ventilation hoses, an effect on the measured weight, as with all other scales, cannot be entirely eliminated. To prevent fluctuations in the weighed result, the hoses should be removed from any fitted ventilation holder and, after lowering the patient on the bed, they should be laid horizontally on the bed with as little tension as possible. As with clothing, the inclusion of parts of the hose supply lines in the weight measurement can lead to a slight increase in the measured weight. However, since these deviations are systematic, they do not affect trend observations.

**NOTE**
To ensure correct determination of patient weight, observe the following instructions:

- Avoid a reduction of the base load on the scale by changing the mechanical structure of the warming bed. **U-LOAD** is displayed on the device.
- Before starting the weighing operation, remove all unnecessary objects from the bed.
- When lifting the patient, do not touch the bed. Leaning on the bed or touching the bed with an overall will distort the weighing result.
- While weighing is in progress, do not place any article on the bed or remove any article from it.
- Always perform the weighing process in exactly the same way. Either lift the hoses with the patient or leave them on the bed each time you carry out a weight measurement.
- Lay the patient to be weighed in the middle of the bed.
- The sensor cable between the weighing unit and display unit must not touch the bed.
- Do not place any heavy objects on the weighing unit.
- Use only scales authorised for use with Babytherm. The serial numbers of the weighing unit and display unit must match.
Configuration mode

Weighing with zero calibration

Preparing for weighing
- Remove all objects that are simultaneously in contact with both the bed and the fixed environment or frame of Babytherm.
- Prepare the patient for lifting.

Switch on the display unit of the scale

1. Plug the power plug into a power supply socket.
2. Press the \textit{Start/Standby} button.
   - 1 beep is emitted. Weighing begins.

Starting the weighing process

1. Press the \textit{Reweigh} button.
   - The patient must be lying on the bed.
   - 1 beep is emitted. Weighing begins.
   - A line of dots is displayed on the screen.
   - 3 beeps are emitted.

   When the Lift Patient \textit{Lift} indicator lights up:
   - Lift the patient off the bed. Take care not to touch the bed and frame of the Babytherm.
   - 2 beeps are emitted. \textit{0.00} is displayed on the screen.

   When the Lower Patient \textit{Lower} indicator lights up:
   - Lay the patient on the bed.
   - The weighing operation is performed.
   - The weighing result is displayed on the screen.
   - A continuous sequence of dashes is displayed on the screen.

   Weighing without zero calibration

   If the last zero calibration was performed no longer than 3 minutes earlier and no object has been removed from the bed or placed on it during this time, repeated calibration is not necessary.

   1. Press the \textit{Reweigh} button.
   - The patient must be lying on the bed.
   - The weighing operation is performed.
   - A line of dots is displayed on the screen.
   - 3 beeps are emitted.

   Weighing is complete.
   - The weighing result is displayed on the screen.
   - After a few minutes, the scale display unit changes to Standby mode.
   - A continuous sequence of dashes is displayed on the screen.
Configuration mode

Shutting down

- Press the On/Off button. The green "On" LED will go out.
Reprocessing

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Reprocessing

WARNING
To avoid risks to hospital staff and other patients, disinfect and clean the device after use. When disinfecting contaminated parts, follow the hospital hygiene regulations (protective clothing, eye protection, etc.).

The Babytherm 8004/8010 infant warmer system must be thoroughly cleaned and disinfected – after each change of patient – at least once a week.

Do not expose device to UV radiation as a means of disinfecting/cleaning. Cracks may be caused in the acrylic glass components.

Only use the recommended cleaning agents and disinfectants. Otherwise, there is a danger of causing cracks in the acrylic glass and macrolon, e.g. when using alcohol.

Clean and disinfect all accessories, e.g. bronchial aspirator, in accordance with their specific Instructions for Use.

Dismantling

- Switch off the device(s). Disconnect the power plug(s) from the mains and switch off all compressed gas supplies used.
- Remove any ancillary equipment installed.
- Remove the ventilation hose clips.
- Remove the hose grommets.
- Swivel the drainage canister bracket upwards and remove from the hole horizontally.

Information on reprocessing

Instructions for reprocessing are based on internationally accepted guidelines, e.g., standard ISO 17664.

Safety information

CAUTION
Risk due to faulty products

Signs of wear, e.g., cracks, deformation, discoloration, or peeling, may occur with reprocessed products.

Check the products for signs of wear and replace them if necessary.
Reprocessing

WARNING
Risk due to inappropriately reprocessed products
Reusable products must be reprocessed, otherwise there is an increased risk of infection.
- Observe the hygiene regulations and reprocessing regulations of the healthcare facility.
- Observe national hygiene regulations and reprocessing regulations.
- Use validated procedures for reprocessing.
- Reprocess reusable products after every use.
- Observe the manufacturer’s instructions for cleaning agents, disinfectants, and reprocessing devices.

CAUTION
Do not wash the mattress in a washing machine. Do not autoclave.

CAUTION
Do not clean the heating elements and reflectors with a moist cloth or brush!

CAUTION
Do not use disinfectants or cleaning products that contain alcohol.

NOTE
Even reusable accessories (e.g. after being prepared) have a limited service life. Wear may be increased and the service life reduced considerably by various factors when handling and preparing them (e.g. disinfectant residues corroding the material when autoclaving them). These parts must be replaced if signs of wear become visible, such as cracks, deformation, discoloration, peeling, etc.

- Before disinfecting/cleaning the radiant warmer, allow it to cool down for about 30 minutes.

Classifications for reprocessing

Classification of medical devices

Medical devices and their components are classified according to the way they are used and the resulting risk.

<table>
<thead>
<tr>
<th>Classification</th>
<th>Explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Non-critical</td>
<td>Components that come only into contact with skin that is intact</td>
</tr>
<tr>
<td>Semi-critical (A, B)</td>
<td>Components that carry breathing gas or come into contact with mucous membranes or pathologically altered skin</td>
</tr>
<tr>
<td>Critical (A, B, C)</td>
<td>Components that penetrate skin or mucous membranes or come into contact with blood</td>
</tr>
</tbody>
</table>
**Classification of device-specific components**

Observe the instructions for use for the components.

The following classification is a recommendation from Dräger.

**Non-critical**
- Bed frame, inside and outside
- Side panels, inside and outside
- Inside walls, inside and outside

**Semi-critical**
- Bed canopy
- Weighing scale
- Stand, including all attachments
- Hose grommets
- Phototherapy filter glass

**Reprocessing list**

<table>
<thead>
<tr>
<th>Components</th>
<th>Surface disinfection with cleaning</th>
<th>Special reprocessing measures</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bed frame, inside and outside</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Side panels, inside and outside</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Inside walls, inside and outside</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>X-ray tray and X-ray tank</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Bed canopy</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Weighing scale</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Stand, including all attachments</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Hose grommets</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Gel mattress (Babytherm 8010)</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>SoftBed foam mattress (Babytherm 8004)</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Phototherapy filter glass</td>
<td>X</td>
<td></td>
</tr>
</tbody>
</table>
Reprocessing

Reprocessing procedures

Validated reprocessing procedures

At the time of product-specific validation, the following reprocessing procedures showed good material compatibility and effectiveness:

<table>
<thead>
<tr>
<th>Procedure</th>
<th>Agent</th>
<th>Manufacturer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surface disinfection with cleaning</td>
<td>Dismozon pur</td>
<td>Bode Chemie</td>
</tr>
</tbody>
</table>

Disinfectants

Use disinfectants that are nationally approved and are suitable for the particular reprocessing procedure.

Surface disinfectants

At the time of the test, the surface disinfectants listed in the following table showed good material compatibility. They can be used in addition to the surface disinfectants listed in the section "Validated reprocessing procedures".

The manufacturers of the surface disinfectants have verified at least the following spectra of activity:
- Bactericidal
- Yeasticidal
- Virucidal or virucidal against enveloped viruses

Observe the specifications of the surface disinfectant manufacturers. Other surface disinfectants are used at one’s own risk.

<table>
<thead>
<tr>
<th>Class of active ingredient</th>
<th>Surface disinfectant</th>
<th>Manufacturer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chlorine-releasing agents</td>
<td>Actichlor plus</td>
<td>Ecolab</td>
</tr>
<tr>
<td></td>
<td>BruTab 6S</td>
<td>Brulin</td>
</tr>
<tr>
<td></td>
<td>Clorox Professional Disinfecting Bleach Cleaner</td>
<td>Clorox</td>
</tr>
<tr>
<td></td>
<td>Dispatch Hospital Cleaner Disinfectant Towels with Bleach</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Klorsept 17</td>
<td>Medentech</td>
</tr>
<tr>
<td>Oxygen-releasing agents</td>
<td>Descogen Liquid</td>
<td>Antiseptica</td>
</tr>
<tr>
<td></td>
<td>Descogen Liquid r.f.u.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Dismozon plus</td>
<td>Bode Chemie</td>
</tr>
<tr>
<td></td>
<td>Dismozon pur</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Oxycide</td>
<td>Ecolab USA</td>
</tr>
<tr>
<td></td>
<td>Perform</td>
<td>Schülke &amp; Mayr</td>
</tr>
<tr>
<td></td>
<td>Virkon</td>
<td>DuPont</td>
</tr>
</tbody>
</table>
Dräger states that oxygen-releasing agents and chlorine-releasing agents may cause color change in some materials. Color change does not indicate that the product is not functioning correctly.

### Surface disinfection with cleaning

1. Remove soiling immediately. Use a cloth dampened with disinfectant to remove soiling.
2. Perform surface disinfection.
3. After the product has been exposed to the disinfectant for the specified contact time, remove residual disinfectant.
4. Wipe with a cloth dampened with water (preferably drinking-water quality). Allow the product to dry.
5. Check the product for visible soiling. Repeat steps 1 to 5 if necessary.
6. Check the product for visible damage and replace if necessary.

### Special reprocessing measures

#### Hose grommets
- Disinfect components in disinfectant bath. After immersing for the prescribed time, rinse with clean water and dry.
- Then wash with a detergent and rinse with clean water; or
- Sterilize at 120 °C (glove program).

#### Gel mattress (Babytherm 8010):
- Wipe gel mattress with disinfectant.

Do not machine wash the mattress and do not autoclave it. The gel mattress may become permanently stained if it comes into contact with liquid, alcoholic, iodine containing or iodine-like disinfectants. This does not impair the function and the mattress can still be used.

#### SoftBed foam mattress (Babytherm 8004):

Clean and wipe disinfect as required and after every change of patient. If it is heavily soiled, the cover may be machine washed at 95 °C. Tumble dry up to 95 °C.

The mattress may become contaminated in exceptional cases. The mattress core (foam) may then be machine washed at 30 °C using mild detergent.
Reprocessing

Phototherapy filter glass

- The lenses should only be cleaned and disinfected with products with a pH value between 7 and 9.

Before next use

- Reassemble components, see page 26 and following pages.
- After wipe-disinfecting, operate the fully assembled unit for a few hours without a patient to eliminate any disinfectant residues:
  - Set the mattress temperature to 37 °C and set the radiant warmer to heating level 3 in man. mode.

Before a patient is next placed in the unit:

- Fit all therapy accessories required.
- Check that the unit is ready for operation, see Page 37 and following pages.
Message – Cause – Remedy

- General device errors ................................................. 90
- System faults – Babytherm 8010 (Mattress heating) ................. 90
- System faults - radiant warmer ................................. 92
- System faults - scale ................................................. 94
Message – Cause – Remedy

General device errors

<table>
<thead>
<tr>
<th>Message</th>
<th>Cause</th>
<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Red LED (\text{\textsuperscript{1}}) is lit, Audible alarm on.</td>
<td>Power failure.</td>
<td>Check that unit is plugged into mains. Check that the mains is</td>
</tr>
<tr>
<td></td>
<td></td>
<td>switched on. Inform internal technical department. Call DrägerService.</td>
</tr>
<tr>
<td>Red \textit{Inop.} LED is lit, continuous tone is emitted.</td>
<td>Malfunction.</td>
<td>Call DrägerService.</td>
</tr>
<tr>
<td>The following message appears on the display: \textit{Battery dis-</td>
<td>Unit has been switched off for a</td>
<td>Battery will be charged automatically when the unit is switched on.</td>
</tr>
<tr>
<td>charged}</td>
<td>relatively long time.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Display disappears after 15 minutes.</td>
</tr>
</tbody>
</table>

System faults – Babytherm 8010 (Mattress heating)

<table>
<thead>
<tr>
<th>Priority</th>
<th>Suppression time</th>
<th>Message</th>
<th>Cause</th>
<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td>!</td>
<td>14</td>
<td>Yellow LED flashing, measured value display flashing, message in text display: \textit{Mattress temp. deviation above 1°C}, audible alarm sounds.</td>
<td>Measured mattress temperature deviates from set value by more than ±1 °C.</td>
<td>If used in combination with radiant warmer: Reduce the output level of the radiant warmer.</td>
</tr>
<tr>
<td>!!!</td>
<td>29</td>
<td>Red LED flashing, Measured value display flashing, message in text display: \textit{Mattress temp. above 40°C}, audible alarm sounds.</td>
<td>Mattress temperature above 40 °C.</td>
<td>If used in combination with radiant warmer: Reduce the output level of the radiant warmer.</td>
</tr>
</tbody>
</table>
### Message – Cause – Remedy

<table>
<thead>
<tr>
<th>Priority</th>
<th>Suppression time</th>
<th>Message</th>
<th>Cause</th>
<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td>!!! 29</td>
<td>10 min</td>
<td>Red LED flashing, the three middle segments of the measured value display are flashing, message in text display: <strong>Mattress temp. sensor fault</strong>, audible alarm sounds.</td>
<td>Temperature sensors in heating surface defective.</td>
<td>Call DrägerService.</td>
</tr>
<tr>
<td>! 2</td>
<td>---</td>
<td>Three dashes at top of measured value display.</td>
<td>Mattress temperature &gt;45 °C.</td>
<td>Wait until mattress temperature is lower than 45 °C.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Three dashes at bottom of measured value display. Message on display: <strong>Mattress temp. below 5°C</strong></td>
<td>Mattress temperature &lt;5 °C.</td>
<td>Wait until mattress temperature is above 5 °C.</td>
</tr>
</tbody>
</table>
## System faults - radiant warmer

<table>
<thead>
<tr>
<th>Priority</th>
<th>Suppression time</th>
<th>Message</th>
<th>Cause</th>
<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td>!!</td>
<td>17</td>
<td>Yellow LED flashing, measured value display flashing, message in text display: <strong>Skin temp. deviation above</strong> X °C, audible alarm sounds. Or <strong>Skin temp. deviation below</strong> X °C, audible alarm sounds.</td>
<td>The measured skin temperature deviates from the set value by more than the set permissible skin temperature deviation (0.3 to 1.0 °C).</td>
<td>If the measured value is higher than the set value: Measure the core temperature! Check the plug connection. Ensure plug is fully inserted. If necessary, contact Dräger-Service. If the measured value is lower than the set value: Check whether both infrared radiators are providing warmth by placing your hand carefully at a safe distance from the radiant warmer. Check that the skin temperature sensor is correctly attached. Call Dräger-Service.</td>
</tr>
<tr>
<td>!!</td>
<td>25</td>
<td>Red LED flashing, the three middle segments of the measured value display are flashing, message in text display: <strong>Plug in skin temp. sensor</strong>, audible alarm sounds.</td>
<td>Skin temperature sensor not connected or sensor defective.</td>
<td>Check the plug connection. Switch to <strong>man.</strong> mode; replace sensor and then switch back to <strong>Skin</strong> mode.</td>
</tr>
<tr>
<td>!!</td>
<td>25</td>
<td>Red LED flashing, the three middle segments of the measured value display are flashing, message in text display: <strong>Skin temp. sensor fault</strong>, audible alarm sounds.</td>
<td>Sensor defective.</td>
<td>Switch to <strong>man.</strong> mode; replace sensor and then switch back to <strong>Skin</strong> mode.</td>
</tr>
<tr>
<td>Priority</td>
<td>Suppression time</td>
<td>Message</td>
<td>Cause</td>
<td>Remedy</td>
</tr>
<tr>
<td>----------</td>
<td>------------------</td>
<td>-------------------------------------------------------------------------</td>
<td>-------------------------------</td>
<td>------------------------------------------------------------------------</td>
</tr>
<tr>
<td>!!! 29</td>
<td>2 min</td>
<td>Red LED flashes, measured value display flashes and the message <strong>Skin temperature above 39°C</strong> appears on the display, audible alarm sounds.</td>
<td>Skin temperature &gt;39 °C.</td>
<td>Check that the peripheral skin temperature sensor is correctly attached. Check that the child is not being heated by additional heat sources, e.g. phototherapy or the sun.</td>
</tr>
<tr>
<td>! 3</td>
<td>---</td>
<td>Three dashes at top of measured value display. Message in text display: <strong>Skin temp. above range.</strong></td>
<td>Skin temperature &gt;42 °C.</td>
<td>Check that the skin temperature sensor is correctly attached.</td>
</tr>
<tr>
<td>! 2</td>
<td>---</td>
<td>Three dashes at bottom of measured value display. Message on display: <strong>Skin temp. below range.</strong></td>
<td>Skin temperature &lt;15 °C</td>
<td></td>
</tr>
<tr>
<td>! 3</td>
<td>---</td>
<td>Three dashes at top of measured value display. Message in text display: <strong>Peripheral temp. above range.</strong></td>
<td>Peripheral temperature &gt;42 °C.</td>
<td>Check that the peripheral skin temperature sensor is correctly attached.</td>
</tr>
<tr>
<td>! 2</td>
<td>---</td>
<td>Three dashes at bottom of measured value display. Message on display: <strong>Peripheral temp. below range.</strong></td>
<td>Peripheral temperature &lt;15 °C.</td>
<td></td>
</tr>
<tr>
<td>!!! 25</td>
<td>14 min</td>
<td>Red LED flashes, the three middle segments of the measured value display are flashing, the message <strong>Peripheral temp. sensor fault</strong> appears on the display and the audible alarm sounds.</td>
<td>Sensor defective.</td>
<td>Replace sensor.</td>
</tr>
</tbody>
</table>
### System faults - scale

<table>
<thead>
<tr>
<th>Fault</th>
<th>Causes</th>
<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Messages on screen:</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>E-FAIL</td>
<td>Device error.</td>
<td>Call DrägerService.</td>
</tr>
<tr>
<td>r-FAIL</td>
<td></td>
<td></td>
</tr>
<tr>
<td>n-FAIL</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ad OuR</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Message O-LOAd and continuous beep</strong></td>
<td>The scale is overloaded.</td>
<td>Observe maximum load of 8 kg.</td>
</tr>
<tr>
<td></td>
<td>Sensor cable not plugged in.</td>
<td>Plug in sensor cable until it engages.</td>
</tr>
<tr>
<td><strong>No display on screen</strong></td>
<td>Display unit not switched on.</td>
<td>Switch on display unit.</td>
</tr>
<tr>
<td></td>
<td>Display unit not connected.</td>
<td>Connect display unit.</td>
</tr>
<tr>
<td><strong>Displayed patient weight is incorrect</strong></td>
<td>Bed or sensor cable mechanically blocked.</td>
<td>Check position of bed and sensor cable.</td>
</tr>
<tr>
<td></td>
<td>Weighing unit and display unit are not compatible with each other.</td>
<td>Check that the serial numbers of the weighing unit and display unit match.</td>
</tr>
<tr>
<td></td>
<td>The scale was not calibrated at the installtion site.</td>
<td>Call DrägerService.</td>
</tr>
<tr>
<td><strong>Long wait before patient weight is displayed</strong></td>
<td>The scale waits until the weighing unit has come to rest.</td>
<td>During the weighing process, the Babytherm must not be exposed to any vibrations. Check the connection and position of the sensor cable.</td>
</tr>
</tbody>
</table>

In the event of any unexpected behaviour of the device, call DrägerService.
Safety information

**WARNING**
**Risk of fire**
Fuses with excessively high nominal values can lead to serious damage. Only use fuses with the specified nominal values.

**WARNING**
**Risk of electric shock**
Before replacing the fuses, disconnect the medical device from the power supply.

**WARNING**
**Risk of infection**
The responsible personnel may be infected by pathogenic germs.
Disinfect and clean device or device parts before any maintenance measures and also before returning the medical device for repair.

**WARNING**
**Risk of electric shock**
There are conducting components under the housing cover.
Do not remove the housing cover.
Maintenance measures must be performed by the personnel responsible.

Overview

This chapter describes the maintenance measures required to maintain the proper functioning of the medical device.

Maintenance measures must be performed by the personnel responsible.

**Definitions of the terms used in maintenance**

<table>
<thead>
<tr>
<th>Concept</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Service</td>
<td>All measures (inspection, preventive maintenance, repair) intended to maintain and restore the functional condition of a medical device</td>
</tr>
<tr>
<td>Inspection</td>
<td>Measures intended to determine and assess the actual state of a medical device</td>
</tr>
<tr>
<td>Maintenance</td>
<td>Recurrent specified measures intended to maintain the functional condition of a medical device</td>
</tr>
<tr>
<td>Repair</td>
<td>Measures intended to restore the functional condition of a medical device after a device malfunction</td>
</tr>
</tbody>
</table>
Inspection

Perform inspections at regular intervals and observe the following specifications.

<table>
<thead>
<tr>
<th>Checks</th>
<th>Interval</th>
<th>Target group</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inspection</td>
<td>Annually</td>
<td>Service personnel</td>
</tr>
<tr>
<td>Safety checks 1)</td>
<td>Annually</td>
<td>Service personnel</td>
</tr>
<tr>
<td>Metrological checks</td>
<td>Every 2 years</td>
<td>Service personnel</td>
</tr>
<tr>
<td>Calibration of the scale (check of the measuring accuracy)</td>
<td>Annually</td>
<td>Specialized service personnel</td>
</tr>
<tr>
<td>Adjustment of the scale</td>
<td>As required</td>
<td>Specialized service personnel</td>
</tr>
<tr>
<td>Verification of the scale</td>
<td>According to official specifications 2) or as required 3)</td>
<td>Officially authorized expert personnel</td>
</tr>
</tbody>
</table>

1) designation applies to the Federal Republic of Germany; corresponds to the “Recurring safety inspection” in the Federal Republic of Austria
2) applies only to scales that are subject to official verification
3) after, e.g., repair, adjustment of scales that are subject to official verification, exceeding the operational error limits

CAUTION
Always disconnect power supply before any maintenance!

Use only Dräger original parts for maintenance.

Performing the inspection

To perform the inspection of Babytherm 8004/8010, use the specifications for the safety checks.

Safety checks

The safety checks are no substitute for service measures indicated by the manufacturer, including the preventive replacement of wear parts.

1. Check the accompanying documents:
   - Latest instructions for use are available

2. Check the following functions according to the instructions for use:
   - Height adjustment
   - The flaps close securely

3. Verify that the device combination is in good condition:
   - Labels are complete and legible
   - No visible damage
   - Fuses which are accessible from the outside are in compliance with the specified values
   - Check the condition of the two heating elements. Check for signs of thermal overload.

4. Check equipment of medical device for completeness according to the instructions for use.

5. Check for electrical safety in accordance with IEC 62353.

6. Check safety features:
   - Power failure alarm
   - LEDs, display, alarm tone

Metrological checks

If required by applicable regulations, the following measurement functions must be checked at the specified intervals:

- Skin temperature (every 2 years)
Service

Conditions:

1. Skin temperature display range is called up.
2. Simulation resistances are used instead of the temperature sensors.
   The following basic values may be used:
   - 33.2 °C (R= 1586 Ω)
   - 36.0 °C (R= 1412 Ω)

37.8 °C (R= 1312 Ω)

Procedure:

- Check the display for 3 different temperature values.

Result:

- The displayed measured value may differ by no more than 0.1 °C from the value simulated by the respective resistance.

Maintenance

**WARNING**
Risk due to faulty components
Device malfunction can occur due to wear and material fatigue of components.
To maintain the function of all components, this device must be inspected and serviced at the intervals specified by the manufacturer.

**WARNING**
Risk of electric shock
Before performing any service work, disconnect all electrical connections and gas connections from power and gas supplies. Only perform service activities when there is no patient in the device.

**CAUTION**
Risk of electric shock
Do not set up or store the device in a dirty environment.
Check the device at regular intervals for dust residues.

**WARNING**
Risk due to faulty heating elements
The heating elements are designed for safe operation within the maintenance intervals. If the maintenance intervals are not adhered to or if the device is used not in accordance with the specified intended use, the patient may be at risk.
Comply with the specified maintenance interval.
Service

Maintenance intervals

<table>
<thead>
<tr>
<th>Component</th>
<th>Interval</th>
<th>Task</th>
<th>Target group</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulb for procedure light/night light</td>
<td>In case of fault</td>
<td>Exchange</td>
<td>Service personnel</td>
</tr>
<tr>
<td>See the “Overview” chapter in the associated instructions for use.</td>
<td></td>
<td>Let the bulbs cool down for at least 15 minutes.</td>
<td></td>
</tr>
<tr>
<td>Heating elements/infrared radiators</td>
<td>After 2 years, see also Page 103.</td>
<td>Exchange in pairs</td>
<td>Specialized service personnel</td>
</tr>
<tr>
<td>Bulb for phototherapy light</td>
<td>After 1000 operating hours</td>
<td>Exchange to preserve the specified therapeutic effect.</td>
<td>Specialized service personnel</td>
</tr>
<tr>
<td>Only use listed bulbs (see &quot;Technical data&quot;).</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Always replace all bulbs.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hose grommets</td>
<td>In case of fault</td>
<td>Replace as soon as the material becomes brittle and/or sticky.</td>
<td>User</td>
</tr>
</tbody>
</table>

Repairs

Dräger recommends that all repairs are carried out by DrägerService and that only authentic Dräger repair parts are used.
 Calibration (Weighing scale option)

Information on the manufacturer's initial set-up and calibration prior to the initial installation on-site are listed on the label "Information on Manufacturer's Initial Calibration". In the listed zone, the weighing scale can be operated within the specified fault tolerances. The calibration is invalid if

- the weighing scale basic settings have been changed,
- the mechanical mounting clip has been manipulated, e.g., due to a repair replacement of the weighing scale elements or changes to the SW settings, or
- the calibration has been changed in the service mode.

In these cases, a new official calibration according to national regulations is required.

If the scales were installed at the factory, the conformity assessment was conducted by the manufacturer. Observe national metrological regulations. Conduct another official calibration according to national regulations before the validity ends.

Calibration intervals (checking measuring accuracy)

<table>
<thead>
<tr>
<th>as required</th>
<th>weekly</th>
<th>every 2 months</th>
<th>every 6 months</th>
<th>once a year</th>
<th>every two years</th>
<th>Target group</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weighing scale</td>
<td>$X^1$)</td>
<td></td>
<td></td>
<td>$X^2$)</td>
<td></td>
<td>Specialized service personnel</td>
</tr>
</tbody>
</table>

1) E.g., after repair of the weighing scale or essential components.
2) Check the measuring accuracy of the measuring scale at regular intervals, commission new adjustment and official calibration as necessary.
Disposal

Disposing of the device

At the end of its service life:

- Have the medical device appropriately disposed of in accordance with applicable laws and regulations.

For countries subject to EU Directive 2002/96/EC

This device is subject to EU Directive 2002/96/EC (WEEE). In order to comply with its registration according to this directive, this device may not be disposed of at municipal collection points for waste electrical and electronic equipment. Dräger has authorized a company to collect and dispose of this device. To initiate collection or for further information, visit Dräger on the Internet at www.draeger.com. Use the Search function with the keyword "WEEE" to find the relevant information. If access to Dräger's website is not possible, contact the local Dräger Organization.
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**Ambient conditions**

For operation

- **Temperature**: 15 °C (59 °F) to 35 °C (95 °F)
- **Atmospheric pressure**: 900 hPa to 1060 hPa (13.1 psi to 15.4 psi)
- **Rel. humidity**: 0 to 75 %, no condensation
- **Air velocity in air-conditioned rooms**: max. 0.3 m/s

For storage/transport

- **Temperature**: –20 °C (–4 °F) to 60 °C (140 °F)
- **Atmospheric pressure**: 700 hPa to 1060 hPa (10.2 psi to 15.4 psi)
- **Rel. humidity**: 0 to 95 %, no condensation
- **Height above sea level (operation)**: Maximum 3000 m

**Radiant warmer**

Radiant power, measured at the nominal value of the mains voltage without additional heat sources, at a distance of 80 cm between the bed and the radiant warmer\(^1\)

- **Heat level 3**: max. 10 mW/cm²
- **Heat level 10**: max. 30 mW/cm²

**Heating elements/infrared radiators**

**Replacement interval**: Calculated based on operating hours and device service life

At an average utilization rate of 30 %, the information in the "Maintenance" chapter corresponds to approx. 5000 hours

**Lights**

- **120 V / 230 V**: max. 16 W
- **Night light**: max. 16 W

**Minimum clearance between top edge of radiant warmer and ceiling**: >50 cm

---

1) measured as per IEC 60601-2-21
Technical data

Skin temperature measurement

Sensor

<table>
<thead>
<tr>
<th>Measuring range/Display range</th>
<th>15 °C (59 °F) to 42 °C (107.6 °F)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Accuracy (sensor)</td>
<td>±0.1 °C</td>
</tr>
<tr>
<td>Accuracy of complete measuring system (not including sensor)</td>
<td>±0.2 °C</td>
</tr>
<tr>
<td>Setting range</td>
<td>35 °C (95 °F) to 37.5 °C (99.5 °F)</td>
</tr>
</tbody>
</table>

Mattress heater (Babytherm 8010 only)

<table>
<thead>
<tr>
<th>Temperature measurement</th>
<th>5 °C (41 °F) to 45 °C (113 °F)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Measuring range/display range accuracy</td>
<td>±0.5 °C between 20 °C (68 °F) and 42 °C (107.6 °F) ±2 °C in all other parts of the range</td>
</tr>
<tr>
<td>Setting range</td>
<td>30 °C (86 °F) to 38.5 °C (101.3 °F)</td>
</tr>
</tbody>
</table>

Height adjustment (optional)

<table>
<thead>
<tr>
<th>Height adjustment range</th>
<th>295 mm (11.6 inches)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Specified operation</td>
<td>Intermittent operation (1 min operation followed by 60 min cooling period)</td>
</tr>
</tbody>
</table>

Bed canopy

<table>
<thead>
<tr>
<th>CO2 concentration with bed canopy fitted</th>
<th>max. 0.5 %</th>
</tr>
</thead>
</table>

Noise emission

<table>
<thead>
<tr>
<th>Operating noise, including phototherapy</th>
<th>54 dB(A)</th>
</tr>
</thead>
</table>

Alarm volume

<table>
<thead>
<tr>
<th>Alarms</th>
<th>Low-priority alarm dB(A)</th>
<th>Medium-priority alarm dB(A)</th>
<th>High-priority alarm dB(A)</th>
<th>Mains power supply failure alarm / device error dB(A)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>min</td>
<td>max</td>
<td>min</td>
<td>max</td>
</tr>
<tr>
<td></td>
<td>-</td>
<td>-</td>
<td>50</td>
<td>75</td>
</tr>
</tbody>
</table>

Measured in front of the device at a distance of 3 m²

Instructions for use Babytherm 8004/8010
Technical data

<table>
<thead>
<tr>
<th>Alarms</th>
<th>Low-priority alarm dB(A)(^1)</th>
<th>Medium-priority alarm dB(A)(^1)</th>
<th>High-priority alarm dB(A)(^1)</th>
<th>Mains power supply failure alarm / device error dB(A)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Measured on the bed, reduced if the hinged hood is used(^2)</td>
<td>-</td>
<td>57</td>
<td>80</td>
<td>&lt;80</td>
</tr>
</tbody>
</table>

1) Initial volume can be configured, see instructions for use
2) Measured in a reverberation chamber in accordance with IEC 60601-2-21

Phototherapy (optional)

Lights

Philips Masterline Plus halogen lamp, Type 13674 12 V / 50 W (6 pcs.)

or

Philips Brilliantline Pro halogen lamp, Type 14619 12 V / 50 W (6 pcs.)

Opening angle

24°

Irradiation power at a distance of 80 cm and a usable surface of 400 x 200 mm

9.5 W/m\(^2\) ± 25 %

The intensity of the bilirubin-affecting irradiation power decreases as the number of operating hours of the bulb increases.

Irradiated useful area on the bed (mm)

- 750
- 400
- 200
- 490
Technical data

Phototherapy (optional)

Measured with LMT radiometer, type POCKET-LUX 2 (P 10 EbiC0), and an integral measurement method with the following relative spectral sensitivity:

![Photo of Relative spectral irradiation power depending on wavelength]

Relative spectral irradiation power depending on wavelength

![Photo of Percentage]

Operating data

Mains voltage

100 V / 110 to 127 V / 120 V / 220 to 240 V (to be specified on order), 50/60 Hz

<table>
<thead>
<tr>
<th>Current consumption (A)</th>
<th>Babytherm 8004</th>
<th>Babytherm 8010</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>100 V</td>
<td>110 to 127 V/120 V</td>
</tr>
<tr>
<td>without height adjustment</td>
<td></td>
<td></td>
</tr>
<tr>
<td>without phototherapy</td>
<td>4.9</td>
<td>5.7</td>
</tr>
<tr>
<td></td>
<td>****</td>
<td>9.3</td>
</tr>
<tr>
<td>with height adjustment</td>
<td></td>
<td></td>
</tr>
<tr>
<td>without phototherapy</td>
<td>8.7</td>
<td>10.3</td>
</tr>
<tr>
<td>with height adjustment</td>
<td></td>
<td></td>
</tr>
<tr>
<td>with phototherapy</td>
<td>****</td>
<td>13.9</td>
</tr>
</tbody>
</table>

Instructions for use Babytherm 8004/8010
Technical data

Maximum power consumption
- Radiant warmer: 600 W
- Mattress heater (Babytherm 8010): 160 W
- Lighting: 32 W
- Height adjustment (optional): 560 W
- Phototherapy (optional): 400 W

Fuses
- 100 V / 120 V: F 15 A UL 248-14, 2 units
- 230 V: F 10 A UL 248-14, 2 units

Height adjustment (optional)
- 100 V / 120 V: T 6.3 A L 250 V; IEC 127-2/III (2 units)
- 230 V: T 3.15 A L 250 V; IEC 127-2/III (2 units)

Standards
- IEC 60601-1
- IEC 60601-2-21, for radiant warmer
- IEC 60601-2-50, for phototherapy
- IEC 80601-2-35, for mattress heater

Electrical protection class
- I

Skin temperature sensor
- Type BF

Heating surface of mattress
- Type BF

Classification
- Class IIb

UMDNS Code
- 15-610

Electromagnetic compatibility (EMC)
- According to EN 60601-1-2

Instructions for use Babytherm 8004/8010
Technical data

Device outlets

WARNING
Only connect devices that meet the requirements of IEC 60950-1 for ungrounded SELV circuits or the requirements of IEC 60601-1 (as of the 2nd edition) for touchable secondary circuits with maximum 24 Vdc nominal voltage.

Serial interfaces COM 1 and COM 2
The two RS 232 interfaces are electrically coupled to each other and to the central alarm input.

<table>
<thead>
<tr>
<th>Protocol</th>
<th>MEDIBUS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Connector</td>
<td>9-pole Sub-D, galvanically isolated 1.5 kV against internal electronics, 0.5 kV against housing.</td>
</tr>
<tr>
<td>Baud rate</td>
<td>9600 baud</td>
</tr>
<tr>
<td>Data bits</td>
<td>8</td>
</tr>
<tr>
<td>Parity</td>
<td>None</td>
</tr>
<tr>
<td>Stop bits</td>
<td>1</td>
</tr>
</tbody>
</table>

Pin assignment

- Pin 1: Coding resistance
- Pin 2: RXD
- Pin 3: TXD
- Pin 4: Open
- Pin 5: SHLD-GND
- Pin 6: Open
- Pin 7: Modem: Internally connected
  MEDIBUS: open
- Pin 8: Modem: Internally connected
  MEDIBUS: open
- Pin 9: Coding resistance
- Housing: SHLD-GND
Technical data

MEDIBUS RS 232 interface
Level measured according to DIN 66020
Monitor cable (Part No. 83 06 488)
Wiring layout diagram

Modem interface RS 232
Connection for external modem for service purposes
Wiring layout diagram

Central alarm
Output for connecting internal paging systems
(Nurse call)
Operating voltage max. 24 V
Current max. 250 mA
Power max. 3 W
Potential-free changeover contact
Technical data

Input
Connection of the Nurse caller from up to 3 devices (the red central alarm LED lights up for every alarm forwarded from these devices to the Babytherm 8004/8010).

Potential-free make contact

Interfaces, optional
All interface signals are electrically isolated from the patient sector.
Electric strength 1.5 kV.

Dimensions

<table>
<thead>
<tr>
<th>Description</th>
<th>Measurement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Length x Width</td>
<td>1315 x 750 mm</td>
</tr>
<tr>
<td>Overall height</td>
<td>1960 mm</td>
</tr>
<tr>
<td>Unit without height adjustment</td>
<td>1896 mm</td>
</tr>
<tr>
<td>Unit with height adjustment</td>
<td>2210 mm</td>
</tr>
<tr>
<td>Working height of bed surface</td>
<td>1025 mm</td>
</tr>
<tr>
<td>without height adjustment</td>
<td>1180 mm</td>
</tr>
<tr>
<td>with height adjustment</td>
<td>220 mm</td>
</tr>
<tr>
<td>Bed surface</td>
<td>750 x 490 mm</td>
</tr>
<tr>
<td>Bed tilting</td>
<td>adjustable in small increments</td>
</tr>
<tr>
<td>Tilt</td>
<td>maximum of 20° front end down</td>
</tr>
<tr>
<td>Height of side panels</td>
<td>150 mm</td>
</tr>
<tr>
<td>Height of inner walls</td>
<td>230 mm</td>
</tr>
<tr>
<td>Size of the X-ray cartridge</td>
<td>Width 500 mm</td>
</tr>
<tr>
<td></td>
<td>Depth 480 mm</td>
</tr>
<tr>
<td></td>
<td>Height 18 mm with X-ray tray</td>
</tr>
<tr>
<td></td>
<td>23 mm without X-ray tray</td>
</tr>
</tbody>
</table>

Instructions for use Babytherm 8004/8010
Technical data

**Weight (with one cabinet)**

Unit without height adjustment

<table>
<thead>
<tr>
<th>With mattress heater</th>
<th>120 kg (265 lbs)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Without mattress heater</td>
<td>110 kg (243 lbs)</td>
</tr>
</tbody>
</table>

Unit with height adjustment

<table>
<thead>
<tr>
<th>With mattress heater</th>
<th>133 kg (293 lbs)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Without mattress heater</td>
<td>120 kg (265 lbs)</td>
</tr>
</tbody>
</table>

**Loading/loads**

<table>
<thead>
<tr>
<th>Lifting columns/standard housing</th>
<th>&lt; 50 kg</th>
</tr>
</thead>
<tbody>
<tr>
<td>of which</td>
<td></td>
</tr>
<tr>
<td>Weight of patient</td>
<td>&lt; 8.0 kg</td>
</tr>
<tr>
<td>Swivel cabinet</td>
<td>&lt; 2.4 kg (observe diagonal mounting position)</td>
</tr>
<tr>
<td>Shelf</td>
<td>&lt; 20 kg (observe maximum mounting height of 140 cm)</td>
</tr>
<tr>
<td>Stand column</td>
<td>&lt; 5.0 kg (attach accessory loads as low and as close to the stand columns as possible and avoid one-sided loading)</td>
</tr>
<tr>
<td>Maximum tilting moment</td>
<td>&lt; 25 Nm</td>
</tr>
<tr>
<td>Individual wheel load</td>
<td>&lt; 100 kg</td>
</tr>
</tbody>
</table>
Technical data

Scale

The measurement precision depends on the local geological and geographical conditions. The specified precision only applies if the device is calibrated at the installation site.

<table>
<thead>
<tr>
<th>Electric power supply</th>
<th>100 to 127 V AC, 50/60 Hz</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>220 to 240 V AC, 50/60 Hz</td>
</tr>
<tr>
<td>Power input</td>
<td>4 W</td>
</tr>
<tr>
<td>Dimensions</td>
<td></td>
</tr>
<tr>
<td>Weighing unit</td>
<td>67.3 x 33 x 3 cm</td>
</tr>
<tr>
<td></td>
<td>(26.5 x 13 x 1.2 inches)</td>
</tr>
<tr>
<td>Display unit</td>
<td>Display unit</td>
</tr>
<tr>
<td></td>
<td>(17 x 14.3 x 2.4 inches)</td>
</tr>
<tr>
<td>Weight</td>
<td></td>
</tr>
<tr>
<td>Weighing unit</td>
<td>4 kg</td>
</tr>
<tr>
<td>Display unit</td>
<td>3.6 kg</td>
</tr>
<tr>
<td>Standards</td>
<td>EC/EN 60601-1</td>
</tr>
</tbody>
</table>

Without "NAWI" option, not verifiable according to 2014/31/EU

| Measuring range        | 300 g to 8 kg             |
| Measuring accuracy     | 10 g                      |
| Resolution             | 10 g                      |

With "NAWI" option, verifiable according to 2014/31/EU

| Measuring range        | 300 g to 8 kg             |
| Measuring accuracy     | (e) 10 g                  |
| Resolution             | (d) 10 g                  |

Classification of the integrated patient weighing scales

Class III non-automatic weighing scale

(in accordance with EC Directive 2014/31/EU)

1) The weighing scale absolute measured values are dependent on the prevailing acceleration due to gravity (g) on-site. The influence of the geographical location (geographical latitude and altitude) must therefore be considered, if necessary, by means of an on-site calibration prior to commissioning if the initial calibration information does not correspond to the respective gravitational zone or no gravitational zone has been designated.

Instructions for use Babytherm 8004/8010
Calibration

NOTE
The scale has been calibrated before delivery. In certain geographical regions, the precision specified in the Technical Data will only be attained if the device is recalibrated at the installation site. Calibration must be performed by authorised personnel only.

EMC declaration

General information
The EMC compliance of the product has been evaluated with the external cables, transducers, and accessories specified in the list of accessories. Other accessories which do not affect EMC compliance may be used if no other reasons forbid their use (see other sections of these instructions for use). The use of noncompliant accessories can result in increased emissions or decreased immunity of the medical device.

The medical device may only be used adjacent to or stacked with other devices if this configuration is approved by Dräger. If adjacent or stacked use of configurations not approved by Dräger is inevitable, verify correct functioning of the medical device in this configuration before it is used. In any case, strictly observe the instructions for use of the other devices.
Electromagnetic emissions

Electromagnetic environment
The medical device is intended for use in an electromagnetic environment as specified below. The user must ensure its use in such an environment.

<table>
<thead>
<tr>
<th>Emissions</th>
<th>Compliance according to</th>
<th>Electromagnetic environment</th>
</tr>
</thead>
<tbody>
<tr>
<td>High-frequency emissions (CISPR 11)</td>
<td>Group 1</td>
<td>The medical device only uses high-frequency energy for its internal functions. Therefore its high frequency emissions are very low and it is unlikely that adjacent electronic devices will be impaired.</td>
</tr>
<tr>
<td></td>
<td>Class A</td>
<td>The medical device is suitable for use in all establishments other than domestic establishments and those directly connected (without transformer) to the public low-voltage power supply network that supplies buildings used for domestic purposes.</td>
</tr>
<tr>
<td>Harmonic emissions (IEC 61000-3-2)</td>
<td>Not applicable</td>
<td></td>
</tr>
<tr>
<td>Transmission of voltage fluctua- tions and flicker (IEC 61000-3-3)</td>
<td>Not applicable</td>
<td></td>
</tr>
</tbody>
</table>
Electromagnetic immunity

The medical device is intended for use in an electromagnetic environment as specified below. The user must ensure that the medical device is operated in such an environment.

<table>
<thead>
<tr>
<th>Immunity against</th>
<th>IEC 60601-1-2/ IEC 80601-2-35/ IEC 60601-2-21/-50</th>
<th>Compliance level (medical device)</th>
<th>Electromagnetic environment</th>
</tr>
</thead>
</table>
| Electrostatic discharge (ESD) (IEC 61000-4-2) | Contact discharge: ±6 kV | ±6 kV | Floors should be wood, concrete, or ceramic tiles. If floors are covered with synthetic material, the relative humidity should be at least 30 %.
| | Air discharge: ±8 kV | ±8 kV | |
| Electrical fast transients/bursts (IEC 61000-4-4) | Power cables: ±2 kV | ±2 kV | Mains voltage quality should be that of a typical commercial or hospital environment.
| | Longer input cables/output cables: ±1 kV | ±1 kV | |
| Impulse voltages/surges (IEC 61000-4-5) | Common mode voltage: ±2 kV | ±2 kV | Mains voltage quality should be that of a typical commercial or hospital environment.
| | Differential mode voltage: ±1 kV | ±1 kV | |
| Magnetic field at supply frequency (50/60 Hz) (IEC 61000-4-8) | 3 A/m | 3 A/m | Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.
| Voltage dips and short-term interruptions of the supply voltage (IEC 61000-4-11) | Dip >95%, 0.5 periods | >95%, 0.5 periods | Mains voltage quality should be that of a typical commercial or hospital environment. If the user of the medical device requires continued operation during mains power supply interruptions, it is recommended that the medical device be powered from an uninterruptible power supply or a battery.
| | Dip 60 %, 5 periods | 60 %, 5 periods |
| | Dip 30 %, 25 periods | 30 %, 25 periods |
| | Dip >95%, 5 seconds | >95%, 5 seconds |
## Technical data

### Immunity against radiated high-frequency disturbance (IEC 61000-4-3)

<table>
<thead>
<tr>
<th>Frequency Range</th>
<th>Compliance Level</th>
<th>Electromagnetic Environment</th>
</tr>
</thead>
<tbody>
<tr>
<td>80 MHz to 2.5 GHz:</td>
<td>10 V/m 3 V/m</td>
<td>Recommended minimum distance between portable and mobile radio frequency transmitters with transmission power PEIRP and the medical device including its lines: (1.84 m (6.04 ft) x (\sqrt{\text{PEIRP}}))</td>
</tr>
</tbody>
</table>

### Conducted high-frequency disturbances (IEC 61000-4-6)

<table>
<thead>
<tr>
<th>Frequency Range</th>
<th>Compliance Level</th>
<th>Electromagnetic Environment</th>
</tr>
</thead>
<tbody>
<tr>
<td>150 kHz to 80 MHz:</td>
<td>10 V</td>
<td>Recommended minimum distance between portable and mobile radio frequency transmitters with transmission power PEIRP and the medical device including its lines: (1.84 m (6.04 ft) x (\sqrt{\text{PEIRP}}))</td>
</tr>
<tr>
<td>150 kHz to 80 MHz:</td>
<td>3 V</td>
<td></td>
</tr>
</tbody>
</table>

1) For PEIRP, insert the highest possible "equivalent isotropic radiated power" of the adjacent radio frequency transmitter in watts. In the vicinity of equipment marked with the symbol ☢, interference can occur. Field strengths from fixed, portable, or mobile radio frequency transmitters at the location of the medical device should be less than 3 V/m within the frequency range from 150 kHz to 2.5 GHz and less than 1 V/m above 2.5 GHz.

2) ISM bands in this frequency range are: 6.765 MHz to 6.795 MHz; 13.553 MHz to 13.567 MHz; 26.957 MHz to 27.283 MHz; 40.66 MHz to 40.70 MHz.
**Technical data**

**Recommended separation distances from portable and mobile high-frequency communication equipment**

The safety clearances listed in the following comply with IEC 60601-1-2.

<table>
<thead>
<tr>
<th>Max. PEIRP (W)</th>
<th>150 kHz to 2.5 GHz</th>
<th>All other frequencies</th>
<th>Examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.03</td>
<td>0.32 m (1.1 ft)</td>
<td>0.96 m (3.2 ft)</td>
<td>WLAN 5250/5775 (Europe)</td>
</tr>
<tr>
<td>0.10</td>
<td>0.58 m (1.9 ft)</td>
<td>1.8 m (5.9 ft)</td>
<td>WLAN 2440 (Europe)</td>
</tr>
<tr>
<td>0.17</td>
<td>0.76 m (2.5 ft)</td>
<td>2.3 m (7.6 ft)</td>
<td>Bluetooth, RFID 2.5 GHz</td>
</tr>
<tr>
<td>0.20</td>
<td>0.82 m (2.7 ft)</td>
<td>2.5 m (8.2 ft)</td>
<td>WLAN 5250 (not in Europe)</td>
</tr>
<tr>
<td>0.25</td>
<td>0.92 m (3.0 ft)</td>
<td>2.8 m (9.2 ft)</td>
<td>UMTS cellular phones</td>
</tr>
<tr>
<td>0.41</td>
<td>1.2 m (3.9 ft)</td>
<td>3.5 m (12 ft)</td>
<td>Cordless DECT telephones</td>
</tr>
<tr>
<td>0.82</td>
<td>1.7 m (5.6 ft)</td>
<td>5 m (16 ft)</td>
<td>RFID 13.56 MHz</td>
</tr>
<tr>
<td>1.00</td>
<td>1.8 m (5.9 ft)</td>
<td>5.5 m (18 ft)</td>
<td>WLAN 5600 (not in Europe)</td>
</tr>
<tr>
<td>1.64</td>
<td>2.4 m (7.9 ft)</td>
<td>7.1 m (23 ft)</td>
<td>GSM 1800/GSM 1900</td>
</tr>
<tr>
<td>3.3</td>
<td>3.3 m (11 ft)</td>
<td>10.00 m (33 ft)</td>
<td>GSM-900 cellular phones, RFID 868 MHz</td>
</tr>
</tbody>
</table>

**Reduced separation distances to portable and mobile radio frequency communication devices**

The following separation distances are based on tests performed by Dräger to determine the minimum separation distances that are absolutely necessary. These reduced separation distances only apply to mobile high-frequency communication equipment that uses the listed standards.

<table>
<thead>
<tr>
<th>Mobile radio frequency communication equipment using ...</th>
<th>Separation distance</th>
</tr>
</thead>
<tbody>
<tr>
<td>GSM 850, GSM 900, RFID 868 MHz (limited to 2 W ERP)</td>
<td>2.60 m (8.53 ft)</td>
</tr>
<tr>
<td>GSM 1800, GSM 1900 (limited to 1 W ERP)</td>
<td>1.83 m (6 ft)</td>
</tr>
<tr>
<td>UMTS, DECT (limited to 0.25 W ERP)</td>
<td>0.91 m (3 ft)</td>
</tr>
<tr>
<td>Bluetooth, WLAN 2450, RFID 2450 (limited to 0.1 W ERP)</td>
<td>0.58 m (1.9 ft)</td>
</tr>
</tbody>
</table>
### Construction and description

**Alarm description** ........................................ 121

**Scale** .......................................................... 122
Babytherm consists of:
- a chassis with infant bed and stand
- a control unit with operating panel
- a radiant warmer and
- a heated gel mattress (for Babytherm 8010).

**Control unit**
The control unit is positioned at the head end underneath the radiant warmer and between the columns of the stand. It has an operating panel and display, and an integral text display with 2 x 40 characters for easier user guidance.

**Radiant warmer**
The radiant warmer contains two infrared ceramic radiating elements and 2 lamps for illuminating the bed surface.

The bed surface lighting can be adjusted for working lighting and night lighting. The good colour reproduction of the lamps ensures easier detection of diagnostically important skin colour nuances.

The radiant warmer can be operated in "manual control mode" or "baby control mode". In manual mode, fixed heater levels of 1 to 10 are set.

In "baby control mode", the skin temperature is constantly adjusted towards a preselected optimum setting. In this mode, the output level of the radiant warmer is automatically adjusted to the needs of the patient.

In both modes, self-check routines and alarm systems provide appropriate system monitoring.

**ThermoMonitoring**
Thermal monitoring is possible with a second skin temperature sensor. Skin temperatures readings are displayed on the controller and can be graphically represented by means of the optional interface (RS 232) and a monitor. This facility improves the diagnostic possibilities of the system.

**Phototherapy**
Light from the halogen lights contributes to photochemical breakdown of the bilirubin in the skin. Dräger halogen lights emit particularly effectively at wavelengths around 460 nm. Unlike the phototherapy units of the past, the colour of the light is not blue, and so the skin colour is accurately visible. Phototherapy is switched on/off from the operating panel on the control unit.

**Mattress heater**
The heated gel mattress consists of the following components: gel mattress, aluminium heating plate and an electronic monitoring and control unit.

The gel mattress consists of a highly heat-conductive gel that does not run or dislocate even when the bed is tilted. The gel is surrounded by a film of soft material compatible with the skin.

The mattress molds itself to the body contours of the patient resting on the mattress. This creates a large contact area that conducts the heat to the patient and avoids pressure points. When the gel mattress is cold, warmth is transferred from the patient to the mattress.

The gel has good heat storage properties. If the mains supply is interrupted, e.g. for in-hospital transport, the patient will be kept warm for about 15 minutes, provided the insulation is sufficient (blanket, bed canopy).

The heating system consists of a thick aluminium plate and a heater element below the plate. This system ensures that heat is distributed evenly over the entire bed surface.

The control and display panel for the heated gel mattress is located on the right-hand side of the control unit.

The mattress temperature is adjusted with reference to a set temperature entered by the user. The set temperature is only attained in the areas where the patient is in contact with the mattress. System monitoring includes self-check routines and alarms.
Configuration mode

Configuration mode is a special operating mode of the control unit. Information on the current software version and on the error protocol can be called up in this mode. The device can be configured to meet the user's specific needs: with maximum permissible skin temperature deviation, volume of soft alarm, language and display contrast. The buttons on the control panel and Nurse call can also be tested.

Description of the skin temperature control

The skin temperature control mode is selected on the control panel. At least the yellow skin temperature sensor (Skin 1) must be connected and correctly attached to the patient. The target value for the skin temperature is set by the user on the control panel.

The patient's current skin temperature is measured by the yellow skin temperature sensor (Skin 1) and compared with the target value that was set. For a target value range between 35 °C and a maximum of 37.5 °C, the device controls the output power of the radiant warmer in such a way that the difference between the target value and the actual value is minimized. If the target value set is greater than the currently measured skin temperature (skin too cold), the radiant heater receives a signal to heat more. The output power of the radiant heater increases, resulting in more heat being supplied to the patient. If the target value set is less than the currently measured skin temperature (skin too warm), the radiant heater receives a signal to heat less. The output power of the radiant heater decreases, resulting in less heat being supplied to the patient.

The patient's skin temperature changes frequently, e.g., as a result of feeding or performing care activities on the child. Deviations of a few tenths of a degree are normal. Therefore: Only change the set value for the skin temperature if the core temperature needs to be corrected. If the current skin temperature deviates by more than ±0.5 °C from the target value, an alarm sounds which can be suppressed by the user.

The alarm ceases as soon as the deviation of the measured skin temperature from the set target value falls below ±0.5 °C (see above) again.

Skin temperature measurement

Two skin temperature sensors can be connected to measure the skin temperature (yellow skin temperature sensor) and the peripheral temperature (white skin temperature sensor, optional). Only the measurement from the yellow skin temperature sensor is used for controlling the radiant heater in the "Skin temperature control" mode.

Safety mode

During skin temperature control there are various situations where the device can no longer control the skin temperature because of a device malfunction and switches to safety mode (fall back). At the same time it generates one of the following alarm messages and switches the heater off.

<table>
<thead>
<tr>
<th>Message</th>
<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Err</strong></td>
<td>Radiant heater is switched off.</td>
</tr>
<tr>
<td></td>
<td>Switch the device off and on again.</td>
</tr>
<tr>
<td></td>
<td>If the fault continues to appear, contact DrägerService.</td>
</tr>
<tr>
<td><strong>Plug in skin temp. sensor</strong></td>
<td>Radiant heater is switched off.</td>
</tr>
<tr>
<td></td>
<td>Connect the skin temperature sensor to the device.</td>
</tr>
<tr>
<td></td>
<td>or</td>
</tr>
<tr>
<td></td>
<td>Select manual operation.</td>
</tr>
<tr>
<td><strong>Skin temp. above range</strong></td>
<td>Radiant heater is switched off.</td>
</tr>
<tr>
<td><strong>Skin temp. below range</strong></td>
<td>Radiant heater is switched off.</td>
</tr>
</tbody>
</table>
The device checks whether the power-off function in case of excess temperature is functional. This is checked during the power-on test and regularly during operation. If this test fails, the device issues an alarm.

### Alarm description

The Babytherm unit distinguishes three alarm levels:

**Warning (high potential risk)**

A continuous audible alarm that cannot be muted is emitted for:
- device malfunction
- mains power supply failure

A sequence of alarms tones that can be muted, combined with a flashing red LED, is triggered for:
- mattress heater sensor failure,
- mattress heating exceeding the temperature limits,
- skin temperature sensor not connected in baby control mode,
- sensor failure of skin temperature sensor,
- skin temperature above 39 °C.

**Caution (medium potential risk)**

An alarm tone sequence that can be muted, combined with a flashing yellow LED, is triggered for:
- deviation between set and actual value of skin or mattress temperature too great,
- 15-minutes alarm (reminder to check the core temperature every 15 minutes with heater levels higher than 4).

**Message (low potential risk)**

Display messages for:
- measured values outside the measuring range,
- battery low.

Any alarm that has not been acknowledged is indicated by the central alarm lamp on the radiant warmer unit.

The text display shows a specific message for the active alarm.

If a second alarm is triggered while the audible alarm is muted, the audible alarm will be reactivated.

<table>
<thead>
<tr>
<th>Message</th>
<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Skin temp. sensor fault</td>
<td>Radiant heater is switched off. Replace the skin temperature sensor. or Select manual operation.</td>
</tr>
</tbody>
</table>

### Table

<table>
<thead>
<tr>
<th>Message</th>
<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Skin temp. sensor fault</td>
<td>Radiant heater is switched off. Replace the skin temperature sensor. or Select manual operation.</td>
</tr>
</tbody>
</table>
Construction and description

Scale

**WARNING**
Only use the integrated scale to determine the weight of the patient. Failure to follow the instructions for use can lead to severe inaccuracies in measuring the patient's weight. To safeguard critical therapeutic decisions based on the patient's weight, the weighing result should be checked against a reference measurement on an external scale.

With the optional fully integrated scale, the weight of the patient can be determined without having to remove the patient from the Babytherm. Even when the scale is installed, the X-ray drawer can still be used. To determine the patient's weight, the patient has to be lifted off the bed once, so that the scale can determine a zero point. The patient can then be placed back on the mattress. The weight is displayed on the screen.

Weighing without lifting the baby a second time (i.e. without repeated zero calibration) is possible. This option can be used when the weight has to be determined again only a short time after the first weighing operation or e.g. if the baby’s weight has to be determined with and then without nappy, or the weight with the full and empty nappy has to be determined. The entire weighing process is accompanied by brief acoustic signals (beeps), so that the nurse’s full attention can be devoted to the patient.

When weighing the patient inside Babytherm, always take care that no hoses or cables are jammed in the bed. Otherwise these components will distort the measurement. If the weight has to be measured with hoses and cables, the best results are obtained if the hoses and cables are removed from the ventilation hose holder of the bed and laid as loosely and as flat as possible on the bed.
<table>
<thead>
<tr>
<th>Name and Description</th>
<th>Part No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Babytherm 8010</td>
<td>2M 22 380</td>
</tr>
<tr>
<td>with mattress heater and baby control mode, 100 V / 120 V / 230 V</td>
<td></td>
</tr>
<tr>
<td>Babytherm 8004</td>
<td>2M 22 370</td>
</tr>
<tr>
<td>with baby control mode and optional X-ray tray, 100 V / 120 V / 230 V</td>
<td></td>
</tr>
<tr>
<td>Modular system, variants to be specified with order:</td>
<td></td>
</tr>
<tr>
<td>– Set of side panels, height 230 mm or set of side panels, height 150 mm</td>
<td></td>
</tr>
<tr>
<td>– Stand with electric height adjustment or stand with fixed column</td>
<td></td>
</tr>
<tr>
<td>– without swivel cabinet, with 1 swivel cabinet or with 2 swivel cabinets (only for stands with electric height adjustment)</td>
<td></td>
</tr>
<tr>
<td>– Interfaces</td>
<td></td>
</tr>
<tr>
<td>– Integral phototherapy</td>
<td></td>
</tr>
<tr>
<td>– Weighing scale</td>
<td></td>
</tr>
<tr>
<td>BiliLux LED phototherapy system</td>
<td>MU20100</td>
</tr>
<tr>
<td>Options (for retrofitting)</td>
<td></td>
</tr>
<tr>
<td>Set of side panels, height 230 mm</td>
<td>2M 30 301</td>
</tr>
<tr>
<td>Set of side panels, height 150 mm</td>
<td>2M 30 242</td>
</tr>
<tr>
<td>Inner panel</td>
<td>2M 20 936</td>
</tr>
<tr>
<td>Swivel cabinet</td>
<td>2M 20 638</td>
</tr>
<tr>
<td>Cabinet fixing</td>
<td>2M 20 868</td>
</tr>
<tr>
<td>Name and Description</td>
<td>Part No.</td>
</tr>
<tr>
<td>Interface option</td>
<td>2M 30 268</td>
</tr>
<tr>
<td>Phototherapy kit</td>
<td>2M 30 450</td>
</tr>
<tr>
<td>MediCable</td>
<td>83 06 488</td>
</tr>
<tr>
<td>Accessories</td>
<td></td>
</tr>
<tr>
<td>Bed canopy</td>
<td>2M 30 392</td>
</tr>
<tr>
<td>Bed canopy bracket</td>
<td>2M 21 342</td>
</tr>
<tr>
<td>Infusion holder</td>
<td>2M 21 514</td>
</tr>
<tr>
<td>Ventilation hose bracket</td>
<td>2M 21 191</td>
</tr>
<tr>
<td>Swivel tray</td>
<td>2M 21 186</td>
</tr>
<tr>
<td>Compact rail</td>
<td>2M 85 337</td>
</tr>
<tr>
<td>Cable clips, bag of 4</td>
<td>G 13 171</td>
</tr>
<tr>
<td>Shelf</td>
<td>2M 21 158</td>
</tr>
<tr>
<td>Shelf bracket</td>
<td></td>
</tr>
<tr>
<td>Centre distance: 37 mm</td>
<td>G 13 822</td>
</tr>
<tr>
<td>Rod for small items</td>
<td></td>
</tr>
<tr>
<td>Ø 25 mm, length 50 cm</td>
<td>G 15 676</td>
</tr>
<tr>
<td>Ø 25 mm, length 100 cm</td>
<td>G 15 677</td>
</tr>
<tr>
<td>Shelf rail</td>
<td>2M 21 161</td>
</tr>
<tr>
<td>Basket 150</td>
<td>M 26 146</td>
</tr>
<tr>
<td>Basket 300</td>
<td>M 26 145</td>
</tr>
<tr>
<td>Basket 600</td>
<td>M 25 121</td>
</tr>
<tr>
<td>Waste bag holder</td>
<td>M 24 695</td>
</tr>
<tr>
<td>Connecting hose clip</td>
<td>2M 85 446</td>
</tr>
<tr>
<td>Bronchial aspiration</td>
<td></td>
</tr>
<tr>
<td>VarioAir P rail</td>
<td>MP 00 545</td>
</tr>
<tr>
<td>VarioVac P rail</td>
<td>MP 00 660</td>
</tr>
<tr>
<td>Vacuum controller VarioVac D rail</td>
<td>MP 00 511</td>
</tr>
<tr>
<td>Ejector VarioAir D rail</td>
<td>MP 00 542</td>
</tr>
<tr>
<td>Clamp for 38 mm tube</td>
<td>2M 85 274</td>
</tr>
</tbody>
</table>
## Order list

<table>
<thead>
<tr>
<th>Name and Description</th>
<th>Part No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single cylinder with lid holder</td>
<td>2M 85 163</td>
</tr>
<tr>
<td>Wall bracket (trolley)</td>
<td>M 26 859</td>
</tr>
<tr>
<td>Cylinder package</td>
<td>2M 85 056</td>
</tr>
<tr>
<td>CS hoses</td>
<td>86 01 697</td>
</tr>
<tr>
<td><strong>Oxygen therapy</strong></td>
<td></td>
</tr>
<tr>
<td>Nebuliser</td>
<td>2M 85 834</td>
</tr>
<tr>
<td>Injector for 230 mm side panels</td>
<td>2M 14 190</td>
</tr>
<tr>
<td>Nebuliser</td>
<td>2M 85 835</td>
</tr>
<tr>
<td>Compact flowmeter</td>
<td>MP 01 200</td>
</tr>
<tr>
<td>O2 flowmeter 4 L, NIST rail</td>
<td>MP 04 552</td>
</tr>
<tr>
<td>O2 flowmeter 4 L, 90° NIST rail</td>
<td>MP 04 553</td>
</tr>
<tr>
<td>O2 flowmeter 16 L, neutral DIN hose</td>
<td>MP 04 575</td>
</tr>
<tr>
<td>O2 flowmeter 16 L, 90°, 90° NIST rail</td>
<td>MP 04 570</td>
</tr>
<tr>
<td>O2 flowmeter 16 L, NIST rail</td>
<td>MP 04 567</td>
</tr>
<tr>
<td>O2 flowmeter 16 L, DIN ISO hose</td>
<td>MP 04 573</td>
</tr>
<tr>
<td>DigiFlow O2, rail, 3 L, 9/16&quot;, NIST</td>
<td>MP 01 212</td>
</tr>
<tr>
<td>DigiFlow O2, rail, 3 L, M34x1.5, NIST</td>
<td>MP 01 213</td>
</tr>
<tr>
<td>DigiFlow O2, rail, 16 L, 9/16&quot;, NIST</td>
<td>MP 01 228</td>
</tr>
<tr>
<td>DigiFlow O2, rail, 3 L, M34x1.5, NIST</td>
<td>MP 01 229</td>
</tr>
<tr>
<td>MX 300-i, O2 monitor</td>
<td>MP 01 440</td>
</tr>
<tr>
<td>MX 300, O2 monitor</td>
<td>MP 01 441</td>
</tr>
<tr>
<td>MX 300, O2 oxygen cell</td>
<td>MP 01 442</td>
</tr>
<tr>
<td>MX 300-i, fastening clamp</td>
<td>MP 01 443</td>
</tr>
<tr>
<td>MX 300-i, replacement cable</td>
<td>MP 01 444</td>
</tr>
<tr>
<td>MX 300-i, T-adapter (22 mm)</td>
<td>MP 01 445</td>
</tr>
</tbody>
</table>

### External ventilation

<table>
<thead>
<tr>
<th>Name and Description</th>
<th>Part No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manual resuscitator children</td>
<td>57 02 321</td>
</tr>
<tr>
<td>Manual resuscitator babies</td>
<td>57 02 322</td>
</tr>
<tr>
<td>PEEP valve, 0 to 10 mbar</td>
<td>84 07 475</td>
</tr>
<tr>
<td>Mask, size 1, round</td>
<td>21 21 026</td>
</tr>
<tr>
<td>Breathing hose, disposable, patient-side, 1 m, 15 pcs.</td>
<td>MX 22 793</td>
</tr>
</tbody>
</table>

### Oxygen therapy

<table>
<thead>
<tr>
<th>Name and Description</th>
<th>Part No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Foam mattress for unheated bed surface</td>
<td>2M 21 012</td>
</tr>
<tr>
<td>Gel mattress for use with mattress heater</td>
<td>2M 20 827</td>
</tr>
<tr>
<td>ThermoTrace™ Skin temperature sensor, yellow, box of 5 (for single use)</td>
<td>MX 11 000</td>
</tr>
<tr>
<td>ThermoTrace™ Skin temperature sensor, white, box of 5 (for single use)</td>
<td>MX 11 001</td>
</tr>
<tr>
<td>ThermoPad™, box of 50</td>
<td>MX 11 002</td>
</tr>
<tr>
<td>Halogen lights, set of 6</td>
<td>2M 30 084</td>
</tr>
<tr>
<td>(Philips Masterline Plus, Type 13674, or Philips Brilliant-line Pro, Type 14619)</td>
<td></td>
</tr>
<tr>
<td>Service kit, lamp, LED, 220 V to 240 V</td>
<td>2M 30 689</td>
</tr>
<tr>
<td>Service kit, lamp, LED, 100 V to 127 V</td>
<td>2M 30 673</td>
</tr>
<tr>
<td>Hose grommet</td>
<td>2M 20 434</td>
</tr>
<tr>
<td>Drainage canister hook</td>
<td>2M 21 293</td>
</tr>
<tr>
<td>Waste bag</td>
<td>M 26 240</td>
</tr>
<tr>
<td>Hug-It positioning aid, size S</td>
<td>MP 01 415</td>
</tr>
<tr>
<td>Hug-It positioning aid, size M</td>
<td>MP 01 416</td>
</tr>
<tr>
<td>Nest positioning aid</td>
<td>MP 01 417</td>
</tr>
<tr>
<td>Positioning aid set (4 small pillows)</td>
<td>MP 01 418</td>
</tr>
<tr>
<td>Name and Description</td>
<td>Part No.</td>
</tr>
<tr>
<td>-------------------------------------------------</td>
<td>------------</td>
</tr>
<tr>
<td>Cuddly toy Cally, light blue or pink</td>
<td>2M 30 462</td>
</tr>
<tr>
<td>Medela APGAR timer</td>
<td>2M 30 466</td>
</tr>
<tr>
<td>Stand for the phototherapy device</td>
<td>2M 21 190</td>
</tr>
<tr>
<td>SoundEar</td>
<td>G 24 925</td>
</tr>
<tr>
<td>SoundEar power supply unit</td>
<td>G 24 926</td>
</tr>
<tr>
<td>Service documentation on request</td>
<td></td>
</tr>
</tbody>
</table>
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Directive 93/42/EWG concerning Medical Devices

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