Caleo®
Neonatal incubator

Instructions for Use
Software 1.n
How to use the Instructions for Use

The headline ... specifies the subject of the main chapter to help you find your way around rapidly.

The page ... contains instructions for use of the unit in a combination of text and illustrations. The information is translated directly into practical actions showing the user how to use the unit.

The left-hand column ... contains text explaining the unit and guiding the user clearly to its uses through concise instructions in ergonomic sequence.

- Dots refer to actions,
- Numbers refer to the illustration next to the text and to the sequence of actions in the case of operations consisting of several steps.

The right-hand column ... contains illustrations as a reference to the text and to guide the user in handling the unit itself. The elements mentioned in the text are highlighted, and unnecessary details are omitted. The user is guided by screens confirming the various steps required for each action.

Preparation

Before using for the first time

1. Check that all packaging materials have been completely removed (see packing slip in the pack).
2. Check that the mains power supply voltage matches the voltage rating specified on the nameplate (see page 123).
3. Check that the altitude above sea level is correctly set (see page 78).
4. Screw the extension pole fully into the base frame and tighten firmly. Check that it is securely held in place.
5. Slide the holder over the pole and secure it at the required working height with the clamping screw.
6. Set the control unit to the desired working position and clamp it securely to the holder with the locking screw.

Make sure that there is sufficient space to swivel and tilt the control unit!

Fitting accessories

Screw in:
- Pole 38 mm/600 (2M 50 691) or alternatively
- Pole 38 mm/310 (2M 50 688)
- Remove the cover plate from the base frame.
- Screw the pole fully into the base frame and tighten firmly. Check that it is securely held in position.

Moving the control unit to the opposite side

4. Remove the control unit = loosen the clamping screw.
5. Remove the holder = loosen the clamping screw.
6. Move the holder to the other pole and set to the desired working height.
7. Fix the holder in position = tighten the clamping screw.
8. Secure the control unit to the holder = tighten the clamping screw.
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For Your Safety and that of Your Patients

Strictly follow the Instructions for Use
Any use of the apparatus requires full understanding and strict observation of these instructions. The apparatus is only to be used for purposes specified here.

Maintenance
The apparatus must be inspected and serviced regularly by trained service personnel at six monthly intervals. Repair and general overhaul of the apparatus may only be carried out by trained service personnel. We recommend that a service contract be obtained with DrägerService and that all repairs also be carried out by them. Only authentic Dräger spare parts may be used for maintenance. Observe chapter "Maintenance Intervals".

Accessories
Do not use accessory parts other than those in the order list.

Not for use in areas of explosion hazard
This apparatus is neither approved nor certified for use in areas where combustible or explosive gas mixtures are likely to occur.

Safe connection with other electrical equipment
Electrical connections to equipment which is not listed in these Instructions for Use should only be made following consultations with the respective manufacturers or an expert.

Liability for proper function or damage
The liability for the proper function of the apparatus is irrevocably transferred to the owner or operator to the extent that the apparatus is serviced or repaired by personnel not employed or authorized by DrägerService or if the apparatus is used in a manner not conforming to its intended use. Dräger cannot be held responsible for damage caused by non-compliance with the recommendations given above. The warranty and liability provisions of the terms of sale and delivery of Dräger are likewise not modified by the recommendations given above.

Dräger Medical AG & Co. KGaA
Intended use

Therapy system providing a controlled supply of warmth, humidity* and O₂ enrichment* in the patient capsule for premature babies and sick neonates up to a body weight of 5 kg or a body length of 55 cm (when treating twins, the total body weight is limited to 5 kg).

Used in

clinical environment, where premature babies or neonates need controlled climate parameters.

The unit may only be used by properly trained personnel under the supervision of qualified medical staff familiar with the currently known risks and benefits of using an incubator.

Possibilities for nursing and therapy

— Heat therapy through control of air temperature or skin temperature
— Humidification
— O₂ therapy through controlled O₂ enrichment
— Normal or intensive care via hand ports or a large front flap
— Bed with pivoting adjustment for raising and lowering the head

With monitoring for

— Air temperature
— Skin temperature
— Relative humidity
— O₂ concentration
— Weight*

* optional equipment feature
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What's what

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4 Handle for transport
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23 Side flap
24 Tubing port (2M 50 412)
25 Feeding grommet, hood (2M 50 352)

* optional equipment feature
Operating concept

Control unit

Hard Keys
these permanently defined keys enable the user to select various functions of Caleo®:

3 Scales
4 Bed tilt
5 Menu selection/configuration
6 Changeover key: air/skin temperature control
7 Trend display
8 Suppress alarm tone
9 Lock key pad function
10 Rotary knob

Visual signals indicate alarm situations
11 Red alarm LED
12 Yellow alarm LED
13 Power failure alarm

Soft Keys
These keys, with variable functions defined by different labels on the screen, guide the user through the specific routines of the unit, from preparing for use to shutting down the unit. Depending on the current menu, different soft keys with varying functions and labels are activated. Only the soft keys required for the current menu actually appear. This precaution keeps the display clear, preventing any confusion for the user.

When a soft key is pressed, its function is activated and the relevant menu is displayed on the screen.

In the standard screen, the soft key labels are as follows:

14 Air/skin temperature
15 Humidity
16 O2

* optional equipment feature
Operating concept

Rotary knob
A single rotary knob is used to select and set parameters.

- Turn knob = select
- Press knob = confirm

Screen
By default, the measured values are displayed as numeric values (standard screen).

- Set values and actual measured values for air temperature/skin temperature
- Set values and actual measured values for relative humidity
- Set values and actual measured values for O2 concentration
- Alarms and warnings

The screen display can also include a trend graph.

* optional equipment feature
Preparation

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Preparation

Before using for the first time

- Check that all packaging materials have been completely removed (see packing slip in the pack).
- Check that the mains power supply voltage matches the voltage rating specified on the nameplate (see page 123).
- Check that the altitude above sea level is correctly set (see page 73).

1. Screw the extension pole fully into the base frame and tighten firmly. Check that it is securely held in place.
2. Slide the holder over the pole and secure it at the required working height with the clamping screw.
3. Set the control unit to the desired working position and clamp it securely to the holder with the locking screw.

Make sure that there is sufficient space to swivel and tilt the control unit!

Fitting accessories

Screw in:
- Pole 38 mm/600 (2M 50 691) or alternatively
- Pole 38 mm/310 (2M 50 688)

- Remove the cover plate from the base frame.
- Screw the pole fully into the base frame and tighten firmly. Check that it is securely held in position.

Moving the control unit to the opposite side

4. Remove the control unit = loosen the clamping screw.
5. Remove the holder = loosen the clamping screw.
6. Move the holder to the other pole and set to the desired working height.
7. Fix the holder in position = tighten the clamping screw.
8. Secure the control unit to the holder = tighten the clamping screw.
Infusion support (2M 21 514)
for pole, 38 mm

- Place the fixing clamp on the stand pillar.
- Push the infusion support into the clip and secure by tightening the clamping screw.

Swivel table (2M 21 186)
For small articles, max. load 3 kg, for mounting on 38 mm pole

- Place the clamp of the swivel table on the stand pillar and tighten the clamping screw.

Make sure that the table has space to swivel freely!

Compact rail (2M 85 337)
Max. load 5 kg, for mounting on 38 mm pole
This rail must only be mounted by qualified technical staff!

For holding accessories, e.g.
- O2 monitor

- Adjust the height of the compact rail to the required height of the mounted accessory.
  1 Fit the compact rail to the pole = push the compact rail over the pole and
  2 fasten with the screws.
**Preparation**

**Fitting accessories**

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**Bronchial aspiration system (2M 85 125)**
Follow the separate Instructions for Use of the bronchial aspirator.

- Fix the bronchial aspirator holder to the standard rail on the wall side or handle side.
- Tighten the clamping lever.
- Establish the hose connections.

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**Basic pole (2M 50 680)**
Maximum load 5 kg

*This pole must only be mounted by qualified technical staff!*

For fixing accessories, e.g.
- additional pole extensions (see page 15).
- swivel table (2M 21 186)
- monitor support plate (2M 50 085)

- Fix to the unit in accordance with the separate Assembly Instructions.

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**Tray 3020 (M 24 678)**
Maximum load must not exceed 2 kg!

- Hang the tray from the standard rail on the wall or handle side and secure in position.
Pole extensions
The following poles can be fixed to the base pole as extension:
- Pole 38 mm/600 (2M 50 691) or
- Pole 38 mm/310 (2M 50 688) or
- Pole 25 mm/600 (2M 50 689).

* Screw the pole into the base pole as far as it will go and tighten securely. Make sure that it is securely held in place.

Notebook holder (2M 22 171)
Maximum load must not exceed 3 kg!

1. Fix the holder to the handle rail of the Caleo®.
2. Align the holder horizontally with the clamping screw.

* Check that the holder is securely fixed in position and that the swivelling mechanism is functioning correctly by turning and tilting the support plate.

Monitor shelf (2M 50 085)
Maximum load must not exceed 20 kg!
Shelf for monitor and ventilation equipment.
**This shelf must only be mounted by qualified technical staff!**
To fix the monitor shelf, fit a second 38 mm pole (see page 12).

* Fit the monitor holder = slide the plate over both 38 mm poles and
3. fix in position with the screws.

The distance between the monitor shelf and the base frame must not exceed 20 cm.
Hose holder for ventilation hoses (84 11 075)

- Open the front flap.
- Raise the bed and pull it out of the incubator.
- Push the mattress slightly to one side.
- Place the hose holder in the mounting hole in the bed and fasten from underneath with the locking screw.
- Replace the bed in the incubator and close the front flap.

The hose holder can be fixed to any of the four corners of the bed.

1. Clip the ventilation hoses and cables into the clips at the end of the ventilation hose holder.

O2 enrichment with O2 control*

2. Screw the O2 connecting hose into the port underneath the incubator.
- Connect the probe to the outlet of the central O2 supply pipeline in the "park" position (see separate Instructions for Use).
  The maximum gas pressure must be 500 kPa.

O2 monitor

To monitor the O2 concentration, use an O2 monitor that has alarm limits:

- Fix the O2 monitor to the handle rail with the holder.
- Place the sensor capsule in Caleo®.
- Route the sensor cable through one of the flexible tubing ports. Where applicable, push the sensor plug into the socket of the O2 monitor (e.g. Oxydig or MiniOx) until it audibly clicks into place (see separate Instructions for Use of O2 monitor).

* optional equipment feature
Vacuum mattress (2M 17 909)
The contour of the vacuum mattress can be altered as required and is then maintained after air evacuation of the mattress. Extreme positions can therefore be obtained for special applications. The foam mattress can remain in the incubator.

- Open the front flap.
- Insert and preform the vacuum mattress.
- Place the patient on the mattress and finally adjust the mattress to the desired shape.
- Connect the vacuum mattress to the hose of the suction system.

1. Open the valve and evacuate the vacuum mattress.
2. Close the valve and disconnect the hose.
3. Close the front flap.

Fitting the drawer (2M 50 565)
Maximum load must not exceed 7 kg.

- Fit the drawer = slide the drawer box into the groove in the base frame.
Doors, ports and bed

Hand ports
To open the hand ports:
1  Press the catch: the relevant hand port swings open.

To close the hand ports:
- Push the hand ports back into place until the catch engages.

Front flap
To open the front flap:
2  Turn the two knobs inwards to the vertical position.
   The red latch becomes visible.

- Lower the front flap until it hangs down vertically towards the floor.

Avoid pinching or jamming hoses and cables in the removable double wall!

To close the front flap:
- Raise the front flap and press into position,
3  Turn the two knobs outwards to the horizontal position until you feel them click into place.

Make sure that both knobs are engaged in position!
The red latches must no longer be visible!
Side flap

- The side flap is opened and closed in the same way as the front flap (see page 18).

Canopy

To open the canopy:
1. Grasp the handle on the canopy and
2. Lift open (approx. 60°).

Do not tilt the canopy forwards!

3. Raise the side support prop, and lower the canopy until the prop is fixed in the slot of the canopy.

To close the canopy:
1. Grasp the handle on the canopy and raise it slightly.
2. Fold down the prop and
3. Close the canopy.

The canopy can be opened from both sides.

To remove the canopy:
4. Grasp the handles on the sides of the canopy with both hands.
5. Lift the canopy horizontally off the pillar elements.

To replace the canopy:
5. Replace the canopy horizontally, so that the guide pins fit into the holes in the pillar element.

Take care with the sensor unit!
**Double wall**

To remove the double wall:
- Remove the canopy and lay it upside down (on a soft, non-abrasive surface).
1. Squeeze the double wall inwards slightly, and
2. release the oblong holes of the double wall from the retaining lugs in the canopy.
- Pull the double wall out of the canopy.

To install the double wall:
- Place the canopy upside down (on a soft, non-abrasive surface).
- Insert the double wall in the canopy.
3. On one longitudinal side, fit the retaining lugs of the canopy into the oblong holes of the double wall.
- Squeeze the double wall inwards slightly and fit the retaining lugs on the second longitudinal side of the canopy into the oblong holes of the double wall.
- Turn the canopy the right way up again. The retaining lugs of the canopy must remain seated in the oblong holes.

**Removing the bed**

Maximum load must not exceed 5 kg
- Open the front flap and fold it down.

Pull the bed out:
4. Set both knobs horizontally to the position marked ,
5. Pull the bed out towards the front as far as it will go, holding it by the recessed handle or by the knobs.

*When the bed is pulled out, monitor the patient constantly to prevent the risk of a fall.*

*Do not lean or rest any weight on the bed when it is pulled out.*
- After completing the care procedure, push the bed back in until it engages. Turn the knobs to the position and close the front flap.

*Push the bed all the way in! Otherwise the hot air duct will be interrupted, and the patient may be warmed or cooled excessively!*

*optional equipment feature*
Using the X-ray drawer

The X-ray drawer can be pulled out when the front flap is either open or closed.

To open the X-ray drawer:
1. Set both knobs vertically to the position marked ,
2. pull the drawer out by the recessed handle or the knobs.
   - Insert or remove the X-ray cassette. Recesses are provided in the X-ray drawer for positioning.

**Do not use the X-ray drawer as a writing support or bed for the patient.**

To close the X-ray drawer:
2. push the drawer inwards under the bed until it tangibly clicks into place.

Push the X-ray drawer in fully! Otherwise the hot air duct will be interrupted, and the patient may be warmed or cooled excessively!

Sealed through-holes
3. Tubing ports (2M 50 412)
4. Tubing grommets (2M 50 385)
5. Feeding grommet, hood (2M 50 352)
   - Route the cables or hoses through the flexible grommets and sleeves.

To route ventilation hoses and cables through the Caleo®, use the ventilation hose holder (page 16).

Drawer (2M 50 565)

Drawer for storing items required for nursing or treatment. The drawer is accessible from both sides.

Open the drawer:
6. grasp the drawer by the handle and pull it out as far as it will go.
   - Place the required material in the drawer.

Close the drawer:
6. push the drawer back in by the handle.
Trolley (base frame) with variable height adjustment *

To use the height adjustment facility:
● Switch on Caleo® (see page 37).
1 Press the left pedal – Caleo® is lowered.
2 Press the right pedal – Caleo® is raised.
● Adjust to a comfortable working height.
When the height stops changing, the frame is at its end position. Release the pedal.

Hoses and cables must be long enough so that they do not kink, tear or become squashed!

Do not store anything under the drawer!

The height adjustment and tilt angle adjustment cannot be operated at the same time!

The height adjustment can only be operated for max. 6 minutes during one hour.

* optional equipment feature
Tilting the bed

To tilt the bed:
- Switch on Caleo® (see page 37).
  1. Press button = raise the bed on the control unit side.
  2. Press button = lower the bed on the control unit side.
- Adjust the bed to the required tilt angle.

The end position is reached when the tilt angle stops changing. Release the button.

The entire housing of the Caleo® incubator is tilted.

Hoses and cables must be carefully routed so that they do not kink, tear or become squashed!

The height adjustment and tilt angle adjustment cannot be operated at the same time!

Do not reach in between the housing and housing support during adjustment. Danger of injury!

Setting the bed to the horizontal position

- Caleo® must be switched on (see page 37).
  1. Press button = the bed will be raised on the control unit side.
  2. Press button = the bed will be lowered on the control unit side.

The spirit levels show whether the bed is horizontal.
  3. Spirit levels for the horizontal position of Caleo® in the transverse axis.
  4. Spirit level for the horizontal position of Caleo® in the longitudinal axis. When using the integrated scales (optional, see "Weighing scale" on page 78), make sure that the unit is on a flat floor before setting.
Using humidifier systems

⚠️ Use only sterilised Aqua dest.!

**Water container (2M 50 040):**
- Disinfect hands.
- 1. Open the water container = lift up the cap.
- 2. Fill the water container with sterilised Aqua dest.

**Do not use any additives!**
Capacity: **2.8 L** (note level marks)
- 1. Close water container = push down the cap.
- 2. Prepare a fresh connection tube (MX 17 018).
- 3. Close the clamp on the connection tube.
- 4. Pierce the silicone nozzle of the water container with the pin of the connection tube.
- 5. Open the clamp on the connection tube.
- 6. Bleed the connection tube (let distilled water drain off).
- 7. Close the clamp on the connection tube.

4. Connect the LuerLock connection to the water connection pipe.
5. Open the clamp on the connection tube.

**Water bag:**

Use only sealed bags with sterilised Aqua dest.!
Do not use any additives!
Must not be confused with infusion solutions!

- Disinfect hands.
- Prepare a new connection tube (MX 17 018) and a water bag with sterilised Aqua dest.
- 6. Close the clamp on the connection tube.
- 7. Insert the pin of the connection tube into the connector of the water bag.
- 8. Open the clamp on the connection tube.
- 9. Bleed the connection tube (let distilled water drain off).
- 10. Close the clamp on the connection tube.
- 11. Connect the LuerLock connection to the water connection pipe.
- 12. Open the clamp on the connection tube.

**Replacing the water bag:**
Distilled water bag empty = the water shortage alarm on Caleo® is triggered.
- Disinfect hands.
- Close the clamp on the connection tube.
- Replace the water bag and reopen the clamp.
Integrated socket strip
The integrated socket strip can be used to connect
– infusion pumps and
– SpO2 measuring equipment.

Connections may only be made by qualified technical staff.

Caleo does not monitor the power supply to external devices.

⚠️ Do not exceed the maximum power input of the connected accessories (all 4 sockets together: max. 2A). The maximum permissible total leakage current must not be exceeded. For the leakage current of Caleo® without socket strip, see "Technical Data" on page 104.
Checking readiness for operation

Before using for the first time

- Check that the mains voltage supply corresponds to the voltage rating on the nameplate (see page 123).
- Check that the altitude above sea level is correctly set (see page 73).

Before each use

- Check that the equipment has been disinfected and cleaned in accordance with the conditions laid down by the hospital (see "Disinfecting/Cleaning/Sterilising" on page 84).
- Check that an adequate gas supply is available for the equipment to be used.
- Check that the required accessories and therapy equipment are available and in perfect condition. Only use parts that have been stripped down and sterilised. Check readiness for operation in accordance with the relevant Instructions for Use.
- Check that there are no cracks or sharp, chipped edges on the incubator canopy.
- Check that the hinges and catches on the canopy are in proper working order.
- Check that the cables and hoses have been routed correctly and safely.
- Check that there is sufficient space for adjusting the tilt and height.
- Connect to the mains supply.
- Check that the slits of the sensor unit are not fouled.

Do not use multiple-plug adapters for the Caleo® power supply!
The patient leakage current may rise above the permitted limits if the protective earth conductor fails when equipment is connected to the socket strip. The risk of electric shocks cannot be excluded in such cases.

Before using the unit, make sure that the following tests have been performed:

Disinfect hands before each test!

Check that the hand ports are secure

1. Open hand port = press catch
2. Close hand port until the catch engages.
3. Pull the edge of the hand port – it must not open.

If the hand port fails to remain engaged:
- Call DrägerService.
Check that the front flap is secure

- Open the front flap and fold it down (see page 18).
  1. Raise and press the flap closed and turn the two knobs outwards until they tangibly engage in the horizontal position.

Make sure that both knobs are engaged!
The red latches must no longer be visible!

If the front flap fails to remain engaged or if the red latches are visible:
- Call DrägerService.

Check that the side flap is secure

- Open the side flap and fold it down (see page 19).
  2. Fold up the side flap and press it closed. Turn the two knobs outwards until they tangibly engage in the horizontal position.

Make sure that both knobs are engaged!
The red latches must no longer be visible.

If the side flap fails to remain engaged or if the red latches are visible:
- Call DrägerService.

Check that the canopy is secure

3. Grasp the handle and open the canopy.
4. Lift open the flap (approx. 60°).

5. Raise the side prop.
- Lower the canopy until the prop is secured in the slot of the canopy.

If the canopy fails to remain open:
- Call DrägerService.
1. Grasp the handles on the sides of the canopy with both hands.
2. Lift the canopy horizontally off the pillar elements.

If the canopy holders are damaged:
- Call DrägerService.

Check that the double wall is securely in place
- All 4 retaining lugs in the canopy must be seated in the oblong holes of the double wall.

If the double wall or the retaining lugs in the canopy are damaged:
- Call DrägerService.

Check that the trough is secure
- Remove the canopy.
- Remove the mattress.
- Remove the bed.
3. Check the catches of the trough.
- Place the bed on the trough.
- Place the mattress on the bed.
- Refit the canopy.

If the catches for the trough are damaged:
- Call DrägerService.

Remove the bed
- Open the front flap and fold it down.
4. Set both knobs horizontally to the position marked ,
5. Grasp the bed by the recessed handle or by the knobs and pull it out towards the front as far as it will go.
5. Push the bed back until it clicks into place,
4. turn the knobs to the position.
- Close the front flap.

If the bed cannot be pulled out or pushed in or if the knobs are damaged:
- Call DrägerService.

If the bed is not fully pushed in, the hot air duct will be interrupted, causing the control system to malfunction! The result may be excessive cooling or overheating of the patient!
Activate the self-test, check audible warning tone

1. Switch on the unit = press the on/off switch until it clicks into position.

During the self-test, the functions of the machine are checked.

The audible signal, alarm beep sequence, screen displays and LEDs must be checked by the user.

- An audible warning signal and an alarm beep sequence are sounded.
  If the warning signal or alarm beeps are not sounded,
  ● Call DrägerService.

- The screen and LEDs initially go dark and are then lit.
  If individual pixels fail to light up or screen images are burnt, or if the LEDs fail to light up,
  ● Call DrägerService.

- The opening screen is displayed.
  If the opening screen is not displayed,
  ● Call DrägerService.

The unit is switched on.
Checking readiness for operation
Before each use

Check the bed tilting mechanism
1 Tilt the bed.

During the tilting process, the entire housing of the Caleo® must move uniformly. If not:
● Call DrägerService.

Do not reach in between the housing and the housing support while the unit is moving! Risk of injury!

2 Return the bed to the horizontal position (see page 23).

● The spirit levels in the bed show whether the bed is horizontal. This check is especially important when using the built-in weighing scales (see page 78)!

If the spirit levels are damaged:
● Call DrägerService.

Check the height adjustment mechanism*
● Operate both foot pedals in succession to raise and lower the Caleo® (see page 22).
● Then adjust to a comfortable working height.

When adjusting the height, the entire housing of the Caleo® must rise or descend uniformly. If not:
● Call DrägerService.

* optional equipment feature
Check the power failure alarm

- Disconnect the unit from the mains.
- The power failure LED must flash.

An intermittent tone must start. The volume of this tone must remain constant for at least 30 seconds.

If the volume decreases too soon:
- Leave the incubator connected to the mains and switched on for 24 hours to recharge the NiCd battery.
- Repeat the test.

If the volume again decreases too soon:
- Call DrägerService.

Check the fresh air filter

- Tilt the unit to remove the fresh air filter more easily (see page 23).

2 Pull the holder for the fresh air filter downwards and remove

If no filter is installed:
- Install the filter.
- Mark the date of first use on the label of the new filter and affix the label to the side of the filter.

If a filter is already installed:
- Remove the filter and check the date of use. The label is on the edge of the filter.
- Replace the filter if fouled or damaged.

If the filter is more than 2 months old:
Separate the filter frame and filter holder at the snap-in fastening and replace the old filter by a new filter.
- Mark the date of first use on the label of the new filter and affix the label to the edge of the filter.

Before operating the unit with skin temperature sensors, it is important that the sensors must be tested before they can be used on the unit (see "Using skin temperature control" on page 42).

The incubator is ready for operation when all checks have been carried out successfully.
Operation

Operation

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Precautions

Before each use, check that the unit is ready for operation (see page 26).

- Make sure that all hoses and cables are routed correctly and safely without obstruction! Otherwise: Risk of extubation! Danger of disconnection!

Lively patients must be observed with particular care.

- When the canopy, front flap, hand ports are opened or when the bed has been pulled out or when the tubing grommets are removed, the patient must be observed constantly to make sure that he or she cannot fall out of the incubator.

Do not lean or apply weight on the bed when it has been pulled out. Maximum load 5 kg

Do not use the X-ray drawer as writing support or bed for the patient.

Allow time for the incubator to warm up before use (page 105).

- Additional external heat sources, such as sunlight, heat lamps, spotlamps and electric cushions should be avoided! They cause the temperature inside the incubator to rise in an uncontrolled manner.

- The baby’s core temperature must be regularly monitored with an independent thermometer.

- The conclusions to be drawn from the measured skin temperature are the responsibility of the attending physicians.

- Skin temperature control mode must not be used on babies who are in shock or who have a high temperature!

- Skin temperature control mode must not be used on twins, since the Caleo® controls only the temperature for one baby. Risk of hypothermia or overheating! For twins, the air temperature control mode must be used.

- Make sure that you do not confuse the positions for the skin temperature sensors. Skin temperature control is regulated by the yellow skin temperature sensor (T1). If this sensor is stuck to the wrong part of the body, the patient may be excessively warmed!

- Do not place any cloths over the hot air duct. The temperature control system would be disrupted, causing a risk of burning if the air from the hot air duct is channelled directly to the patient.

When treating larger babies, the higher heater output may cause the air temperature in the Caleo® incubator to rise. In this case the double wall should be removed.

The room temperature must be at least 3 °C lower than the air temperature in the Caleo® incubator.

When treating twins, pay attention to the risk of cross-infection!

Fire hazard from supply of O₂

- No naked lights or smoking. Textiles, oil and plastics can very easily catch fire and burn rapidly in an O₂ enriched atmosphere!
- All fittings and seals in contact with oxygen must be kept free of oil and grease!
- Open valves on O₂ cylinders slowly!
- Do not use Caleo® in the presence of flammable anaesthetic gases or disinfecting agents. Risk of explosion!
- Do not use or keep flammable liquids, such as alcohol, ether or acetone, in Caleo®!
- Do not use any electrical equipment in the patient capsule, unless this equipment is expressly designed for use in explosion-hazard areas.

Note the physiological risks from O₂.

During O₂ therapy, monitor the O₂ concentration with a separate O₂ monitor.

The air in the incubator may only be enriched with O₂ when prescribed by a doctor. O₂ enrichment must be controlled on the basis of the arterially measured oxygen partial pressure in the patient’s blood. Otherwise, there is a danger of hyperoxaemia (damage to the eyes) or hypoxaemia (brain damage).

Medicaments and similar substances must not be atomised in the patient capsule. Correct functioning of the incubator may be impaired by the precipitation of atomised substances.

Do not cover the sensor unit and do not hang anything from the slits in the sensor unit. Keep the slits of the sensor unit free of impurities.

In the event of a power failure, the CO₂ level in the patient capsule may rise due to an insufficient supply of fresh air. Pay attention to the risk of CO₂ poisoning!

The central alarm light can be deactivated (see page 122). Always check the alarm LEDs on the control panel and the associated alarm beep signals.
Front flaps
When closing the front flaps, make sure that the patient is not lying in the closing path.
*The front flaps are not properly shut until the red latches are no longer visible.*
When opening and closing the front flaps, make sure that the hoses and cables are not caught in the moving double wall!

Side flaps
When opening and closing the side flaps, make sure that the hoses and cables are routed safely and clear of any obstructions!
*The side flaps are not properly shut until the red latches are no longer visible.*

Canopy
The canopy must not be used as a shelf for laying clothing, instruments etc.
Before moving the canopy, make sure that nothing has been laid on top of it.
When fitting and removing the canopy cover hold it firmly in your hand.
The canopy installation catch must engage correctly.
Do not lift the canopy when engaged.
Do not tilt the canopy forwards.
When closed, make sure that the canopy cover sits firmly in place!

Double wall
To fit and remove the double wall, remove the canopy from Caleo®.

Tilting/height adjustment
Ensure that the cables and hoses are routed correctly and safely without obstruction. Hoses and/or cables risk being trapped when tilting the Caleo® and when opening and closing the front flap.

Kangaroo mode
The patient’s core temperature must be monitored constantly, because it is outside the controlled climate.
Particular attention must be paid to critical patients vital parameters.

When using Phototherapy
Absorption of light through the baby’s skin will supply heat, which may increase the baby’s core temperature.
For this reason:
- Roughly 15 minutes before starting phototherapy, decrease the air temperature setting for the incubator air by about 2 °C.
- Decrease the set value for humidity.
- Room temperature must be at least 3 °C lower than the air temperature in Caleo®.
  This value applies to Dräger Type 4000 phototherapy units. When using other phototherapy units, especially units without a built-in fan, the temperature in the Caleo® incubator may rise even more sharply.
- Phototherapy may only be used with a stand!

The core temperature of the baby must be monitored with particular care during phototherapy!

The supply of fluid to the baby must be increased, e.g. by parental infusion, to compensate for the increased loss of water during phototherapy.

The phototherapy lamp and Caleo® canopy must not be covered with cloths, aluminium foil or other materials in order to boost the phototherapeutic effect – risk of heat build-up, as the incubator cannot be adequately cooled with ambient air.
Danger of overheating the patient.

In-house transport
An even floor surface must be ensured when moving the Caleo® incubator. The incubator must not be used outside the hospital building, as the castors may be damaged or dislodged.
The Caleo® may only be moved when empty.

High noise levels
Excessive noise levels that can disturb the patient may be caused by:
- using O₂ head boxes and delivering pressurised gases,
- wear on the bearings of the fan motor,
- placing objects on the canopy.
  • Observe the specified maintenance intervals – see page 94.

Cleaning mode may only be used when Caleo® is empty.
After use, allow Caleo® to cool down before dismantling.
Risk of burning when touching the heater!
Electrical safety

Do not use multiway adapters to connect the power supply of the Caleo®!

The patient leakage current may rise above the permitted limits if the protective earth conductor fails while equipment is connected to the socket strip. In such cases, the risk of electric shock cannot be excluded.

Only use electromedical ancillary equipment conforming to IEC / EN 60601-1.

When using the integrated socket strip: take into account the total leakage current and total current consumption!

See "Technical Data" on page 104.

Caleo® does not monitor the power supply to external devices.

Mobile telephones must not be used within 10 metres of the incubator.

Mobile telephones can interfere with the functioning of electromedical equipment and therefore endanger the patient!*

* Dräger medical equipment conforms to the interference immunity requirements laid down in product-specific standards or in EN 60601-1-2 (IEC 60601-1-2). However, depending on the design of mobile phone and the application situation, field strengths exceeding the values laid down in the specified standards may be generated in the immediate vicinity of mobile phones, thereby causing interference and malfunctions.
Switching on Caleo®

- Connect the unit to the mains.

1. Switch on the unit = press the on/off switch until it clicks into place.

An audible signal is emitted.

- The opening screen is displayed.

The incubator performs a self-test. During the self-test, the functions of the machine are checked. The audible signal and screen displays must be checked by the user, see page 29.

- After the self-test, the standard screen for air temperature control is displayed.

- The currently activated function is always highlighted by a light background.

With air temperature control, the unit takes 20 minutes to warm up. During this period the alarm for "Air temp. deviation above 1.5 °C" is suppressed.
Operation
Using air temperature control

The baby’s core temperature must be measured regularly with an independent thermometer!

Do not leave the canopy open for any length of time, otherwise the air temperature in the Caleo® will drop.

Adjusting the set value for air temperature

1. Adjust the set value = press the button.

   — The current actual measured value and the desired set value are displayed on the screen both as bar graphs and as numerical values.

   — The message »set value with rotary knob« appears at the top of the screen.

2. Increase the set value = turn the rotary knob clockwise.
3. Decrease the set value = turn the rotary knob counterclockwise.
4. Confirm the new setting = press the rotary knob.

If you do not wish to change the settings: press = the new settings are cancelled
The display returns to the standard screen. The previous set value is retained.

Or
- Wait for 7 seconds: Caleo® emits 4 short beeps to prompt the user to press the rotary knob. If the rotary knob is not pressed, the display automatically returns to the standard screen after 7 seconds, and the previous set value is retained.

Standard set value range 28 °C to 37 °C
Extended set value range 37.1 °C to 39 °C
20 °C to 27.9 °C
Default setting 33 °C

---

---
When using the extended set value range, particular care must be taken to monitor the baby’s core temperature!

If the specified set value exceeds the standard set value range
- The following message is displayed at the top of the screen:
  »confirm extended range with rotary knob«.

- Confirm the extended range = press the rotary knob.
- Continue increasing the set value = turn the rotary knob clockwise.

- The advisory message »>37.0 °C« appears on the screen.
- The following message appears at the top of the screen:
  »set value with rotary knob«.

- Confirm the set value = press the rotary knob.

If the specified set value is below the standard range
- The following message is displayed at the top of the screen:
  »confirm extended range with rotary knob«.

- Confirm the extended range = press the rotary knob.
- Continue decreasing the set value = turn the rotary knob counter-clockwise.
Operation
Using air temperature control

— The advisory message \( \Delta <28.0 \, ^\circ \text{C} \) appears on the screen.
— The following message appears at the top of the screen: "set value with rotary knob".

* Confirm the set value = press the rotary knob.

— The display returns to the standard screen. The measured values are displayed.
— The set value and \(" <28.0 \, \Delta \) are displayed alternately.

Reducing the air temperature in Caleo®

The cooling rate depends on the incubator design and can be increased by:
— removing the double wall
— reducing the outside temperature (if possible)
— reducing the set humidity value.

The cooling rate is not accelerated by:
— reducing the air temperature setting below the actually desired value.

If you do not wish to confirm the new set value:

1. Cancel adjustment of the set value = press button.

The screen returns to the standard display mode. The previous set value is retained.

Or
— Wait for 7 seconds: Caleo® emits 4 short beeps to prompt the user to press the rotary knob. If the rotary knob is not pressed, the display automatically returns to the standard screen after 7 seconds, and the previous set value is retained.

In urgent cases: open the canopy, front flap or hand ports.
In this case, the baby must be monitored constantly to ensure that he or she cannot fall out of the incubator.
Alarms
Alarm limits can be changed in the configuration (see page 71).

Example: if the deviation between the set and measured air temperature exceeds 1.5 °C:
– The screen displays the warning message «Air temp. deviation above 1.5 °C»,
– The alarm tone sequence (3 beeps) is sounded,
  1 The central alarm indicator lights up**,
  2 The measured value flashes,
  3 The yellow alarm LED flashes.

The intermittent alarm tone sequence can be muted for 15 minutes.
  4 Suppress intermittent alarm tone = press button or
  5 Press the rotary knob.
– The warning message remains on the screen,
– The intermittent alarm tone is muted,
  1 The central alarm indicator goes out,
  2 The measured value continues to flash,
  3 The yellow alarm LED continues to flash.

When the measured value returns within the range ±1.5 °C:
– The warning message disappears,
– The intermittent alarm tone is muted,
  1 The central alarm indicator goes out,
  2 The measured value remains on-screen but does not flash,
  3 The yellow alarm LED goes out.

If the air temperature is over 38 °C (or over 40 °C in the case of the extended set value range):
– The screen displays the warning message: «Air temperature too high»,
– The intermittent alarm tone sequence (5 beeps) is sounded,
  1 The central alarm light is lit**,
  2 The measured value flashes,
  6 The red alarm LED flashes.

The alarm tone can be muted for 5 minutes.
Caleo® heats up if necessary to attain the desired interior air temperature setting.
  2 The measured value continues to flash,
  6 The red alarm LED continues to flash.

When the air temperature again drops below the alarm value:
  4 Press the button to cancel the alarm.

For other alarms, see "Troubleshooting – Error Messages” on page 98. See "Alarm description” on page 121.

* The numerical values in this description are examples.
** See “Setting alarms” on page 74.
*** The central alarm light can be switched off.
  See “Setting system parameters” on page 73.
Using skin temperature control

Immediately before using the yellow skin temperature sensor or the white peripheral sensor, insert it in the yellow or white socket and wait for the measurement signal to appear on screen.

1. Measurement signal from the yellow skin temperature sensor (T1)
2. Measurement signal from the white peripheral temperature sensor (T2)

If no measurement signal appears, the sensor must be replaced (page 100).

Using skin temperature measurement in air temperature control mode or skin temperature control mode.

Connect the temperature sensors for measuring the skin temperature and peripheral temperature:
- Plug the yellow skin temperature connector into the yellow socket on the sensor unit (skin temperature, T1). When using skin temperature control, the control functions refer to this sensor.
- Plug the connector of the white peripheral temperature sensor into the white socket on the sensor unit (peripheral temperature, or skin temperature of the twin in the case of air temperature control mode, T2).
- Route the sensor cable through one of the flexible tubing grommets.
- Remove the protective foil from the adhesive pad and place the skin temperature sensor on the pad.
- Using the adhesive pad, attach the sensor tip to the appropriate part of the baby's skin.

Positioning the skin temperature sensor (yellow):
If the baby is lying on his/her back:
- Attach the yellow sensor to the abdomen, near the liver.
If the baby is lying on his/her belly:
- Attach the yellow sensor to the back, preferably near the kidneys.

Positioning the peripheral temperature sensor (white):
- Attach the white sensor to the extremities, preferably the foot or arm.

Do not use the skin temperature sensor (yellow) or peripheral temperature sensor (white) to measure rectal temperature!

Regularly check that the skin temperature sensor is correctly attached to the patient's skin. If the skin temperature sensor has fallen off, it will measure the air temperature, resulting in possible overheating of the patient (although the air temperature would not rise above 39 °C).

When a skin temperature sensor is attached but “air temperature control” is activated, the measured skin temperature is displayed.
However, in this case, the temperature is not controlled as a function of skin temperature!
Changing over between air/skin temperature control

Do not use skin temperature control mode on babies in shock, as their skin temperature is much lower than normal. Skin temperature control would increase the air temperature by too much and would endanger the patient. Use air temperature control – see page 38.

Do not use skin temperature control mode on babies with a high temperature, as their skin temperature is much higher than normal. The control function would overcool the incubator, resulting in the risk of hypothermia.

Do not use skin temperature control mode on twins, because Caleo® only controls the temperature for one baby. Danger of hypothermia or overheating!

1 Change temperature control mode = press button.
1 The LED for the control mode requiring confirmation flashes.

– The activated control mode is displayed on the screen. The actual and set values are displayed as bar graphs and numeric values.

– The upper part of the screen displays the message: »set value with rotary knob«.
Operation
Changing over between air/skin temperature control

After the new mode has been activated, you can set the desired value with the rotary knob.

1. The LED of the activated mode flashes.
2. Adjust the set value = turn the rotary knob.
2. Confirm the new set value = press the rotary knob.

— The upper part of the screen displays the advisory message «confirm new mode with rotary knob».

2. Confirm the new mode = press the rotary knob.

The display returns to the standard screen.

1. The LED of the activated mode is lit.

If you do not wish to change the settings:
3. press = the new settings are cancelled.

The display returns to the standard screen. The previous set value is retained.

Or
— Wait for 7 seconds: Caleo° emits 4 short beeps to prompt the user to press the rotary knob. If the rotary knob is not pressed, the display automatically returns to the standard screen after 7 seconds, and the previous set value is retained.
Using skin temperature control

Do not use skin temperature control mode on babies in shock, as their skin temperature is much lower than normal. Skin temperature control would increase the air temperature by too much and would endanger the patient. Use air temperature control mode – see page 38.

Do not use skin temperature control mode on babies with a high temperature, as their skin temperature is much higher than normal. The control function would overcool the incubator, resulting in the risk of hypothermia.

Do not use skin temperature control mode on twins, because Caleo® only controls the temperature for one baby. Danger of hypothermia or overheating!

Make sure that you do not confuse the positions for the skin temperature sensors. Skin temperature control is regulated by the yellow skin temperature sensor (T1). If this sensor is stuck to the wrong part of the body, the patient may be excessively warmed.

Do not attach the sensor under the baby, because otherwise the core temperature, not the skin temperature, will be measured and used as control parameter!

- The baby’s core temperature must be measured regularly with an independent thermometer!

Do not leave the canopy open for any length of time, because the air temperature in Caleo® will drop!

Adjusting the set value

<table>
<thead>
<tr>
<th>Standard set value range</th>
<th>34 °C to 37 °C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Extended set value range</td>
<td>37.1 °C to 38 °C</td>
</tr>
<tr>
<td>Default setting</td>
<td>36.5 °C</td>
</tr>
</tbody>
</table>

1 Adjust the set value = press the button.
Operation
Using skin temperature control

- The actual measured value and the set value are displayed on the screen both as bar graphs and as numerical values.
- The following message is displayed in the top part of the screen »set value with rotary knob«.

1 Increase the set value = turn the rotary knob clockwise.
1 Decrease the set value = turn the rotary knob counterclockwise.
1 Confirm the new setting = press the rotary knob.

If you do not wish to change the settings:
← press = the new settings are cancelled
The display returns to the standard screen. The previous set value is retained.
Or
- Wait for 7 seconds: Caleo® emits 4 short beeps to prompt the user to press the rotary knob. If the rotary knob is not pressed, the display automatically returns to the standard screen after 7 seconds, and the previous set value is retained.

⚠ When using the extended set value range, particular care must be taken to monitor the baby’s core temperature!

If the specified set value exceeds the standard set value range,
- the following message is displayed »confirm extended range with rotary knob«.

1 Confirm the extended range = press the rotary knob.
1 Continue increasing the set value = turn the rotary knob clockwise.
— The advisory message »Δ >37.0 °C« appears on the screen.
— The following message appears at the top of the screen: »set value with rotary knob«.

Confirm the set value = press the rotary knob.

— The display returns to the standard screen. The measured values are displayed.
— The set value and »set: >37.0 Δ « are displayed alternately.

If you do not wish to confirm the set value:

Cancel adjustment of the set value = press button.
The screen returns to standard display mode. The previous set value is retained.

Or
— Wait for 7 seconds: Caleo® emits 4 short beeps to prompt the user to press the rotary knob. If the rotary knob is not pressed, the display automatically returns to the standard screen after 7 seconds, and the previous set value is retained.
Alarms

Alarm limits can be changed in the configuration (page 74).

Example: if the deviation between the set and measured skin temperature exceeds ±0.5 °C:
— The screen displays the warning message
  »Skin 1 temp. deviation above 0.5 °C«,
— The alarm tone sequence (3 beeps) is sounded,
  1 The central alarm indicator lights up**,
  2 The measured value flashes,
  3 The yellow alarm LED flashes.

The intermittent alarm tone sequence can be muted for 5 minutes:
4 Suppress intermittent alarm tone = press button, or
5 Press the rotary knob.
— The warning message remains on the screen,
— The intermittent alarm tone is muted,
  1 The central alarm indicator light goes out,
  2 The measured value continues to flash,
  3 The yellow alarm LED continues to flash.

When the measured value returns within the range ±0.5 °C:
— The warning message disappears,
— The intermittent alarm tone is muted,
  1 The central alarm indicator light goes out,
  2 The measured value remains on-screen but does not flash,
  3 The yellow alarm LED goes out.

If the sensor plug is disconnected:
— Instead of the measured value, 3 dashes are displayed.

After 3 seconds:
— The screen displays the warning message
  »Connect skin 1 sensor«.
— The intermittent alarm tone sequence (5 beeps) is sounded,
  1 The central alarm indicator lights up**,
  2 The measured value flashes,
  3 The red alarm LED flashes.
Then:
● Immediately plug in the sensor

If the sensor is defective:
— The screen displays the warning message
  »Skin 1 sensor fault«.
— The intermittent alarm tone sequence (5 beeps) is sounded,
  1 The central alarm indicator lights up**,
  2 The measured value flashes,
  3 The red alarm LED flashes.
Then:
● Change the skin temperature sensor.

As long as 3 dashes appear on screen, Caleo® does not heat. Danger of hypothermia for the patient!

* The numerical values in this description are examples.
  See "Setting alarms" on page 74.
** The central alarm light can be switched off.
  See "Setting system parameters" on page 73.
The alarm tone can be muted for 5 minutes:
1  Suppress intermittent alarm tone = press button or
2  Press the rotary knob.
   — The warning message remains on the screen,
   — The intermittent alarm tone is muted,
   — The central alarm indicator light goes out,
3  The measured value continues to flash,
4  The yellow alarm LED continues to flash.

For other alarms, see "Troubleshooting – Error Messages" on page 98.
See "Alarm description" on page 121.
Using humidity control

- Connect the humidifier system
  (see "Using humidifier systems" on page 24).

1. Set humidity control = press button.

- The actual value and current setpoint of the humidity control are displayed as bar graphs and numerical values.

Soft key assignments:

2. ↔ = Do not activate new settings.
3. off = Switch off humidity control.
4. manual = Activate manual humidity control mode.
5. auto = Activate automatic humidity control mode (see page 119).

- On activating the humidity control, AUTO humidity is proposed as default.
Using humidity control

Setting AUTO humidity
In AUTO humidity, the humidity setpoint is automatically calculated and set by the system as a function of the air temperature setting (see page 119). The maximum relative humidity depends on the ambient temperature and incubator air temperature at max. 75 % relative humidity.

1 Switch humidity control over to AUTO humidity = press button (soft key).
2 Activate AUTO humidity = press the rotary knob.

- The display returns to the standard screen. The measured value and the automatic set value are displayed.

Manually adjusting the set value

Standard set value range 30 % to 99 %
Default value 50 %

3 Switch humidity control over to manual mode = press button (soft key).
4 Increase set value = turn rotary knob clockwise.
4 Decrease set value = turn rotary knob counter-clockwise.
4 Confirm set value = press rotary knob.

The maximum relative humidity attained is between 85 % and 99 % r.h., depending on the ambient temperature and the incubator air temperature.
Operation
Using humidity control

- The actual value and the current set value of the humidity control are displayed as bar graphs and numerical values.

- The display returns to the standard screen. The measured values and set values are displayed.
Alarms

In the event of water shortage
– The following warning message appears on the screen
  »Water empty, please refill«,
– The alarm tone sequence (3 beeps) is sounded,
– The central alarm indicator lights up,
  1 The measured value flashes,
  2 The yellow alarm LED flashes.

Fill the water container or replace the water bag – see page 24 and the following pages.

The intermittent alarm tone sequence can be muted for 15 minutes:
  3 Suppress intermittent alarm tone = press button,
  or
  4 Press the rotary knob.
– The warning message remains on the screen,
– The intermittent alarm tone is muted,
– The central alarm indicator light goes out,
  1 The measured value continues to flash,
  2 The yellow alarm LED continues to flash.

When the cause of the alarm has been remedied,
– The warning message disappears,
– The intermittent alarm tone is muted,
– The central alarm indicator light goes out,
  1 The measured value remains on the screen but does not flash,
  2 The yellow alarm LED goes out.

For other alarms, see "Troubleshooting – Error Messages" on page 98.
See "Alarm description" on page 121.

* The central alarm light can be switched off.
See "Setting system parameters" on page 73.
Using O2 control

Note the physiological risks from O2!

The air in the incubator should only be enriched with O2 when prescribed by a doctor. O2 enrichment must be controlled on the basis of the arterially measured oxygen partial pressure in the patient's blood. Otherwise, there is a danger of hyperoxaemia (damage to the eyes) or hypoxaemia (brain damage).

- Connect the probe of the O2 hose to the outlet of the central medical gas pipeline supply (page 16).

1. Set O2 control = press soft key button.

- The actual value and the current set value of the O2 control are displayed as bar graphs and numerical values.

Soft key assignments:

2. \( \rightarrow \) = Do not activate new settings.

3. off = Switch off O2 control.

4. on = Switch on O2 control.

- After switching on, automatic O2 control is active.
- The following message is displayed at the top of the screen: »set value with rotary knob«.

The current measured value is displayed after a few seconds.

* optional equipment feature
**Adjusting the set value**

- **Standard set value range**: 21 Vol.% to 40 Vol.%
- **Extended set value range**: 40.1 Vol.% to 75 Vol.%
- **Default setting**: 21 %

1. **Increase set value**: turn rotary knob clockwise.
2. **Decrease set value**: turn rotary knob counter-clockwise.
3. **Confirm set value**: press rotary knob.

---

The current actual measured value and the set value are displayed on the screen both as bar graphs and as numerical values.
**Note the dangers of high O₂ concentrations (see page 34)!**

If the set value exceeds the standard set value range

- The following message is displayed at the top of the screen: »confirm extended range with rotary knob«.

1. Confirm the extended set value range = press the rotary knob.
2. Continue increasing the set value = turn the rotary knob clockwise.

- The advisory message » **>40 %**« appears on the screen.
- The following message appears at the top of the screen »set value with rotary knob«.

1. Confirm the set value = press the rotary knob.

- The display returns to the standard screen. The measured values are displayed.

- The set value and »set: >40 ** « are displayed alternately.
Alarms

Alarm limits can be changed in the configuration (see page 74).

Example: if the deviation between the set and measured O₂ concentration exceeds ±5 %:
- The screen displays the warning message "Oxygen deviation above 5 %",
- The alarm tone sequence (5 beeps) is sounded,
- The central alarm indicator lights up**,
1 The measured value flashes,
2 The red alarm LED flashes.

The intermittent alarm tone sequence can be muted for 2 minutes:
3 Suppress intermittent alarm tone = press button or
4 Press the rotary knob.
- The warning message remains on the screen,
- The intermittent alarm tone is muted,
- The central alarm indicator goes out,
1 The measured value continues to flash,
2 The red alarm LED continues to flash.

When the measured value returns within the range ±5 Vol.%:
- The warning message disappears,
- The intermittent alarm tone is muted,
- The central alarm indicator goes out,
1 The measured value remains on-screen but does not flash,
2 The red alarm LED goes out.

For other alarms, see "Troubleshooting – Error Messages" on page 98.
See "Alarm description" on page 121.

* The numerical values in this description are examples.
** See "Setting alarms" on page 74.
See "Setting system parameters" on page 73.
### Selecting menus

1. Select menu = press »menu« button.

The required mode can be selected from the menu displayed:

2. Select item = turn the rotary knob.
2. Confirm selection (activate item) = press the rotary knob.

---

**Kangaroo mode**

See page 116 for a description of Kangaroo mode.

The patient’s core temperature or skin temperature must be constantly monitored, because it is outside the controlled climate. Particular attention must be paid to critical patients vital parameters. Ensure that all hoses and cables are routed correctly and safely without obstruction!

In **Kangaroo mode the incubator is operated in air temperature control mode**. If Caleo® was previously operated in air temperature mode, the set value for air will be activated in Kangaroo mode. If Caleo® was previously operated in skin temperature mode, the last air temperature value will be loaded as set value. The yellow skin temperature sensor and the peripheral temperature sensor can be used in Kangaroo mode to monitor the skin temperature of the baby. The alarm limits for monitoring must be redefined (see page 74).

The previously set values for
- Humidity (page 51) and
- O2 (page 55) are retained in Kangaroo mode.

The previous set value for skin temperature control is stored in buffer memory.
Activating Kangaroo mode

1. Display the main menu = press »menu« button.

Select "Kangaroo mode" from the menu.

2. Select item = turn the rotary knob.
2. Confirm and activate item = press the rotary knob.

— The screen displays the following advisory message:
   »You are leaving the current mode. Please confirm the new Kangaroo Mode with rotary knob«.

2. Confirm Kangaroo mode = press the rotary knob.

— When activated, Kangaroo mode is highlighted on screen by a light background.
— The duration of Kangaroo mode can be displayed on the screen in minutes and seconds (mm:ss).

Alarm muting is automatically activated, i.e. all alarms occurring within the next 5 minutes for the following parameters
— Air temperature deviation
— Kangaroo mode
— O2 deviation
are automatically displayed as "acknowledged" for the relevant period. See "Alarm suppression" on page 77.

● The tubing grommets can be removed from the pillar elements so that the hoses and cables remain well ordered during Kangaroo mode.
Alarms

Alarm limits can be changed in the configuration (page 74).

If the skin temperature of the yellow skin temperature sensor (skin 1) falls below the alarm limit set in the configuration:

- The screen displays the warning message
  »Skin 1 temperature below 36.0 °C«.
- The alarm tone sequence (3 beeps) is sounded,
  1 The central alarm indicator lights up,
  2 The measured value flashes,
  3 The yellow alarm LED flashes.

The intermittent alarm tone sequence can be muted for 15 minutes.
  4 Suppress intermittent alarm tone = press button or
  5 Press the rotary knob.

- The warning message remains on the screen,
- The intermittent alarm tone is muted,
  1 The central alarm indicator goes out,
  2 The measured value continues to flash,
  3 The yellow alarm LED continues to flash.

When the measured value returns above the alarm limit:
- The warning message disappears,
- The intermittent alarm tone is muted,
  1 The central alarm indicator goes out,
  2 The measured value remains on-screen but does not flash,
  3 The yellow alarm LED goes out.

For other alarms, see "Troubleshooting – Error Messages" on page 98.
See "Alarm description" on page 121.

* The numerical values in this description are examples.
See "Setting alarms" on page 74.
** The central alarm light can be switched off.
See "Setting system parameters" on page 73.
Ending Kangaroo mode

1. Display menu list = press »menu« button.
   Select the item »Back to air mode« or »Back to skin mode« from the displayed menu.

2. Select item = turn the rotary knob.
2. Activate item = press the rotary knob.

   - The screen displays the advisory message »You are leaving the current mode. Please confirm the new Air/Skin mode with rotary knob«.

2. Exit Kangaroo mode = press the rotary knob.

   The former operating status with the previous set values is reactivated. The display returns to the standard screen.

After quitting Kangaroo mode, reinsert the tubing grommets in the pillar element.

- To maintain Kangaroo mode = press « key.
Or
   - Wait for 7 seconds: Caleo® emits 4 short beeps to prompt the user to press the rotary knob. If the rotary knob is not pressed, the display automatically returns to the standard screen after 7 seconds, and the previous set value is retained.
Trend display

The trend screen is used for the graphical and numerical display of the measurement parameters. The data window always shows the last data in the selected time interval. In addition, the current measured values and set values are numerically displayed.

Switching over between standard/trend screen

1. Display trend = press » « button.

Selecting trend display

To set the trend you wish to display:

2. Open the menu = press the »menu« button.

Select the "Trend Main Page" item from the displayed menu.

3. Select menu item = turn the rotary knob.

3. Activate menu item = press the rotary knob.

Select the desired trend display from the displayed list.

Defaults:

– Air temperature trend
– Zoom factor 3 hours

3. Select trend = turn the rotary knob.

3. Activate trend = press the rotary knob.

* Back to menu = press » « key.
— The trend for the air temperature over the last 3 hours is displayed. The current air and skin temperature values are displayed on the right next to the trend.

— T1: Yellow skin temperature sensor (Skin 1)
— T2: White peripheral sensor (Skin 2)

- To exit trend display mode = press » « button.

The display returns to the standard screen.

**Setting the time interval (Zoom)**
1. Display menu = press «menu» button.

Select the “Trend Main Page” item from the displayed menu.

2. Select menu item = turn the rotary knob.
2. Activate menu item = press the rotary knob.
Select the desired trend from the displayed list.

1. Display the zoom menu = press button.
2. Select the desired zoom from the menu.
   2.1 Select zoom = turn the rotary knob.
   2.2 Activate zoom = press the rotary knob.

- Back to menu = press » button (soft key).
- Back to Trend Main Page = press the »Trend« button (soft key).
— In the illustrated example, the trend for the air temperature over the last 3 hours will be displayed.

— T1: Yellow skin temperature sensor (skin 1)
— T2: White peripheral sensor (skin 2)

Trend analysis

Trend analysis is used for the graphical and numerical display of measurement parameters and their associated set values. The data window can be selected at will to any period over the last 7 days. Trend analysis can therefore be used to evaluate thermo-monitoring data.

The following parameters can be set:
— Skin temperature 1
  (yellow skin temperature sensor, skin 1, T1)
— Peripheral temperature or skin temperature 2 (for twins)
  (white temperature sensor, skin 2, T2)
— Air temperature
— Relative humidity (rel. %)*
— O2 concentration (Vol %)*.

While trend analysis is in progress, no current measured values (air, skin, etc.) are displayed. The numerical values displayed are values from past readings.

If no button is pressed for 2 minutes, the display automatically reverts to the standard screen.

1 To activate trend analysis = hold down button for 4 seconds.

* optional equipment feature
The trend graph for the selected measured value is displayed on the screen.

Default values:
- Trend 1: Skin temperature
- Trend 2: Air temperature
- Zoom: 3 hours

The selected value is displayed as a trend graph. In this graph, the trend curve of the measured value is overlaid on the corresponding set value curve.

Time cursor:
The time cursor is displayed as a vertical dotted line marking a precise point of time on the graph’s time axis.

- To move the time curve on the time scale = turn the rotary knob.

The time marked by the cursor line is specified underneath the dotted line by the date and time. The start time and end time of the current time window are specified to the right and left underneath the trend graph.

- If the time cursor is moved beyond the displayed time range, the screen adapts automatically and displays the relevant time range:
  - Less recent time range = turn the rotary knob counterclockwise.
  - More recent time range = turn the rotary knob clockwise.

Data window:
The time cursor is associated with a data window situated to the right of the displayed trend. This data window shows the numerical values valid at the time marked on the time axis by the time cursor.

The following values are displayed in the data window:
1. Name of the selected parameter
2. Set value of this parameter at the marked time
3. Measured value at the marked time
**Trend selection**
A trend graph can be selected with the soft keys.

Soft Key assignments:

1. \( \leftarrow \) = Cancel. New settings will not be activated.
2. trend 1 = Select trend 1.
3. trend 2 = Select trend 2.
4. zoom = Select time interval.

Select trend 1:
2. Display the trend 1 menu = press soft key.
The following parameters can be selected as "trend 1":
   - air
   - skin
   - humidity*
   - O2*
   - weight*
3. Select trend 1 = turn rotary knob.

The newly selected trend will be displayed on the screen.

Select trend 2:
3. Display the trend 2 menu = press soft key.
The following parameters can be selected as "trend 2":
   - air
   - humidity*
   - oxygen*
   - weight*
   - disable
4. Select trend 2 = turn rotary knob.
5. Confirm (activate) selection = press rotary knob.

* = optional equipment feature
The newly selected trend is displayed on the screen underneath trend 1. In addition, a second data window containing the relevant parameters is opened next to trend 2. The time cursor and time range (zoom) are the same for both trend displays. The «disable» option removes trend 2 from the screen, so that only trend 1 is displayed.

Selecting the time interval (zoom)

1. Open the Zoom menu = press soft key.

The following intervals of time can be selected for the time window (zoom) function:
- 3; 6; 12; 24; 48 hours or 7 days

2. Select zoom = Turn rotary knob.
3. Activate zoom = Press rotary knob.

The selected trends will then be displayed in the newly selected time window.

All data going back a maximum of 7 days can be displayed. The individual measuring points together with the associated data for the data window are displayed. When displaying the measured weight values, up to 30 values can be stored and displayed as a trend. Values between the individual measured values are interpolated.

Ending trend analysis

To exit trend analysis
- Back to menu = press «soft key button
- Terminate trend analysis = press button
- or
- if no soft key on the screen is pressed for 2 minutes, the display will automatically revert to the standard screen.
**Cleaning mode**

Cleaning mode is only available if Caleo® is equipped with humidity control.

Cleaning mode may only be used when Caleo® is **empty**.

In cleaning mode the water heater is boiled empty and then cooled. Cleaning mode lasts about 60 minutes.

1. Open menu = press the »menu« button.
2. Select item = turn the rotary knob.
3. Confirm (activate) item = press the rotary knob.

---

Select "Cleaning Mode" from the displayed menu.

- The required operating steps are specified on the screen:
  - Disconnect Luerlock connector from the apparatus.

---

* optional equipment feature
Operation
Selecting menus

- Start Cleaning Mode = press both soft keys simultaneously.

The water heater is boiled dry.
The heater is then cooled.

Risk of burns on contact with the heater!
Do not disassemble Caleo® during cleaning mode.

- The screen indicates when cleaning mode is complete.

- Confirm end of cleaning mode = press rotary rotary knob.

After ending cleaning mode:
- Disinfect and clean Caleo® (see "Disinfecting/Cleaning/ Sterilising" on page 84).
Configuration

In configuration mode, you can set
- Language, date and time
- System parameters and
- Alarm parameters
and
you can obtain information on
- O2 sensors
- the software version.

Activating configuration mode

1. Activate configuration mode = hold down the »menu« button for 4 seconds.

2. Select configuration parameters = turn the rotary knob.
2. Activate the configuration parameters = press the rotary knob.

Cancel selection = press « key.
- The display returns to the standard screen.
Language/date/time

1 Select language/date/time = Turn and press the rotary knob.

— The language selection box is highlighted by a bright border.

1 Select language = turn the rotary knob.
1 Confirm language = press the rotary knob.

— The selected language is displayed on the screen.

The date and time are set by the same procedure.

1 Save settings = turn the rotary knob to «back» and then press the control button.

• Cancel selection = press » « soft key button.

— The display returns to the configuration parameter selection menu (see page 71).
## Setting system parameters

1. Select the temperature unit = turn and press the rotary knob.

   - The temperature unit is highlighted by a light background.

2. Select unit = turn the rotary knob.
3. Confirm unit = press the rotary knob.

   - The selected unit is displayed on the screen.

The weight unit, dT/T2 display, central alarm indicator and altitude above sea level* can be selected by the same procedure,

- The weight unit can only be set if the optional weighing scale is integrated in Caleo®.
- If using 2 skin temperature sensors, the screen can display T1 and either T2 or the difference between T1 and T2 (dT).
- If the central alarm light is deactivated, alarm situations are only indicated by the flashing "Alarm" LED on the control panel and the alarm beep signal.
- The altitude above sea level* can only be set if O2 control is integrated in Caleo®.

If the local altitude is entered incorrectly, the measurement accuracy of the O2 sensors will be reduced (e.g. 1.5 % additional error for 1000 m difference in altitude).

- Cancel selection = press « soft key button.

   - The display returns to the configuration parameter menu (see page 71).

---

* Altitude above sea level.
Setting alarms

1 Select alarms = turn and push the rotary knob.

— The menu that is then displayed allows the user to select Kangaroo mode, the initial volume of the audible alarms and the alarm limits for temperature control and O2 control.

1 Select menu item = turn rotary knob.
1 Confirm menu item = press rotary knob.

• Cancel selection = press » « soft key button.
— The display returns to the configuration parameter selection menu (see page 71).

Kangaroo mode:

Adjustment ranges:
Skin alarm T1 min 33 °C to 37 °C and off
Skin alarm T2 min 33 °C to 37 °C and off
dT alarm min –2 °C to 2 °C and off
dT alarm max 2 °C to 5 °C and off
Default values: Same as last setting

The text items in the description that follows are examples:
1 Select skin alarm $T_{1\ min}$ (lower alarm limit for skin temperature) = turn and press the rotary knob.
— The $T_{1\ min}$ alarm selection box is highlighted by a bright border.
1 Select alarm = turn the rotary knob.
1 Confirm alarm = press the rotary knob.
The display returns to the alarm settings menu (page 74).

The other alarms are set by the same method.

• Cancel selection = press » « soft key button.
— The screen returns to the configuration parameters menu (see page 71).
For a description of Kangaroo mode alarms please refer to page 117.
Initial volume (sound intensity) of the alarms:

Adjustment range: 1 to 8
Default: 1

- The initial sound intensity is highlighted by a light background.

- Set an initial sound intensity = turn rotary knob.
- Confirm initial sound intensity = press rotary knob.
The screen returns to the alarm settings menu (page 74).

- Cancel selection = press » « soft key button.
- The display returns to the configuration parameter menu (see page 71).

Alarm limits:

Adjustment ranges:
Air temperature deviation –1.5 or –2.5 °C
Skin temperature deviation ±0.3 to 1.0 °C
O₂ deviation* ±3 % or ±5 %

Default values:
Air temperature deviation –1.5 °C
Skin temperature deviation ±0.5 °C
O₂ deviation* ±5 %

- Select the alarm limit for air/skin temperature and O₂* = turn and push the rotary rotary knob.
- The alarm limit setting for the deviation in air temperature is highlighted by a bright border.

- Select air temperature deviation = turn the rotary knob.
- Confirm air temperature deviation = press the rotary knob.
The screen returns to the alarm settings menu (page 74).

The other alarm limits are set in the same way.

- Cancel selection = press » « soft key button.
- The display returns to the configuration parameter menu (see page 71).

* optional
O2 sensor information *

The screen displays the following information for the O2 sensors used:
- Date of manufacture
- Date of last calibration
- Date of next calibration.

Viewing software information

- The software version and the number of operating hours are displayed on the screen. Where applicable, this screen contains additional information on service intervals. Further information on this subject is provided in the service documentation.

1. Return to configuration parameter menu = press soft key button
   or
2. Press the rotary knob.
   - The display returns to the configuration parameter menu (see page 71).

* Optional equipment feature
Lock key pad functions

1. To lock further on-screen setting = press button.
   1. The LED on the button lights up.

   – After 4 seconds, all screen functions are locked, except for:
     1. Lock key pad function
     2. Alarm suppression
     3. Rotary knob

   1. The LED in the button remains lit.

Enabling

1. Change setting on screen = press button.
   1. The LED in the button goes out.
   – After 4 seconds, the screen functions can be changed.
     The LED in the button remains off.

Alarm suppression

2. Pressing this key if an alarm is active:
   – The audible alarm is muted
   – The central alarm indicator goes out*.

2. The LED in the button is not lit.

The permitted duration of alarm suppression depends on the type of alarm. When the cause of the alarm is removed, alarm suppression is automatically ended.

2. Pressing this key if no alarm is active (alarm muting):
For all alarms occurring in the next 4 minutes, of the type
   – Air temperature deviation
   – Skin temperature deviation
   – O₂ deviation,
no alarm tone will be sounded, and the central alarm indicator will not be lit.

However: the alarm message is displayed on the screen, and the LEDs for the measured value and alarm flash.
2. The LED in the button will be lit.

* The central alarm light can be switched off.
See “Setting system parameters” on page 73.
Weighing scale

The weighing scale is located directly underneath the bed. In the weighing process, the entire bed and the items placed on top of it are weighed with the baby. However, these extras are deducted again as soon as the baby is lifted off the bed (tare weight), so that precise weighing of the baby is possible. Before weighing, check that the bed is fully pushed in and is in the horizontal position.

The scales must be regularly checked by a test weight.

The spirit levels show whether the bed is aligned horizontally.
1 Spirit levels for the horizontal alignment of Caleo® in the transverse axis.
2 Spirit level for the horizontal alignment of Caleo® in the longitudinal axis.

To align the bed horizontally see page 23.

During the weighing process, Caleo® must not be exposed to any vibrations.

During the weighing process, no objects may be placed on the bed surface.

During the weighing process, no objects may be placed between the bed and the housing.

The weighing accuracy may be adversely affected when using the ventilation hose holder.

Starting the weighing procedure
- Remove supply lines, supply line brackets or other articles from the bed.
- 3 Activate the weighing process = Press button.

If the scale is not ready for operation, this function cannot be activated.

* optional equipment feature
During weighing, the user is guided through the sequence of operating steps by the following prompts:

- Please adjust Caleo® to horizontal position (see page 23).
- Turn knobs to X-ray position (see page 21).
- Lift baby clear of mattress.

The system waits until at least 250 g have been taken off the scale and the scale has been at rest for 3 seconds. The tare is determined.
- 1 beep is sounded.

- Place patient on bed.

The system waits until at least 250 g have been added to the scale and the scale has been at rest for 3 seconds. Weighing then proceeds.
- 1 beep is sounded.

The weighing procedure is complete.
- The current and last weighing results are displayed on the screen.

After 30 seconds, the display reverts to the standard screen. or
- Display the standard screen = press the rotary knob.

The screen displays the last weight as advisory text for the next 10 minutes.

**Weighing process cancelled**
The weighing process is cancelled if the scales are not unloaded or loaded within 30 seconds.
- 3 short beeps are emitted.
**Weighing without tare**

If the last tare weight was obtained no longer than 60 minutes earlier, and if no objects have since been removed from the bed or placed on it, recalibration with a new tare weight is not necessary.

1. Activate the weighing process = press soft key.

Caleo® proposes reweighing directly without obtaining a new tare.

2. Weigh without tare = press button.

- Lay the patient on the bed.

The system waits until the weighing scale has remained at rest for 3 seconds. Weighing then proceeds.

- 1 beep is emitted.
The weighing procedure is ended.

- The present and last weights are displayed.

After 30 seconds, the display reverts automatically to the standard screen.

or

Display the standard screen = press the rotary knob.

The screen displays the last weight as advisory text for the next 10 minutes.

---

**Ending operation**

**Switch off incubator**

1. Switch off the incubator = press the on/off switch.

The incubator is switched off.

If the O₂ hose is connected:

- Disconnect the probe from the outlet of the central O₂ supply pipeline and place it in the "Park" position (strictly follow the separate Instructions for Use).

---

* optional equipment feature
Care

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Care list ............................................................................................ 93
Maintenance intervals ................................................................. 94
Disposal ......................................................................................... 95
Disinfecting/Cleaning/Sterilising

**Risk of burns on contact with the heater**
When the incubator is closed, the heater is still hot enough to cause serious burns for a long time after switching off (still approx. 70 °C after 1 hour).

Clean and disinfect the incubator thoroughly:
- after each change of patient
- at least once a week.

Clean and disinfect the accessories, such as the suction equipment and the flowmeter in accordance with their separate Instructions for Use.

**Disinfecting/cleaning/sterilising**
Use surface disinfectants for disinfecting.
For reasons of material compatibility, use disinfectants based on
- aldehydes
- quaternary ammonium compounds.

Due to the chemical composition and direct effects on material compatibility, the following products are only suitable to a limited extent:
- Halogen-releasing compounds
- Strong organic acids
- O2 releasing compounds.

Use only cleaning agent in autoclaves and other cleaning and disinfecting machines. Do not use any alkaline or chlorine releasing disinfectants, due to the risk of corrosion. As a general rule, the manufacturer's recommendations must be taken into account when choosing a cleaning agent. Liability for the specified use and for any material damage that may be caused by the product rests with the manufacturer.

**Do not use any disinfectants or cleaning agents containing alcohol.**

We recommend the use of disinfectants based on the current DGHM list (DGHM: German Society for Hygiene and Microbiology). The DGHM list (published by mhp Verlag GmbH, Wiesbaden) also gives the composition of each disinfectant. For countries where the DGHM list is not familiar, we recommend the types of disinfectant specified above.
The following surface disinfectants are recommended:

<table>
<thead>
<tr>
<th>Surface disinfectants</th>
<th>Manufacturer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dismozon® pur</td>
<td>Bode Chemie GmbH &amp; Co., Germany</td>
</tr>
<tr>
<td>Incidur®</td>
<td>Henkel-Ecolab Deutschland GmbH</td>
</tr>
<tr>
<td>Trichlorol</td>
<td>Lysoform, Germany</td>
</tr>
<tr>
<td>Virkon</td>
<td>Tetenal, Germany</td>
</tr>
<tr>
<td>Bacillol 25</td>
<td>Bode Chemie GmbH &amp; Co</td>
</tr>
<tr>
<td>Seculyse</td>
<td>Paragerm (Henkel Ecolab), France</td>
</tr>
<tr>
<td>Sekupoudre</td>
<td>Paragerm (Henkel Ecolab), France</td>
</tr>
<tr>
<td>Vaposeptol</td>
<td>Paragerm (Henkel Ecolab), France</td>
</tr>
<tr>
<td>Cidex</td>
<td>Johnson &amp; Johnson, Taiwan</td>
</tr>
<tr>
<td>Habitane</td>
<td>Zeneca Limited, Norway</td>
</tr>
<tr>
<td>Kloramin</td>
<td>Norsk Medisinal Depot A/S, Norway</td>
</tr>
<tr>
<td>Sactiv</td>
<td>Diversey Lever, Finland</td>
</tr>
<tr>
<td>Viraclean</td>
<td>Whiteley, Australia</td>
</tr>
</tbody>
</table>

Note the manufacturer’s directions for use.

**Do not treat Caleo® and its components in formaldehyde vapour or ethylene oxide.**

The following materials are used in the patient capsule:

<table>
<thead>
<tr>
<th>Components</th>
<th>Material</th>
</tr>
</thead>
<tbody>
<tr>
<td>Canopy, flaps</td>
<td>Polycarbonate</td>
</tr>
<tr>
<td>Support pillar units</td>
<td>Styrene-butadiene thermo foam rubber</td>
</tr>
<tr>
<td>Bed</td>
<td>Styrene-butadiene thermo foam rubber</td>
</tr>
<tr>
<td>Housing</td>
<td>Polystyrene</td>
</tr>
<tr>
<td>Caleo® mattress, standard</td>
<td>Polyurethane</td>
</tr>
<tr>
<td>Caleo® SoftBed™</td>
<td>Polyurethane/polyester</td>
</tr>
</tbody>
</table>
Stripping down

Checking the fresh air filter
To make the filter easier to remove, set the incubator to the tilted position.
- Connect to the mains power supply.
- Switch on the incubator = press the on/off switch until it clicks into position.
- Tilt the bed (see page 23).

1 Pull the fresh air filter unit downwards and out of the unit.
- Check the installation date marked on the label fixed to the filter rim.

If the filter is fouled, damaged or older than 2 months, separate the filter frame and filter holder at the snap-on fitting and replace the old filter with a new filter.
- Write down the installation date of the new filter, and stick the label to the filter rim.

Removing the water supply
- Close the clamp on the connection tube.
- Remove and dispose of the water bag and connection tube or
2 Remove the water container from the holder.
3 Detach the connection tube from the water container and water connection pipe.
- Dispose of the connection tube.

- Clean the water container in the cleaning and disinfecting machine at 93 °C*
or
- sterilise at 134 °C*.

* Observe national and international standards on cleaning, disinfecting and sterilisation procedures (e.g. EN 285, EN 554, EN 556).
Removing the water connection pipe
- Tilt the incubator to make the water connection pipe and fresh air filter easier to remove (see page 23).

1  Turn the water connection pipe 90° clockwise and pull it out of the guide channel.

- Clean the water connection pipe in the cleaning and disinfecting machine at 93 °C* or
  - sterilise at 134 °C*.

- Place Caleo® horizontal (see page 23).
- Switch on cleaning mode** (see page 69).

After cleaning mode** is complete or if cleaning mode is not provided:
- Switch off the incubator and disconnect from mains.
- Remove any auxiliary equipment installed (for care instructions see the specific Instructions for Use of the equipment concerned).

Risk of burns on contact with the heater. Allow Caleo® to cool down before stripping down further.

Remove the canopy.
2  Grasp the handles on the sides of the canopy with both hands.
3  Lift the canopy horizontally off its supports (pillar elements).

- Remove visible soiling with a disposable cloth and detergent.
- Wipe-disinfect the surfaces.
- After allowing the disinfectant time to take effect (see manufacturer’s specifications), wipe the surface with a clean damp cloth and then rub dry.

* Observe national and international standards on cleaning, disinfecting and sterilisation procedures (e.g. EN 285, EN 554, EN 556).
** optional equipment feature
To remove the double wall:
- Remove the canopy and lay it upside down (on a soft, non-abrasive surface).
1. Squeeze the double wall inwards slightly, and
2. Release the oblong holes of the double wall from the retaining lugs in the canopy.
- Pull the double wall out of the canopy.
- Remove visible soiling with a disposable cloth and detergent.
- Wipe-disinfect the surfaces.
- After allowing the disinfectant time to take effect (see manufacturer’s specifications), wipe the surface with a clean damp cloth and then rub dry.

Open the front flap:
3. Turn the two locking knobs inwards as far as they will go and fold down the front flap.
- Fold out the adjustable double walls to clean them.
- Open the side flaps in the same way.
- Remove visible soiling with a disposable cloth and detergent.
- Wipe-disinfect the surfaces.
- After allowing the disinfectant time to take effect (see manufacturer’s specifications), wipe the surface with a clean damp cloth and then rub dry.

- Remove all through-hole grommets, sleeves and liners etc.
- Clean them at 93 °C in the cleaning and disinfecting machine*.

- Remove the mattress from the bed.
4. Remove the bed.
**Take care not to damage the sensor unit.**
- Remove visible soiling with a disposable cloth and detergent.
- Wipe-disinfect the surfaces.
- After allowing the disinfectant time to take effect (see manufacturer’s specifications), wipe the surface with a clean damp cloth and then rub dry.

---
* Observe national and international standards on cleaning, disinfecting and sterilisation procedures (e.g. EN 285, EN 554, EN 556).
Remove the X-ray drawer:
- Using the recessed handle, pull out the X-ray drawer as far as it will go.
1. Tilt the drawer upwards and pull it out of the unit.
- Remove visible soiling with a disposable cloth and detergent.
- Wipe-disinfect the surfaces.
- After allowing the disinfectant time to take effect (see manufacturer's specifications), wipe the surface with a clean damp cloth and then rub dry.

Remove trough:
2. Press both catches inwards and pull the trough upwards.
3. Lift out the trough.
4. Remove the fan impeller.
- Remove visible soiling with a disposable cloth and detergent.
- Wipe-disinfect the surfaces.
- After allowing the disinfectant time to take effect (see manufacturer's specifications), wipe the surface with a clean damp cloth and then rub dry.

5. Turn the trough over.
5. Lift the air guide plate to the side to disinfect and clean.
- Remove visible soiling with a disposable cloth and detergent.
- Wipe-disinfect the surfaces.
- After allowing the disinfectant time to take effect (see manufacturer's specifications), wipe the surface with a clean damp cloth and then rub dry.
Disinfecting/Cleaning/Sterilising

Stripping down

Base frame/body

- Remove visible soiling with a disposable cloth and detergent.
- Wipe-disinfect the surfaces.
- After allowing the disinfectant time to take effect (see manufacturer's specifications), wipe the surface with a clean damp cloth and then rub dry.
- Remove any impurities near the slits of the sensor unit.

Do not allow any moisture to enter the sensor.
Do not disinfect the sensor unit by immersion or spraying.

Control panel

- Pull off the turn-and-press rotary knob.
- Remove visible soiling with a disposable cloth and detergent.
- Wipe-disinfect the surfaces.
- After allowing the disinfectant time to take effect (see manufacturer's specifications), wipe the surface with a clean damp cloth and then rub dry.

Do not allow any moisture to enter the control panel.
Do not disinfect the control panel by immersion or spraying.

Use only the recommended cleaning agents and disinfectants. Otherwise, there is a risk of cracks forming in the transparent parts, e.g. due to the use of alcohol.

Do not use UV radiation on the incubator, because this treatment may cause cracks in the transparent components.
Before reusing for a patient

- Check that the system has been cleaned and disinfected in conformity with the regulations of the hospital.
- Reassemble the equipment with disinfected hands.
- Refit all equipment – see "Stripping down", page 86.

When inserting the trough, make sure that both holders click into place.

If the holders of the trough are damaged:
- Call DrägerService.

Fitting the water connection pipe:

1. Check the O-ring (2M 50 346) for damage and replace if necessary.
2. Check the locking valve for damage.

To make the water connection pipe easier to install, tilt the incubator.

- Connect the incubator to the mains power supply.
- Switch on the incubator = press the on/off switch until it clicks into position (see page 37).
- Tilt the bed (see page 23).

3. Push the water connection pipe into the guide duct, holding the LuerLock connection vertically at the top.

- After overcoming a spring force,

4. Turn the water connection pipe counter-clockwise $90^\circ$ and secure the locking valve in the holder.
Check that the incubator is ready for operation – see “Before each use” on page 26.

To remove possible disinfectant residue, we recommend running the incubator in standby mode.
- Switch on Caleo® (see page 37).
- Activate air temperature control (see page 38).
- Run Caleo® at 37 °C with opened flaps.

If using a water container:
Do not refill the water container until just before placing the patient in the incubator (see page 24)!

If using a water bag:
Do not connect the water bag until just before placing the patient in the incubator (see page 24)!
<table>
<thead>
<tr>
<th>Reusable components</th>
<th>Care intervals</th>
<th>How often</th>
<th>Wipe&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Cleaning and disinfecting machine&lt;sup&gt;b&lt;/sup&gt; 93 °C&lt;sup&gt;c&lt;/sup&gt;</th>
<th>Sterilise 134 °C&lt;sup&gt;c&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Connection tube</td>
<td>Replace weekly and with each change of patient</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Water container</td>
<td>Change of patient/weekly</td>
<td></td>
<td>yes</td>
<td></td>
<td>yes</td>
</tr>
<tr>
<td>Water connection pipe</td>
<td>Change of patient/weekly</td>
<td></td>
<td>yes</td>
<td></td>
<td>yes</td>
</tr>
<tr>
<td>Canopy</td>
<td>Change of patient/weekly</td>
<td></td>
<td>yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Double wall</td>
<td>Change of patient/weekly</td>
<td></td>
<td>yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Front flap</td>
<td>Change of patient/weekly</td>
<td></td>
<td>yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Side flap</td>
<td>Change of patient/weekly</td>
<td></td>
<td>yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Double walls</td>
<td>Change of patient/weekly</td>
<td></td>
<td>yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Through-hole sleeves etc.</td>
<td>Change of patient/weekly</td>
<td></td>
<td>yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mattress</td>
<td>Change of patient/weekly</td>
<td></td>
<td>yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>X-ray drawer</td>
<td>Change of patient/weekly</td>
<td></td>
<td>yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Trough</td>
<td>Change of patient/weekly</td>
<td></td>
<td>yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fan bar</td>
<td>Change of patient/weekly</td>
<td></td>
<td>yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fan impeller</td>
<td>Change of patient/weekly</td>
<td></td>
<td>yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Base frame</td>
<td>Change of patient/weekly</td>
<td></td>
<td>yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Control panel</td>
<td>Change of patient/weekly</td>
<td></td>
<td>yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rotary knob (control panel)</td>
<td>Change of patient/weekly</td>
<td></td>
<td>yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fresh air filter</td>
<td>Replace every 2 months</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Care list**

- **a.** Use surface disinfectants based on aldehydes and quaternary ammonium compounds.
- **b.** Use only detergent. Do not use disinfectants that release alkali or chlorine. Risk of corrosion!
- **c.** Observe national and international standards on cleaning, disinfecting and sterilisation procedures (e.g. EN 285, EN 554, EN 556).
# Maintenance intervals

Disinfect and clean the incubator or the relevant parts before each maintenance operation, even when returning the equipment for repair purposes.

**Disconnect from the mains before each maintenance operation. Risk of electric shock.**

## Replaceable parts:

<table>
<thead>
<tr>
<th>Part</th>
<th>Intervals</th>
<th>Who?</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fresh air filter</td>
<td>when necessary weekly every 2 months every 6 months once a year every two years</td>
<td>Medical and technical personnel</td>
</tr>
<tr>
<td>Grommets, sleeves etc.</td>
<td>X^a</td>
<td>Medical and technical personnel</td>
</tr>
<tr>
<td>O-ring, for water connection pipe</td>
<td>X</td>
<td>Medical and technical personnel</td>
</tr>
<tr>
<td>Fan motor</td>
<td></td>
<td>X^b Authorised technicians</td>
</tr>
<tr>
<td>Lithium battery</td>
<td></td>
<td>X Authorised technicians</td>
</tr>
<tr>
<td>Skin temperature sensors</td>
<td>X^c</td>
<td>Medical personnel</td>
</tr>
<tr>
<td>Adhesive pads</td>
<td>X^d</td>
<td>Medical personnel</td>
</tr>
<tr>
<td>Foam mattress</td>
<td>X</td>
<td>Medical personnel</td>
</tr>
</tbody>
</table>

## Maintenance:

Device servicing and maintenance | X | Authorised technicians

## Calibration:

O₂ sensors | X | Authorised technicians

---

a. Replace if the material becomes brittle or sticky or if strips of material have become detached
b. Replace after 18000 – 20000 operating hours
c. At the latest when changing patient
d. At the latest when changing patient
Disposal

Disposal of the connection tube and fresh air filter
— with household waste.

Disposal of O2 sensors and battery
— Do not throw into the fire. Risk of explosion.
— Do not force open: risk of chemical burns.
— Do not recharge the battery.
Special waste:
— Dispose of special waste in accordance with national waste disposal regulations.

Further information can be obtained from national and local environmental and legislative authorities and suitable waste disposal companies.

Disposal of incubator
At the end of its useful life.
— Dispose of the incubator in conformity with national waste disposal regulations.
or
— Hand the incubator over to a suitable waste disposal company for disposal.
Further information can be obtained from national and local environmental and legislative authorities.

All plastic components weighing more than 50 g must be marked with their material code in order to ensure environmentally sound disposal.
Troubleshooting – Error Messages

Contents

Troubleshooting – Error Messages ............................................. 98

Troubleshooting – Faults ......................................................... 102
## Troubleshooting – Error Messages

All error messages are displayed on the screen. They are listed in the table below in alphabetical order. See also "Alarm description" on page 121.

<table>
<thead>
<tr>
<th>Message</th>
<th>Cause</th>
<th>Remedy</th>
<th>Duration</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Air heater inoperable</strong></td>
<td>Air heater defective.</td>
<td>Call DrägerService.</td>
<td>1 min.</td>
</tr>
<tr>
<td>Measured value flashes on screen.</td>
<td>Red alarm LED on the control panel flashes.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Central alarm indicator lights up.*</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Intermittent audible alarm is sounded (5x).</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Air temp. deviation above 1.5 °C</strong></td>
<td>The specified tolerance limit for deviation from the set value has been exceeded by a value either above the maximum or below the minimum value</td>
<td>Reduce/increase humidity.</td>
<td>15 min.</td>
</tr>
<tr>
<td>(see &quot;Configuration&quot; on page 71.)</td>
<td>Measured value flashes on screen.</td>
<td>Close canopy, front flap and hand ports.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Yellow alarm LED on the control panel flashes.</td>
<td>Check the set value and the configuration (page 71).</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Central alarm indicator lights up.*</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Intermittent audible alarm is sounded (3x).</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Air temp. sensor inoperable</strong></td>
<td>Air temperature sensors are defective.</td>
<td>Call DrägerService.</td>
<td>1 min.</td>
</tr>
<tr>
<td>Red alarm LED on the control panel flashes.</td>
<td>Central alarm indicator lights up.*</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Intermittent audible alarm is sounded (5x).</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Air temperature too high</strong></td>
<td>Incubator temperature above 38 °C.</td>
<td>Remove external temperature sources.</td>
<td>5 min.</td>
</tr>
<tr>
<td></td>
<td>Incubator temperature above 40 °C.</td>
<td>Call DrägerService.</td>
<td></td>
</tr>
<tr>
<td>Measured value flashes on screen.</td>
<td>Red alarm LED on the control panel flashes.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Central alarm indicator lights up.*</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Intermittent audible alarm is sounded (5x).</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Battery discharged</strong></td>
<td>Flat battery.</td>
<td>Connect the incubator to the power supply for about 30 minutes</td>
<td>1 min.</td>
</tr>
<tr>
<td>Yellow alarm LED on the control panel flashes.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Connect skin 1 sensor</strong></td>
<td>Probe for skin temperature (yellow) is not connected.</td>
<td>Check connection and correct if necessary.</td>
<td>5 min.</td>
</tr>
<tr>
<td>Three dashes flash on screen in place of the measured value.</td>
<td>Red alarm LED on the control panel flashes.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Central alarm indicator lights up.*</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Intermittent audible alarm is sounded (5x).</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Fan inoperable</strong></td>
<td>Fan defective.</td>
<td>Check fan impeller.</td>
<td>5 min.</td>
</tr>
<tr>
<td>Red alarm LED on the control panel flashes.</td>
<td>Central alarm indicator lights up.*</td>
<td>Call DrägerService.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Intermittent audible alarm is sounded (5x).</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Heater temp. sensor inoperable</strong></td>
<td>Air heater temperature sensors defective.</td>
<td>Call DrägerService.</td>
<td>1 min.</td>
</tr>
<tr>
<td>Red alarm LED on the control panel flashes.</td>
<td>Central alarm indicator lights up.*</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Intermittent audible alarm is sounded (5x).</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* The central alarm light can be switched off. See "Setting system parameters" on page 73.
## Troubleshooting – Error Messages

<table>
<thead>
<tr>
<th>Message</th>
<th>Cause</th>
<th>Remedy</th>
<th>Duration</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Humidifier inoperable</strong></td>
<td>Measured value (closed-loop humidity control) or set value (open-loop humidity control) flashes on the screen. Red alarm LED on the control panel flashes. Central alarm indicator lights up.* Intermittent audible alarm is sounded (5x).</td>
<td>Water heater is defective. Switch off humidity module. Call DrägerService.</td>
<td>1 min.</td>
</tr>
<tr>
<td><strong>Humidity deviation above 10%</strong></td>
<td>Yellow LED lights up on the control unit.</td>
<td>Canopy, front flap or hand ports are open. Sensor defective.</td>
<td>Close canopy, front flap or hand ports. Check water supply connection. Call DrägerService.</td>
</tr>
<tr>
<td><strong>Humidity sensor inoperable</strong></td>
<td>Measured value on screen flashes. Red alarm LED on the control panel flashes. Central alarm indicator lights up.* Intermittent audible alarm is sounded (5x).</td>
<td>Humidity sensor is defective</td>
<td>Switch off humidity control. Call DrägerService.</td>
</tr>
<tr>
<td><strong>Key pad locked</strong></td>
<td>Intermittent audible alarm is sounded (3x) when a button is pressed. Button functions (set value input/weighing/menu button) disabled</td>
<td></td>
<td>Activate push button functions (page 77).</td>
</tr>
<tr>
<td><strong>Oxygen concentration below 18%</strong></td>
<td>O2 concentration less than 18 Vol.%</td>
<td></td>
<td>Check that the correct gas is connected. Call DrägerService.</td>
</tr>
<tr>
<td><strong>Oxygen deviation above 5%</strong></td>
<td>Measured value flashes on screen. Red alarm LED on the control panel flashes. Central alarm indicator lights up.* Intermittent audible alarm is sounded (5x).</td>
<td>Difference in measured O2 concentration is greater than 3 or 5%.</td>
<td>Close the canopy, front flap and hand ports. Check O2 connection. Check O2 supply via central medical gas pipeline or O2 cylinder. Check configuration (page 71). Call DrägerService.</td>
</tr>
<tr>
<td><strong>Oxygen module inoperable</strong></td>
<td>Display flashes on screen. Red alarm LED on the control panel flashes. Central alarm indicator lights up.* Intermittent audible alarm is sounded (5x).</td>
<td>Controller of O2 module is defective</td>
<td>Switch off oxygen module. Call DrägerService</td>
</tr>
<tr>
<td><strong>Oxygen sensor deviation above 3%</strong></td>
<td>Measured value flashes on screen. Red alarm LED on the control panel flashes. Central alarm indicator lights up.* Intermittent audible alarm is sounded (5x).</td>
<td>Sensor 1 or sensor 2 is defective.</td>
<td>Switch off oxygen module. Call DrägerService</td>
</tr>
</tbody>
</table>

* The central alarm light can be switched off. See “Setting system parameters” on page 73.
| Message                                      | Cause                                          | Remedy                                  | Duration | audible
|----------------------------------------------|-----------------------------------------------|-----------------------------------------|----------| alarm muted |
| Oxygen sensor 1 inoperable                   | Sensor for oxygen measurement is defective.   | Switch off oxygen module. Call DrägerService | 1 min.   |              |
| Measured value flashes on screen.            | Sensor 1 is defective.                        | Call DrägerService                      |          |              |
| Red alarm LED on the control panel flashes.  |                                               |                                         |          |              |
| Central alarm indicator lights up.*          |                                               |                                         |          |              |
| Intermittent audible alarm is sounded (5x)   |                                               |                                         |          |              |
| Oxygen sensor 2 inoperable                   | Sensor for oxygen measurement is defective.   | Switch off oxygen module. Call DrägerService | 1 min.   |              |
| Measured value flashes on screen.            | Sensor 2 is defective.                        | Call DrägerService                      |          |              |
| Red alarm LED on the control panel flashes.  |                                               |                                         |          |              |
| Central alarm indicator lights up.*          |                                               |                                         |          |              |
| Intermittent audible alarm is sounded (5x)   |                                               |                                         |          |              |
| Skin 1 less than 0.5 °C above skin 2         | Kangaroo mode alarm: temperature difference   | Check patient's heat exchange.          | 15 min.  |              |
| (see "Configuration" on page 71.)            | between the skin temperature (yellow) and     | Check configuration (page 71).           |          |              |
| The measured values for skin temperatures    | peripheral temperature (white) too low        |                                         |          |              |
| flash on-screen.                             |                                               |                                         |          |              |
| Yellow alarm LED on the control panel        |                                               |                                         |          |              |
| flashes.                                     |                                               |                                         |          |              |
| Central alarm indicator lights up.*          |                                               |                                         |          |              |
| Intermittent audible alarm is sounded (3x).  |                                               |                                         |          |              |
| Skin 1 more than 4.0 °C above skin 2         | Kangaroo mode alarm: temperature difference   | Check patient's heat exchange.          | 15 min.  |              |
| (see "Configuration" on page 71.)            | between the skin temperature (yellow) and     | Check configuration (page 71).           |          |              |
| The measured values for skin temperatures    | peripheral temperature (white) too high       |                                         |          |              |
| flash on-screen.                             |                                               |                                         |          |              |
| Yellow alarm LED on the control panel        |                                               |                                         |          |              |
| flashes.                                     |                                               |                                         |          |              |
| Central alarm indicator lights up.*          |                                               |                                         |          |              |
| Intermittent audible alarm is sounded (3x).  |                                               |                                         |          |              |
| Skin 1 sensor fault                          | Sensor for core temperature is defective.     | Replace sensor                          | 5 min.   |              |
| Measured value flashes on screen.            |                                               |                                         |          |              |
| Red alarm LED on the control panel flashes.  |                                               |                                         |          |              |
| Central alarm indicator lights up.*          |                                               |                                         |          |              |
| Intermittent audible alarm is sounded (5x).  |                                               |                                         |          |              |
| Skin 1 temp. deviation above 0.5 °C          | The specified limit value for deviation       | Check that the sensor is correctly      | 5 min.   |              |
| (can be set between 0.3 and 1.0 °C –        | from the set value has been exceeded.         | attached to the patient.                |          |              |
| see "Configuration" on page 71.)             |                                               | Close the canopy, front flap or hand    |          |              |
| Measured value flashes on screen.            |                                               | ports.                                  |          |              |
| Yellow alarm LED on the control panel        |                                               | Switch off external heat sources.       |          |              |
| flashes.                                     |                                               | Remove double walls.                    |          |              |
| Central alarm indicator lights up.*          |                                               | Change configuration (page 71).          |          |              |
| Intermittent audible alarm is sounded (3x).  |                                               |                                         |          |              |
| Skin 1 temperature above 39 °C               | Peripheral temperature too high.              | Check that sensor is correctly          | 2 min.   |              |
| Measured value on screen flashes.            |                                               | attached to the patient.                |          |              |
| Red alarm LED on the control panel           |                                               | Switch off external heat sources.       |          |              |
| flashes.                                     |                                               | Check whether double walls can be       |          |              |
| Central alarm indicator lights up.*          |                                               | removed.                               |          |              |
| Intermittent audible alarm is sounded (5x).  |                                               |                                         |          |              |

* The central alarm light can be switched off. See "Setting system parameters" on page 73.
<table>
<thead>
<tr>
<th>Message</th>
<th>Cause</th>
<th>Remedy</th>
<th>Duration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Skin 1 temperature below 36.0 °C</td>
<td>Kangaroo mode alarm: Skin temperature is falling below the alarm limit.</td>
<td>Increase heat supply to the patient. Check configuration.</td>
<td>15 min.</td>
</tr>
<tr>
<td>(See “Configuration” on page 71)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Measured value on screen flashes.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yellow alarm LED on the control panel flashes.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Central alarm indicator lights up.*</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intermittent audible alarm is sounded (3x).</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Skin 2 sensor fault</td>
<td>Sensor for peripheral temperature measurement is defective.</td>
<td>Replace sensor</td>
<td>5 min.</td>
</tr>
<tr>
<td>Measured value flashes on screen.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Red alarm LED on the control panel flashes.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Central alarm indicator lights up.*</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intermittent audible alarm is sounded (5x).</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Skin 2 temperature above 39 °C</td>
<td>Peripheral temperature too high</td>
<td>Check that sensor is correctly attached to the patient. Switch off external heat sources. Check whether double walls can be removed.</td>
<td>2 min.</td>
</tr>
<tr>
<td>Measured value on screen flashes.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Red alarm LED on the control panel flashes.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Central alarm indicator lights up.*</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intermittent audible alarm is sounded (5x).</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Skin 2 temperature below 34.0 °C</td>
<td>Kangaroo mode alarm: Peripheral temperature is falling below the alarm limit.</td>
<td>Increase heat supply to the patient. Check configuration.</td>
<td>15 min.</td>
</tr>
<tr>
<td>(see “Configuration” on page 71.)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Measured value on screen flashes.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yellow alarm LED on the control panel flashes.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Central alarm indicator lights up.*</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intermittent audible alarm is sounded (3x).</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Water empty, please refill</td>
<td>Water container is empty</td>
<td>Refill water container. Replace infusion bottle.</td>
<td>15 min.</td>
</tr>
<tr>
<td>Measured value flashes on screen.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yellow alarm LED on the control panel flashes.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Central alarm indicator lights up.*</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intermittent audible alarm is sounded (3x).</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wrong oxygen sensor 1</td>
<td>Wrong sensor in O2 module. No measurement can be performed after replacing the oxygen sensor.</td>
<td>Call DrägerService. Switch off O2 control.</td>
<td>1 min.</td>
</tr>
<tr>
<td>Message on screen flashes.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Red alarm LED on the control panel flashes.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Central alarm indicator lights up.*</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intermittent audible alarm is sounded (5x).</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wrong oxygen sensor 2</td>
<td>Wrong sensor in O2 module. No measurement can be performed after replacing the oxygen sensor.</td>
<td>Call DrägerService. Switch off O2 control.</td>
<td>1 min.</td>
</tr>
<tr>
<td>Message on screen flashes.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Red alarm LED on the control panel flashes.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Central alarm indicator lights up.*</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intermittent audible alarm is sounded (5x).</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* The central alarm light can be switched off. See “Setting system parameters” on page 73.
Troubleshooting – Faults

All faults are listed in the table below in alphabetical order. See also "Alarm description" on page 121.

<table>
<thead>
<tr>
<th>Fault</th>
<th>Cause</th>
<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Height adjustment does not switch off.</td>
<td>Switch defective or loose contact. Motor defective.</td>
<td>Call DrägerService.</td>
</tr>
<tr>
<td>Red alarm LED on control panel is lit. Continuous audible alarm tone. Audible alarm cannot be muted.</td>
<td>Serious device fault.</td>
<td>Call DrägerService.</td>
</tr>
<tr>
<td>Red LED at symbol flashing, intermittent alarm tone is sounded.</td>
<td>Power failure alarm.</td>
<td>Check power supply. — Connect power supply to mains. Call DrägerService.</td>
</tr>
<tr>
<td>Short beep signal (3x) is sounded.</td>
<td>While setting a set value, the rotary knob is not pressed within 20 seconds.</td>
<td>Press rotary knob or cancel input.</td>
</tr>
<tr>
<td>The yellow LED in the alarm button is lit.</td>
<td>Alarm muting is active.</td>
<td>Deactivate alarm muting.</td>
</tr>
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</table>

If the machine does not operate as expected please call DrägerService.
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Environmental conditions ................................................ 104
Operating data ................................................................. 104
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Measurement and control parameters ................................. 105
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Technical Data

Environmental conditions

In normal operation
- Temperature: 20 °C to 35 °C
- Atmospheric pressure: 600 hPa to 1060 hPa
- Relative humidity: 10 to 95 %, no dew formation
  
  Applies to hospitals at altitudes of 0 to 4000 m above sea-level

During storage/transport
- Temperature: –20 °C to 60 °C
- Atmospheric pressure: 210 hPa to 1060 hPa
- Relative humidity: 10 to 95 %, no dew formation
  
  Applies to transport by air (10,000 m altitude)

Operating data

Electrical power supply
- 100 V / 120 V / 127 V / 230 V to 240 V AC
- (please specify in your order) 50 Hz / 60 Hz
- Max. power consumption at 100 V: 10 A*
- Max. power consumption at 120 V: 8.7 A*
- Max. power consumption at 127 V: 9.1 A*
- Max. power consumption at 230 V: 5.5 A*
- Max. power consumption at 240 V: 5.6 A*
- Leakage current 100 – 127 V / 50 Hz / 60 Hz: 150 µA
- Leakage current 230 – 240 V / 50 Hz: 250 µA

Heater power
- Air heater: 500 W
- Water heater: 140 W

Built-in socket strip
- Max. overall current consumption of all sockets at all power supply voltages: 2 A

⚠️ Max. leakage current:
By connecting devices to the socket strip the overall leakage current may be increased to an unacceptably high level. Where applicable, national limits must be observed. The operator is responsible for adhering to the specified maximum overall leakage current.

Europe (IEC / EN 60601-1):
- Permissible overall leakage current: 500 µA
- Max. leakage current of the socket strip: 250 µA

USA (UL 2601-1):
- Permissible overall leakage current: 300 µA
- Max. leakage current of the socket strip: 150 µA

Height adjustment and tilting
- Duty cycle: 10 %
- Shut-off mode: 6 minutes ON, 54 minutes pause

* This value takes into account the current consumption of the built-in socket strip.
Performance characteristics

Warm-up time
20 minutes from 20 °C to 31 °C (at 20 °C ambient temperature)

Increase in O₂ concentration from 21 to 60 Vol.%
<10 min

Humidification
Heating of sterilised Aqua dest. or demineralised water

Air velocity over the bed
<8 cm / second

Fresh air supply
30 L/min

CO₂ flush, conforming to IEC / EN 60 601-2-19 / 105.1
Max. available CO₂ concentration in the incubator <0.5 Vol.%

Bed tilting
infinitely variable up to 15° tilt angle on both sides.

Operating noise inside the canopy
≤50 dB(A) at 50 Hz
≤50 dB(A) at 60 Hz

Particle filter
Particle class P 2 conforming to EN 149 FFP2 pass volume 2 %

Measurement and control parameters

The specified values depend on the environmental conditions.

Air temperature control

- Measuring principle: NTC, 2 x
- Measuring range: 13 °C to 42 °C
- Measurement uncertainty: ±0.8 °C
- Set value range: 20 °C to 39 °C in 0.1 °C increments*
  <28 °C and >37 °C (extendable with confirmation)

Skin temperature control

- Measuring principle: NTC
- Measuring range: 13 °C to 43 °C
- Measurement uncertainty: ±0.3 °C
- Set value range: 34 °C to 38 °C in 0.1 °C increments
  >37 °C (extendable with confirmation)

O₂ control

- Measuring principle: Electrochemical sensor (capillary)
- Measuring range: 18 Vol.% to 99 Vol.%
- Measurement uncertainty: ±3 Vol. %
- Influencing factors: Atmospheric humidity <1.5 %
- Set value range: 21 Vol.% to 75 Vol.% in 1 Vol.% increments
  >40 Vol. % (extendable with confirmation)
  In rare cases the maximum actual value attained can be less than 75 vol.% but never less than 65 vol.%. The set value must be at least 3 °C above the ambient temperature. At lower ambient temperatures, the increased heat loss may prevent high set values (39 °C) being fully attained. Use the double wall.
### Relative humidity control
- **Measuring principle**: Capacitive
- **Measuring range**: 10 % r.h. to 99 % r.h.
- **Measurement uncertainty**: ±10 %
- **Set value range**: 30 % r.h. to 99 % r.h. in increments of 1 %

### Weighing scale
- **Measuring range**: 0 kg to 10 kg
- **Measurement uncertainty**: ±2 g (0 kg to 2 kg), ±5 g (2 kg to 10 kg)
- **Resolution**: 1 g

### BabyLink interface (optional)
- 2 serial interfaces for the output of incubator status data (actual values, set values and alarms).
- All signals are electrically isolated from the patient.
- Dielectric strength 1500 V

### Central alarm
- **Output for connection to in-house P.A. systems (nurse call)**
  - **Operating voltage**: max. 24 V
  - **Current**: max. 250 mA
  - **Power**: max. 3 W
  - **Potential-free changeover contact**

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* The maximum attainable humidity depends on the air temperature and relative humidity. At high air temperature or low relative humidity, the humidity attainable in the Caleo® is reduced.
**Dimensions**

Unit (Width x Depth) 1090 mm x 680 mm
Height of overall unit with variable pillar 1220 mm to 1520 mm
Height of overall unit with fixed pillar 1270 mm / 1370 mm / 1470 mm
Height of mattress surface (variable height) 800 mm to 1100 mm
Height of mattress surface (fixed height) Optionally 850 mm, 950 mm or 1050 mm
Bed (width x depth) 645 mm x 500 mm

**Weight**

Overall weight 137 kg (basic equipment)
Overall load capacity 60 kg
Standard rail each 10 kg
Tube extensions each 14 kg
Monitor support/drawer/basic tube overall 32 kg

**Overview of max. loads**

- max. 5 kg
- max. 3 kg
- max. 3 kg
- max. 20 kg
- max. 5 kg
- max. 5 kg
- max. 3 kg
- max. 2 kg

**Classification**

according to EC Directive 93/42/EEC Appendix IX Class IIb
UMDNS Code 12-113
Universal Medical Device Nomenclature System

Standards
- Device conforms to IEC / EN 60601-1, IEC / EN 60601-2-19
- Enclosure protection class I
- IEC / EN 60601-2-19 (36.202.2.1) 10 V/m
- Electromagnetic compatibility
- Skin temperature sensor Type BF
# Description

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Description

Operating principle
Caleo® is an incubator for premature children and sick neonates up to a body weight of 5 kg and a body length of 55 cm. Inside the Caleo® patient "capsule", the patients are supplied with a controlled amount of heat and, if necessary, humidity* and oxygen*. The user/operator can adapt the incubator climate to suit the needs of the patient, by adjusting the air temperature, relative humidity* and oxygen content*. The patient capsule acts as a specially protected zone for the patient. The ambient air is therefore filtered before it enters the interior.

Accessibility
Caleo® provides excellent accessibility to the patient for all normal and intensive care requirements: for this purpose, the four hand ports (two on each longitudinal side) are designed with especially large dimensions (Caleo® JumboPorts™). The two longitudinal sides can be completely folded down. In addition, two smaller side flaps at the head and foot ends of the patient can be folded down. If necessary, the canopy cover can also be propped up from two different sides or can be fully removed in an emergency – in order to provide free accessibility to the patient from above.

A total of ten generously dimensioned hose through-hole sleeves ensure clear organisation and routing of hoses and cables through the incubator. Each corner has two such sleeves, which can be easily removed, especially during "Kangarooing" (where the patient rests in direct contact on the mother's or father's chest), to ensure clearly manageable cable and hose routing even when the patient is outside the patient capsule. Two other large tubing ports are located in the side flaps at the head and foot end. In addition, the canopy cover contains an aperture, e.g. for feeding the patient.

Bed and mattress
The bed is especially large in order to provide sufficient space for the patient under a variety of conditions – such as hose installation, the use of supports and storage aids, the treatment of twins in a single incubator and turning over the patient, e.g. for reintubation in the incubator. The patient lies on a well-insulated foam mattress that is surrounded by foil for easier cleaning and has extra-efficient thermal insulation. To prevent decubitus even more effectively, an extra-soft mattress can be used as alternative (SoftBed™ Caleo®**). The bed can be pulled out when the front flap is open. It can be electrically tilted in order to obtain either a head-up position (Trendelenburg position) or head-down positon (±15°).

* optional equipment feature
** see Caleo® Accessories range.
**Twins in the Caleo® (Caleo® Twincubator™)**

Twins can be placed together in Caleo® if there are no medical objections and if their total combined weight does not exceed 5 kg. When treating twins in an incubator, Caleo® must be operated in air control mode.

The treatment of twins together in a single incubator can help to prevent post-natal separation trauma. Direct skin contact between the twins can have positive effects on the development of the patient, as is the case with "Kangaroo" mode (see page 116). If necessary, the incubator air temperature may have to be reduced, because the patients mutually warm each other by direct contact and would therefore risk overheating.

During operation in air temperature mode, the skin temperature of the first patient can be monitored with the yellow temperature sensor, whilst the second temperature channel (white probe) can be used to monitor the skin temperature of the second patient.

Possible dangers from the treatment of twins in a single incubator are derived, e.g. in the case of infections, from the risk of contagion or confusion between the two patients when administering medicines or foods. Also, if the twins require different ambient temperatures or an ambient air with different oxygen or humidity saturation, they should be placed in two separate incubators.

**X-rays**

With the X-ray drawer, which is accessible from outside, the patient can be X-rayed in the incubator without having to be removed or lifted. The X-ray drawer can be pulled out without having to open the Caleo® front flap. Unnecessary disturbance of the patient is therefore avoided. A grid is provided on the X-ray drawer to help align the X-ray cassette.
Weighing scale*

With the optionally fully integrated incubator weighing scale, the weight of the patient can be determined without having to remove the patient from the protective climate of the incubator. Even with the built-in scales, the use of the X-ray drawer remains possible. To weigh, the patient must be lifted once in order to reset the scales to a zero point. The patient can then be placed back on the mattress. The current and previous weight are displayed for information. The results of the last 30 weighs can be graphically represented in the trend display.

Weighing without lifting the patient again (i.e. without new tare) is possible. This option is useful when the weight has to be taken again for checking purposes shortly after the first weigh, or if e.g. the weight with full and empty nappies has to be determined. The entire weighing procedure is accompanied by short audible signals, so that the operator's full attention can be paid to the patient.

When weighing the patient inside an incubator, take care to ensure that no hoses or cables cramp the bed or can distort the weight measurement. If possible, all hoses or cables should be lifted together with the patient. When the patient is placed back on the mattress, the hoses and cables should still be held slightly raised until the new weight appears on the screen (to ensure that the weight of the hoses and cables is not also weighed during the weighing procedure).

The weighing scale in the Caleo® comprises four weighing elements located underneath the bed, an electronic measuring and analysis unit and a special page on the control monitor. In normal mode, the entire bed rests on these four weighing elements under the bed. A safety system prevents the weighing elements being damaged if loads of more than 10 kg are applied. When removing the bed, it should be raised slightly by turning two fixing knobs so that it can then be pulled out smoothly.

* optional equipment feature
Airflow Routing

The heated and humidified air flows into the canopy from both sides. It is guided up the inside of the front flap, along the canopy cover and then down the two transverse sides by suction. The air from the interior is mixed with fresh ambient air by an air filter and is circulated by a fan impeller. Along this path, the air is channelled past an electrically powered heater and is humidified if necessary.

The patient lies in a calm zone with low airflow speeds. Heat loss due to flow is kept to a minimum. On opening the large flaps or hand ports (Caleo® JumboPorts™), an efficient hot air curtain is formed to prevent cooling of the patient capsule.

Air temperature control

The user sets the desired air temperature in the patient capsule by means of the control panel (set value for air temperature in air temperature mode).

The current air temperature is measured by the air temperature sensor in the patient capsule (at the patient's head end of the incubator) and is then compared to the set value. If the set value is greater than the actual measured air temperature, the heater receives the signal to apply more heat. The air temperature inside Caleo® therefore increases.

If the set value is lower than the actual measured air temperature, the heater receives the signal to apply less heat. The air temperature inside Caleo® drops.

If the current air temperature deviates from the set value by more than ±1.5 °C an alarm is triggered.

The audible signal of this alarm can be muted by the user. As soon as the deviation in the measured air temperature is within ±1.5 °C of the set value (see above), the alarm is cancelled.

* optional equipment feature
** other configurations possible
Caleo® temperature control characteristics
The desired temperature increase is achieved rapidly due to the high heating power. A temperature reduction takes longer, due to the good thermal insulation of the incubator.

Note on setting the air temperature set value in the Caleo®:
The patient has limited
— heat loss due to airflow, because the airflow rate above the mattress is low.
— heat loss through the mattress, because the foam mattress is well insulated.
— heat loss due to evaporation, provided that humidity* is set relatively high (greater than 60%).
— heat loss due to radiation, provided that the double wall** is installed.

Skin temperature measurement
Two skin temperature sensors can be connected to measure the skin temperature (yellow skin temperature sensor) and peripheral temperature (white skin temperature sensor). The measured value of the yellow skin temperature sensor is used to regulate the incubator heater in "skin temperature control mode".

ThermoMonitoring™
The term ThermoMonitoring™ refers to the continuous measurement and display of a central temperature and a peripheral temperature. Instead of the central temperature, a near-core skin temperature can be used, as is the case e.g. during skin temperature control at the incubator.

The continuous display of the difference between these two temperatures permits early detection of the occurrence of cold stress. Heat stress, thermoregulation problems and e.g. infections can also be more rapidly detected by displaying the two temperature values and evaluating their difference.

Consequently, Caleo® provides the possibility of switching between a standard screen with large numeric digits and a trend screen showing the trend graph of a maximum of two temperatures. In this way, the difference between near-core skin temperature and peripheral temperature required by ThermoMonitoring™ can be displayed continuously.

In addition, by means of trend analysis, values from the past can be called up for subsequent explanation, e.g. the onset of sickness symptoms or the development of hypothermal stress. Values going back a maximum of seven days in the past can be accessed. In addition, the desired time window can be set (between 3 hours and seven days).

* optional equipment feature
** see Caleo® Accessories range
With the ‘Trend Main Page’ option selected from the main menu, the user can preconfigure which trend to display when switching over between numerical display and trend display.

In addition to displaying the two skin temperatures, the display can show the air temperature, humidity*, oxygen* and the weight’ for up to 30 weighs.

**Skin temperature control**

Caleo® is operated in skin temperature mode, this mode can be set on the control unit. At least the yellow skin temperature sensor (skin 1) must be plugged in and correctly attached to the patient. The user sets the set value for the skin temperature on the control panel. The actual skin temperature of the patient is measured by the yellow skin temperature sensor (skin 1) and compared with the set value.

The difference between the set value and the measured actual value is used to control the air temperature in Caleo® between a minimum of 20 °C and a maximum of 39 °C.

If the set value is greater than the currently measured skin temperature (skin too cold), the heater receives a signal to supply more heat. The air temperature in Caleo® rises, thereby also increasing the skin temperature of the patient.

If the set value is lower than the actual measured air temperature (skin too hot), the heater receives a signal to apply less heat. The air temperature inside Caleo® drops, thereby also reducing the skin temperature of the patient.

The longer the deviation between the set value and the actual measured value persists, the more powerfully heat is supplied by the heater (if the skin is too cold) or the more the air temperature in Caleo® is reduced (if the skin is too hot).

**Waiting for the controller to settle.**

The patient’s skin temperature varies frequently, e.g. due to food intake or medical care. Deviations of a few tenths of a degree are normal. Therefore:

Only change the set value of the skin temperature if the core temperature needs to be changed.

If the actual temperature deviates from the set skin temperature by more than ±0.5 °C**, an alarm is sounded. This audible alarm can be muted by the user. As soon as the measured value deviates from the set value by less than ±0.5 °C** (see above), the alarm is deactivated again.

---

* optional equipment feature
** other configurations possible
Kangaroo Mode

Kangaroo mode (Caleo® KangarooMode™) simplifies the operation of the incubator when the baby is removed to have direct skin contact with the mother or father ("Kangarooing"). This mode provides the user with extended monitoring functions in order to detect overheating and overcooking of the patient even when the patient is outside the patient capsule.

Other Caleo® features designed to ensure easier removal of the patient from Caleo® for "Kangarooing":
- the removable tubing grommets (sleeves) in the corners of the incubator,
- the facility to lower the patient's bed by up to 80 cm (with the variable "height adjustment" option*)
- the minimal space requirement of the fold-down front flap of the Caleo®.

The incubator is switched over to Kangaroo mode on the control panel after pressing the "Menu" button. Once Kangaroo mode is activated, the following functions are automatically activated:

- **Switchover to "Standby" mode**
  Since the baby is no longer inside Caleo® during "Kangarooing", the baby's skin temperature should no longer be used as a measure for controlling the air temperature inside the incubator. Instead, the incubator should be set up so that after terminating "Kangarooing" and putting the patient back inside Caleo®, the incubator should already be heated to the same temperature and climate as when the baby was taken out of the incubator. Consequently, during "Standby" mode, the following logic is applied:
  If Caleo® was previously operated in skin temperature mode, it is switched over to air temperature mode for the duration of Kangaroo mode. The set value for the air temperature is automatically set to the average air temperature over the last three minutes. The previous set value for the skin temperature is stored in buffer memory.
  If Caleo® was previously operated in air temperature mode, the setting remains unchanged.

- **Automatic alarm muting**
  Since, generally, after switching over to Kangaroo mode the user opens the flap and takes the baby out of the incubator, alarms triggered by opening the flaps are no longer meaningful. Consequently, all alarms that would normally be activated by opening the large flaps are automatically muted for the next 4 minutes. The following alarms are involved:
  - Air temperature deviation
  - Oxygen deviation

* optional equipment feature
- **Activation of special Kangaroo mode alarms**

During Kangarooing, the skin (and core) temperature of the patient is frequently found to rise. However, in some cases, the patient is cooled. Consequently, the core or skin temperature patient must be regularly monitored. In order to perform this monitoring with as little nuisance as possible to mother/father and baby, Caleo® provides the facility for activating special Kangaroo mode alarms during operation in Kangaroo mode. These alarms are set in the configuration. The following alarm limits are set:

- Lower alarm limit for the skin temperature (skin 1; T1) = Skin T1 min
- Lower alarm limit for the peripheral temperature (skin 2; T2) = Skin T2 min
- Lower alarm limit for the difference between T1 and T2 = Delta T min
- Upper alarm limit for the difference between T1 and T2 = Delta T max

These Kangaroo alarms have the following significance:

**Skin T1 min:**
The alarm is triggered as soon as the skin temperature (T1, yellow skin temperature sensor) falls below the set value.

**Skin T2 min:**
The Skin T2 min alarm is triggered as soon as the peripheral temperature (T2, white temperature sensor) falls below the set value.

**ΔT min:**
This alarm is triggered if the difference between Skin T1 and Skin T2 is less than the set value (risk of hyperthermia).
∆T max:
This alarm is triggered if the difference between Skin T1 and Skin T2 is above the set value (risk of hypothermia).

Each of these alarms can either be set or switched in advance to "OFF". However, immediately after switching over to Kangaroo mode, automatic alarm muting is activated for the first four minutes, including for all special Kangaroo alarms, in order to give the patient time to adapt to the new environment.

**O2 enrichment**
When adjusting the oxygen concentration in the patient capsule, the additional oxygen supply is metered by a microprocessor-controlled valve. The oxygen is thereby channelled into the air routing system, so that it is heated and humidified with the air.
Closed-loop humidity control*

Caleo® provides the possibility for hygienic humidification of the incubator air. In this system, water is supplied from a separate water container (water bag or water tank) and heated. With closed-loop humidity control, the desired relative humidity is set, and Caleo® then automatically regulates the evaporator output of the humidifier.

Caleo® can control the humidity in two ways:
- By manual input of the desired relative humidity
- Automatic humidity control (AUTO humidity)

Manual humidity control
A value between 30 % and 99 % relative humidity can be set. The actual humidity concentration is measured by the humidity sensor in the patient capsule (integrated in the sensor module at the patient's head end of the capsule).
If the set value is higher than the actual measured relative humidity (air too dry), the humidifier receives a signal to allow more water vapour into the patient capsule. The relative humidity inside Caleo® therefore rises.
If the set value is lower than the actual measured relative humidity (air too humid), the humidifier receives a signal to allow less water vapour into the patient capsule. The relative humidity inside Caleo® therefore falls.

AUTO humidity
When AUTO mode is set, the set value for the relative humidity is calculated and set automatically by the system as a function of the air temperature (see graph).
This function is based on the observation that small and relatively immature patients require both a higher air temperature and higher relative humidity than larger patients. Consequently, in AUTO mode, the set value for relative humidity is calculated and set as a function of the air temperature. The lower the air temperature setting, the lower the set value for relative humidity.

* optional equipment feature
Cleaning Mode*

Cleaning mode is only available if Caleo® is equipped with humidity control. Cleaning mode (Caleo® CleanSwitch™) simplifies the task of cleaning Caleo® and must be used after ending operation of Caleo® (see page 69), after the water supply has been disconnected and the water supply connection pipe has been removed (see page 87).

Cleaning mode may only be used if Caleo® is empty.

Cleaning mode is activated on the control panel after pressing the "Menu" button and selecting the "Cleaning Mode" option.

In Cleaning mode, the water heater is made to run dry. For this purpose, the humidifier is heated to above 100 °C, so that all remaining water in the humidifier is evaporated. After the residual water has been evaporated, the temperature in the humidifier is maintained at a temperature higher than 100 °C for approximately 10 minutes longer. Then the humidifier is allowed to cool down. When there is no longer any danger of burns for the user, a message is displayed on the screen to inform the user that Cleaning Mode is complete and that the incubator can be dismantled for disinfecting and cleaning (see "After completing cleaning mode" on page 87).

Safety systems

After switching on the incubator, a self-test is performed to test all memory addresses of the microprocessor control system and the fault-free running of the program segments. The functions of the control elements and feedback messages are tested by switching on and off. This test is also performed at ten-minute intervals during operation. In this test, all modules installed in the incubator are tested. Any error message will be displayed even when the defective module is switched off. Any unauthorised operating state will cause Caleo® to switch off the heater or water heater as a safety precaution. An additional temperature sensor in the hot air duct restricts the heat output in cases were the control loop would otherwise operate the heater at full power for a long period. Typical situations include e.g. flaps opened for a long period, high set values (39 °C) at low ambient temperatures (<22 °C) or a partly covered hot air duct. This safety system considerably reduces the risk of burns caused by excessive heating of the surfaces next to the air outlet or by the hot air stream itself.

---

* optional equipment feature
Alarm description

Visual signals on the control panel
1. Red alarm LED
2. Yellow alarm LED
3. Red LED next to the \[\text{\textregistered}\] symbol

Caleo® distinguishes between 4 different alarm priorities:

**Serious device faults**

Device fault:
- Continuous tone that cannot be muted, and
- the red alarm LED lights up.

**Power failure:**
- Intermittent alarm tone that cannot be muted, and
- the red alarm LED next to the \[\text{\textregistered}\] symbol lights up.

**Warning (highest danger level)**
- Alarm tone sequence (5 beeps), which can be muted,
- Red alarm LED flashes,
- Central alarm indicator lights up* and
- Measured value flashes.
For example:
  - Skin 1 temperature above 39 °C

**Caution (medium danger level)**
- Alarm tone sequence (3 beeps), which can be muted,
- Yellow alarm LED flashes,
- Central alarm indicator lights up and
- Measured value flashes.
For example:
  - Air temp. deviation above 1.5 °C

**Advisory (low danger potential)**
- Yellow alarm LED lights up.
For example:
  - Humidity deviation above 10 %

Information messages on the active alarm are displayed on screen.

If another alarm occurs while the alarm tone is muted, the alarm tone will be reactivated. See “Alarm suppression” on page 77.

* The central alarm light can be switched off.
  See “Setting system parameters” on page 73.
Central alarm indicator

1 The central alarm indicator lights up when an alarm occurs.

1 The central alarm indicator does not light up with an alarm if
   — the alarm muting button is pressed, or
   — the central alarm indicator has been deactivated in the
     Configuration. See “Setting system parameters” on
     page 73.

Key to the symbols used

- Alarm muting
- Cancel, stop setting procedure
- Weighing scale
- Bed tilt
- Bed can be pulled out
- Radioscopy, X-ray drawer can be removed
- Caution: please note special information on this
  function in the Instructions for Use.
- Type BF
- Disable keypad
- Waiting for input from rotary control
Labelling

1. Connections for skin temperature sensors, page 42

2. auxiliary power sockets, page 25, Technical Data, page 104
   Type designation for fuses for auxiliary power sockets
   Type designation for system fuses

   Equipotential bonding
   On/off switch, page 26

   Nameplate

3. page 86

4. page 16

5. page 24
1. Never leave baby unattended when doors are open! page 26

2. Do not tilt canopy forwards, page 27

3. WARNING!
   Never block or obstruct air vents! Risk of burning!
   page 9 and page 113

4. Always close X-ray tray tightly! page 21

5. max 5 kg, 11lbs page 107

6. max 7 kg, 14lbs page 107

7. Use max. 6 min. within 60 min.
   Do not place any objects on the base plate, page 22

8. page 23

9. Aquadest page 24

10. Internal battery, page 94

11. Interface connections, page 106

(inside)

No spray disinfection! Wipe disinfection only! page 93

Switch off device and allow heater to cool down for 90 min. before touching the surface! page 35 page 87
**Order list**

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<td><strong>Caleo®</strong></td>
<td></td>
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<tr>
<td>Caleo incubator with air and skin servo modes, ThermoMonitoring™ (temperature probes optional) With large front and rear access panels, 6 extra large hand ports (Jumbo Ports™), detachable hood which opens to either side, electronic bed inclination (±15°); 10 extra large tubing ports, integrated X-ray tray, integrated mains distribution box (4 sockets), large graphic EL display with Dräger rotary knob, trend display and large numerical display, central alarm light.</td>
<td>2M 50 000 / 2M 50 555</td>
</tr>
</tbody>
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**Options**

- Trolley with fixed height  Feature
- Trolley with variable height  Feature
- Standard humidity  Feature
- O₂ control  Feature
- Double wall  Feature
- Bed scales, integrated  Feature
- Drawer  Feature
- Interface  Feature
  (2x RS232, 1x sister call)

**Accessories for O₂ enrichment**

- Oxygen flowmeter, 32 L/min, 5 bar  2M 85 506
- Silicone hose, connecting hose between 12 03 606
- Oxygen flowmeter 2M 85 506 and Caleo® 2M 29 323
- Oxygen connecting hose 3M (black)  M 29 233
- Oxygen connecting hose 5M (black)  M 29 253
- CS hose O₂ DN6, 3/6M w/o connect.  M 30 873
- MiniOx  2M 22 464
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</tr>
<tr>
<td>O2 central supply system hose 5M NIST EN-F</td>
<td>86 02 515</td>
</tr>
<tr>
<td>CS hose O2 3M, DIN-ST</td>
<td>M 34 402</td>
</tr>
<tr>
<td>CS hose O2 5M, DIN-ST</td>
<td>M 34 403</td>
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<tr>
<td>CS hose O2 5M without connector</td>
<td>M 34 416</td>
</tr>
<tr>
<td>Oxygen hose 5M NIST, SW, w/o connect.</td>
<td>M 32 037</td>
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<tr>
<td>Adapter O2 DIN/NIST</td>
<td>M 32 366</td>
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<tr>
<td>Oxygen CS hose 3M DIN BL</td>
<td>M 29 231</td>
</tr>
<tr>
<td>Oxygen CS hose 5M DIN BL</td>
<td>M 29 251</td>
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<td>Basic pole</td>
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<tr>
<td>Infusion support, 38 mm pole</td>
<td>2M 21 514</td>
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<td>Rail-mounted infusion holder</td>
<td>2M 20 719</td>
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<td>Compact rail</td>
<td>2M 85 337</td>
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<td>M 24 678</td>
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<td>Notebook Holder</td>
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<tr>
<td>Caleo® SoftBed™</td>
<td>MX 17 012</td>
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<td>Fixation Set Caleo® (France only)</td>
<td>2M 50 080</td>
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<td>O₂ – Air connection hose, 5M (black)</td>
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<td>Holder for litter bags</td>
<td>M 24 695</td>
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<tr>
<td>Set of 100 litter bags</td>
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<td><strong>Accessories for phototherapy</strong></td>
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<td>Phototherapy 4000 (220-240 V)</td>
<td>2M 21 000</td>
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<tr>
<td>Phototherapy 4000 (110-127 V)</td>
<td>2M 21 700</td>
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<tr>
<td>Phototherapy 4000 (100 V)</td>
<td>2M 22 090</td>
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<tr>
<td>Stand for phototherapy unit 4000</td>
<td>2M 21 190</td>
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<td>Water container set, complete</td>
<td>2M 50 040</td>
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<tr>
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<td>2M 50 042</td>
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<tr>
<td>Nozzle for the water container</td>
<td>2M 50 039</td>
</tr>
<tr>
<td>Transfer set</td>
<td>MX 17 018</td>
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<tr>
<td>O-ring, for water connection pipe</td>
<td>2M 50 346</td>
</tr>
<tr>
<td>Caleo Fresh air filter (20 pcs)</td>
<td>MX 17 015</td>
</tr>
<tr>
<td>ThermoTrace™, core (set of 5), yellow</td>
<td>MX 11 000</td>
</tr>
<tr>
<td>ThermoTrace™, peripheral (set of 5), white</td>
<td>MX 11 001</td>
</tr>
<tr>
<td>ThermoPad™ (set of 50)</td>
<td>MX 11 002</td>
</tr>
<tr>
<td>Oxy-Trace™ Incu.</td>
<td>MX 01 050</td>
</tr>
<tr>
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</tr>
<tr>
<td>Tubing port</td>
<td>2M 50 412</td>
</tr>
<tr>
<td>Feeding grommet, hood</td>
<td>2M 50 352</td>
</tr>
<tr>
<td>Caleo® standard mattress</td>
<td>2M 50 556</td>
</tr>
<tr>
<td>Caleo® SoftBed™</td>
<td>MX 17 012</td>
</tr>
<tr>
<td>Vacuum mattress</td>
<td>2M 17 909</td>
</tr>
<tr>
<td>Lithium batteries, 3V/1400 mAh</td>
<td>18 35 343</td>
</tr>
<tr>
<td>Mattress cover (blue-nose kitten)</td>
<td>2M 21 272</td>
</tr>
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### Parts List

As an alternative to the part numbers listed in the Order List, the following parts and devices, which are no longer supplied by Dräger, may be used.

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<tr>
<td>Servo controlled oxygen</td>
<td>2M 50 740</td>
</tr>
<tr>
<td>Bed scale, integrated</td>
<td>2M 50 745</td>
</tr>
<tr>
<td>Double wall</td>
<td>2M 50 421</td>
</tr>
<tr>
<td>Drawer</td>
<td>2M 50 565</td>
</tr>
<tr>
<td>Interface, RS 232</td>
<td>2M 50 750</td>
</tr>
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</table>

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</tr>
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<td>2M 85 041</td>
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<td>Hose holder</td>
<td>2M 19 630</td>
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<tr>
<td>Stand for phototherapy</td>
<td>2M 18 780</td>
</tr>
<tr>
<td>O2 distributor with cylinder connection</td>
<td>2M 18 828</td>
</tr>
<tr>
<td>Oxydig, O2 measuring and warning device, inc. sensor capsule</td>
<td>83 04 411</td>
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These Instructions for Use apply only to Caleo® with Serial No.:
If no Serial No. has been filled in by Dräger these Instructions for Use are provided for general information only and are not intended for use with any specific machine or device

Directive 93/42/EGG concerning Medical Devices

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