

User Manual

Careclave® 618

Combination steam sterilizer

from software version 21.0.3



EN

Dear customer,

We thank you for your confidence demonstrated by the purchase of this MELAG product. As an owner-run and operated family concern founded in 1951, we have a long history of successful specialization in hygiene products for practice-based use. Our focus on innovation, quality and the highest standards of operational reliability has established MELAG as the world's leading manufacturer in the instrument reprocessing and hygiene field.

You, our customer are justified in your demand for the best products, quality and reliability. Providing **“competence in hygiene”** and **“Quality – made in Germany”**, we guarantee that these demands will be met. Our certified quality management system is subject to close monitoring: one instrument to this end is our annual multi-day audit conducted in accordance with EN ISO 13485. This guarantees that all MELAG products are manufactured and tested in accordance with strict quality criteria.

The MELAG management and team.

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1 General guidelines

Please read this user manual carefully before commissioning the device. The manual includes important safety instructions. Make sure that you always have access to digital or printed version of the user manual.

Please read this user manual carefully before starting to use the accessories. The user manual includes important safety information. Make sure that you always have access to digital or printed version of the user manual.

Should the manual no longer be legible, is damaged or has been lost, you can download a new copy from MELAG download centre at www.melag.com.

Symbols used

Symbol	Explanation
	Indicates a dangerous situation, which if not avoided, could entail slight to life-threatening injuries.
	Indicates a dangerous situation, which if not avoided, could lead to burns.
	Draws your attention to a situation, which if not avoided, could result in damage to the instruments, the practice fittings or the device.
	Draws your attention to important information.

Formatting rules

Example	Explanation
Universal- Program	Words or phrases appearing on the display of the device are marked as display text.
	Prerequisites for the following handling instruction.
	Refer to the glossary or another text section.
	Information for safe handling.

Disposal

MELAG devices are synonymous with high quality and a long life-span. When you eventually need to decommission your MELAG device, we offer a special disposal service. Simply contact your stockist.

Dispose of accessories and consumption media which you no longer require in the appropriate manner. Comply with all relevant disposal specification in terms of possibly contaminated waste.

The packaging protects the device against transport damage. The packaging materials have been selected for their environmentally-friendly disposability and can be recycled. Returning the packaging to the material flow reduces the amount of waste and saves raw materials. Dispose of all non-required packaging materials at the collection points of the dual system.

Dispose of spare parts that are no longer used, e.g. seals, properly.

2 Safety



When operating the device, comply with the following safety instructions as well as those contained in subsequent chapters. Use the device only for the purpose specified in these instructions. Failure to comply with the safety instructions can result in injury and/or damage to the device.

Qualified personnel

- Both instrument reprocessing and the sterilization of instruments and textiles with Careclave may only be performed by qualified personnel.
- The operator must ensure that the users have been trained in the operation and safe handling of the device.
- The operator must ensure that users are regularly trained in the operation and safe handling of the device.

Set-up, installation and commissioning

- Check the device after unpacking for any damage suffered during transport.
- The device should only be setup, installed and commissioned by MELAG authorized persons.
- Have the electrical connection, the connections for supply and wastewater and compressed air set up only by a specialist.
- The device is not suitable for operation in explosive atmospheres.
- Install and operate the device in a frost-free environment.
- The device is conceived for use outside the patient area. The device should be located a minimum of 1.5 m radius away from the treatment area.

Power cable and power plug

- Only the power cable included in the scope of delivery may be connected to the device.
- The power cable may not be replaced by a cable determined to be insufficient.
- Comply with all legal requirements and locally-specified connection conditions.
- Never operate the device if the plug or power cable are damaged.
- The power cable or plug should only be replaced by ►authorized technicians.
- Never damage or alter the power plug or cable.
- Never unplug by pulling on the power cable. Always take a grip on the plug.
- Ensure that the power cable does not become jammed in.
- Never lead the cable along a source of heat.
- Never fix the power cable with sharp objects.
- The mains socket must be freely accessible after installation so that the device can be disconnected from the electrical mains at any time if necessary by pulling the mains plug.

Spring safety valve

- The free movement of the valve rod must be guaranteed. For example, the spring safety valve must not be taped or blocked. Install the device in such a way that the proper functioning of the spring safety valve is guaranteed.

Reprocessing and sterilization

- Follow the manufacturer's instructions of your textile articles and instruments regarding their reprocessing and sterilization.
- Comply with the relevant standards and directives applicable to the reprocessing and sterilization of textiles and instruments (in Germany e.g. from the ►RKI and ►DGSV).

Normal operation

- The door area and the area of the cooler and the safety valves at the rear of the device can become hot while the device is switched on and remain hot for a long time after it is switched off.
- The sterile filter is no longer effective if it has become wet. Stop using the sterile filter and replace it.
- Do not replace the sterile filter during a program run.

Program termination

- Please observe that opening the door following a program abort can lead to hot steam leaving the sterilization chamber.
- Observe the instructions shown on the display of the device. Sterilize the affected ▶equipment after re-wrapping.

Maintenance

- Maintenance should only be performed by ▶authorized technicians.
- Maintain the specified servicing intervals.
- Only original MELAG spare parts may be used.

Repair

- Never open the device housing. Incorrect opening and repair can compromise electrical safety and pose a danger to the user. The device may only be opened by an ▶authorised technician who must be a qualified electrician (according to VDE 0105-100 or IEC 60050).

Malfunctions

- Should the device issue the same malfunction message repeatedly, turn off the device and if necessary, inform your stockist.
- The device may only be serviced by ▶authorized technicians.

Notification requirement in the event of serious accidents in the European Economic Area

- Please note that all serious accidents which occur in connection with the medical product (e.g. death or serious deterioration in the state of health of a patient) which were presumably caused by the product, must be reported to the manufacturer (MELAG) and the relevant authority of the member state, in which the user and/or patient resides.

Personal protective equipment

- The removal of the Carebox should only be carried out with an aid, e.g. a protective glove.
- The sterilized equipment should only be removed using an aid, e.g. protective glove or tray lifter.

3 Performance specifications

Intended use

Careclave is intended for use in the medical field, especially in dentistry. This device is not intended to be used in a patient environment. In accordance with ►EN 13060 this device is a steam sterilizer (autoclave) with type B cycles. As a universal steam sterilizer, it is suited to highly-demanding sterilization tasks. It can be used to sterilize, e.g., narrow lumen instruments – both wrapped or unwrapped – and textiles.

In addition, Careclave can be used to reprocess compatible connectible dental instruments (e.g. transmission instruments) in a container (Carebox) provided for this purpose. The internal and external cleaning as well as the subsequent thermal disinfection comply with the specifications of ►EN ISO 15883-1 and -2. Optionally, sterilization with type S cycles can be carried out instead of thermal disinfection, or automatic care can be carried out with care oil.



WARNING

Any attempt to sterilize liquids can result in a ►delay in boiling. This can result in burns and damage to the device.

- Never use this device to sterilize fluids. It is not licensed for the sterilization of fluids.

User benefits

With the help of Careclave, you can fully comply with the reprocessing recommendations of the German Commission for Hospital Hygiene and Infection Prevention at the ►RKI, as well as the normative requirements from ►EN 13060 and ►EN ISO 15883-1 and -2.

Reprocessing of semi-critically classified dental transmission instruments

The simultaneous reprocessing of a maximum of eight dental transmission instruments takes place exclusively in the Carebox. For successful cleaning and disinfection, the transmission instruments must be connected to the appropriate adapters. Cleaning and disinfection take place both in the inner area of the dental instrument (interior cleaning) and on the outer surfaces (exterior cleaning). During subsequent care with care oil, only the drive areas of the dental transmission instruments are specifically treated. Depending on the program selected, either thermal disinfection is carried out before care or sterilization after care.

Reprocessing of semi-critically classified connectible hollow bodies

The simultaneous reprocessing of a maximum of eight ultrasonic and air scaler tips (e.g. ZEG tips) takes place exclusively in the Carebox. For successful cleaning and disinfection, the tips must be connected to the appropriate adapters. Cleaning and disinfection take place both in the inner area of the tips (internal cleaning) and on the outer surfaces (external cleaning). Depending on the program selected, either thermal disinfection or sterilization is then carried out.

Combined sterilization

In the Care-B program, reprocessing in the Carebox is possible in combination with type B sterilization. This allows you to sterilize instruments or tips in the Carebox as well as critically classified (wrapped) and narrow instruments in the sterilization chamber in one program run.

Program runs for the Carebox

Cleaning & disinfection

Careclave can simultaneously clean and thermally disinfect up to eight dental transmission instruments or ultrasonic and air scaler tips. For successful cleaning and disinfection, the instruments in the Carebox must be connected to the adapters provided.

Cleaning and disinfection takes place both in the inner area of the dental instrument (internal cleaning) and on the external surfaces (external cleaning). Cleaning is done with demineralised water, partly with the support of compressed air pulses and without the use of chemicals.

During the program run, the following steps (phases) are completed:

Carebox detection

Carebox detection (*Carebox-Connect*) determines the type of Carebox used (Blue/Green) and activates (Carebox Blue) or deactivates (Carebox Green) the care for the program run.

Pre-cleaning

Pre-cleaning is done with cold water. The transmission instruments soiled and contaminated by proteins or coarse organic adhesions are cleaned mechanically to avoid denaturation (coagulation) due to excessively high water temperatures. At the end of pre-cleaning, the rinse liquor is drained to remove the proteins and other contaminants from the system.

Intermediate cleaning

In intermediate cleaning, a new rinse liquor is used and further soiling is dissolved. During this process, the temperature of the rinse liquor rises slightly. At the end of intermediate cleaning, the rinse liquor is drained again to further lower the residual contamination.

Final cleaning

In final cleaning, the newly introduced rinse liquor is continuously heated. At a temperature of 55 °C, any remaining organic deposits and soiling are dissolved from the instruments.

Disinfection (Program Care-Therm)

Immediately after final cleaning, ►A0-value controlled thermal disinfection takes place at a temperature above 92 °C. The rinse liquor from the final cleaning is reused for thermal disinfection. The disinfection phase is conceived so as to reach an A0 value of at least 3000. This eliminates vegetative bacteria and fungi or fungal spores and inactivates viruses (incl. HBV, HCV) and achieves the effective range AB according to the specifications of the ►Robert Koch Institute.

The transmission instruments prepared with this method are suitable for use in treatments with the risk classification semi-critical B.

Care (optional)

During care, the care oil is specifically dosed only into the drive areas of the transmission instruments. The dosing system is designed so that each instrument, regardless of its type, receives a sufficient amount of the care oil.

Sterilization

Sterilization S (Program Care-S)

Instead of disinfection in the Carebox, sterilization with type S cycles (according to ►EN 13060) can also be carried out. This is specially designed for the steam penetration of dental transmission instruments. Sterilization is carried out at a temperature of 134 °C and with a sterilization time of 3:30 min. The transmission instruments prepared in this way are suitable for use in treatments with the risk classification semi-critical B.

Sterilization B (Program Care-B)

Instead of sterilization exclusively in the Carebox, sterilization of the Carebox including the entire chamber can also be carried out with type B cycles (according to ►EN 13060). This is additionally designed for the steam penetration of product with narrow lumen (hollow body A), both wrapped and unwrapped. Sterilization takes place at a temperature of 134 °C and with a sterilization time of 5:30 min. The wrapped transmission instruments prepared in the sterilization chamber are suitable for use in treatments with a risk classification of critical B.

Drying

Drying before the end of the program not only serves to intensively dry the inner lumen of the instruments, but also to empty the Carebox. It is carried out both as vacuum drying and in the form of compressed air drying. Apart from conformity with ►EN 13060, drying also contributes to maintaining the value of your instruments. In the Care-B program, drying is also designed for wrapped loads in the sterilization chamber.

Program runs for the sterilization chamber

A reprocessing program runs in three main phases: the air removal and heating up phase, the sterilization phase and the drying phase. After program start, you can follow the program run on the display. It shows the chamber temperature and pressure as well as the time until the end of drying.

Program phases of a standard sterilization program

Program phase	Description
1. Air removal and heating up phase	Air removal The air removal phase comprises the conditioning and the fractionating phase. During conditioning, steam is repeatedly injected into and removed from the ►sterilization chamber. This generates over-pressure and the residual air is removed. Then, during fractionation, the mixture of air and steam is evacuated from the sterilization chamber and steam is injected. This procedure is also called the fractionated vacuum procedure.
	Heating The continued steam injection into the sterilization chamber leads to an increase in pressure and temperature which continues until the program-specific sterilization parameters have been reached.
2. Sterilization phase	Sterilizing If the pressure and temperature correspond to the program-dependent nominal values, the sterilization phase begins. The corresponding process parameters (pressure and temperature) are held at sterilization level.
3. Drying phase	Pressure release The sterilization phase is followed by pressure release from the sterilization chamber.
	Drying The sterile material is dried using a vacuum (vacuum drying).
	Ventilation Upon program end, the sterilization chamber is filled with sterile air via the sterile filter and adjusted to the ambient pressure.

Program phases of the vacuum test

Program phase	Description
1. Evacuation phase	The sterilization chamber is evacuated until the pressure for the vacuum test has been reached.
2. Equilibration time	An equilibration time of five minutes will follow.
3. Measurement time	The measurement time amounts to ten minutes. The pressure increase within the sterilization chamber is measured during the measurement time. The evacuation pressure and the equilibration time or measurement time are shown on the display.
4. Ventilation	The chamber is ventilated after the end of the measuring time.
5. Test end	The display shows the test result, the batch number, the total number of batches and the leakage rate.

Care procedure

Careclave offers the possibility of caring for transmission instruments with the help of an integrated function for distributing and dosing care oil. Furthermore, chuck care can be done manually with the ADDcare integrated care station.

The Carebox Blue is used for the reprocessing of instruments that must be maintained with maintenance oil.

The amount of care oil applied to the transmission instruments is sufficient for the next treatment. In addition, observe the instructions of the instrument manufacturer to prevent damage to the transmission.

See also:

- [Oiling the chucks](#) ► page 38]

Sterilization procedure

The steam sterilizer sterilizes on the basis of the ▶[fractionated vacuum procedure](#). This guarantees the complete and effective wetting/penetration of the sterilization material with saturated steam.

The steam sterilizer uses the double jacket technology to generate the sterilization steam, i.e. the steam sterilizer is fitted with a separate steam generator combined with a double-walled sterilization chamber. After heating, steam is held constantly available in the double jacket. This gives the walls of the sterilization chamber a defined temperature and protects the sterilization chamber from overheating. This especially effective procedure supports the quick ▶[evacuation](#) of the air from the sterilization chamber, the sterilization packaging and instrument cavities. This permits the sterilization of large quantities of instruments or textiles directly one after each other, thereby achieving excellent drying results.

Type of the feed water supply

The steam sterilizer works with a one-way ▶[feed water](#) system. This means that it uses fresh feed water (demineralized or distilled water) for every sterilization procedure. The quality of the feed water is subject to permanent monitoring via integrated ▶[conductivity measurement](#). If combined with careful preparation of the instruments, this serves largely to prevent stain accretion on the instruments and soiling of the steam sterilizer.

Safety equipment

Internal process monitoring

A ▶[process evaluation system](#) is integrated in the electronics of the steam sterilizer. It compares the process parameters (such as temperature, time and pressure) during a program run. It monitors the parameters in terms of their threshold values during control and regulation and guarantees safe and successful sterilization. A monitoring system checks the device components of the steam sterilizer for their functionality and their plausible interaction. If one or more parameters exceeds pre-determined threshold values, the steam sterilizer issues warnings or malfunction messages and if necessary, aborts the program. In the case of a program abort, follow the instructions on the display.

The steam sterilizer uses an electronic parameter control. This enables the steam sterilizer to optimise the total operating time of a program in dependence on the load.

Internal logic monitoring

The electronics of the steam sterilizer monitor the successful program run by means of two separate test processes. When a program has been successfully completed, it is shown on the display as a successful program. In addition, the status LED below the display illuminates green.

Door mechanism

Careclave constantly checks pressure and temperature in the sterilization chamber and prevents the door from being opened when over-pressure has built up in the sterilization chamber. The motor-driven automatic door locking mechanism opens the door slowly by turning the locking spindle. This also holds the door whilst it opens. Pressure equalisation will have been performed by the time that the door is completely open, even following pressure differences.

Automatic monitoring of the feed water

The quantity and quality of the ▶[feed water](#) is automatically checked before every program start.

Overview of sterilization programs

The results in this table show which inspections were performed on the steam sterilizer. The marked fields demonstrate compliance with all the applicable sections of the standard ▶EN 13060.

Type tests	Universal-B	Quick-S	Prion-B	Gentle-B	Care-B	Care-S
Program type in accordance with ▶EN 13060	Type B	Type S	Type B	Type B	Type B	Type S
▶Dynamic pressure test of the sterilization chamber	X	X	X	X	X	X
▶Air leakage	X	X	X	X	X	X
▶Empty chamber test	X	X	X	X	X	X
▶Solid load	X	X	X	X	X	X
▶Porous partial load	X	--	X	X	X	--
▶Porous full load	X	--	X	X	X	--
▶Simple hollow body (▶Hollow body B)	X	--	X	X	X	X
▶Product with narrow lumen (▶Hollow body A)	X	--	X	X	X	--
▶Single wrapping	X	--	X	X	X	--
▶Multiple wrapping	X	--	X	X	X	--
Drying ▶solid load	X	X	X	X	X	--
Drying, porous load	X	--	X	X	X	--
Sterilization temperature	134°C	134°C	134°C	121°C	134°C	134°C
Sterilization pressure	2.1 bar	2.1 bar	2.1 bar	1.1 bar	2.1 bar	2.1 bar
Sterilization time	05:30 min	03:30 min	20:30 min	20:30 min	05:30 min	03:30 min
X = Complies with all applicable sections of the standard ▶EN 13060						

4 Description of the device

Scope of delivery

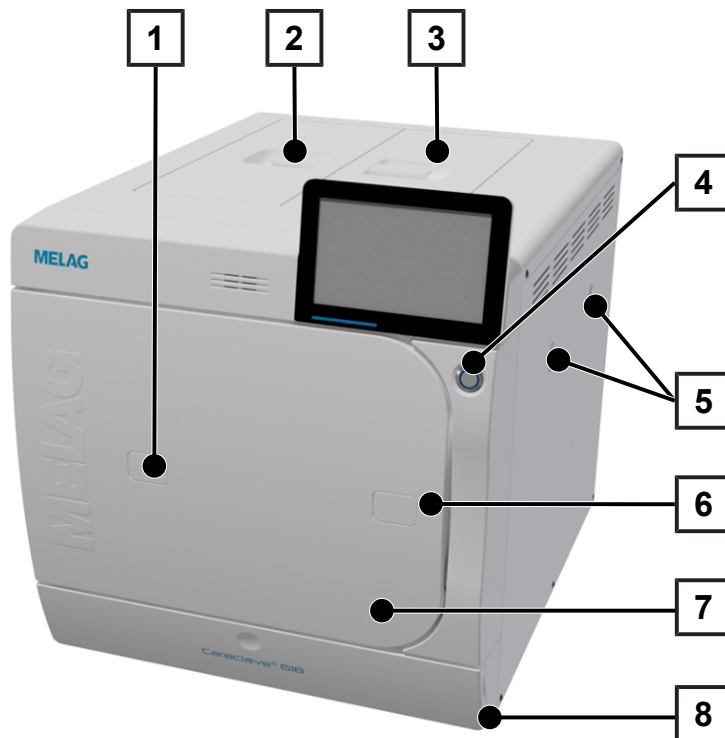
Please check the scope of delivery before setting up and connecting the device.

Standard scope of delivery

- Careclave
- Careclave user manual
- Technical manual
- User manual accessories for Careclave
- Manufacturer's inspection report and declaration of conformity
- Warranty certificate
- Record of installation and setup
- Instruction protocol
- Instrument logbook
- Power cable
- 2x carrying strap
- Installation set consisting of
 - Double-chamber siphon
 - 3x hollow screw G 1/4"
 - 3x SVS-E swivel
 - 6x Cu seal 13.5x18.5
 - 2x Cu seal 13.5x20
 - 5 m PUR hose, black
 - 5 m PTFE plastic pipe
 - 2x double hose nozzle for siphon
 - Cold water adapter MELAdem 40
 - Coupling plug Compressed air
 - 2x fitting
 - 2x SVS-E straight
- 4x cover caps for niches for mounts in the side wall
- Tray lifter
- Heat protection gloves
- Allen key with which to open the door in an emergency
- Screwdriver (TX 6)
- Drain hose
- Oil for door lock bushing
- MELAG USB flash drive
- 2x can of MELAG Care Oil (1x installed and 1x stock)
- Carebox (optional)
- Short/long mount for trays and cassettes (optional)

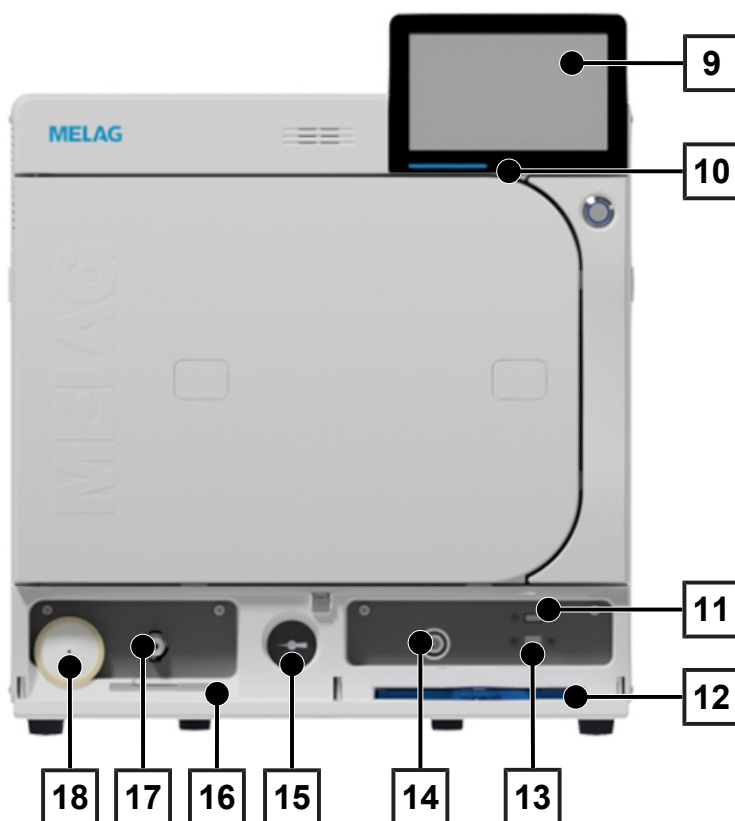
Views of the device

View from front

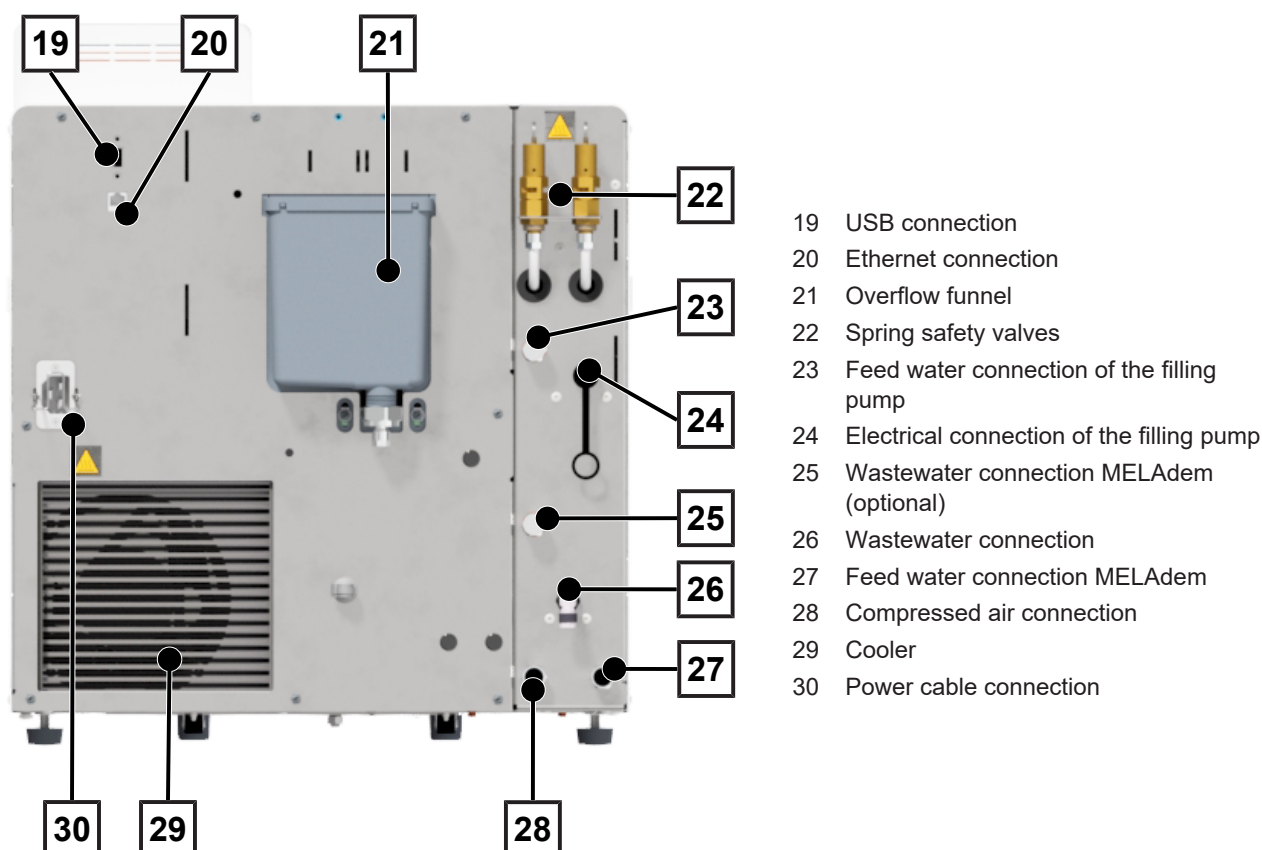


- 1 Access to the validation fitting
- 2 Feed water tank cover
- 3 Accessories compartment cover
- 4 Power switch
- 5 Optional: Mounts for Carebox
- 6 Opening for door opening in an emergency
- 7 Door
- 8 Service hatch

Front view, with open service hatch



- 9 Smart touch display
- 10 LED status bar
- 11 USB connection
- 12 Dust filter
- 13 Service connection
- 14 Overheat protection reset button
- 15 Manometer for pressure display on the double jacket steam generator
- 16 Allen key with which to open the door in an emergency
- 17 Drain valve for emptying the feed water tank
- 18 Sterile filter

Rear view, without cover**Service hatch**

The service hatch is magnetic and can be opened by pulling on any side.

**Symbols on the device****Type plate**

Manufacturer of the medical device



Date of manufacture of the medical device



Identifies a medical device



Medical device serial number from the manufacturer



Article number of the medical device



Information about the chamber volume



Operating temperature of the device



Indicates the permitted temperature range (min./max.) of the water supply



Operating pressure of the device



Flow pressure at the connected compressed air connection from min. to max.



Flow pressure at the connected water inflow from min. to max.



Electrical connection of the device: AC current



The user manual includes important safety information. Failure to comply with these instructions can result in injury and material damage.



Read this user manual carefully before commissioning the device.



In affixing this CE mark, the manufacturer declares that this medical product fulfils the basic requirements of the Medical Device Directive. The four-digit number confirms that this is monitored by an approved certification agency.



In affixing this CE mark, the manufacturer declares that this product fulfils the basic requirements of the Pressure Equipment Directive. The four-digit number confirms that this is monitored by an approved certification agency.



The device may not be disposed as domestic waste. The vendor is responsible for appropriate disposal of the device - it must be delivered to the vendor to be disposed of.
MELAG devices are synonymous for long-term quality. When you eventually need to decommission your MELAG device, we offer a special disposal service. Simply contact your stockist.

Warning symbols



This symbol indicates that the marked area becomes hot during operation. Contact with it during or shortly after operation can pose the danger of burns.

Device symbols - front

Symbol	Description	Symbol	Description
	Sterile filter		Dust filter/device filter
	Drain connection		Overheat protection reset button
	USB connection		Service connection

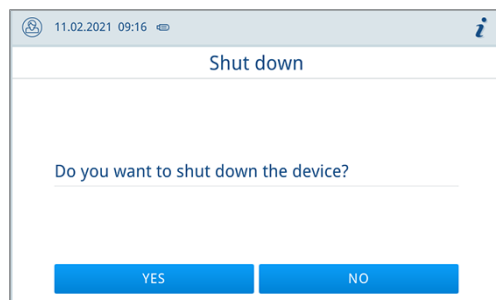
Device symbols - rear

Symbol	Description	Symbol	Description
Aqua dem 	Feed water connection, water treatment unit	Osmosis drain 	Wastewater connection, water treatment unit
Pump aqua dem 	Connection, filling pump	Drain 	Wastewater connection
Air 	Compressed air connection	Pump power 	Electrical connection of the filling pump

Power switch**PLEASE NOTE**

The device cannot be shut down during a running program.

By pressing the power switch, you can call up the shutdown dialogue.



By pressing the power switch again, you switch the device on again.

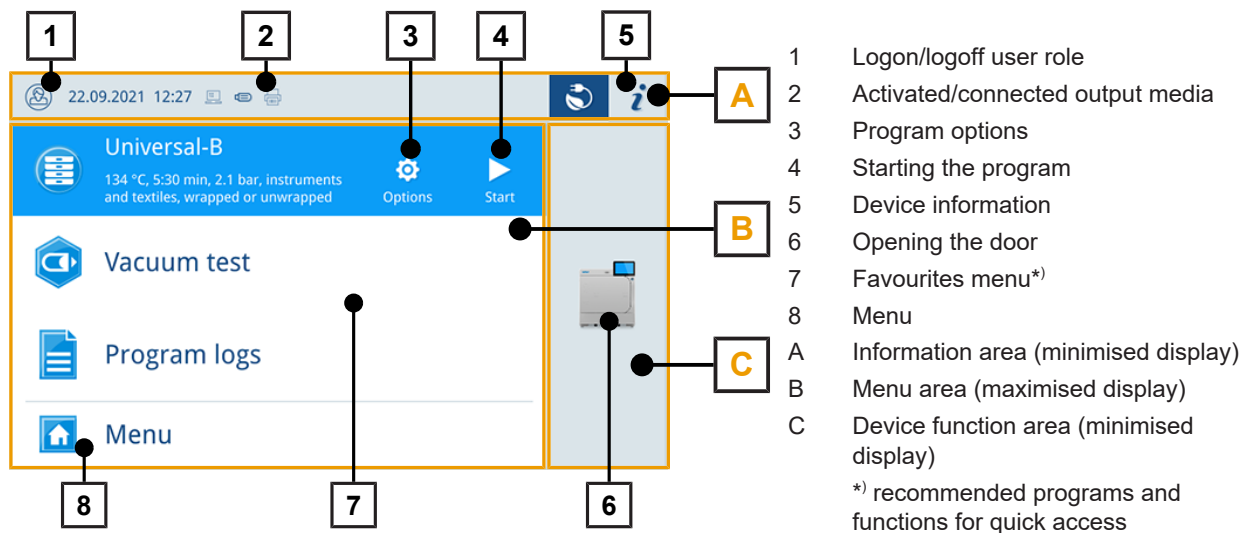
State	Description
illuminated	The device is shut down.
not illuminated	The device is in standby or a program is running.
vibrating	The device is powering up.

Smart touch display

The user interface consists of a colour 7-inch touch display.

The selected menu item is always highlighted in colour.

The display of the areas (A, B, C) is dynamic and can change depending on the device status.



Due to the dynamic display, the display and position of the buttons on the device may differ from the illustrations shown.

Buttons in the program selection

Button	Description
	Starting the program
	Select program options and start program
	Abort program

Buttons in the information area

Buttons	Description
	Show or hide Device status
	Open or close Device status
	Malfunction message present Show or hide Malfunction message
	Warning message present Show or hide warning message
	Energy-saving activated Show or hide energy-saving dialogue

Chuck care buttons

Symbol	Description	Button
	Chuck care is switched off	Switch on chuck care
	Chuck care is switched on	Switch off chuck care
	Chuck care venting is switched on	Switch off chuck care

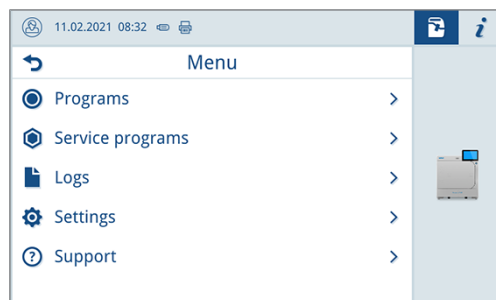
LED status bar

The LED status bar on the lowest edge of the display indicates different situations with various colours.

Colour of the LED	Description
Blue	Device is in operation, no program active Program in progress
Green	Program successfully completed Drying in progress
Red	Malfunction message Program abort in progress Program not successfully completed
Yellow	Warning message

Menu

The Menu gives you access to the programs available in the device mode, to various settings and to the log output.



The **Support** menu item contains the contact details of your service technician and the License information.

Device mode**Careclave mode**

When Careclave mode is activated, the Carebox symbol is displayed in the information area.

Symbol	Meaning
	Careclave mode active

The device can be operated in Careclave mode (with Carebox) or in Vacuclave mode (as an steam sterilizer).

**PLEASE NOTE**

Careclave mode can only be activated by inserting a Carebox.

Door mode

When door mode is activated, the door mode symbol is displayed in the information area.

Symbol	Meaning
	Door mode active

The device can be operated in door mode to switch on the device and open the door. No device processes are activated in the background.

The following functions are available in door mode:

- Door mode
- Change settings
- Subsequent log issue

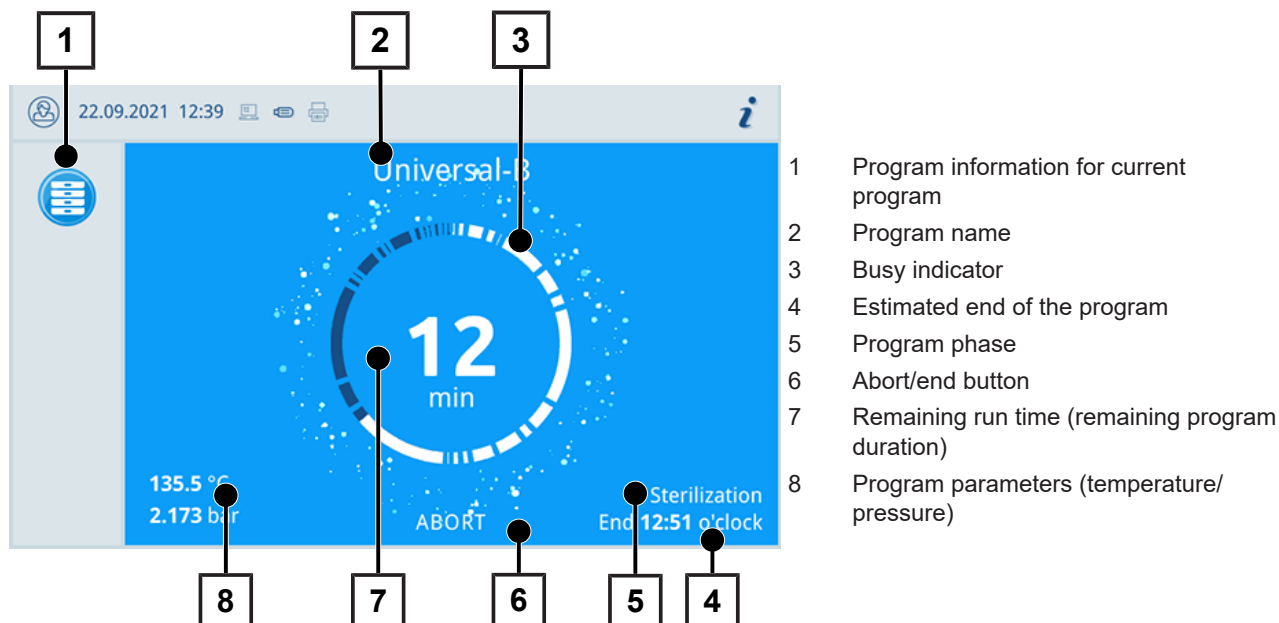
See also:

- [Activating/deactivating door mode](#) [► page 23]

Program run

During a program run, all important information is shown on the display.

If no input is made on the display, the program display maximises and overlays the menu. Touch the display to show the menu.



The display indicates whether the sterilization phase has been completed successfully. The busy indicator and the LED status bar both change from blue to green as soon as the drying phase is initiated.

5 Opening and closing the door

The device is fitted with a motor-driven automatic door locking mechanism with a threaded spindle.

Opening the door



PLEASE NOTE

The door is to be left open only whilst loading and unloading the device. Keeping the door closed saves energy.

Please observe the following when opening the door:

- Never use force to open the door.
- Do not pull vigorously at the door to open it. The door opens automatically.
- 1. Open the door by pressing **OPEN DOOR**.
The button is displayed when the menu area is minimised.
→ The door opens automatically.
- 2. Open the door completely until it snaps into place.

See also:

- [Oiling the door spindle](#) [▶ page 69]

Closing the door

- ▶ To close the door, press it firmly inwards until the automatic door lock engages.



- After the door has been closed, the display returns to the default view. The door is locked pressure-tight upon program start.

When closing the door, comply with the following instructions to guarantee faultless operation of the door locking mechanism:

- Never slam the door.
- Press the door closely to the housing.
- Hold the door closed for min. 3 s until the door lock engages.



PLEASE NOTE

The door can only be closed with the oil can inserted.

For safety reasons, the automatic door lock is deactivated when no oil can is present.

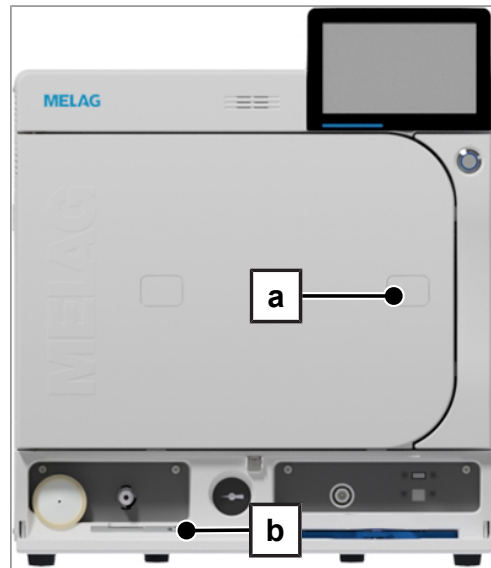
See also:

- [Replacing the oil can](#) [▶ page 68]

Manual door emergency-opening

In emergency situations e.g. power failure, the door can be opened in the following fashion:

1. Switch off the steam sterilizer and remove the power plug from the socket.
2. Remove the cover cap (pos. a) in order to enable the emergency door opening by pressing the cover cap in on one side.



3. Insert the Allen key (5 mm) included in the scope of delivery into the opening. The Allen key can be stored in the specially designed bracket behind the service hatch (pos. b).



CAUTION

When the door is opened in an emergency, hot water vapour may escape and hot water may be present in the sterilization chamber and in the Carebox.

This could result in scalding.

- Never touch the load, the sterilization chamber or the door with unprotected hands. The components are hot.
- The water that may be present can be seen through the transparent lower section of the Carebox.

4. Turn the Allen key to open the door in a clockwise direction.
 - ➔ The door opens a crack.
5. Remove the Allen key.
6. Open the door and return the cover cap.

Activating/deactivating door mode

1. Switch on the device.
2. Touch the bottom two corners of the display during device startup.
 - ➔ The symbol for the door mode appears in the information area.
3. Press the symbol for the door mode to deactivate it.

See also:

- [Device mode](#) [▶ page 20]

6 First steps

Setup and installation



PLEASE NOTE

Comply with the specifications of the technical manual during setup and installation. This contains all building-side requirements.

Record of installation and setup

The record of installation and setup is to be completed by the responsible stockist and a copy sent to MELAG as proof of the correct setup, installation and initial commissioning. This is a constituent part of any guarantee claim.

Feed water supply

Steam sterilization requires the use of [▶distilled](#) or [▶demineralized water](#), known as [▶feed water](#). Annex C of [▶EN 13060](#) specifies the guideline values to be observed.

The steam sterilizer requires approx. 4 L feed water for the first filling of the steam generator.

The supply with [▶feed water](#) is effected either via an external water storage container, which must be filled periodically with water of the corresponding quality, or automatically via a water treatment unit (e.g. MELAdem 53/53C or MELAdem 47).

Compressed air supply

According to [▶EN 13060](#), the device must not be operated without the sterile filter integrated in the compressed air hose.

The requirements for the compressed air are described in the chapter [Technical data](#) [▶](#) page 92].

The device cannot be operated in Careclave mode without a compressed air supply. If the device needs to be operated in Vacuclave mode without compressed air, the **Carebox detection** must be deactivated.

See also:

- [Technical data](#) [▶](#) page 92]
- [Program options](#) [▶](#) page 60]

Adapter installation

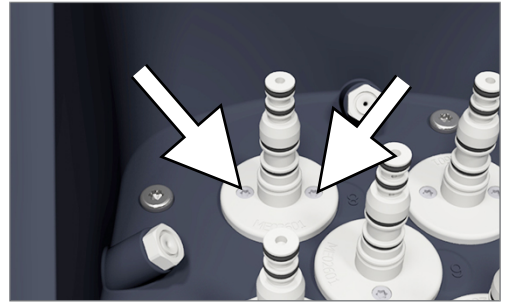
Comply with the following for safe handling:

- The Carebox Blue is used for the reprocessing of instruments that must be maintained with maintenance oil.
- The Carebox Green is used for the reprocessing of instruments that must not be maintained with maintenance oil.
- Use only original accessories.

The following must be fulfilled or present:

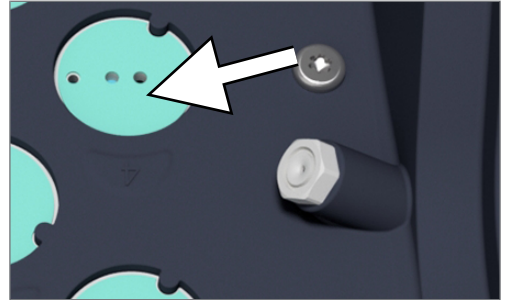
- ✓ The adapters for transmission instruments, hollow-body instruments or unused connections.
- ✓ A new sealing washer, a new O-ring and screws for each adapter.
- ✓ A TX6 screwdriver.

1. Loosen and remove the screws of the existing adapter.

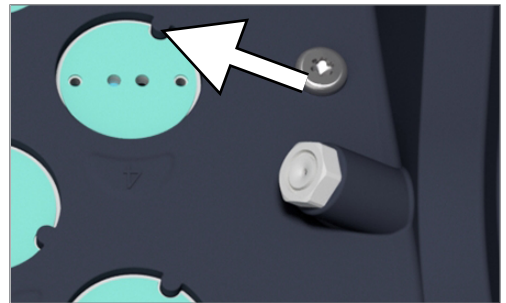


2. Remove the adapter.

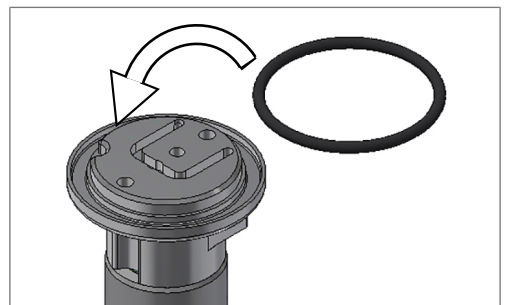
3. Remove the sealing washer and any residue from the seal. Be careful not to scratch the seal face of the Carebox.



4. Insert the new sealing washer by aligning it with the recess and hole pattern.



5. Insert the new O-ring into the adapter.



6. Insert the new adapter by aligning it with the recess.

7. Screw the adapter tight.



8. Perform the **Carebox test**.

**PLEASE NOTE**

Further information on the installation of special adapters, e.g. ME22407, can be found in the user manual "Accessories for Careclave".

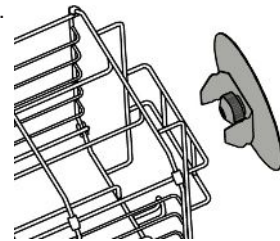
See also:

- [Carebox test](#) [▶ page 56]

Load mounts

Detailed information regarding the various mounts, their combinability with various load holders and their application can be found in the user manual "Accessories for Careclave".

A spring clip is located on the rear panel of the sterilization chamber to fix the mount.



Switching on the device

The following must be fulfilled or present:

- ✓ The device is connected to the power supply.
 - ✓ The feed water supply is secure.
 - ✓ The wastewater disposal is connected.
 - ✓ The compressed air supply is secure.
1. Switch on the device at the power switch.
 - ↳ The double jacket steam generator is ventilated, whereby the air flow can cause a hissing noise.
 2. The welcome screen is displayed. The display then switches to the Favourites menu.
 - ↳ The feed water level is checked and pre-heated immediately after activation.

After device activation, a ▶**heating up time** of a maximum of 15 minutes is required. This time is required for the pre-heating of the double jacket steam generator.

**PLEASE NOTE**

You can start a program immediately without waiting for the heating up time.

See also:

- [Views of the device](#) [▶ page 15]

7 Important information regarding routine operation

Manufacturer's recommendation for the routine operation of type B steam sterilizers ¹⁾

When is it necessary to make checks?	How should the checks be made?
Once per working day	- Visual inspection of the Carebox (sieve, cover and media seals, O-rings) - Visual check of the door seal and the door lock for damage - Check the operating media (electricity, ►feed water and water connection if necessary) - Check the documentation media (printer paper, computer, network) We recommend performing the steam penetration test with MELAcontrol / MELAcontrol Pro in the Universal-Program (test system in accordance with ►EN 867-5).
Once a week	- Vacuum test Tip: In the mornings before starting work – the steam sterilizer must be cold and dry. - Carebox test (with all Careboxes)
Batch-related tests	With "Critical B" instruments: - MELAcontrol / MELAcontrol Pro must be used as ►batch control with every sterilization cycle. With "Critical A" instruments: - The process indicator (type 5 in accordance with ►EN ISO 11140) must be used as batch control with every sterilization cycle. With "Critical A + B" instruments: - MELAcontrol / MELAcontrol Pro must be used as batch control with every sterilization cycle. This simplifies the working procedure and increases security. You can omit the daily steam penetration test with MELAcontrol / MELAcontrol Pro (see above). The use of another test system in accordance with ►EN 867-5 is possible. The number of the available test systems means that MELAG is not able to provide technical support when using a different system.



PLEASE NOTE

The results of the tests must be documented.

- The indicator test strips used need not be stored.

¹⁾ in accordance with the current recommendations from the Robert Koch Institute

8 Careclave mode

Preparation and pre-cleaning

When preparing and pre-cleaning the instruments, comply with the following:

- If instruments are to be subject to manual preparation for cleaning, ensure that no media or tools/resources are deployed which could damage their surface. Never use any aggressive cleaning agents, wire or brass wire brushes or metal scourers. Information regarding correct instrument reprocessing is available from your instrument manufacturer.
- Water-insoluble treatment substances (e.g. dental cement, root canal disinfectants, alginates, silicone etc.) must be cleaned manually directly after use. Consult the product data sheets of the treatment substances.
- Other substances can also necessitate manual pre-cleaning. These include ultrasound gels and other auxiliary substances.
- Hollow bodies (transmission instruments, cannulae etc.) are to be checked for passage.
- Where applicable, instruments must be dismantled for reprocessing in accordance with the manufacturer's specifications.
- Corroded or defective instruments must be separated. Crusted instruments must be subject to a basic cleaning/repair.
- ►KRINKO►BfArM (2012) recommend that instruments of the risk class "Semi-critical B" and "Critical B" are subjected to pre-cleaning directly after use.

Loading the Carebox



WARNING

Danger from contaminated transmission instruments.

Observe the following when loading the Carebox:

- The instruments are contaminated after treatment and have sharp edges and tips. Use suitable protective gloves when loading the Carebox.
- Do not use the protective gloves used for loading for further operation of the device.
- Check the instruments for free passage before loading to prevent residues from impairing the disinfection.
- Only dental transmission instruments or hollow bodies may be reprocessed inside the Carebox. These must be attached to the adapters provided for this purpose. It is not possible to reprocess instruments or hollow bodies that are not attached to adapters.

Transmission instruments should be reprocessed immediately after use.



NOTICE

Correct instrument care

Observe the type of Carebox (Blue/Green) when loading.

- Instruments which may not be cared for (oiled) belong in the Carebox Green.
 - Instruments which require care belong in the Carebox Blue.
- ⇒ This will prevent damage to the instrument due to incorrect care.

The following must be fulfilled or present:

- ✓ There are no tools (drills) in the chucks.



PLEASE NOTE

If necessary, adapters can remain unfitted.

With the Carebox Blue, this can lead to increased oil leakage into the Carebox.

1. Insert the transmission instruments onto the adapters as far as they will go.
 - ↳ The heads point sideways to the edge.
 - ↳ The instrument engages audibly and noticeably.
2. For Sirona T1 Classic:
Align the transmission instrument so that the connections of the spray channels match the receptacles provided in the adapter.
3. Check that the transmission instrument and adapter are correctly connected by gently pulling on the instrument.
4. For tips:
Screw the tip onto the adapter using the torque wrench provided by the instrument manufacturer.
 - ↳ Reprocessing with a torque wrench connected is possible, as far as this has been approved by the manufacturer for automatic decontamination.
5. Check in the lower section whether the Carebox sieve is inserted and fixed from below with the safety spring.



6. Check in the lower section whether the lateral media filters are inserted.



7. Place the upper section on the lower section of the Carebox.
8. **NOTICE! If the latches are very difficult to close, check the cover to ensure that it is correctly seated.** Close the latches on both sides until you hear and feel them engage.

See also:

- [Adapter installation](#) [► page 24]
- [Changing the filter](#) [► page 73]

Loading Careclave

The following must be fulfilled or present:

- ✓ The Carebox has been cleaned and disinfected from the outside before it is moved to the clean side of the reprocessing room.
 - ✓ The sterilization chamber is dry.
 - ✓ The outside of the Carebox is dry and cooled down.
Please note: A Carebox that can only be handled with heat protection gloves is too hot and must be cooled down before it is processed in Careclave.
1. Open the door completely until it snaps into place.
 2. Place the Carebox on the upper mount in the inside of the door with a slight incline.



3. Slowly guide the Carebox to the lower mount until the magnetic connection engages.
 - ➔ The display shows **Carebox installed** and switches to Careclave mode.
4. Close the door.
 - ➔ The door is automatically pulled towards the device.
 - ➔ Careclave is ready for program selection.



PLEASE NOTE

Avoid incrustation of impurities.

- Immediately make a program selection and start Careclave.
 - ⇒ If no program start has occurred after twenty seconds, the warning "Carebox warming" is displayed and a warning tone is emitted.

Program selection



WARNING

Danger due to subsequent contamination

- Do not use potentially contaminated gloves to operate the device.
- If in doubt, clean and disinfect the affected surfaces with a surface disinfectant.



WARNING

In the Care-S and Care-Therm program, no reprocessing takes place in the sterilization chamber!

Program loading

Program		Packaging	Especially suitable for	Load
Care-S		Carebox ^{*)}	Transmission instruments semicritical B	Carebox with intended loading (max. 8 pcs.)
Care-Therm ^{**)}		Carebox ^{*)}	Transmission instruments semicritical B ^{***)}	Carebox with intended loading (max. 8 pcs.)
Care-B		Carebox	Transmission instruments semicritical B ^{***)}	Carebox: with intended loading (max. 8 pcs.)
		Single and multiple wrapped instruments	Instruments critical B narrow-lumen hollow bodies A	Sterilization chamber: short mount for trays with solid load max. 5 kg

^{*)} Reprocessing takes place exclusively in the Carebox. No reprocessing takes place in the sterilization chamber.
^{**)} There must be a pause of four minutes between the execution of two Care-Therm programs.
^{***)} The transmission instruments are heated with steam and must be temperature-resistant up to 135 °C.



PLEASE NOTE

If you are reprocessing loads in the sterilization chamber in the Care-B program, observe the sections on preparation, load, removal and storage in the chapter [Vacuclave mode](#) [► page 40].

See also:

- [Preparing the sterilization material](#) [► page 40]
- [Loading the steam sterilizer](#) [► page 41]
- [Removing the sterile material](#) [► page 47]
- [Storing sterile material](#) [► page 48]

Program parameters

Program	Temperature	Pressure	Disinfection/ sterilization time	Operating time ^{*)}	Drying
Care-S	134 °C	2.1 bar	03:30 min	23:24 min	03:04 min
Care-Therm	--	--	A0 > 3000	18:25 min	03:04 min
Care-B	134 °C	2.1 bar	05:30 min	43:30 min	15 min

^{*)} The operating times with Carebox Green are approx. two minutes shorter.



PLEASE NOTE

If the feed water or ambient temperature is too high, the operating time can be extended by up to eight minutes.

Starting the program

The door locks pressure-tight upon program start. The device performs automatic tests, e.g. a quantity check and conductivity measurement of the feed water.

1. Press **START PROGRAM**.
2. Confirm the subsequent dialogue window with **START PROGRAM**.
3. With enabled **Authentication at Reprocessing program start**, authenticate by entering a PIN.



During the program run, the display shows the current program duration, the current parameters and the expected end of the program.

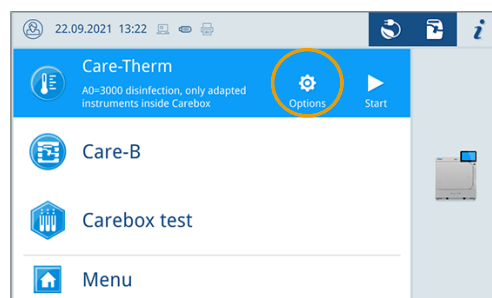
If no input is made on the display, the program display maximises and overlays the menu. Touch the display to show the menu.

Program options

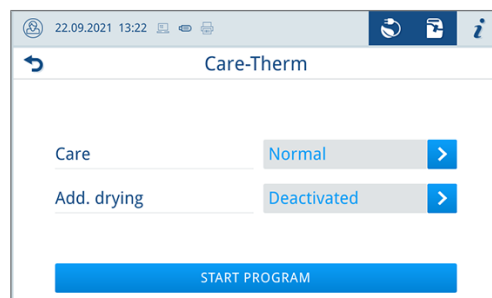
The **Options** button can be used to change settings one time for the selected program.

The options available in the menu depend on the device mode, the selected program and the user role.

1. Press the **Options** button.



2. Select the desired options.



3. Start the program with **START PROGRAM**.
4. With enabled **Authentication at Reprocessing program start**, authenticate by entering a PIN.
5. Confirm the subsequent dialogue window with **START PROGRAM**.

See also:

- [Program options](#) [▶ page 60]

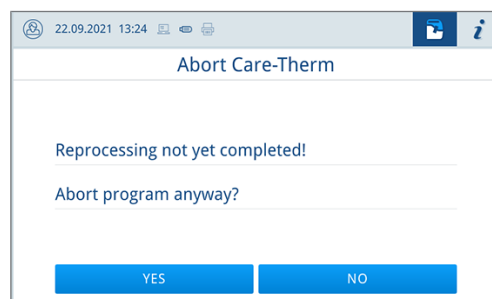
Manual program abort

You can abort the program at any time. If you abort the program before the end of the sterilization phase, the load is **not** sterile.

1. Press **ABORT** to abort a program.



2. Confirm the security query with **YES**.



- ➔ The load is not processed.
- ➔ Aborting the program may take a few minutes as steam and condensate are removed from the chamber.



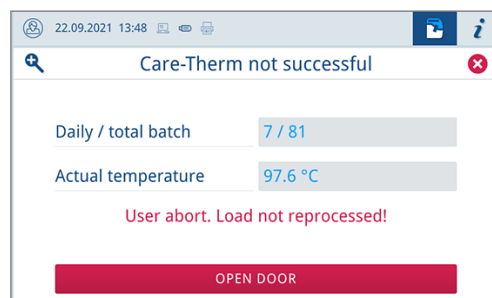
CAUTION

When the door is opened after a program abort, hot steam may escape and hot water may be present in the sterilization chamber and the Carebox.

This could result in scalding.

- Never touch the load, boiler or door with unprotected hands. The components are hot.
- The water that may be present can be seen through the transparent lower section of the Carebox.

3. Press **OPEN DOOR** to remove the load.



PLEASE NOTE

After the program has been aborted, the Carebox must cool down and be dry on the outside before reprocessing can be performed in Careclave.

Ending the program prematurely

You can end the program prematurely. If you abort the program before the end of the drying phase, the load is not completely dried and should be used immediately.

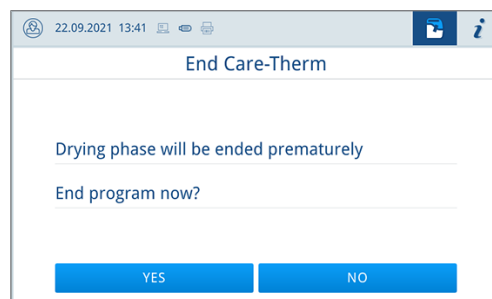
The following must be fulfilled or present:

- ✓ The drying phase has been reached.

1. To end the program prematurely, press **END**.



2. Confirm the termination of the drying process with **YES**.



→ The program will be aborted prematurely.

Program is ended

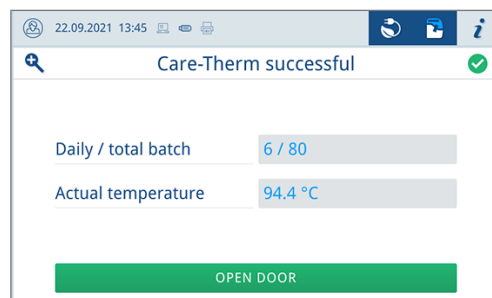


PLEASE NOTE

If the program has been carried out successfully, a corresponding message appears on the display and the status LED below the display illuminates green.

- If the display indicates that the program was not successful or the LED does not illuminate green, the program must be repeated.

1. Press **OPEN DOOR** to remove the load.



2. With enabled **Authentication at Reprocessing program end**, authenticate by entering a PIN.

Removing the Carebox



CAUTION

Danger of burns from hot parts and hot condensate.

Comply with the following specifications when removing the Carebox:

- Never touch the Carebox, the inside of the device or the inside of the door with bare hands. The components are hot.
- Use suitable protective gloves to remove the Carebox. The Carebox and the instruments may be over 100 °C hot after the program run.
- After the program run, there may still be a small amount of hot condensate in the Carebox. The condensate may drip out when the Carebox is removed or carried. Any condensate that may be present can be seen through the transparent lower section of the Carebox.
- When opening the Carebox, make sure that you do not come into direct contact with hot condensate.

The following must be fulfilled or present:

- ✓ End of the program has been reached
- 1. Unlock the door by pressing **OPEN DOOR**.
- 2. Open the door completely until it snaps into place.
- 3. **CAUTION! Danger of burns from hot parts and hot condensate.** Remove the Carebox.



- 4. Close the door.
 - ➡ The door is automatically pulled towards the device.



PLEASE NOTE

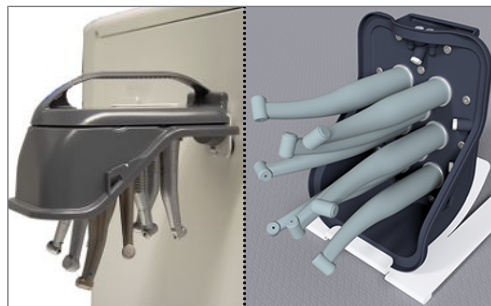
If the transmission instruments are to be used later, you can leave the Carebox closed during cooling.

In this case, it takes at least thirty minutes for the transmission instruments to cool down.

- 5. **CAUTION! Danger of burns from hot condensate.** Separate the upper section of the Carebox from the lower section of the Carebox to allow the transmission instruments to cool down.



6. Place the upper section of the Carebox in the Carebox cover holder or hang it on the side wall of the Careclave.



7. Allow the instruments to cool down.



PLEASE NOTE

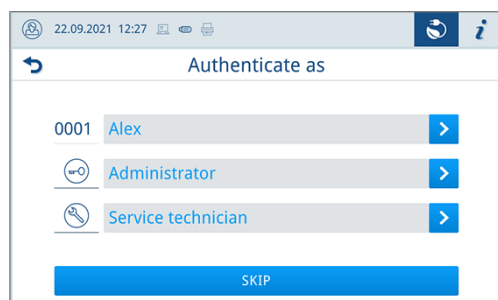
For fast and safe cooling of the instruments, MELAG recommends the Cooling Box.

See [Cooling with the Cooling Box](#) [► page 38].

The Approval process

According to ►RKI "Hygiene requirements for the treatment of medical products", instrument reprocessing ends with the documented approval for storage and application of the ►sterile material. The approval process consists of ►batch indication and batch approval and must be performed by authorised and expert personnel.

Batch approval comprises the checking of the process parameters using the sterilization results on the device and the sterilization log as well as checking of the individual packaging for damage and residual moisture. The sterilization log records the approval of the ►batch and any indicators. Depending on the setting in the user administration, approval for the ►sterile material requires the user PIN of the person who provides approval for the batch and the indicators.



See also:

- [Logging](#) [► page 49]
- [Administrative settings](#) [► page 62]

Removing instruments and hollow objects



CAUTION

Danger of burns from hot parts and hot condensate.

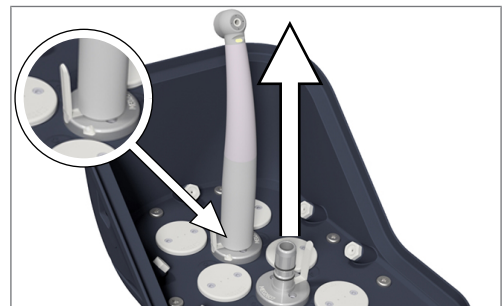
Comply with the following for safe handling:

- To ensure safe removal of the instruments and hollow objects, check whether they have cooled down sufficiently.
- Reprocess any instruments and hollow bodies that have fallen off adapters. They are not sterile.
- If necessary, take additional measures to cool down the instruments and hollow bodies:
 - Using the Cooling Box: Cooling time approx. four minutes
 - Cool down in the ambient air with the Carebox open: Cooling time at least fifteen minutes
 - Cool down in the closed Carebox: Cooling time at least thirty minutes

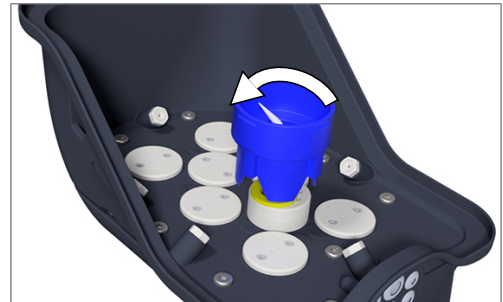
1. Check that the transmission instrument is firmly seated.
 ↳ Should the connection work loose after reprocessing, the instrument must be reprocessed again.
2. Hold the upper section of the Carebox with one hand.
3. When using an adapter for Sirona T1 Classic:
 Pull the transmission instrument forcefully and in a straight direction from the adapter with your other hand.



4. When using an adapter with an ISO coupling:
 Operate the latch with your other hand and then pull the transmission instrument from the adapter forcefully and in a straight direction.



5. When using an adapter for tips:
 Unscrew the adapter using the torque wrench.



6. Keep the Carebox in the clean area.



PLEASE NOTE

The transmission instruments are dried by compressed air/vacuum drying of the inner channels.

The result of the drying process strongly depends on the type and design of the instruments.

- Check the drying result of the transmission instruments and, if necessary, dry them with medical compressed air after removal.

Cooling with the Cooling Box

The following must be fulfilled or present:

- ✓ The Cooling Box is switched on.
- ✓ The upper section of the Carebox has been separated from the lower section.



NOTICE

Danger of short circuit

- Allow any wet instruments and hollow bodies to drip off before placing the upper section of the Carebox on the device.

1. Place the upper section of the Carebox on the cooling chamber of the device.



2. Check whether the upper section of the Carebox is correctly positioned on the seal of the cooling chamber.
3. Start a program run by pressing the corresponding program selection button.

Oiling the chucks

Chuck systems must be regularly maintained with suitable care products and oils. Comply with the information provided by the instrument manufacturer.



PLEASE NOTE

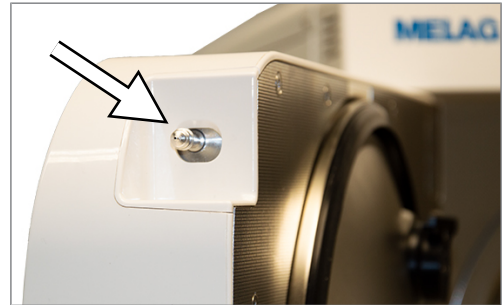
To ensure proper functioning of the collets, MELAG recommends oiling them once a week.

Chuck care must be performed after reprocessing with the Carebox.

- Perform the chuck care in addition to the care of the instruments by the Care programs.
- Comply with the information provided by the instrument manufacturer.

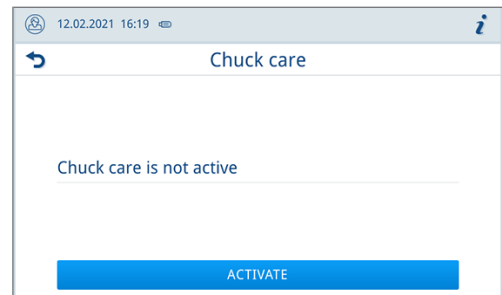
The integrated ADDcare care station can be used to care for the instruments. Proceed as follows:

1. Open the door completely until it snaps into place.
2. Start chuck care by briefly pressing (1 s) on the nozzle or by pressing the chuck care button.



→ The dialogue for chuck care appears on the user interface.

3. Press **ACTIVATE**.



4. Briefly press the instrument onto the nozzle with the chuck unlocked to perform chuck care.



5. Press the instrument briefly onto the nozzle repeatedly until oil emerges from the instrument head.
6. Then carefully clean the nozzle with a dry, lint-free cloth.
7. Press the chuck care button to switch off chuck care.

→ The dialogue for chuck care appears on the user interface.

8. Press **DEACTIVATE**.



PLEASE NOTE

If chuck care is not performed, the dialogue is automatically closed.

9 Vacuclave mode

Preparing the sterilization material

Cleaning and disinfection must always have been performed correctly and appropriately before sterilization. Only in this way is it possible to guarantee the subsequent sterilization of the [sterilization material](#). The materials used, cleaning agents and reprocessing procedure are of decisive significance.

Reprocessing the instruments

Unwrapped sterile material loses its sterility on contact with ambient air. If you intend to store your instruments sterilely, wrap them in suitable packaging before sterilization.

Ensure the following when reprocessing used and brand-new instruments:

- Observe both the instrument manufacturer's instructions regarding reprocessing and sterilization as well as the relevant standards and directives (in Germany, for example, from [RKI](#), [DGSV](#) and [DGUV Regulation 1](#)).
- Clean the instruments exceptionally thoroughly e.g. using an ultrasonic device or washer-disinfector.
- Rinse the instruments after cleaning and disinfecting, where possible with de-mineralized or distilled water and then dry the instruments thoroughly with a clean, non-fuzzing cloth.
- Use only those care agents suitable for steam sterilization. Consult the manufacturer of the care agents. Do not use any water repellent agents or oils impermeable to steam.
- When using ultrasonic devices, care equipment for handpieces and washer-disinfectors, comply with the manufacturer's reprocessing instructions.



NOTICE

The presence of residual disinfection and cleaning fluids results in corrosion.

This could result in increased maintenance requirements and a restriction of the steam sterilizer function.

Reprocessing textiles

Comply with the following points when reprocessing textiles and putting the textiles in sterilization containers:

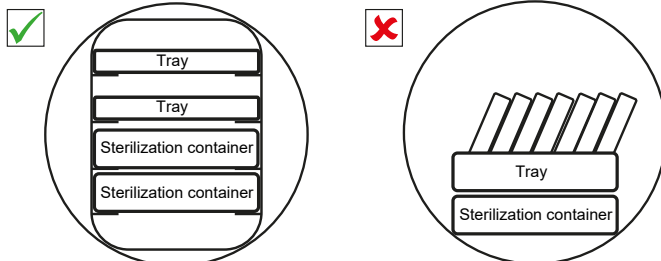
- Comply with both the manufacturer's instructions of the textiles regarding reprocessing and sterilization as well as the relevant standards and directives (in Germany e.g. from the [RKI](#) and [DGSV](#)).
- Arrange the folds in the textiles parallel to each other.
- Stack textiles vertically wherever possible and not too closely together in the sterilization container. This enables the development of flow channels.
- If textile packages do not remain together, wrap the textiles in sterilization paper.
- Only ever sterilize dry textiles.
- The textiles may not be permitted to come into direct contact with the sterilization chamber; otherwise they will become saturated with [condensate](#).

Loading the steam sterilizer

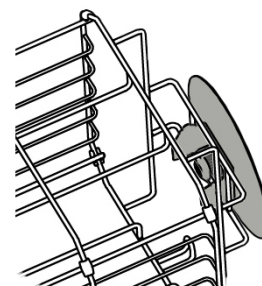
Effective sterilization and good drying is only possible if the steam sterilizer has been loaded correctly.

Ensure the following during loading:

- Insert trays or cassettes in the chamber only with their appropriate mount.



- Slide the mount into the sterilization chamber to its fullest extent. The holder must engage audibly and noticeably in the spring clip.



- Use perforated trays such as those from MELAG. Only in this way can **condensate** drain off. Non-perforated bases or half-shells for holding the **sterilization material** lead to poor drying results.
- The use of paper tray inserts can result in poor drying results.
- Wherever possible, ensure the separate sterilization of textiles and instruments in separate sterilization containers or sterilization packaging. This leads to better drying results.



Packaging

Only ever use packaging materials and systems (**sterile barrier systems**) which fulfil the standard **EN ISO 11607-1**. The correct use of suitable packaging is important in achieving successful sterilization results. You can use re-usable rigid packaging systems or soft packaging such as transparent sterilization packaging, paper bags, sterilization paper, textiles or fleece.

Closed sterilization containers



WARNING

Risk of contamination due to insufficient steam penetration or poor drying.

- Use only suitable sterilization containers.
- Do not cover the perforations when stacking the sterilization containers so that the condensate can drain off.

Please comply with the following when using closed sterilization containers:

- Use aluminium sterilization containers. Aluminium retains and conducts heat and thus accelerates drying.
- Closed sterilization containers must be either perforated or have a valve on at least one side. MELAG sterilization containers (e.g. MELAstore Boxes) fulfil the requirements for successful sterilization and drying.
- Wherever possible, ensure that sterilization containers are only stacked on top of those of identical size, so that the condensate can run down their sides.
- Ensure that the perforations are not covered when stacking the containers.

Soft sterilization packaging



WARNING

Risk of contamination due to insufficient drying

To improve the drying results for full loads of soft sterilization packages, the setting **Drying: Intelligent** must be activated.

► **Soft sterilization packaging** can be used in both sterilization containers and on trays. Please comply with the following when using soft sterilization packaging e.g. MELAfol:

- Arrange soft sterilization packaging in a perpendicular position and at narrow intervals.
- Place transparent sterilization packages on their edge wherever possible and with the paper side facing downwards.
- Do not place multiple soft sterilization packages flat on top of each other on a tray or in a container.
- If the seam seal tears during sterilization, this could be caused by the choice of undersized packaging. Re-pack the instruments with larger packaging and perform sterilization again.
- Should the seam seal rip during sterilization, extend the sealing pulse on the film sealing device or make a double seam.

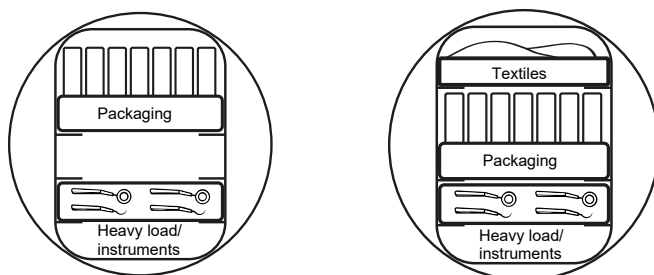
Multiple wrapping

The steam sterilizer works with a fractionated vacuum procedure. This permits the use of ► **multiple wrapping**.

Mixed loads

Observe the following when sterilizing ► **mixed loads**:

- Always place textiles at the top.
- Place the sterilization containers at the bottom.
- Place unwrapped instruments at the bottom.
- Place the heaviest loads at the bottom.
- Place transparent sterilization packaging and paper bags at the top - except in combination with textiles. In this case, place them at the bottom.



Selecting the program

Now select the sterilization program according to how and whether the ►sterilization material is packed. It is also necessary to take into account the temperature resistance of the sterilization material. All sterilization programs are displayed in the **Programs** menu. The following table shows which ►sterilization material fits which program and which additional programs are available.

Program loading

Program		Packaging	Especially suitable for	Load ^{*)}
Universal-B		Single and multiple wrapped	Mixed loads Long narrow-bore hollow bodies A	- Textiles (multiple wrapped) 2 kg - Instruments (single and multiple wrapped) 6 kg - Solid load (container^{**) / unwrapped instruments) 11 kg} - mixed load (textile/solid) 1.5 kg / 5.5 kg
Quick-S		Unwrapped	Unwrapped loads	Solid load 7 kg
Prion-B		Single and multiple wrapped	Instruments under suspicion of carrying the danger of infection through abnormally altered proteins (e.g. Creutzfeldt–Jakob, BSE)	- Textiles (multiple wrapped) 2 kg - Instruments (single and multiple wrapped) 6 kg - Solid load (container^{**) / unwrapped instruments) 11 kg} - Mixed load (textile/solid) 1.5 kg / 5.5 kg
Gentle-B		Single and multiple wrapped	Textiles; thermo-unstable items (e.g. plastic, rubber articles)	- Textiles (multiple wrapped) 2 kg - Instruments (single and multiple wrapped) 6 kg - Solid load (container^{**) / unwrapped instruments) 7 kg} - Thermo-unstable items / unwrapped instruments 7 kg - mixed load (textile/solid) 1.5 kg / 5.5 kg

^{*)} The max. weight per component is 2 kg for instruments or textiles.

^{**) checked with sterilization containers / MELAstore Box. Other load configurations must be checked individually and locally.}

Program parameters

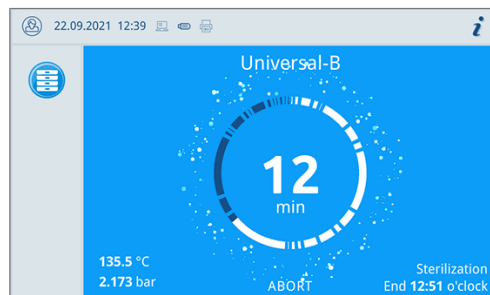
Program	Sterilization temperature	Sterilization pressure	Sterilization time	Operating time ^{*)}	Intelligent drying	Time-controlled drying
Universal-B	134 °C	2.1 bar	5:30 min	13-20 min	5-25 min	15 min
Quick-S	134 °C	2.1 bar	3:30 min	9-15 min	5-25 min	15 min
Prion-B	134 °C	2.1 bar	20:30 min	28-35 min	5-25 min	15 min
Gentle-B	121 °C	1.1 bar	20:30 min	27-37 min	5-25 min	15 min

^{*)} without drying, with a full load and dependent on the load and set-up conditions (such as e.g. water temperature and mains voltage)

Starting the program

With the start of the program, the door closes pressure-tight and the device checks the quantity of feed water and its conductivity.

1. Press **START PROGRAM**.
2. Confirm the subsequent dialogue window with **START PROGRAM**.
3. With enabled **Authentication at Reprocessing program start**, authenticate by entering a PIN.



During the program run, the display shows the current program duration, the current parameters and the expected end of the program.

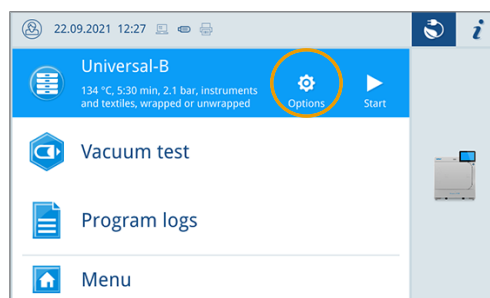
If no input is made on the display, the program display maximises and overlays the menu. Touch the display to show the menu.

Program options

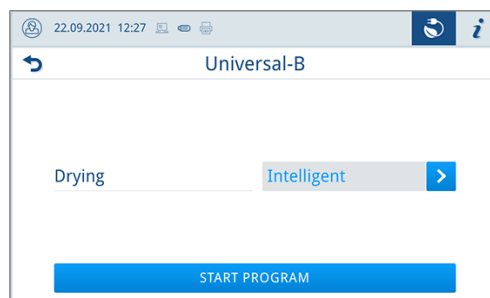
The **Options** button can be used to change settings one time for the selected program.

The options available in the menu depend on the device mode, the selected program and the user role.

1. Press the **Options** button.



2. Select the desired option.



3. Start the program with **START PROGRAM**.
4. With enabled **Authentication at Reprocessing program start**, authenticate by entering a PIN.
5. Confirm the subsequent dialogue window with **START PROGRAM**.

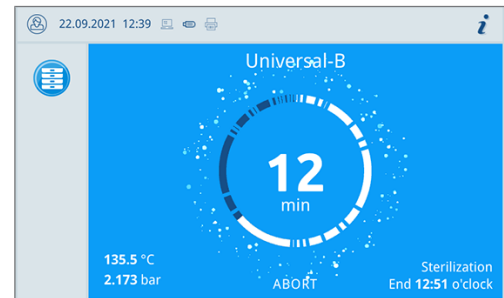
See also:

- [Program options](#) [► page 60]

Manual program abort

You can abort the program at any time. If you abort the program before the end of the sterilization phase, the load is **not** sterile.

1. Press **ABORT** to abort a program.



2. Confirm the security query with **YES**.



- The load is not sterile.
- Aborting the program may take a few minutes as steam and condensate are removed from the chamber.



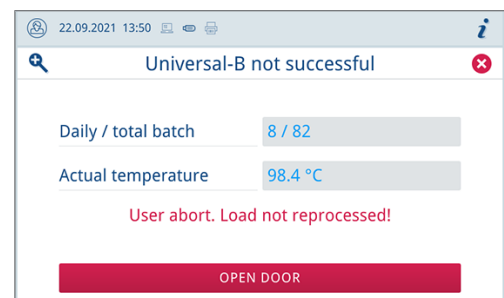
CAUTION

When the door is opened after a program abort, hot steam may escape and hot water may be present in the sterilization chamber.

This could result in scalding.

- Never touch the load, the sterilization chamber or the door with unprotected hands. The components are hot.

3. Press **OPEN DOOR** to remove the load.



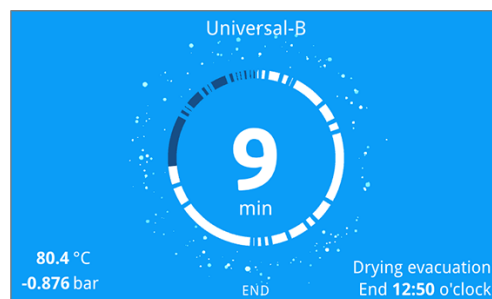
Ending the program prematurely

You can end the program prematurely. If you abort the program before the end of the drying phase, the load is not completely dried and should be used immediately.

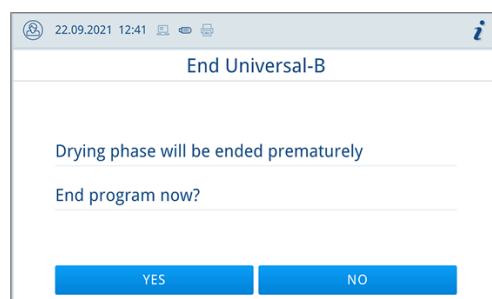
The following must be fulfilled or present:

- ✓ The drying phase has been reached.

1. To end the program prematurely, press **END**.



2. Confirm the termination of the drying process with **YES**.



➔ The program will be aborted prematurely.

See also:

- [Program options](#) [▶ page 60]

Program is ended

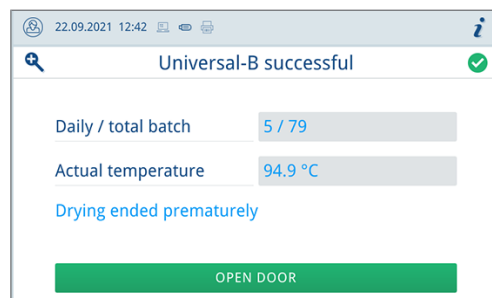


PLEASE NOTE

If the program has been carried out successfully, a corresponding message appears on the display and the status LED below the display illuminates green.

- If the display indicates that the program was not successful or the LED does not illuminate green, the program must be repeated.

1. Press **OPEN DOOR** to remove the load.

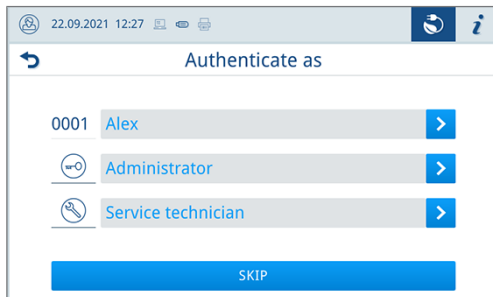


2. With enabled **Authentication at Reprocessing program end**, authenticate by entering a PIN.

The Approval process

According to ►RKI "Hygiene requirements for the treatment of medical products", instrument reprocessing ends with the documented approval for storage and application of the ►sterile material. The approval process consists of ►batch indication and batch approval and must be performed by authorised and expert personnel.

Batch approval comprises the checking of the process parameters using the sterilization results on the device and the sterilization log as well as checking of the individual packaging for damage and residual moisture. The sterilization log records the approval of the ►batch and any indicators. Depending on the setting in the user administration, approval for the ►sterile material requires the user PIN of the person who provides approval for the batch and the indicators.



See also:

- [Logging](#) [► page 49]
- [Administrative settings](#) [► page 62]

Removing the sterile material



CAUTION

Danger of burns from hot loads

- Use a tray lifter or heat protection gloves.



CAUTION

Unsterile instruments resulting from damaged or burst packaging. This endangers the health of your patients and practice team.

- Should the packaging be damaged or have burst after sterilization, wrap the sterilization material again and re-sterilize it.

If you remove the ►sterile material from the device directly after the end of the program, it is possible that the instruments can be partially damp. According to the red brochure of the Working Group Instrument Reprocessing (►AKI), the tolerable residual moisture is - in practice - a few drops of water capable of evaporating within 15 min, but actual pools of water are not acceptable.

Comply with the following specifications when removing the sterile material:

- Never use force to open the door. This could damage the device or result in the emission of hot steam.
- Hold the mount level when removing it from the device. Otherwise, the load could slide off.
- Hold the tray level when removing it from the steam sterilizer. Otherwise, the load could slide off.
- When removing the load from the device separately, ensure that the mount does not slide out unintended.
- Use a tray lifter to remove the tray.
- Use suitable protective gloves to remove the trays.
- Never touch the sterilized equipment, the chamber, the mount or the inside of the door with bare hands. The components are hot.
- Check the packaging of the sterile material for damage when removing it from the device. Should the packaging be damaged, wrap the sterilization material again and re-sterilize it.

Storing sterile material

The maximum storage time is dependent on the packaging and the storage conditions. For standard-conform wrapped ▶sterile material – (if protected from dust) it can amount to up to six months. Comply with the provisions of ▶DIN 58953, part 8 and the criteria specified below for the storage of sterile material:

- Comply with the maximum storage duration in accordance with the packaging type.
- Do not store the ▶sterile material in the reprocessing room.
- Store the sterile material in a dust-protected environment e.g. in a closed instrument cabinet.
- Store the sterile material in an environment protected against moisture.
- Store the sterile material in an environment protected against excess temperature variations.

10 Logging

Batch documentation

The batch documentation serves as proof of the successful conclusion of the program and represents an obligatory part of quality assurance. The device internal log memory saves such data as the program type, [▶batch](#) and process parameters of all the programs completed.

To obtain the batch documentation, you can output the internal log memory and transfer its data to various output media. This can be performed immediately at the end of every program or at a later point, such as at the end of the day.

If authentication is activated, the user ID and the result of the approval process are documented in the log header and, if required, on a label.

See also:

- [User administration](#) [▶ page 62]
- [Authentication](#) [▶ page 63]
- [Batch approval](#) [▶ page 63]

Logs menu

The **Logs** menu provides you with the following options:

- Display and output of program logs
- Display and output of malfunction logs
- Display and output of status logs
- Display and output of system logs
- Printing of labels

You can issue logs subsequently and independently of the time of a program end. You can set the output media yourself.

Log types

Log type	Description
Program log	Log of a program
Malfunction logs	Log with malfunctions that occurred outside a program run
Status log	Summary of all important settings and system statuses
System log	List of all the malfunctions and changes to the system in order of time (log book)

See also:

- [Log output](#) [▶ page 52]

Log list

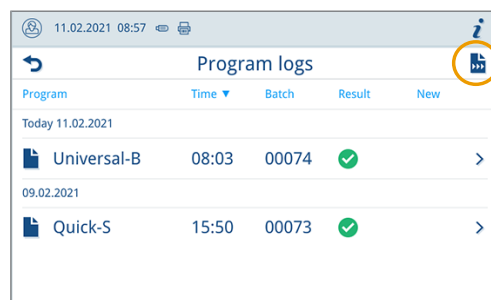
In the log list you can view all logs in detail. It displays all the logs present in the memory. You can sort the list by pressing the column headings.

The **Result** column shows whether the program ended successfully or not.

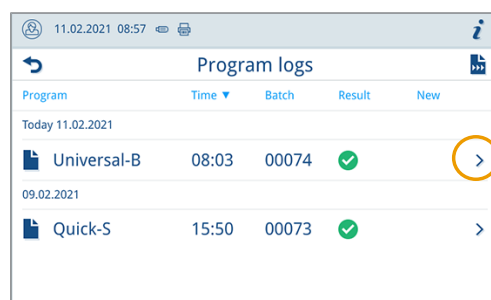
Symbol	Description
Green checkmark	Program successfully completed
Red cross	Program not successfully completed

Logs that have not yet been output are marked with a dot in the **New** column.

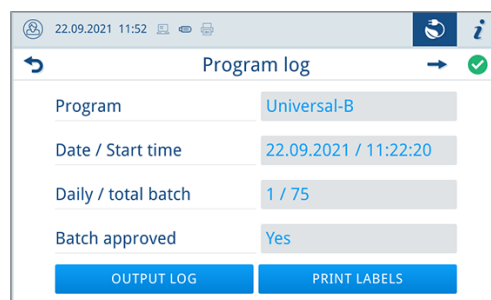
1. Press the button at the top right to customise the **Log output options** and output multiple logs.



2. Press the button with the arrow to view and output a log.



3. Press **OUTPUT LOG** to enter the **Log output options** and output the displayed log.



4. Press **PRINT LABELS** to open the label printing dialogue.
5. Press the button with the arrow to change the **Quantity** or the **Storage duration**.
 ➡ Confirm the changes with **OK**.
6. Press **PRINT LABELS** to print labels for the displayed log.

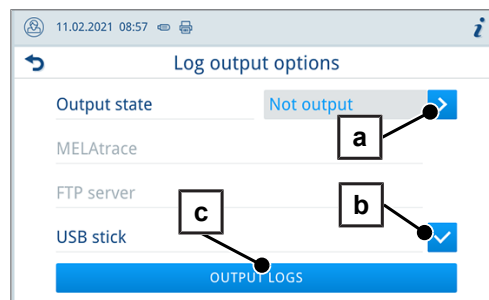
See also:

- [Log output](#) [▶ page 52]

Log output options

In the **Log output options** menu, you can set which logs are to be output and then output the logs.

1. Press the button with the arrow to select the desired output status (pos. a).



2. Activate the desired output medium (pos. b).
 ↳ Unavailable output media are displayed greyed out.
3. Press **OUTPUT LOGS** (pos. c).
 ↳ The output takes place on the selected output media.

Output status

The following settings are possible:

Output status	Description
Not output	All logs that are not output will be output.
Last	The log of the last successful completed program is outputted.
All	The logs of all successfully completed programs will be outputted.

Output media

The following output media can be activated and configured in the menu **Settings > Log output**:

- MELAt race
- FTP
- USB on USB stick

For activated output media, the symbol in the information area is displayed faintly.

For activated and connected output media, the symbol is displayed in full.

Output media that are not activated are not displayed, even if they are connected.



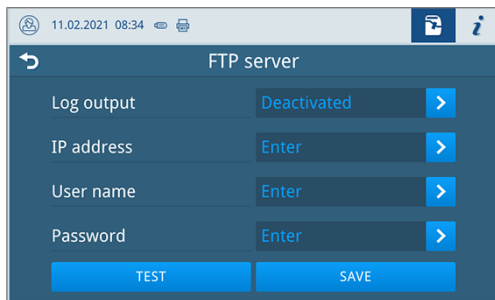
NOTICE

You can only connect one USB storage medium.

Symbol	Output media	Description
	USB flash drive	Output to a USB stick connected to the USB port
	FTP	Output to an FTP server
	MELAt race	Output to MELAt race
	Label printer	Output to a connected label printer

FTP server configuration

Under the FTP menu item, the configuration of the FTP server is also carried out via the IP address, the user name and the password.



The **TEST** button can be used to test the set configuration.

See also:

- [Views of the device](#) [► page 15]
- [Smart touch display](#) [► page 19]
- [Log output](#) [► page 52]

Log output

In the **Settings > Log output** menu, you can set how the log should be output by default for each output medium. You can activate the log output for several output media at the same time.

For each output medium, one of the following options can be selected for the **Log output**:

Option	Description
deactivated	No log output
Manual	Manual log output
Automatic (immediately after program run)	Automatic log output after end of the program for the defined programs

For the option **Automatic** a dialogue follows for the definition for which programs the automatic log output should take place.

See also:

- [Log list](#) [► page 50]
- [Log output options](#) [► page 51]
- [Output media](#) [► page 51]

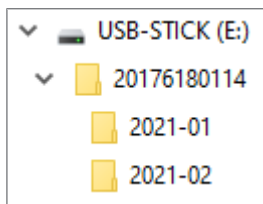
Finding logs

Storage location for logs

When transferring the logs to a USB stick, they will be stored in a separate folder in the main directory. Direct transfer of the logs to a computer via the network and using the MELAG **FTP** server allows you to work directly in the FTP server to determine directly where on your computer the device directory with log files is to be saved. With output via MELAtrace, you can work directly in the program to determine the folder in which they are to be saved.

Log directory

A folder is created on all memory media (USB stick or computer) after log output containing the serial number of the issuing device. This folder contains sub-folders with the month of log generation e.g. 2021-01 for January 2021. This contains all logs generated by the device this month.



Displaying logs on the computer

The log files are generated in html format and can be displayed and printed on the computer with a web browser or in MELAtrace/MELAvi.

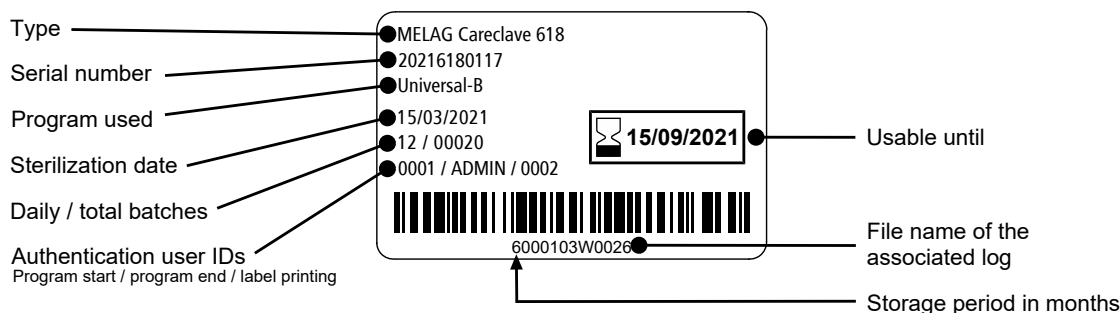
The program, malfunction and status logs contain a file entry for each line. The program logs contain graphic data and can be displayed as graphic logs in MELAvi.

Protocol					
file:///F:/2021-01-29_00001_20186180124_UNI_OK_000103N0001.html					
000	Device ID	1001000124	000	Ident informations of the device	
010	File name	2021-01-29_00001_20186180124_UNI_OK_000103N0001	010	File name of the log	
020	Device type	Careclave 618	020	Device type	
030	Program name	Universal-B Program	030	Program name	
035	Program type	134 °C wrapped	035	Program type	
040	Date	29.01.2021	040	Creation date of the log	
045	Daily / total batch	01 / 00001	045	Daily batch number and total batch number	
050	User program start	0001	050	User ID at program start	
055	User program end	0002	055	User ID at program and batch approval	
060	Indicator changed	Yes	060	Indicator assessment	
065	Batch approved	Yes	065	Status batch approval	
070	Program result	Program successfully completed	070	Program result	
141	Sterilization temperature	135.5 ±0.09/-0.21 °C	141	Sterilization temperature with max. deviation	
143	Sterilization pressure	2.17 ±0.00/-0.02 bar	143	Sterilization pressure with max. deviation	
144	Plateau time	5 min 30 s	144	Sterilization time	
150	Conductivity	0 µS/cm (0 ml : 0.01 µS/cm)	150	Conductivity of feed water and feed quantity	
155	Start time	07:42:17	155	Time at program start	
156	End time	08:01:11 (18:54 min)	156	Time at program end and program duration	
160	Device serial number	20186180124	160	Serial number of the device	

Step	Start [m:s]	End [m:s]	Dauer [m:s]	P [mbar]	T [°C]	
Program start						
SP-S	00:00	00:00	00:00	c 0	c 0.0	Program start

Label printer as output medium

The use of a label printer facilitates batch traceability. Using the sterilization date, the storage duration, batch number, user ID of the person approving the application for use, the steam sterilizer used and the file name it is possible to assign the sterilized instruments to the patient and sterilization batch. Faultless packages containing sterile material are marked with labels after sterilization. As such, the preconditions for correct "Approval" by the person conferred with the task of decontamination are given. All information regarding the correct sterilization procedure can be attributed to the instruments used in patient records.



See also:

- [Label print](#) [▶ page 61]

11 Function tests

Service programs

Service programs in Vacuclave mode

Program	Program name	Operating time	Use/function
	Vacuum test	18 min	For measuring the leakage rate, test with a dry and cold device (test without load)
	Bowie & Dick test	15 min	Steam penetration test with special test package (available from specialist stockists)
	Draining	5 min	For emptying and pressure release of the double jacket steam generator, e.g. for service, decommissioning or before transport

Service programs in Careclave mode

Program	Program name	Operating time	Use/function
	Carebox test	3 min	For checking the Carebox without load
	Oil dosing venting ^{*)}	22 min	For diagnosis and maintenance of the oil dosing system
*) Administrator login required			

See also:

- [Vacuum test](#) [▶ page 55]
- [Bowie & Dick test](#) [▶ page 56]
- [Emptying](#) [▶ page 77]
- [Carebox test](#) [▶ page 56]

Vacuum test

The device can be checked for leakages in the steam system using the ▶ **vacuum** test. This determines the leakage rate at the same time.

Perform a vacuum test in the following circumstances:

- Once a week in routine operation
- During commissioning
- Following longer operating pauses
- In the case of a corresponding malfunction (e.g. in the vacuum system)



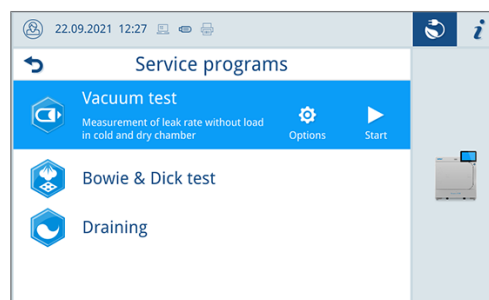
PLEASE NOTE

Perform the vacuum test with the device in a cold and dry state.

The following must be fulfilled or present:

- ✓ There is no Carebox in the Careclave.

1. Switch on the device.
2. Working in the **Service programs** menu, select **Vacuum test** and press **Start**.



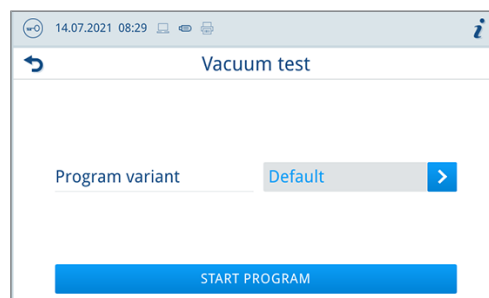
➡ The vacuum test is started in the **Default** program version.

➡ The evacuation pressure and the equilibration time or measuring time are shown on the display. The chamber is ventilated after the end of the measuring time. Then the message will be shown on the display with an indication of the leakage rate. Should the leakage rate be too high i.e. over 1.3 mbar, a corresponding message will be issued on the display.

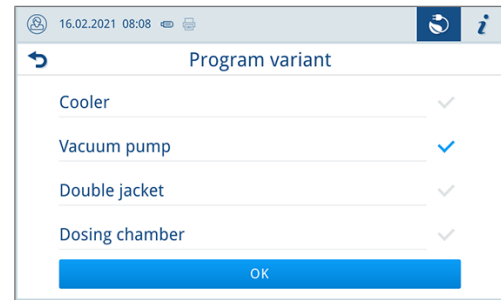
Options for the vacuum test

Under **Options**, you can extend the vacuum test to areas that are connected to the sterilization chamber. In this way, you can also evaluate their leak tightness.

1. Press the button with the arrow to select another version of the vacuum test.



2. Select the desired version and accept it with **OK** at the end of the list.



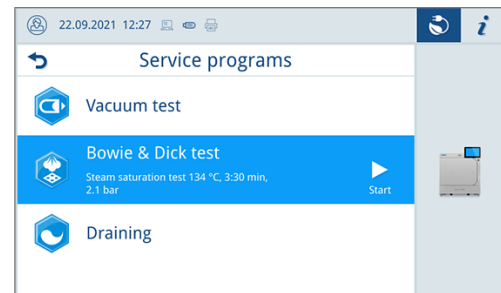
3. Start the vacuum test with **START PROGRAM**.

Bowie & Dick test

The **Bowie & Dick-Test** serves as proof of steam penetration of **porous materials** such as e.g. textiles. You can perform a routine function check for proof of steam penetration. Use the Bowie & Dick test program to this end. Specialist stockists provide various test systems for the Bowie & Dick test. Perform the test according to the test system manufacturer's specifications.

The following must be fulfilled or present:

- ✓ A new test system.
 - ✓ There is no Carebox in the Careclave.
 - ✓ The sterilization chamber is empty.
1. Place the test system in the sterilization chamber according to the manufacturer's instructions.
 2. Close the door.
 3. Working in the **Service programs** select **Bowie & Dick test** and press **Start**.



Carebox test

Use the **Carebox test** to check that the Carebox and its supply channels are functioning correctly.

Perform a **Carebox test** in the following circumstances:

- Once a week during routine operation, after the vacuum test
- During initial start-up of a Carebox or device
- After each adapter exchange
- Following longer operating pauses
- In the case of a corresponding malfunction (e.g. Internal cleaning or Carebox detection)



PLEASE NOTE

The distribution of the rinse liquor directly affects the cleaning performance of Careclave. Blocked channels in the upper section of the Carebox or in the instrument adapters can prevent successful cleaning.



PLEASE NOTE

If several Careboxes are used, all Careboxes must be checked.

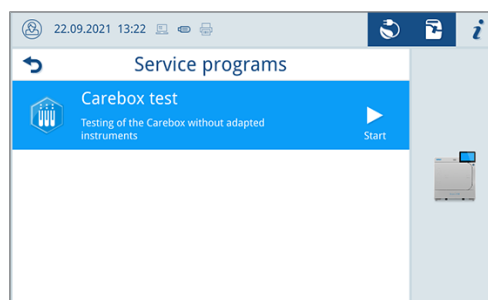
The following must be fulfilled or present:

- ✓ The measuring device.
- ✓ There are no transmission instruments or hollow bodies in the Carebox.

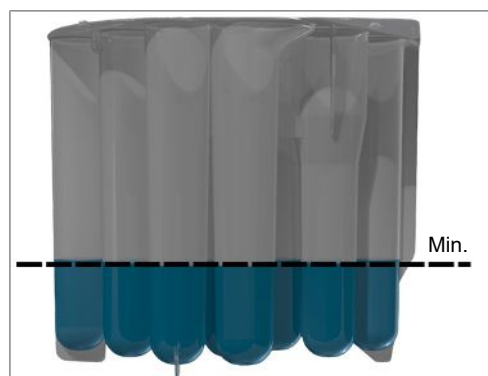
1. Slowly and carefully place the measuring device in the lower section of the Carebox, making sure that it is correctly aligned.



2. Close the Carebox.
3. Load the device with the Carebox.
4. Start the **Carebox test** service program.



5. Remove and open the Carebox.
6. **CAUTION! Danger of burns.** Remove the measuring device.
7. Check that all filling levels are at or above the minimum level line.



8. Answer the following question according to the result.
9. If a filling level is below the line for the minimum level, repeat the **Carebox test**.
10. If the minimum level is still not reached, please contact your stockist or MELAG customer service.

12 Settings

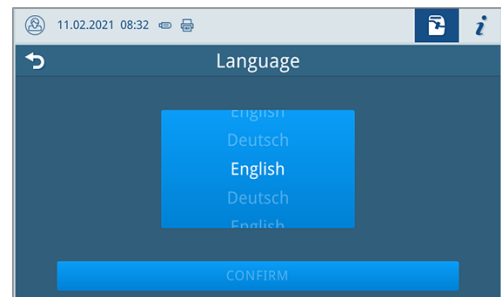
General settings

General settings can be changed by any user.

Language

In the **Settings** > **Language** menu, you can switch between the enabled languages.

1. Set the desired language.



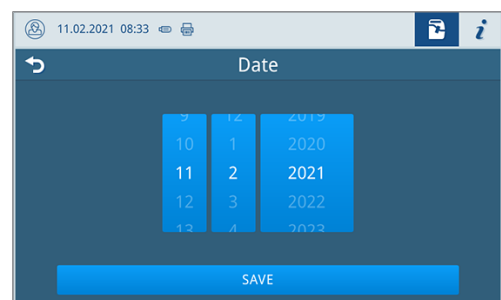
2. Press **CONFIRM** to accept the changes.

→ The dialogues on the display and the log texts are changed to the selected language.

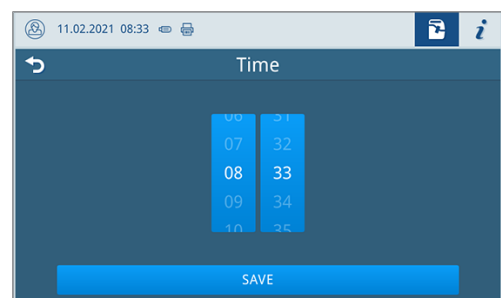
Date and time

Date and time of the device must be correctly set for proper batch documentation. Ensure that you take into account any clock change in autumn and summer, as this is not adjusted automatically. Set the date and time as follows:

1. Open the **Settings** menu.
2. Select the **Date** menu item.
3. Set the date.



4. Press **SAVE** to accept the changes.
5. Select the **Time** menu item.
6. Set the time.



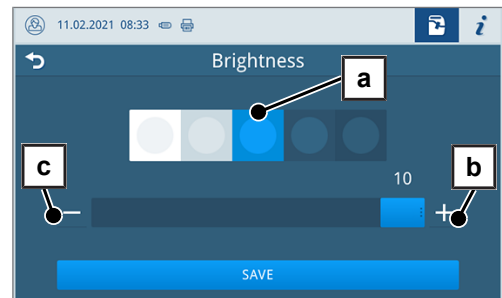
7. Press **SAVE** to accept the changes.

Display brightness

In the **Settings** > **Brightness** menu, you can individualise the brightness of the display.

The display brightness is adjusted immediately. The colour bar (pos. a) gives you an impression of the colour contrast.

1. Move the slider to the left or right or press the plus (pos. b) or minus (pos. c) buttons.



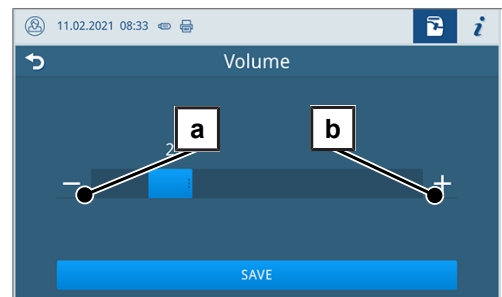
→ The display brightness can be adjusted in ten steps.

2. Press **SAVE** to accept the changes.

Volume

In the **Settings** > **Volume** menu, you can individualise the sound output.

1. Move the slider to the left or right or press the minus (pos. a) or plus (pos. b) buttons.



→ The volume can be adjusted in ten steps.

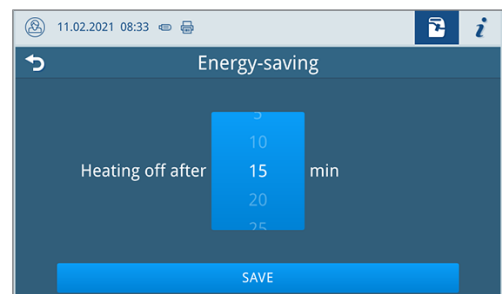
→ At level 0, the sound is switched off.

2. Press **SAVE** to accept the changes.

Energy-saving

In the **Settings** > **Energy-saving** menu, you can set after how long the device is inactive the heating is switched off.

1. Set the desired switch-off time in minutes on the number wheel.

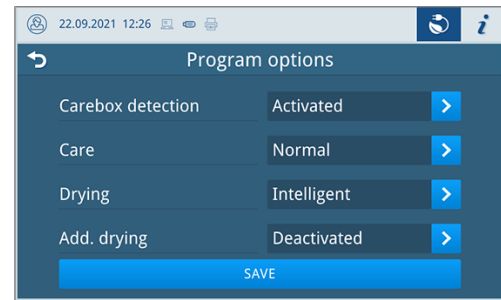


2. Press **SAVE** to accept the changes.

Program options

In the **Settings > Program options** menu, you can set default settings for program options.

1. Press the button with the arrow to make changes.



2. Activate or deactivate the desired setting by selecting or deselecting it.
3. Confirm the changes with **OK**.
4. Press **SAVE** to accept the changes.

The following settings are possible:

Device mode	Designation	Short description
General	Carebox detection	Checks the Carebox holder of the inside of the door for a Carebox and the Carebox inserted.
Careclave	Care: Normal	Cares for the instruments connected to the instrument adapters with care oil.
	Care: Intensive	Cares for the instruments connected to the instrument adapters with ample care oil.
	Care: Off	Does not perform any care. Comply with the information in the cleaning instructions from the instrument manufacturer.
Careclave	Add. drying	Performs additional drying in the Carebox to optimise the drying of the Carebox.
Vacuclave	Drying: Intelligent	Automatically monitors and ends the drying phase once the load is dry.
	Drying: Time-controlled	Ends the drying phase after a specified duration.

Carebox detection

Before the program starts, the device checks whether there is a Carebox on the suspension in the sterilization chamber. If a Carebox is detected, the version (Blue/Green) is checked as well as whether all channels to be cleaned are accessible.

Drying

You have the option to change the preset drying mode once at the program start via **Program options**.

Time-controlled drying

In the case of timed drying, the duration of the drying phase is determined by the program.

Intelligent drying

In contrast to a conventional time-controlled drying process, the duration of the intelligent drying is automatically calculated using the residual moisture in the sterilization chamber. The drying phase is ended as soon as the load is dry. A number of factors play a role in this process including e.g. the type of the load, wrapped or unwrapped, the load quantity, the distribution of the load in the sterilization chamber etc.

**PLEASE NOTE**

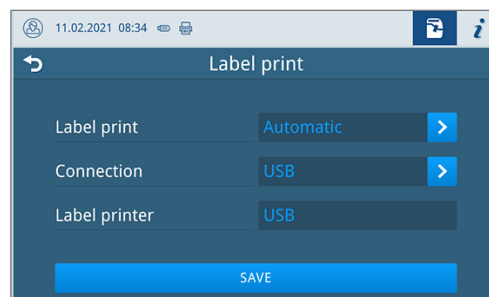
Intelligent drying is activated in the delivery state.

Label print

In the **Settings > Label print** menu, you can configure the label printer and set default settings.

The label printer can be connected via USB or via integration into a local network (LAN). If several devices access the label printer, it must be integrated via a local network (LAN).

1. Press the button with the arrow to make changes.



2. Activate or deactivate the desired setting by selecting or deselecting it.
 ➔ For the option **Automatic** a dialogue follows for the definition for which programs the automatic label print should take place.
3. Confirm the changes with **OK**.
4. Press **SAVE** to accept the changes.

The following settings are possible:

Option	Description
deactivated	No label printing
Manual	Manual label printing
Automatic (immediately after program run)	Automatic label printing after the end of the program for the defined programs

**PLEASE NOTE**

The last storage time set during label printing is taken over individually for each program as a presetting for the next label printing.

See also:

- [Label printer as output medium](#) [▶ page 53]

Water supply

In the **Settings > Water supply** menu, you can switch to a manual water supply.

The following settings are possible:

Designation	Description
Automatic	The feed water is automatically supplied via the <i>MELAdem feed water connection</i> .
Manual	Before starting the program, the feed water tank must be filled manually. The required amount is about two litres. PLEASE NOTE: The fresh water tank must be filled before each program start.

Logging

Further information on the logging settings and the output media can be found in the chapter [Logging](#) [▶ page 49].

Administrative settings

To make administrative settings, such as changes to the user administration, you must log in as an Administrator or a Service technician.

See also:

- [Logging on user role](#) [▶ page 65]

User administration

An individual ID and user PIN can be issued to every user to facilitate reliable traceability via the approval process after the end of a sterilization program. With the user PIN, the user can authenticate himself before the batch is approved.

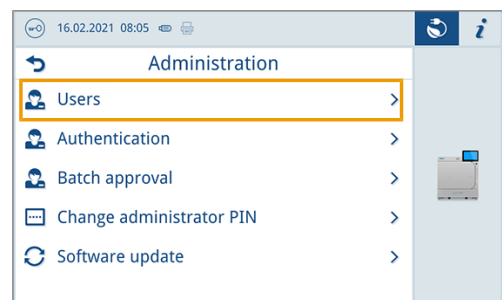
Only created users are authorised to approve and can approve a batch with their user PIN.

In the **Settings > Administration** you can create or edit users.

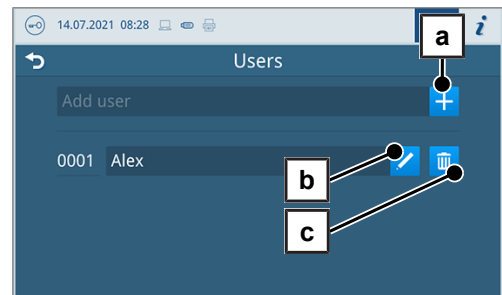
The following must be fulfilled or present:

- ✓ The logged-in user role is: Administrator or Service technician.

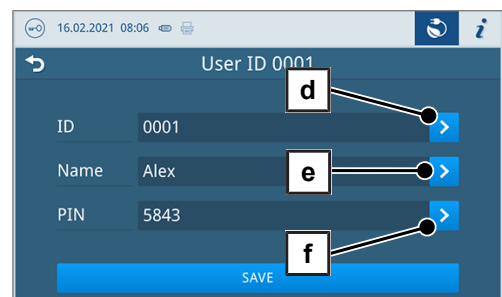
1. Select the **Users** menu.



2. Press the plus button (pos. a) to create a new user.



3. Edit (pos. b) or delete (pos. c) the user using the buttons next to the user name.
4. Press the buttons with the arrow to change the ID (pos. d), the user name (pos. e) or the PIN (pos. f).



5. Confirm the changes with **OK** and apply the changes with **SAVE**.

**PLEASE NOTE**

You can determine the necessity of user authentication via a PIN in the **Authentication** menu.

See also:

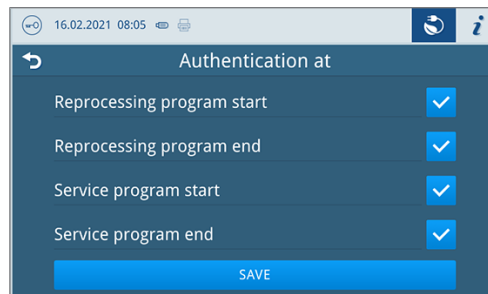
- [Authentication](#) [▶ page 63]
- [Batch approval](#) [▶ page 63]

Authentication

In the **Settings > Administration** menu, you can activate authentication (PIN entry) for the start or end of the program.

The following must be fulfilled or present:

- ✓ The logged-in user role is: Administrator or Service technician.
1. Select the **Authentication** menu.
 2. Activate or deactivate the desired setting by selecting or deselecting it.



3. Press **SAVE** to accept the changes.

The following settings are possible:

Designation	Description
Reprocessing program start	PIN entry required to start a program
Reprocessing program end	PIN entry required to open a door
Service program start	PIN entry required to start a service program
Service program end	PIN entry required to open door after a service program

**PLEASE NOTE**

All options are disabled in the factory settings.

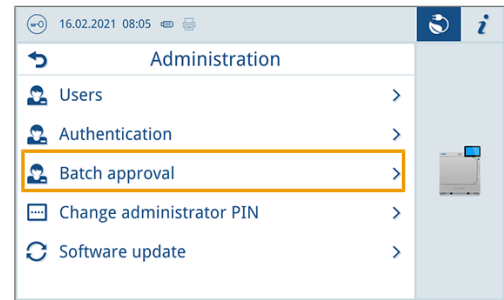
Batch approval

In the **Settings > Administration** menu you can activate the batch approval after the program has ended successfully and the indicator assessment.

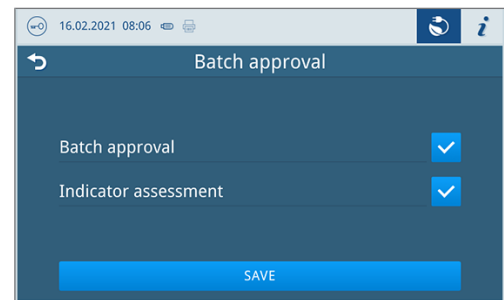
The following must be fulfilled or present:

- ✓ The logged-in user role is: Administrator or Service technician.

1. Select the **Batch approval** menu.



2. Activate or deactivate the desired setting by selecting or deselecting it.



3. Press **SAVE** to accept the changes.

The following settings are possible:

Log type	Description
Batch approval	Batch approval after successful program end
Indicator assessment	Indicator assessment approval after successful program end

Software update

In the **Settings > Administration** menu, you can perform a software version update.



NOTICE

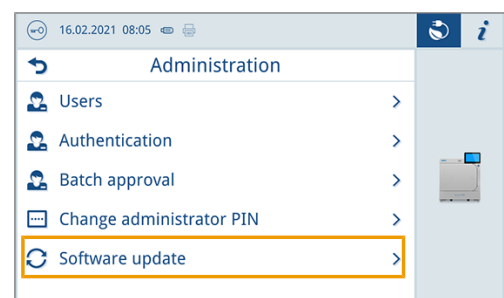
During a software update, all program logs are deleted.

- Check whether all required logs have been output to an output medium.

The following must be fulfilled or present:

- ✓ The logged-in user role is: Administrator or Service technician.
- ✓ A USB stick in FAT 32 format with installation data.
- ✓ All required logs have been output.

1. Select the **Software update** menu.



2. Insert a USB stick with the installation data in any USB connection.

3. Press **NEXT** to perform the software update.
 - ➔ During the software update, the device independently performs one or more restarts.

See also:

- [Log list](#) [▶ page 50]

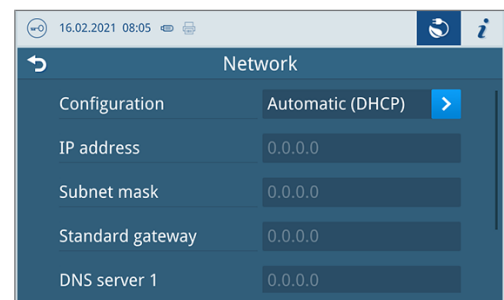
Network

In the **Settings > Network** menu, you can select an automatic configuration via DHCP or enter the required address details manually.

The following must be fulfilled or present:

- ✓ The logged-in user role is: Administrator or Service technician.

1. Press the button with the arrow to make changes.

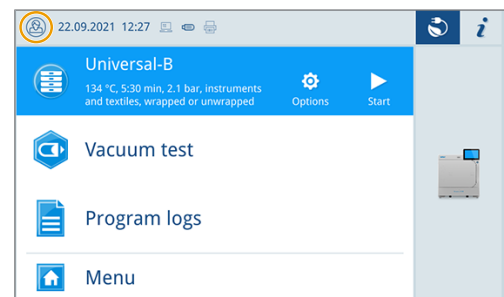


2. Press **SAVE** to accept the changes.

Logging on user role

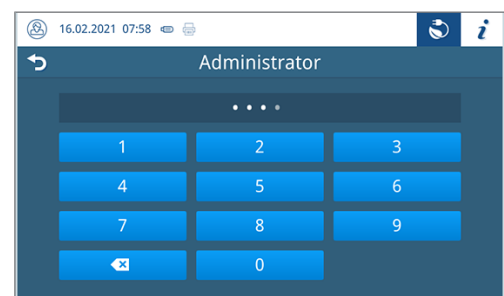
To log on a user role, proceed as follows:

1. Press the User Role button.



2. Select the desired role, e.g. Administrator.

3. Enter the associated PIN.

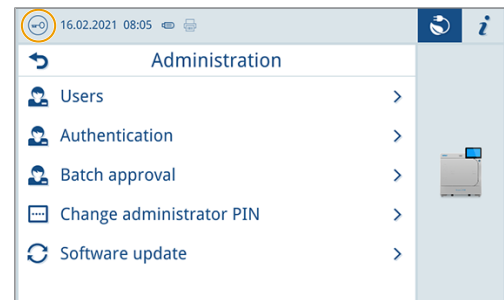


- ➔ The symbol of the user role button changes.
- ➔ Further setting options are now available in the menu.

Logging off a user role

In order to log off a user role, proceed as follows:

1. Press the User Role button.



2. Press **LOGOUT**.

→ The symbol of the user role button changes.

Service settings

To make service settings, such as a **Software reinstallation**, you must log in as a Service technician. Only ►[authorised technicians](#) have access to the further service documents required for this purpose.

See also:

- [Logging on user role](#) [► page 65]

13 Maintenance



PLEASE NOTE

The maintenance work described below can be performed by the user as part of in-house maintenance.

All maintenance activities beyond this must only be carried out by an [authorized technician](#).

Servicing intervals

Interval	Measure	Device components
Daily	Check for soiling, deposits or damage	Chamber inc. door seal and chamber sealing face, door lock, mount for the load
Daily	Check the Carebox for contamination and clean if necessary	Carebox
	Check the Carebox sieve for contamination and clean if necessary	Carebox lower section
	Check O-rings on the adapters for presence, wear or damage and replace if necessary	Carebox upper section
	Visual inspection of the seals (housing seal and media seals) and replace if necessary	Carebox
Weekly	Cleaning	all device components and the Carebox
	Carebox test	Carebox
After 2 months	Oil the door spindle	Door mechanism
Upon display request	Changing the oil can	Door
	Changing the media filter	Carebox
After 600 cycles (recommended)	Change adapter seals (O-rings)	Carebox
1x annually or following 1000 cycles	Replace the dust filter	behind the service hatch
After 24 months or 3000 cycles	Maintenance	By the authorized customer services working in accordance with the maintenance instructions
As required	Cleaning the surfaces	Housing parts

Accessories

Also comply with the maintenance intervals of the optional accessories.

Interval	Measure	Device component
After 24 months	Change the HEPA filter	Cooling Box
After 24 months	Replace the housing seal	Cooling Box

See also:

- [Cleaning](#) [► page 70]

Replacing the oil can



NOTICE

Danger due to incorrect cleaning and care of the transmission instruments

Can result in damage to the transmission instruments.

- Use only original accessories.



PLEASE NOTE

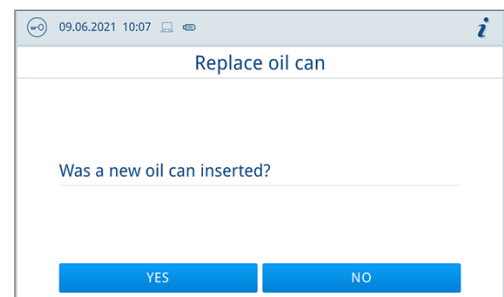
After changing the oil can, carry out a filter change on all Careboxes.

The following must be fulfilled or present:

- ✓ A new oil can with adequate best-before date.
 - ✓ The device is switched on.
1. Open the door completely until it snaps into place.
 2. Screw the cover in the side of the door.



3. Remove the oil can.
4. Insert the new oil can into the door.
5. Screw the cover cap back in up to the locking points.
6. Reset the counter by answering the question with YES.



7. Close the door.
8. After changing the oil can, carry out a media filter change on all Careboxes.

See also:

- [Changing the filter](#) [▶ page 73]

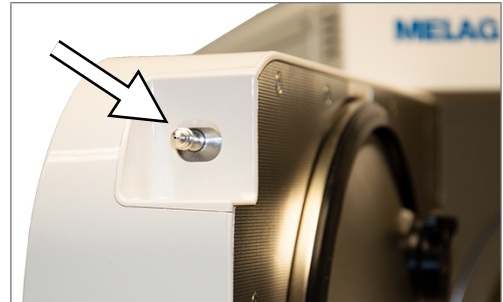
Venting the chuck care system



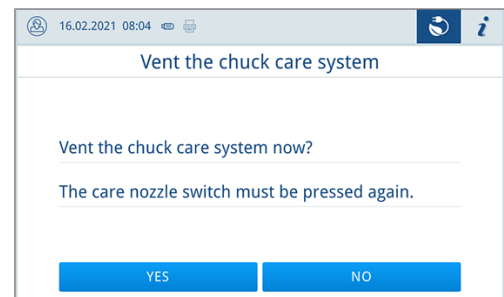
PLEASE NOTE

Venting of the chuck care system is only necessary in case of malfunction.

1. Press an instrument or lint-free cloth onto the care nozzle for more than five seconds to vent the chuck care system.



2. Activate venting by answering the question with **YES**.



3. Press the care nozzle until you hear an acoustic signal.

→ The chuck care system has been vented.

See also:

- [Oiling the chucks](#) [▶ page 38]

Oiling the door spindle

Oil the door spindle **every two months** as follows:

1. Clean the spindle with a non-fuzzing cloth.
2. Place two drops of oil in the bushing on the door from the oil can included in the scope of delivery (art. no. ME27515).

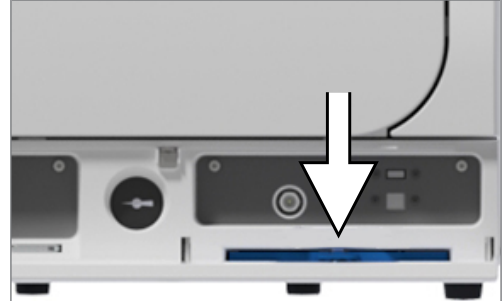
Instructions for oiling the door spindle are printed on the inside of the door.

Changing the dust filter

The following must be fulfilled or present:

- ✓ A new and dry dust filter.
- ✓ The device is switched on.

1. Open the service hatch.



2. Press down the centre of the grip and pull out the dust filter.



3. Insert the new dust filter until it snaps into place.
The latch nose of the grip must point upwards.
4. Close the service hatch.
5. Answer the following question with YES.

Cleaning



NOTICE

Inappropriately performed cleaning can lead to the scratching of and damage to surfaces and the development of leaks in sealing surfaces.

This also favours the development of soiling deposits and [corrosion](#) in the [sterilization chamber](#).

- Comply with all information regarding cleaning of the part affected.

Sterilization chamber, chamber sealing face, mount, trays

To maintain the value of your device and to avoid stubborn contamination and deposits, MELAG recommends weekly cleaning of the surfaces (e.g. with the MELAG chamber cleaning set).

Cleaning the sterilization chamber, chamber sealing face, mount and trays



NOTICE

Damage to the surfaces

Do not use any hard objects for cleaning such as a metal saucepan cleaner or a steel brush.

The following must be fulfilled or present:

- ✓ The device has been switched off and the power plug has been unplugged from the socket.
- ✓ The device has been completely cooled.
- ✓ Trays or cassettes and the associated mount have been removed from the sterilization chamber.
- 1. Wet the surfaces to be cleaned completely with the cleaning fluid.
Please note: You should not allow cleaning fluid to enter the piping coming from the sterilization chamber.
- 2. Spread the cleaning fluid evenly with a non-fuzzing cloth.
- 3. Allow the cleaning fluid to act and evaporate for a sufficient time.
- 4. Using a new non-fuzzing cloth, spread ample amounts of demineralized water over the cleaned surfaces.
- 5. Wipe the surfaces thoroughly to remove cleaning residues. Repeat this process as necessary after wringing out the cloth.
 - ➡ Residues of cleaning fluids can ignite or cause deposits on the instruments.
- 6. Allow the cleaned areas to dry completely. This may take a few minutes.
- 7. Finally, wipe the cleaned surfaces with a dry, non-fuzzing microfibre cloth.

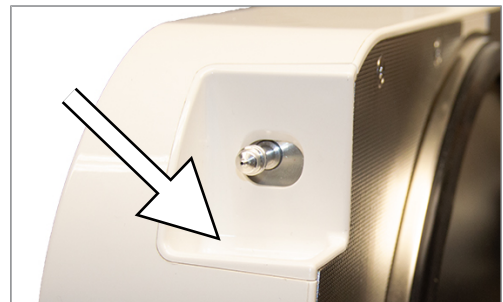
Housing parts

Where necessary, clean the housing parts with a neutral fluid cleaner or spirit.

Floor trough - chuck care system

The following must be fulfilled or present:

- ✓ A dry and non-fuzzing cloth.
- ▶ Wipe out the floor trough with a cloth.



Clean the feed water tank

Drain the feed water tank

The following must be fulfilled or present:

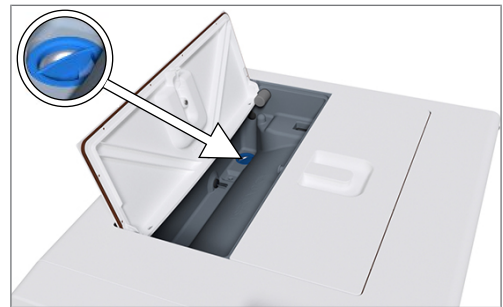
- ✓ A drain hose (included in the scope of delivery).
- ✓ A collection container (with a capacity of up to 3 l, normally 150 ml flows out).
- ✓ The device is not in operation. The device should cool down for approx. 15 min after switching off.
- 1. Switch off the device.
- 2. Open the service hatch.
- 3. Place the collection container in front of the device and the end of the drain hose in the collection container.
- 4. Push the drain hose onto the drain valve until it noticeably snaps into place. The knob must be horizontal.

5. Open the drain valve by turning the knob 1/4 in an anti-clockwise direction.
6. Drain the water into the collection container.
7. To remove the drain hose, turn the drain valve into the vertical position again.
8. Close the service flap.



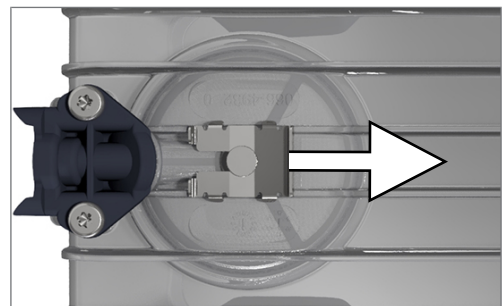
Clean the feed water tank

1. Open the cover cap on the top of the device.
2. Put the cover aside.
3. Check the tank for contamination and, if necessary, clean it with a sponge and solvent-free, non-alkaline cleaner (e.g. washing-up liquid).
4. Remove the tank filter from the base of the feed water tank by pulling it out.
5. Clean the tank filter under running water or with the MELAjet spray pistol for MELAdem 40.
6. Check the cleaning result against the light.
7. Reinsert the tank filter.
8. Please reinsert the cover cap correctly and close it.

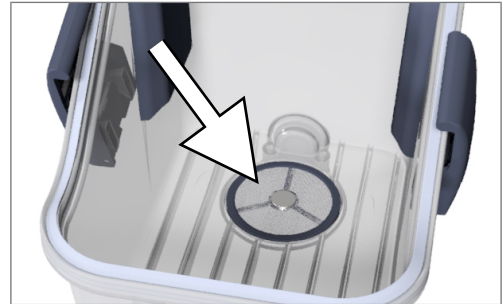


Clean Carebox

1. On the underside of the Carebox, release the fixing clamp of the Carebox sieve by pulling it out in the direction of the arrow.



2. Press against the pins from below and remove the sieve from inside the Carebox base.



3. Clean the sieve under running water with a soft brush or with the MELAjet spray gun for MELAdem 40.
4. Check the cleaning result against the light.
5. Reinsert the sieve.
6. Reinsert the fixing clamp.

Carebox

Changing the filter

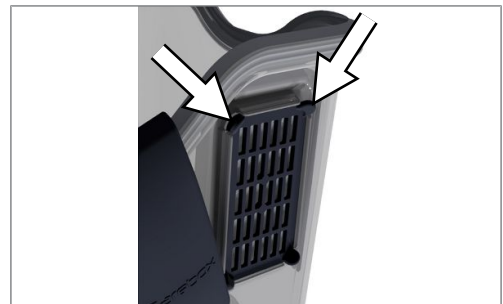


PLEASE NOTE

After changing the oil can, carry out a filter change on all Careboxes.

The following must be fulfilled or present:

- ✓ Two new filters.
1. Perform a detailed visual inspection of the new filters to check for damage, e.g. holes.
 2. Remove the cover of the filter by pulling on the two upper plastic tabs.



3. Remove the filter using one of the plastic tabs on the cover.

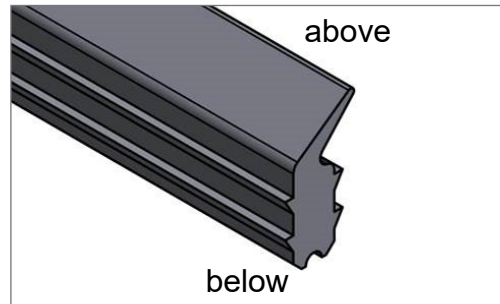


4. Insert the new filter.
5. Replace the cover by placing it at the bottom first.
6. Change the filter on the opposite side.

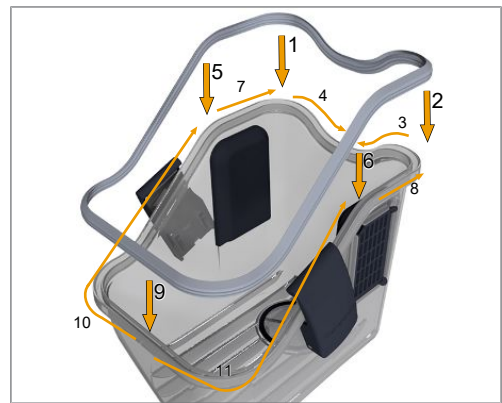
Replace the Carebox housing seal

The following must be fulfilled or present:

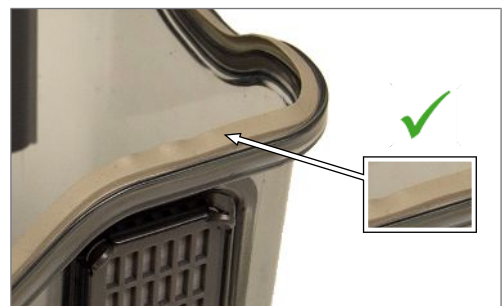
- ✓ A new Carebox housing seal.
- 1. Pull out the housing seal upwards using the fingers or with the help of tweezers.
- 2. When inserting the housing seal, note the correct orientation of the curvature.



- 3. Insert the new housing seal by hand. Fix it in the sealing groove in the order shown. Start with the upper part and press the edges going downwards into the sealing groove at the end.



- 4. **NOTICE! Do not pull on the housing seal.** Press the housing seal into the sealing groove circumferentially.
- 5. Check the housing seal for correct position.



- The sealing lip may have small wave after insertion. However, these waves should be avoided to avoid leakage that may occur.

Changing the media seals

The following must be fulfilled or present:

- ✓ A new set of media seals (five small, two large).
- 1. Remove the six media seals (1-6) on the rear side of the Carebox upper section, e.g. using tweezers.



- 2. Remove the media seal (7) on the rear side of the Carebox lower section, e.g. using tweezers.
- 3. Insert the new media seals by pressing them in.
- 4. Check the media seals for correct position.

Change adapter seals



PLEASE NOTE

Perform a detailed visual inspection of the new adapter seal to check for damage, e.g. cracks.

The following must be fulfilled or present:

- ✓ A new adapter seal (O-ring).
- ✓ If necessary, use aids such as a pointed object.
- 1. Remove the damaged adapter seal by disconnecting it or cutting it open.
Do not damage the adapter in the process.



- 2. Insert the new adapter seal.



3. Check the adapter seal for correct position.

See also:

- [Adapter installation](#) [► page 24]

Maintenance



NOTICE

Continuing operation beyond the maintenance interval can result in malfunctions in the device.

- Maintenance should only be performed by trained and authorized service technicians or stockist technicians.
- Maintain the specified servicing intervals.

Regular maintenance is vital to ensure reliable operation and value retention of the device. All function and safety-relevant components and electrical units must be checked during maintenance and replaced where necessary. Maintenance must be performed in accordance with the pertinent maintenance instructions of the device.

Maintenance work is to be performed regularly after 3000 program cycles but must be performed after 24 months. The steam sterilizer will issue a maintenance message at the relevant time.

14 Pause times

Duration of the operating pause

Duration of the operating pause	Measure
Short pauses between two sterilization processes	▪ Keep the door closed to save energy ▪ Set Energy-saving as required
Pauses which last longer than an hour	▪ Switching off device
Longer pauses e.g. over night or the weekend	▪ Leave the door ajar to prevent premature wear and the sticking of the door seal ▪ Switching off device ▪ If present, shut off the water inflow of the water treatment unit
Longer than two weeks	▪ Perform the Draining service program ▪ Perform a Vacuum test. ▪ After a successful vacuum test, perform an empty sterilization in a reprocessing program

See also:

- [Energy-saving](#) [► page 59]
- [Service programs](#) [► page 54]

Starting after long pauses

After a long period of non-use, the device will heat up after being switched on.

Decommissioning

When decommissioning the device for a long pause (e.g. due to holiday), proceed as follows:

1. Empty the double jacket steam generator, see [Emptying](#) [► page 77].
2. Switch on the device at the power switch.
3. Disconnect the power plug from the socket and if necessary, allow the device to cool.
4. Empty the internal storage tank via the drain hose.
5. Shut off if present, the water inflow of the water treatment unit.

Emptying

You have the option of emptying the water in the double jacket steam generator easily via the Draining program. In order to do so, the device is heated once, building up pressure in the double jacket so that the water can be emptied fully from the double jacket steam generator.

1. Working in the **Service programs** menu, select the **Draining** program and press **Start**.
2. Confirm the dialogue window.
 - ➡ The double jacket steam generator is emptied.
3. Confirm the message **Draining successful**.
 - ➡ The device switches to door mode.
4. Switch off the device.

Transport



CAUTION

Danger of injury from incorrect carrying.

Lifting and carrying too heavy a load can result in spinal injury. Failure to comply with these provisions can result in crushing.

- The device should always be carried by two people.
 - Use the correct carrying straps to carry the device.
-

On-site transport

To transport the device within a room or floor, proceed as follows:

1. Decommission the device, see: [Decommissioning](#) [► page 77].
2. Disconnect the connection hoses connected on the rear of the device.
3. Install the carrying aid.

Off-site transport

To transport the device over longer distances, to different floors or for shipping, proceed as follows:

1. Decommission the device, see: [Decommissioning](#) [► page 77].
2. Pack the device so that it is protected from mechanical hazards (e.g. blows) and moisture.
3. Observe the transport and storage conditions, see: [Technical data](#) [► page 92].

15 Malfunctions

Not all notifications on the display are malfunction messages. Warnings and malfunction messages are issued on the display with an event number. This number serves identification purposes.

	Nature of the display notification	Description
	Messages	Many messages are notifications. They serve to inform and provide assistance in the operation of the device.
	Warnings	Warning messages are displayed when necessary. Warning messages contain instructions that help to ensure smooth operation and to recognise undesirable conditions. Comply with these warnings early in order to avoid malfunctions.
	Malfunction messages	Malfunction messages are issued when it is not possible to ensure safe operation or safety of sterilization. These can appear on the display shortly after activating the steam sterilizer or during a program run. If a malfunction occurs during a program run, the program will be aborted.

Troubleshooting online

All messages with current descriptions can be found in the Troubleshooting portal on the MELAG website (<https://www.melag.com/en/service/troubleshooting>).



Before contacting customer services

Ensure that you have complied with all instructions relating to a warning or malfunction message issued on the display of the steam sterilizer. The following table contains a summary of the most important events. Should you be unable to find the relevant event in the table below, or your efforts do not redress the problem, you can contact your nearest stockist or an authorized MELAG customer service provider. To enable us to give the best possible service, please have your steam sterilizer serial number and a detailed description of the malfunction to hand.

Malfunction logs

In the **Logs > Malfunction logs** menu, you can view malfunction logs and output them to a USB stick.

See also:

- [Logs menu](#) [▶ page 49]
- [Log list](#) [▶ page 50]

General events

The following tables indicate possible causes for certain events and the corresponding operating information for their remedy. Should you be unable to find the relevant event, or your efforts do not redress the problem, you can contact your nearest stockist or authorised MELAG customer service provider.

Event	Possible cause	What you can do
Poor cleaning outcome	Encrusted soiling on the instruments.	Do not allow soiling to dry on. Rinse off soiling immediately. Immediately make a program selection and start Careclave after inserting a Carebox.
Excessive care oil in the Carebox	Unnecessary oil leakage at unused adapters	Load all adapters with instruments or replace unused adapters with dummy adapters as standard.

Event	Possible cause	What you can do
Insufficient drying in the program Care-Therm	Unsuitable loading of the Carebox Green	Activate Add. drying. Use the program Care-S instead of the program Care-Therm.

Warning and malfunction messages

Event	Possible cause	What you can do
10026	The oil can is empty.	Please replace the oil can. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
10062	The lack of water in the fresh water tank could not be remedied within the monitoring time.	Ensure water supply (main valve) or fill in case of supply from canisters
10063	The manual feed water supply is activated. The device must be filled with at least 1.5 l of demineralised water.	Please supply the device with sufficient demineralised water before starting the program or ensure an automatic water supply via a water treatment unit. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
10082	In the program start, the counter for the feed quantity is evaluated. The limit value is exceeded. Therefore, starting the program is not possible.	The automatic emptying was skipped several times because the drying was ended manually. Starting the program is not possible before the "Empty double jacket" service program has been carried out. Please start the "Empty double jacket" service program. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
10093	In the program start, the counter for the rinse value is evaluated. The limit value is exceeded. Therefore, starting the program is not possible.	The automatic emptying was skipped several times because the drying was ended manually. Starting the program is not possible before the "Empty double jacket" service program has been carried out. Please start the "Empty double jacket" service program. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
10094	In the program start, the system checks whether the vacuum test can be performed successfully at the current ambient temperature.	The ambient temperature of the device is very high. Allow the device to cool down. Observe the set-up conditions. Ensure that the device is sufficiently ventilated. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
10098	A failure of the supply voltage was detected in the program run.	Connect the device to a specially fused power supply to which no other electrical device is connected. Check the power cable on the rear of the device to ensure that it is firmly seated. Attach the safety latch.
10099	A failure of the supply voltage was detected in the program run of a service program.	Connect the device to a specially fused power supply to which no other electrical device is connected. Check the power cable on the rear of the device to ensure that it is firmly seated. Attach the safety latch.

Event	Possible cause	What you can do
10101	The float switch (S13) in the overflow funnel detects a short-term impermissible water level, which indicates a blockage in the wastewater system.	Short-term wastewater blockage - please check wastewater hose for kinks or, if applicable, a closed shut-off. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
10102	The float switch (S13) in the overflow funnel detects a permanently impermissible water level, which indicates a blockage in the wastewater system.	Permanent wastewater blockage - please check wastewater hose for kinks or, if applicable, a closed shut-off.
10104	Contact Carebox K4. The status of the Carebox (inserted/not inserted) changes when the door is closed.	The Carebox is detected by a magnetic switch on the door of the Carebox. If you sterilize magnetic loads in the Carebox, please place them at the back of the chamber so as not to interfere with the magnetic detection.
10109	Door process. The limit current for the door motor was exceeded during opening. Apparently, the door locking mechanism or the door motor is blocked.	The automatic opening of the door is disturbed. Please allow the device to cool down and open the door using the tool in the media board. Please ensure regular oil maintenance of the door spindle and door nut. If this occurs repeatedly, please contact the technical service.
10117	Door process. The monitoring time when opening the door has expired and both the door contact switch K1 and K2 signal a closed door.	The automatic opening of the door is disturbed. Please allow the device to cool down and open the door using the tool in the media board. Please ensure regular oil maintenance of the door spindle and door nut. If this occurs repeatedly, please contact the technical service.
10120	Door process. The limit current for the door motor is exceeded when opening from the pressure-tight status (Z4) to the vapour-tight status (Z3). Apparently, there is a blockage of the door.	The automatic opening of the door is disturbed. Please allow the device to cool down and open the door using the tool in the media board. Please ensure regular oil maintenance of the door spindle and door nut. If this occurs repeatedly, please contact the technical service.
10130	Double jacket feeding. The maximum feed quantity or feed duration when feeding fresh water into the double jacket has been exceeded.	Remove and clean the filter in the fresh water tank.
10134	Vacuum system cooling. The temperature at the cooler cannot be reduced sufficiently within the monitoring time. Apparently, the cooling system has a fault.	Please allow the device to cool down. Please observe the set-up conditions. Please ensure sufficient ventilation of the device.
10137	Evacuation/test during vacuum test. The maximum permissible pressure was exceeded in the waiting or test phase of the vacuum test.	Please allow the device to cool down. Check the door seal for visible defects. Please clean the door seal with a damp cloth.

Event	Possible cause	What you can do
10145	Monitoring time only runs when evacuation is started in negative pressure. Malfunction is triggered if evacuation cannot be completed within the monitoring time.	Please check the dust filter for contamination and replace it if necessary. The intake area of the cooling system under the device must be free; check for paper or similar items underneath the device which could stop the air flow. Please ensure that the device is sufficiently ventilated. Heat dissipation must be able to take place freely, ensure ventilation; conversion cabinets are not recommended. Please observe the set-up conditions (e.g. ambient temperature). Check the loading of the device for compliance with the permissible loading quantities. Check the pressure release filter in the chamber for blockage.
10165	Double jacket emptying. The maximum runtime for emptying has expired.	Please allow the device to cool down. Please observe the set-up conditions. Please ensure sufficient ventilation of the device. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
10168	Abort routine. The Carebox could not be emptied successfully during the abort routine.	The Carebox tank could not be emptied. CAUTION! There may be hot water in the Carebox. Please clean the Carebox daily under running water. Please clean the sieve in the lower section of the Carebox. If this occurs repeatedly despite cleaning, please contact the technical service. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
10169	Please contact technical services	Please contact technical services. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
10179	Chamber pressure release. The pressure release valve had to be opened several times before a pressure drop occurred.	Check the chamber to see if any debris from loading or packing is clogging the fittings. Remove and check the coarse filter at the back bottom of the chamber for blockages. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
10187	Circulation emptying. The maximum permissible duration of the circulation emptying process was exceeded without the Carebox being emptied.	Check the sieve in the lower section of the Carebox and clean if necessary. Check the pressure drain line on the back of the device (hatched drop) for kink-free installation.
10194	Start-up time of the circulation pump. The intended circulation pressure (S2) for pulsed circulation is not reached. There may be blockage in the circulation circuit.	Please check the seals of the Carebox and clean the Carebox sieve.
10195	Pre-cleaning test. The temperature of the fresh water is too high. Pre-cleaning is not possible under these conditions.	The temperature of the incoming water is too high for a cold wash; ensure compliance with the installation requirements.

Event	Possible cause	What you can do
10196	Care. The dosing chamber for the care oil could not be emptied within the blow-out limit time.	Please check the compressed air supply (compressor, main valve). Please check the media seals of the Carebox.
10198	Care. The dosing chamber for the care oil could not be blown out within the runtime.	Please check the compressed air supply (compressor, main valve). Please check the media seals of the Carebox.
10199	Care. The dosing of the care oil has failed.	The oil can is completely empty. Please insert a new oil can. Replace the media separation filter on the side of the Carebox at the same time.
10200	Carebox pressure reduction. The pressure reduction in the Carebox takes too long.	Check the sieve in the lower section of the Carebox and clean if necessary. Check condition of media seals and replace if necessary. Check the pressure drain line on the back of the device (hatched drop) for kink-free installation.
10201	Final cleaning. The maximum time for heating up the rinse liquor has expired.	Make sure that at least one instrument has spray channels as these are subjected to flow for heating. Alternatively, an adapter slot can also be left empty to enable heating.
10202	Wash liquor heating. The temperature increase is smaller than expected. The wash liquor is being heated up too slowly.	Make sure that at least one instrument has spray channels as these are subjected to flow for heating. Alternatively, an adapter slot can also be left empty to enable heating. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
10203	Carebox water which is detected by a pressure increase in the chamber. As this is a warm wash, the protein load of the wash liquor is already so low that only a warning message is issued at this point.	Please make sure that the Carebox is correctly seated. If this occurs repeatedly, replace the small seals on the back of the Carebox. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
10204	Final cleaning. The maximum time that the limit pressure in the chamber may be exceeded during the heating of the rinse liquor has expired.	Check condition of media seals and replace if necessary. Check the status of the Carebox housing seal and replace if necessary. Check installation of splash guard and media separation filter (PTFE) on the sides of the Carebox base and correct if necessary. Check the pressure drain line on the back of the device (hatched drop) for kink-free installation. Check the pressure release filter in the chamber for blockage.
10207	Pre-cleaning. The maximum permitted temperature for pre-cleaning has been exceeded; pre-cleaning is not possible.	The Carebox must cool down between cycles. Use the Cooling Box. If continuous operation is desired, Carebox can be operated with two or more Careboxes in alternation. A pause time of 4 min between cycles must be observed.
10208	External cleaning. The circulation pressure when circulating the rinse liquor is too low.	Please check the Carebox seals and clean the Carebox sieve.
10212	Spray channel cleaning. A leakage of the Carebox was detected during the cleaning of the spray channels.	Please make sure that the Carebox is correctly seated. If this occurs repeatedly, replace the small seals on the back of the Carebox.

Event	Possible cause	What you can do
10213	Drive channel cleaning. A leakage of the Carebox was detected during the cleaning of the drive channels.	Please make sure that the Carebox is correctly seated. If this occurs repeatedly, replace the small seals on the back of the Carebox.
10214	Reverse feeding. A leakage of the Carebox was detected during reverse feeding.	Please make sure that the Carebox is correctly seated. If this occurs repeatedly, replace the large seals on the back of the Carebox lower section.
10215	External cleaning. A leakage of the Carebox was detected during external cleaning.	Please check the condition of the media seal of the Carebox and replace it if necessary. Please check the condition of the Carebox housing seal and replace it if necessary. Please check the chamber for contamination. If this occurs repeatedly, replace the seals on the back of the Carebox.
10216	Steam inlet drive channels. The limit time for the pressure build-up at the steam inlet drive channels has been exceeded.	Make sure that at least one instrument has drive channels as these are subjected to flow for heating. Alternatively, an adapter slot can also be left empty to enable heating.
10217	Steam inlet spray channels. The limit time for the pressure build-up at the steam inlet spray channels has been exceeded.	Make sure that at least one instrument has spray channels as these are subjected to flow for heating. Alternatively, an adapter slot can also be left empty to enable heating.
10228	Door process. Closing the door of the Carebox is only possible when an oil can is inserted.	Please insert an oil can. If you do not have a full oil can available, you must insert the (possibly empty) oil can that has been removed before you can close the door. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
10230	The pneumatic Carebox detection delivers illogical values. Although a Carebox is inserted (contact switch), the pressure increase suggests a missing Carebox.	The Carebox must be dry on the outside before it is inserted in the Carebox; falling drops can cause an excessively high pressure increase. Please check the media seals of the Carebox.
10231	The pneumatic Carebox detection delivers illogical values. Although a Carebox is inserted (contact switch), the pressure increase suggests a missing Carebox. There may be very impermeable instruments at positions 5 and 8 or at positions 6 and 7.	The Carebox must be dry on the outside before it is inserted in the Carebox. Falling drops can cause an excessively high pressure increase. Please check the media seals of the Carebox. Please change the instrument position in the Carebox. Insert an instrument from position 5 or 8 into position 1 to 4 and an instrument from position 6 or 7 into position 1 to 4.
10233	The pneumatic Carebox detection delivers illogical values. Although no Carebox is inserted (contact switch), the pressure increase suggests a Carebox is inserted.	Please check the compressed air supply (compressor, main valve). Please check the media seals of the Carebox.
10234	The continuity check of the Carebox delivers impermissible values.	Flow through the Carebox is not possible. Please check the compressed air supply as well as the instruments and adapters for free passage. Run the Carebox test service program.
10235	The continuity check of the Carebox delivers impermissible values. No pressure increase is detected when flow passes through V20.	Flow through the Carebox is not possible. Please check the compressed air supply as well as instruments and adapters at position 5 and 8. Run the Carebox test service program.
10236	The continuity check of the Carebox delivers impermissible values. No pressure increase is detected when flow passes through V16.	Flow through the Carebox is not possible. Please check the compressed air supply as well as instruments and adapters at position 6 and 7. Run the Carebox test service program.

Event	Possible cause	What you can do
10237	The continuity check of the Carebox delivers impermissible values. No pressure increase is detected when flow passes through V19.	Flow through the Carebox is not possible. Please check the compressed air supply as well as instruments and adapters at position 2 and 3. Run the Carebox test service program.
10238	The continuity check of the Carebox delivers impermissible values. No pressure increase is detected when flow passes through V18.	Flow through the Carebox is not possible. Please check the compressed air supply as well as instruments and adapters at position 1 and 4. Run the Carebox test service program.
10239	The continuity check of the Carebox delivers impermissible values. No pressure increase is detected when flow passes through V22.	Make sure that at least one instrument has spray channels as these are subjected to flow for heating. Alternatively, an adapter slot can also be left empty to enable heating.
10241	Pressure-controlled evacuation. This is triggered when the termination gradient is violated in a gradient-monitored process, which leads to an abort of the program run (in the context of pressure-controlled evacuation/pressure gradient monitoring). The vacuum performance is insufficient.	Please check the dust filter for contamination and replace it if necessary. The intake area of the cooling system under the device must be free; check for paper or similar items underneath the device which could stop the air flow. Please ensure that the device is sufficiently ventilated. Heat dissipation must be able to take place freely, ensure ventilation; conversion cabinets are not recommended. Please observe the set-up conditions (e.g. ambient temperature). Check the loading of the device for compliance with the permissible loading quantities. Check the pressure release filter in the chamber for blockage.
10242	This is triggered when the termination gradient is violated in a gradient-monitored process, which leads to an abort of the program run (in the context of clock-controlled evacuation VT/pressure gradient monitoring). The vacuum performance is insufficient.	Please check the dust filter for contamination and replace it if necessary. The intake area of the cooling system under the device must be free; check for paper or similar items underneath the device which could stop the air flow. Please ensure that the device is sufficiently ventilated. Heat dissipation must be able to take place freely, ensure ventilation; conversion cabinets are not recommended. Please observe the set-up conditions (e.g. ambient temperature). Check the loading of the device for compliance with the permissible loading quantities. Check the pressure release filter in the chamber for blockage.
10253	This is triggered when the Carebox is hooked in and the door is closed without starting a program or performing other activities on the GUI. The aim is to prevent a contaminated box from being stored in the warm device, which would lead to unnecessary drying of the contamination.	Remove the Carebox or start a reprocessing program. This is only a warning. The reprocessing result is not affected. You can continue to use the device.

Event	Possible cause	What you can do
10256	Monitoring pressure gradient during evacuation. The pressure change at pressure sensor S1 is too low during evacuation.	Please check the dust filter for contamination and replace it if necessary. The intake area of the cooling system under the device must be free; check for paper or similar items underneath the device which could stop the air flow. Please ensure that the device is sufficiently ventilated. Heat dissipation must be able to take place freely, ensure ventilation; conversion cabinets are not recommended. Please observe the set-up conditions (e.g. ambient temperature). Check the loading of the device for compliance with the permissible loading quantities. Check the pressure release filter in the chamber for blockage.
10257	Monitoring pressure gradient during evacuation in the vacuum test. The pressure change at pressure sensor S1 is too low during evacuation.	Please check the dust filter for contamination and replace it if necessary. The intake area of the cooling system under the device must be free; check for paper or similar items underneath the device which could stop the air flow. Please ensure that the device is sufficiently ventilated. Heat dissipation must be able to take place freely, ensure ventilation; conversion cabinets are not recommended. Please observe the set-up conditions (e.g. ambient temperature). Check the loading of the device for compliance with the permissible loading quantities. Check the pressure release filter in the chamber for blockage.
10262	Chamber support pressure. The support pressure could not be built up within the specified time.	Please check the compressed air supply (compressor, main valve). Please check the media seals of the Carebox.
10263	Monitoring pressure gradient during pressure release of General Carebox detection. The pressure change at pressure sensor S1 is too low during pressure release.	The Carebox must be dry on the outside before it is inserted in the Carebox; falling drops can cause an excessively high pressure increase. There should be no puddles in the chamber, e.g. from a previous program stop. If necessary, remove residual moisture from the chamber or allow it to evaporate. Check the pressure drain line on the back of the device (hatched drop) for kink-free installation. Check the pressure release filter in the chamber for blockage.
10264	Monitoring pressure gradient during pressure release after continuity check. The pressure change at pressure sensor S1 is too low during pressure release.	The Carebox must be dry on the outside before it is inserted in the Carebox; falling drops can cause an excessively high pressure increase. Check the pressure drain line on the back of the device (hatched drop) for kink-free installation. Check the pressure release filter in the chamber for blockage.

Event	Possible cause	What you can do
10266	Pressure-controlled evacuation. The pressure change is less than expected; the vacuum performance is decreasing.	Please check the dust filter for contamination and replace it if necessary. The intake area of the cooling system under the device must be free; check for paper or similar items underneath the device which could stop the air flow. Please ensure that the device is sufficiently ventilated. Heat dissipation must be able to take place freely, ensure ventilation; conversion cabinets are not recommended. Please observe the set-up conditions (e.g. ambient temperature). Check the loading of the device for compliance with the permissible loading quantities. Check the pressure release filter in the chamber for blockage. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
10267	Clock-controlled evacuation. The pressure change is less than expected; the vacuum performance is decreasing.	Please check the dust filter for contamination and replace it if necessary. The intake area of the cooling system under the device must be free; check for paper or similar items underneath the device which could stop the air flow. Please ensure that the device is sufficiently ventilated. Heat dissipation must be able to take place freely, ensure ventilation; conversion cabinets are not recommended. Please observe the set-up conditions (e.g. ambient temperature). Check the loading of the device for compliance with the permissible loading quantities. Check the pressure release filter in the chamber for blockage. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
10268	Steam intake. The pressure change is less than expected; the steam intake performance is decreasing.	Check the loading of the device for compliance with the loading quantities. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
10269	Ventilation. The volume flow during ventilation is lower than expected.	Check the sterile air filter in the media board; if it is heavily worn or blocked, replace it. This is only a warning. The reprocessing result is not affected. You can continue to use the device.

Event	Possible cause	What you can do
10270	Pressure release. The pressure change is less than expected; the pressure release speed is decreasing.	Please check the dust filter for contamination and replace it if necessary. The intake area of the cooling system under the device must be free; check for paper or similar items underneath the device which could stop the air flow. Please ensure that the device is sufficiently ventilated. Heat dissipation must be able to take place freely, ensure ventilation; conversion cabinets are not recommended. Please observe the set-up conditions (e.g. ambient temperature). Check the loading of the device for compliance with the permissible loading quantities. Check the pressure release filter in the chamber for blockage. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
10271	Conductivity monitoring. The warning value for poor conductivity was exceeded in the program start. A program start is still possible	Have a regenerated cartridge ready for your water treatment unit. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
10272	Conductivity monitoring. The warning value for insufficient conductivity was exceeded in the cleaning. A program start is still possible	Have a regenerated cartridge ready for your water treatment unit. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
10273	Conductivity monitoring. The limit value for insufficient conductivity was exceeded in the program start. A program start is not possible	Ensure the supply of DI water of suitable quality. Insert a regenerated cartridge into your water treatment system.
10274	Conductivity monitoring. The limit value for insufficient conductivity was exceeded in the cleaning. A program start is not possible	Ensure the supply of DI water of suitable quality. Insert a regenerated cartridge into your water treatment system.
10275	This is triggered when the measuring turbine (S9) of the feed pump (P1) indicates that the volume flow is too low.	Please remove and clean the filter in the storage tank.
10276	Reverse feeding. The measuring turbine (S9) of the feed pump (P1) indicates that the volume flow is too low.	Please remove and clean the filter in the storage tank.
10277	Program start. The filling level of the oil can is low, so it can be assumed that an oil care operation can no longer be performed successfully in the foreseeable future (~20 ml).	Please have a can of MELAG Care Oil ready for replacement. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
10278	The manual feed water supply is activated. The device must be filled with at least 1.5 l of demineralised water.	Please supply the device with sufficient demineralised water before starting the program or ensure an automatic water supply via a water treatment unit.

Event	Possible cause	What you can do
10279	Carebox pressure release. The pressure change is less than expected; the pressure release speed is decreasing.	Please clean the Carebox daily under running water. Please clean the sieve in the lower section of the Carebox. If this occurs repeatedly despite cleaning, please contact the technical service. Wastewater blockage - please check wastewater hose for kinks or, if applicable, a closed shut-off. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
10280	Monitoring pressure gradient during pressure release of General Carebox detection. The pressure change at pressure sensor S1 is too low during pressure release.	The Carebox must be dry on the outside before it is inserted in the Carebox; falling drops can cause an excessively high pressure increase. There should be no puddles in the chamber, e.g. from a previous program stop. If necessary, remove residual moisture from the chamber or allow it to evaporate. Check the pressure drain line on the back of the device (hatched drop) for kink-free installation. Check the pressure release filter in the chamber for blockage.
10281	Monitoring pressure gradient during pressure release. The pressure change at pressure sensor S1 is too low during pressure release.	Please check the dust filter for contamination and replace it if necessary. The intake area of the cooling system under the device must be free; check for paper or similar items underneath the device which could stop the air flow. Please ensure that the device is sufficiently ventilated. Heat dissipation must be able to take place freely, ensure ventilation; conversion cabinets are not recommended. Please observe the set-up conditions (e.g. ambient temperature). Check the loading of the device for compliance with the permissible loading quantities. Check the pressure release filter in the chamber for blockage.
10282	Monitoring pressure gradient during pressure release of General Carebox detection. The pressure change at pressure sensor S1 is too low during pressure release.	The Carebox must be dry on the outside before it is inserted in the Carebox; falling drops can cause an excessively high pressure increase. There should be no puddles in the chamber, e.g. from a previous program stop. If necessary, remove residual moisture from the chamber or allow it to evaporate. Check the pressure drain line on the back of the device (hatched drop) for kink-free installation. Check the pressure release filter in the chamber for blockage.

Event	Possible cause	What you can do
10283	Pressure-controlled evacuation. This is triggered when the termination gradient is violated in a gradient-monitored process, which leads to an abort of the program run (in the context of pressure-controlled evacuation/pressure gradient monitoring). The vacuum performance is insufficient.	Please check the dust filter for contamination and replace it if necessary. The intake area of the cooling system under the device must be free; check for paper or similar items underneath the device which could stop the air flow. Please ensure that the device is sufficiently ventilated. Heat dissipation must be able to take place freely, ensure ventilation; conversion cabinets are not recommended. Please observe the set-up conditions (e.g. ambient temperature). Check the loading of the device for compliance with the permissible loading quantities. Check the pressure release filter in the chamber for blockage.
10286	Pressure-controlled evacuation. This is triggered when the termination gradient is violated in a gradient-monitored process, which leads to an abort of the program run (in the context of pressure-controlled evacuation/pressure gradient monitoring). The vacuum performance is insufficient.	Please check the dust filter for contamination and replace it if necessary. The intake area of the cooling system under the device must be free; check for paper or similar items underneath the device which could stop the air flow. Please ensure that the device is sufficiently ventilated. Heat dissipation must be able to take place freely, ensure ventilation; conversion cabinets are not recommended. Please observe the set-up conditions (e.g. ambient temperature). Check the loading of the device for compliance with the permissible loading quantities. Check the pressure release filter in the chamber for blockage.
11000	The log output was aborted due to a connection error.	Please check the connection of the device to the practice network via the network interface on the back of the device. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
11001	Several USB sticks are directly connected to the device	Please connect only one USB stick to the device. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
11002	The USB storage medium is not plugged in although write access to the USB stick has been requested.	Please insert the USB stick in the media board. If necessary, use the USB socket on the rear of the device. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
11003	The USB storage medium does not have enough free space to store the required log data.	Please save the log data on the USB stick in the practice network and then empty the USB stick to output the new logs. This is only a warning. The reprocessing result is not affected. You can continue to use the device.

Event	Possible cause	What you can do
11004	The writing of the log data to the USB storage medium has failed.	Please insert the USB stick in the media board. If necessary, use the USB socket on the rear of the device. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
11006	The maximum number of program logs not output was reached. The oldest log will be overwritten the next time the program is run.	Please output the internally stored logs to a USB stick or to your practice network. The log output can also be done automatically; this must be configured in the settings menu. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
11007	The printer cover is open while a print job was being sent.	Please close the printer cover. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
11009	A printer is configured but not connected.	Please connect the printer via the network interface on the back of the device. Please restart the printer. First start the device and then the printer. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
11011	Several printers are directly connected to the device	Please connect only one printer to the device. Please restart the printer. First start the device and then the printer. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
11012	The paper of the printer will be used up soon.	Please have a new roll ready. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
11013	General printer error	Please restart the printer. First start the device and then the printer. This is only a warning. The reprocessing result is not affected. You can continue to use the device.
11100	The log output was aborted due to a connection error.	Please check the connection of the device to the practice network via the network interface on the back of the device. This is only a warning. The reprocessing result is not affected. You can continue to use the device.

16 Technical data

Device type	Careclave 618
Device dimensions (H x W x D)	56.2 x 48.0 x 65.3 cm
Empty weight	69 kg
Operating weight	82.5 kg
Sterilization chamber	
Chamber diameter	250 mm
Chamber depth	350 mm
Chamber volume/steam generator	17.8 l / 4.4 l
Carebox	
Volume Carebox	2.4 l
Electrical connection	
Power supply	220-240 V 50 Hz
Max. voltage range	Fluctuations of the mains supply voltage up to $\pm 10\%$ of the nominal voltage
Max. power consumption in operation	3000 W
Max. power consumption in idle mode	0.5 W
Building fuse protection	16 A, RCD with rated residual current = 30 mA
Length of the power cable	2 m
Overvoltage category	transient overvoltage up to the values of overvoltage category II
Degree of contamination (in accordance with EN 61010)	2
Ambient conditions	
Installation location	interior of a building
Noise emission LP(a) in 1 m distance	70 dB(A)
Waste heat (with maximum load)	1.7 kW
Ambient temperature	5-40 °C (ideal range 16-26 °C)
Relative humidity	max. 80 % at temperatures of up to 30 °C, max. 50% at 40 °C (decreasing in linear fashion in-between)
Degree of protection (in accordance with IEC 60529)	IP20
Transport and storage conditions	Temperature: -5 to +40 °C, air humidity: < 80%
Max. altitude	2000 m
Feed water	
Max. water consumption	5 l/h
Max. water consumption in the Care-Therm program	1.0 l
Max. water consumption in the Care-S program	1.2 l
Max. water consumption in the Care-B program	1.3 l
Average water consumption	2.5 l/h
Water quality (feed water)	distilled or demineralized water in accordance with EN 13060, Appendix C
Water temperature	1 °C - 35 °C
Cold water (for connection of the water treatment unit)	
Min. static water pressure (WTU) ²⁾	2 bar
Max static water pressure (WTU)	10 bar
Water quality (water treatment unit)	potable water

²⁾ When using a MELAdem 47, the minimum static water pressure is 3.0 bar. If the static water pressure is less than 3.0 bar, the booster pump (art. no. ME22500) must be used on the MELAdem 47.

Compressed air

Min. pressure	4 bar relative
Max. pressure	8 bar relative
Min. compressed air supply	55 NI/min
Average compressed air supply	50 NI/cycle
Compressed air quality	dried, condensate-free, bacteria-free, oil-free and filtered (filter fineness $\leq 2 \mu\text{m}$)

Wastewater

Max. throughflow volume (wastewater)	2 l/min
Max water temperature (wastewater)	90 °C for 30 seconds, maximum 98 °C for 1 second

Working and operating pressures

Permissible operating pressure chamber	-1 bar to + 3 bar relative
Permissible operating pressure jacket	-1 bar to + 3 bar relative
Working pressure chamber/jacket	2.2 bar relative

17 Accessories and spare parts

You can obtain the specified articles and an overview of further accessories from your stockist.

Accessories for the device

Category	Article	Art. no.
Mounts	Mount C "Plus" for 6 trays or 3 MELAstore Boxes 100	ME81370
	Mount D "Plus" for 2 MELAstore Boxes 200 or 2 MELAstore Boxes 100 and 2 narrow trays	ME82640
	Mount E "Plus" for 6 trays (standard) and 2 narrow trays	ME82400
	Mount F "Plus" for 3 MELAstore Boxes 100 and 2 narrow trays	ME82660
	Careclave 4+2 tray mount for 4 trays (short) and 2 narrow trays (short)	ME21778
	Careclave 6+2 tray mount for 6 trays (short) and 2 narrow trays (short)	ME22346
Trays	Tray	ME00280
	Tray, narrow	ME01320
	Tray for Careclave 4+2 and 6+2 mounts	ME21774
	Narrow tray for Careclave 4+2 and 6+2 mounts	ME21776
Package holder	Package holder	ME22410
Sterilization container with disposable paper filter according to EN 868-8	15K (18 x 12 x 4.5 cm)	ME01151
	15M (35 x 12 x 4.5 cm)	ME01152
	15G (35 x 12 x 8 cm)	ME01153
	17K (20 x 14 x 5 cm)	ME01171
	28M (32 x 16 x 6 cm)	ME01284
	28G (32 x 16 x 12 cm)	ME01285
MELAstore System	MELAstore Tray 50 (18 x 11.8 x 3 cm)	ME01180
	MELAstore Tray 100 (27.5 x 17.6 x 3 cm)	ME01181
	MELAstore Tray 200 (27.5 x 17.6 x 4.3 cm)	ME01182
	MELAstore Box 100 (31.2 x 19 x 4.6 cm)	ME01191
	MELAstore Box 200 (31.2 x 19 x 6.5 cm)	ME01192

General accessories

Category	Article	Art. no.
Test body system	MELAcontrol Helix consisting of a Helix test body and 250 indicator strips	ME01080
	MELAcontrol Pro consisting of a Helix test body and 40 indicator strips	ME01075
	MELAcontrol Pro refill pack	ME01076
Water treatment	MELAdem 47 reverse osmosis unit	ME01049
	MELAdem 53/MELAdem 53 C	ME01038/ME01036
Water supply	Filling pump P10	ME65010
	Booster pump for MELAdem 47	ME22500
Cleaning	Chamber cleaning set	ME01081

Category	Article	Art. no.
For the documentation	USB flash drive	ME19901
	MELAprint 60 label printer	ME01160
	Network cable (cross-over), 2 m	ME15813
	Network cable (cross-over), 5 m	ME15814
	Network cable (cross-over), 10 m	ME15815
Other	Water stop	ME01056
	Surface-mounted siphon	ME37410
	Carrying system	ME21121
	Cooling Box	ME11000

Carebox accessories

Category	Article	Art. no.
Carebox	Carebox Green	ME10704
Carebox	Carebox Blue	ME10708
Carebox	Table mount for Carebox upper section	ME22161
Carebox Blue	Adapter for KaVo turbines (MULTIflex)	ME02601
Carebox Blue	Adapter for Sirona turbines	ME02602
Carebox Blue	Adapter for unused connections	ME02603
Carebox Blue	Adapter for Sirona T1 Classic	ME02604
Carebox Blue	Adapter for W&H turbines (Roto Quick)	ME02605
Carebox Blue	Adapter for KaVo/BienAir WS-heads	ME02606
Carebox Blue	Adapter for ISO connection (INTRA coupling)	ME02607
Carebox Blue	Adapter for BienAir turbines	ME02608
Carebox Blue	Adapter for Midwest connection (4/5 hole)	ME02609
Carebox Blue	Adapter for exterior spray channels	ME21914
Carebox Blue	O-rings for ISO adapter (10 pcs)	ME02627
Carebox Blue	O-rings for Sirona T1 Classic adapter (5 pcs)	ME02624
Carebox Blue	O-rings for Sirona turbine adapters (9 pcs)	ME02622
Carebox Blue	O-ring set for KaVo turbine adapters (Multiflex) (set for 1 adapter)	ME02621
Carebox Blue	O-ring set for W&H turbine adapter (Roto Quick) (set for 1 adapter)	ME02625
Carebox Blue	O-rings for KaVo/BienAir head adapter (5 pcs)	ME02626
Carebox Blue	O-ring for set of BienAir turbine adapters (set for 1 adapter)	ME02628
Carebox Green	Adapter for tips (M3.0 x 0.5 mm), external thread	ME80750
Carebox Green	Adapter for tips (M3.6 x PH1.5 P0.5), internal thread	ME80751
Carebox Green	Adapter for tips (M3.0 x 0.35 mm), external thread	ME80752
Carebox Green	Adapter for tips (M3.5 x 0.35 mm), internal thread	ME80755
Carebox Green	Adapter for tips (M3.0 x 0.6 mm), external thread	ME80756
Carebox Green	Adapter for tips (M3.5 x 0.6 mm), internal thread	ME80760
Carebox Green	Adapter for tips (M3.0 x 0.5 mm), internal thread	ME80790
Carebox Green	Adapter M8x1 eccentric	ME22407

Spare parts

Category	Article	Art. no.
Careclave	Oil for door lock bushing	ME27515
Careclave	Oil can with care oil - MELAG Care Oil	ME84740
Careclave	Dust filter	ME82260

Category	Article	Art. no.
Careclave	Sterile filter	ME84790
Careclave	Feed water tank filter	ME21358
Careclave	Power cable with hot appliance plug	ME21301
Careclave	Nozzle, chuck care	ME80016
Carebox	Carebox measuring device	ME21273
Carebox	Carebox holder for the device side	ME22162
Carebox upper section	Screwdriver TX6 for adapters	ME21867
Carebox upper section	Sealing set for adapter foot	ME21328
Carebox upper section Carebox lower section	Set - Media seals for Carebox: 2x media seal - big 5x media seal - small	ME21465
Carebox lower section	Filter (12 pieces)	ME21412
Carebox lower section	Carebox sieve	ME10701
Carebox lower section	Housing seal	ME21404
Carebox lower section	Fixing clamp for the sieve	ME21692

Glossary

A0-value

The A0 value represents a standard for the elimination of microorganisms and the deactivation of viruses in the disinfection procedure with damp heat. The A0 value depends on temperature and time.

Air leakage

An air leakage is a location through which air can pass in and out without this being desired. Verification of the leakage serves to prove that the volume of air ingress in the chamber during the vacuum phase does not exceed a value which would prevent steam penetration of the sterilizer load and that the air leakage does not cause the possible contamination of the sterilizer load during the drying phase.

AKI

AKI is the abbreviation for "Arbeitskreis Instrumentenaufbereitung"

Authorised technician

The term "authorised technician" refers to an employee of a customer service provider or stockist who has been trained and authorised by MELAG to perform maintenance and installation work on MELAG devices. Only they may carry out this work.

Batch

The batch is the composition of items which has been subject to the same reprocessing procedure.

BfArM

Federal Institute for Drugs and Medical Devices (Bundesinstitut für Arzneimittel und Medizinprodukte, BfArM) in Germany.

Bowie & Dick test

Steam penetration test with a standard test package; described in EN 285; the test is recognized in the large-scale sterilization industry.

Condensate

Fluid (e.g. water) produced by the cooling of and resultant separation from the vaporous state.

Conductivity

Conductivity refers to the ability of a conductive chemical material or mixture to conduct or transfer other materials or particles.

Corrosion

The chemical alteration or destruction of metal materials by water and chemicals

Delay in boiling

This refers to the phenomenon that it is possible under certain circumstances to heat a fluid beyond its boiling point without them boiling. This represents an unstable state; even low-level agitation can produce a large bubble within the shortest period, which expands explosively.

Demineralised water

Water without the minerals usually found in normal spring or tap water; is produced through ion exchange of normal tap water. It is used here as feed water.

DGSV

Abbr.: Deutsche Gesellschaft für Sterilgutversorgung (German Society German Society for Sterile Supply). The DSGV training centres are specified in DIN 58946, part 6 as "Requirements of personnel".

DGUV Regulation 1

DGUV is the abbreviation for "Deutsche Gesetzliche Unfallversicherung" (German Statutory Accident Insurance). The regulation 1 governs the principles of prevention.

DIN 58953

Standard – sterilization, sterile equipment supply

Distilled water

From the Latin aqua destillata; also referred to as aqua dest; water which to a great extent is free from salts, organic material and micro-organisms, is produced from normal tap water or pre-cleaned water through the process of distillation (evaporation and subsequent condensation). It is used here as feed water.

Dynamic pressure test of the sterilization chamber

Serves to prove that the rate of pressure variations during a sterilization cycle does not exceed a particular value which could result in the damage of the packaging material. [see also EN 13060]

Empty chamber test

Test run without a load, performed to assess the performance of a sterilizer without the influence of a load; facilitating verification of the temperatures maintained in comparison to the temperatures set. [see EN 13060]

EN 13060

Standard – small steam sterilizers

EN 867-5

Standard – non-biological systems for use in sterilizers – part 5: The determination of indicator systems and test bodies for the performance inspection of type B and type S small sterilizers.

EN ISO 11140-1

Standard – the sterilization of products for use in medical treatment – chemical indicators – part 1: General requirements

EN ISO 11607-1

Standard - packaging for medical devices to be sterilized in the final packaging - Part 1: Requirements placed on materials, sterile barrier systems and packaging systems

EN ISO 15883

Standard - Washer-disinfectors

Evacuation

Creation of a vacuum in a vessel

Feed water

Feed water is required to produce steam for sterilization. Guide values for water quality in accordance with EN 285 / EN 13060 – Appendix C

Fractionated vacuum procedure

A technical procedure in steam sterilization; the repeated evacuation of the sterilization chamber in alternation with steam injection.

FTP

Abbr.: (File Transfer Protocol) is a data transmission procedure serving to transport data from the internet. This data can include programs, files or even information. Special FTP programs (FTP clients) serve to load the data onto a server.

Heat-up phase

The time required after the steam sterilizer has been switched on / after the start of a sterilization program, to heat the double-jacket steam generator before the sterilization procedure starts. The duration is dependent on temperature at which sterilization takes place.

Hollow body A

See product with narrow lumen

Hollow body article B

See simple hollow body

KRINKO

Abbreviation for "Kommission für Krankenhaushygiene und Infektionsprävention" (Commission for Hospital Hygiene and Infection Prevention) at the Robert Koch Institute in Germany.

Mixed loads

wrapped and unwrapped sterilization material within a single load

Multiple wrapping

e.g. wrapped instruments sealed in a double layer of film or wrapped in film and placed in an additional container or a container wrapped in textiles.

Porous

Pervious to liquids and air, e.g. textiles

Porous full load

Serves to prove that the values set on the control satisfy the necessary sterilization conditions in porous loads with a maximum mass for which the sterilizer is designed in accordance with EN 13060 [see also EN 13060].

Porous partial load

Serves to prove that the values set on the control allow steam to enter the pre-determined test package quickly and equally [see also EN 13060]

Process evaluation system

Also known as the self-monitoring system – this observes itself and compares the various sensors during a current program.

Product with narrow lumen

An article open on one side to which the following applies: $1 \leq L/D \leq 750$ and $L \leq 1500$ mm or an article with an opening on both sides which is: $2 \leq L/D \leq 1500$ and $L \leq 3000$ mm and which does not correspond to a hollow body article B; L...length of hollow body article; D...Diameter of hollow body article [see also EN 13060]

RKI

Abbreviation "Robert Koch Institute". It is one of the most important bodies for the safeguarding of public health in Germany.

Simple hollow bodies

An article open on one side to which the following applies: $1 \leq L/D \leq 5$ and $D \geq 5$ mm or an article with an opening on both sides which is: $2 \leq L/D \leq 10$ and $D \geq 5$; L...hollow body article length; D... hollow body article diameter [see also EN 13060]

Single wrapping

Wrapped once e.g. instruments sealed in foil – in opposition to: Multiple wrapping

Soft sterilization packaging

e.g. a paper bag or transparent sterilization packaging

Solid

Without hollows or gaps, solid, compact, closed

Solid load

Serves to prove that the necessary sterilization conditions have been reached within the entire load with the values set in the control. The load must represent the largest weight of massive instruments designed for sterilization in a sterilizer in accordance with EN 13060. [see also EN 13060]

Sterile barrier system

A closed minimum packaging which prevents the entrance of micro-organisms e.g. through sealing bags, sealed and re-usable containers and folded sterilization towels etc.

Sterile material

Also referred to as a batch: a load which has already been sterilized, i.e. is sterile

Sterilization chamber

The interior of a sterilizer accommodates the sterilization material

Sterilization material

Unsterile, sterilizable material which is still to be sterilized.

Vacuum

In common parlance, an area devoid of all material. In the technical sense: volumes with a reduced gas pressure (at least air pressure)



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Original instructions

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Your stockist