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Further information, accessories, consumables and spare parts are available from:

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info@atmosmed.de
www.atmosmed.com
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1.0 Introduction

1.1 Notes on operating instructions

These operating instructions contain important notes on how to operate the ATMOS C 341 Battery safely, correctly and effectively. These operating instructions serve not only for new operating personnel to be instructed in its use, but also for use as a reference manual. Any reprint - even in extracts - only after written permission from ATMOS.

These operating instructions must always be kept available near the device.

Care, period tests, regular cleaning and proper application are indispensable. They guarantee the operational safety and usability of the ATMOS C 341 Battery. Maintenance, repairs and period tests may only be carried out by persons who have the appropriate technical knowledge and are familiar with the product. To carry out these measures the person must have the necessary test devices and original spare parts.

Prior to start-up please peruse chapter „2.0 Hints for your safety“ on page 11, in order to be prepared for any possible dangerous situations. This helps you avoid potentially dangerous situations.

The product ATMOS C 341 Battery bears CE marking CE 0124 according to the EC Directive of the council for medical products 93/42/EEC and meets the basic requirements of Appendix I of the directive.

The product ATMOS C 341 Battery complies with all applicable requirements of the Directive 2011/65/EC restricting the use of certain hazardous substances in electrical and electronic equipment ("RoHS").

The declaration of conformity and our general standard terms and conditions can be obtained on our website at www.atmosmed.com.

The quality management system at ATMOS has been certified according to international standards EN ISO 13485.

These operating instructions are valid for the following devices:

- ATMOS C 341 Battery REF 318.0000.0
- ATMOS C 341 Battery / DDS REF 318.1300.0
- ATMOS C 341 Battery / Serres® REF 318.1400.0
- ATMOS C 341 Battery / Medi-Vac® REF 318.1600.0
- ATMOS C 341 Battery / Universal bracket REF 318.1700.0

Some figures show the ATMOS E 341 Battery. However, the devices do not differ in their functionality described.
1.2 Explanation of pictures and symbols

In these operating instructions

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="danger.png" alt="Danger" /></td>
<td>Warning of a danger which causes immediate death or serious injury. Observe the necessary measures.</td>
</tr>
<tr>
<td><img src="warning.png" alt="Warning" /></td>
<td>Beware of a danger which can cause death or serious injury. Observe the necessary measures.</td>
</tr>
<tr>
<td><img src="caution.png" alt="Caution" /></td>
<td>Beware of a danger which can easily hurt you. Observe the necessary measures.</td>
</tr>
<tr>
<td><img src="notice.png" alt="Notice" /></td>
<td>Indication of a danger where the product or other items can be damaged. Observe the necessary measures.</td>
</tr>
</tbody>
</table>

Warning of a danger which can cause death or serious injury.

- Information regarding possible material damage which can be caused.
- Useful information on the handling of the device.

   - Numeration.
   - Result of an action.
   - Move, plug in this direction.
   - Engage, check correct fit.

On device and type plate

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="blue.png" alt="Follow operating instructions" /></td>
<td>Follow operating instructions (blue)</td>
</tr>
<tr>
<td><img src="book.png" alt="Observe operating instructions" /></td>
<td>Observe operating instructions</td>
</tr>
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<td><img src="manufacturer.png" alt="Manufacturer" /></td>
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<tr>
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<td>Serial number</td>
</tr>
<tr>
<td><img src="degree.png" alt="Degree of protection" /></td>
<td>Degree of protection</td>
</tr>
<tr>
<td><img src="order.png" alt="Order number" /></td>
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<tr>
<td><img src="short.png" alt="Short term operation" /></td>
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<td><img src="disposal.png" alt="Professional disposal" /></td>
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<tr>
<td><img src="patient.png" alt="PATIENT" /></td>
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</tr>
<tr>
<td><img src="connection.png" alt="Connection suction hose/patient" /></td>
<td>Connection suction hose/patient (Serres® canister system)</td>
</tr>
<tr>
<td><img src="single.png" alt="For single use only" /></td>
<td>For single use only (Symbol is on the consumables)</td>
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This product complies with the relevant requirements of the EU Directives
On the recharging accessories

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<td>This product complies with the relevant requirements of the EU Directives</td>
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<td><img src="image" alt="Manufacturer" /></td>
<td>Manufacturer</td>
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<tr>
<td><img src="image" alt="Serial Number" /></td>
<td>Serial number</td>
</tr>
<tr>
<td><img src="image" alt="Output Voltage" /></td>
<td>Output voltage (13.8 V / 3.5 A)</td>
</tr>
<tr>
<td><img src="image" alt="Input Voltage" /></td>
<td>Input voltage (100 - 240 V / 50 - 60 Hz / 1.1 A)</td>
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<td><img src="image" alt="Alternating Current" /></td>
<td>Alternating current</td>
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<td><img src="image" alt="Protection Class" /></td>
<td>Protection class II</td>
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<thead>
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<th>Information</th>
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<tbody>
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<td>REF Order number</td>
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<tr>
<td><img src="image" alt="Certificate" /></td>
<td>Professional disposal</td>
</tr>
<tr>
<td><img src="image" alt="Certificate" /></td>
<td>IP40 Degree of protection</td>
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<tr>
<td><img src="image" alt="Heart" /></td>
<td>Application part type CF</td>
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<tr>
<td><img src="image" alt="House" /></td>
<td>For indoor use only</td>
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</table>

**On the battery**

<table>
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</tr>
</thead>
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<td><img src="image" alt="Certificate" /></td>
<td>China RoHS: Environmental friendly use period (EFUP) in years</td>
</tr>
<tr>
<td><img src="image" alt="Contains Lead" /></td>
<td>Contains lead, recyclable</td>
</tr>
<tr>
<td><img src="image" alt="Contains Lead" /></td>
<td>Contains lead, dispose of professional</td>
</tr>
</tbody>
</table>

1.3 Intended use and side effects

**Intended use**

**Name:** ATMOS C 341 Battery  
**Main functions:** Temporary and spontaneous suction of secretion, blood and body fluids and also liquid, viscous and solid pieces of food in the medical sector. To evacuate vacuum mattresses and inflatable splints.

**Medical indications / application:** Suction of the upper and lower respiratory tract.

**Specification of the main function:** Drainage and temporary collection of body fluids. By means of an electrical suction pump a negative pressure will be created. The integrated suction canister allows a temporarily collection of the derived body fluids.

**Application organ:** Upper respiratory tract (nose, nasal cavity, throat), lower respiratory tract (larynx, trachea, bronchial system)

**Application time:** Temporary use on the patient (< 60 min.)

**Application site:** The application site is the clinic, the practice, the accident & emergency department, the nursing and home care sector, as well as for outdoor application and during transport. The application of the device may only be performed by medically trained and introduced staff.
**Contraindications:** Not suitable for
- The continuous operation by drainages in the low vacuum range (e.g. thorax drainages or wound drainages).
- Permanent endoscopic use.
- Suction in medical rooms where a potential equalization is necessary (e.g. heart surgery).
- Use outside the medical sector.
- Suction of flammable, corrosive or explosive substances.
- Suction in explosion-hazardous areas.

**The product is:** active

**Sterility:** No sterile product

**Single use product / reprocessing:** The device and part of the accessories are reusable. For information on reprocessing, cleaning and disinfection please see the operating instructions.

**Possible side effects during suction**
- Bleeding in the nasal pharyngeal area
- Injury to the vocal cords
- Tracheal injury
- Hypoxemia
- Cardiovascular instability
- Bradycardia, Arrhythmia and Asystole (caused by vagus stimulation)
- Tachycardia (caused by stress)
- Choking, nausea, vomiting and coughing
- Nosocomial infection of the respiratory tract
- Seizures by patients who tend to develop cramps

Attention must be paid to these operating instructions in order to keep the side effects as minimal as possible.
1.4 Function

The ATMOS C 341 Battery is a mobile, portable, mains operated medical device for the temporary application on adults, children and babies. The device is operated with an electronically controlled, maintenance-free diaphragm pump.

The pump can be optionally operated by rechargeable battery or via an external DC voltage source (12 V).

During operation the pump generates a vacuum within the hose system and collection canister, thus sucking off secretion, blood and body fluids as well as liquid, viscous and solid pieces of food. The fluid is gathered in the collection canister.

With the help of the vacuum control the target vacuum (-0.05 bar up to -0.8 bar) and with it the suction performance can infinitely be adjusted. The set value can be read at the vacuum display.

**DDS secretion canister:**

The DDS secretion canister is affixed laterally to the device and is plugged via Direct Docking onto the suction connection at the support for the DDS canister system. Therefore there is no intermediate hose. Now the user can/must only connect the suction hose. A hydrophobic bacterial filter located in the lid of the canister prevents bacteria and liquids from entering the pump.

A mechanical oversuction stop (float ball) is integrated in the canister lid. This prevents an accidental absorption of secretion into the pump head. The float ball rises to the top of the secretion until it blocks the outlet.

**Disposable suction canister:**

The disposable suction canister is comprised of an external canister, disposable suction bag, vacuum hose and the disposable suction hose.

The disposable suction canister is affixed laterally to the device. The vacuum hose of the canister is connected to the suction connection of the device. The secretion is transported to the disposable suction bag via the suction hose. The disposable suction bag is a single use product. As soon as the suction bag is full it is removed from the external canister and disposed of. The disposable suction bag and the disposable suction hose must not be reused.

A bacterial filter is integrated in the disposable suction bag. This prevents secretion, liquid and bacteria from seeping into the device.

1.5 Intended operators

The ATMOS C 341 Battery may only be used by persons who were medically trained, and were trained in the medical suction. Prior to application the user must be familiar with the device. Please note the country-specific requirements and regulations.

ATMOS recommends: Instruction on the operation of the device must be performed by an authorized person.
1.6 Scope of delivery

1. Please compare the contents on completeness immediately upon receipt (see delivery note).

**Basic device**

- ATMOS C 341 Battery with device base
- Hose reel (mounted)
- Power supply and recharging unit 318.0035.0
- 2-pin mains connection cable 008.0920.0

**DDS canister system (mounted)**

- Support for DDS canister system
- Reusable suction hose, Ø 10 mm L 1.3 m
- Secretion canister 1 l with canister lid, filter holder, sealing ring
- 10 x Bacterial filter
- 10 x Fingertip

**Serres® canister system (mounted)**

- Support for Serres® canister system
- Vacuum hose with angled connection
- Serres® external canister 1 l
- 10 x disposable suction hose with fingertip, Ø 6 mm

**Medi-Vac® canister system (not mounted)**

- Support for Medi-Vac® canister system
- Vacuum hose with angled connection
- Medi-Vac® external canister 1 l
- 10 x disposable suction hose with fingertip, Ø 6 mm
Universal bracket (not mounted)

Support for canister system
Vacuum hose with angled connection

Hydrophobic bacterial and viral filter is not included in the scope of delivery and must be ordered separately for use with a canister system without an integrated bacterial filter.

Not included in the scope of delivery:

- Suction catheter
- Adapter for vacuum mattresses
- Serres’ suction bag 1 l
- Medi-Vac® suction bag 1 l
- Wall and device support

1.7 Transport and storage

Transport the device only in a shipping carton, which is padded and offers sufficient protection.

If you notice any damage:
1. Document and report the transport damage.
2. Fill in the form QD 434 “customer complaint/return shipment”. This form is enclosed to each delivery and can be found at www.atmosmed.com.
3. Send in the device to ATMOS (chapter „6.3 Sending in the device“ on page 40).

Ambient conditions during transport and storage:

- Temperature: - 40...+ 70 ° C
- Relative air humidity: 5...95 % without condensation
- Air pressure: 540...1100 hPa
2.0 Hints for your safety

The safety of the ATMOS C 341 Battery complies with all the recognized rules of technology and the guidelines of the Medical Products Law.

Please read and pay attention to the safety instructions prior to using the product.

2.1 General safety information

Make yourself familiar with the device at an early stage, so you can use it even in hectic situations.

Never operate the unit, if it shows any obvious safety defects. Check the unit at regular intervals for safety and function.

2.2 Danger for users, patients and third parties

Take care that the device is always functional and ready for use.

Your patient may suffocate.

- Ensure that the device is always ready for use in an emergency.
- Position the unit in an easily accessible location and keep access free.
- Make sure that the charging accessories are functional. Replace defective charging accessories immediately.
- Recharge the battery at the latest after 3 months, even if you do not use the device.
- Carry out a function check after each use. Carry out a function check every 4 weeks in case you do not use the device for a longer period.
- ATMOS recommends always having another suction device ready to hand in case of any device failure. So you can suck even in the event of equipment failure.
- Please observe the notes on electromagnetic compatibility (EMC) of the device.

Avoid misapplication.

Your patient may be seriously injured.

- The ATMOS C 341 Battery may only be used by persons who were medically trained, and were trained in the medical suction.
- Please select the vacuum according to the patient and the application.
- Observe the valid guidelines.

Reduce the risk of infection for you and your patients!

Deadly diseases can be transmitted.

- Always wear disposable gloves, if you could come into contact with secretion.
- Never use components marked with \(\text{\ding{116}}\) more than once. These components are intended for single use only.
- Only use sterile packaged parts, when the packaging is undamaged.
- Do not operate the device without a bacterial filter.
Protect yourself against an electric shock.
Burns, cardiac arrhythmias and even death are possible.

- Do not operate the device if it has been dropped. In this case please clean the device and send it in to ATMOS for repair.
- Disconnect the device from the mains power supply prior to cleaning or disinfection.
- Prior to each use, please check whether the device or the recharging accessories are damaged. Never operate the device if you detect any failure. In this case please clean the device and send it in to ATMOS for repair.
- Take care that no liquid penetrates the device. In case that liquid has penetrated the device it may no longer be operated. In this case please clean the device and send it in to ATMOS for repair.
- The ATMOS C 341 Battery cannot be sterilized.
- Use the recharging accessories in dry surroundings. The surroundings must be non-conductive.
- Only use the recharging accessories according to the operating instructions.
- Only use original accessories and original spare parts from ATMOS. This specifically applies to the recharging accessories and the battery.
- Please pay attention to the period tests in chapter „6.0 Maintenance and service“ on page 39.
- Assembly, repairs, modifications and period tests may only be carried out by authorized persons.
- Do not modify the device without permission of the manufacturer.

Explosion and fire hazard!
Burns and injuries are possible.

- Never suction any explosive, flammable or corrosive gases or liquids. Please observe the intended use in chapter „1.3 Intended use and side effects“ on page 6.
- Never operate the device in explosion-hazardous areas or areas which are oxygenated.
- Only use original accessories and original spare parts from ATMOS. This specifically applies to the recharging accessories and the battery.

Danger of suffocation for children through accessories!
Children can strangle themselves or be suffocated by small parts.

- Keep children away from hoses and connection cables.
- Keep children away from swallowable small parts. Small parts are, e.g. fingertip and sealing ring.

Tripping hazard by cables.
Injuries and fractures are possible.

- Lay connecting cables properly.

Only a fully functional product will meet the safety requirements of users, patients and third parties. Please therefore read the following instructions carefully.
2.3 Damage to the device

Please observe the ambient conditions regarding transport, storage, operation and recharging of the battery.

Take care that no liquid penetrates the device. In case that liquid has penetrated the device it may no longer be operated. In this case please clean the device and send it in to ATMOS for repair.

Always place the device on firm, level surface. The device must always be in a vertical position, when you use it. Otherwise secretions may enter the unit.

Only use proper power connections and extension cords.

If possible, avoid a transport at temperatures below \(-5\) °C. After transport in temperatures below \(-5\) °C: The device must be acclimatized for up to 6 hours at room temperature before you continue with the next steps.

The device may only be connected to the mains power supply when mains voltage and frequency of device and mains power supply correspond.
3.0 Setting up and starting up

Please observe that insufficient battery charge can result in damage to the battery.
1. The battery must be fully charged prior to first use.

3.1 Device overview

3.1.1 Front and rear view

Basic device

DDS canister system
With Serres® canister system

- Angle (connection disposable suction hose)
- Serres® suction bag
- Serres® external canister
- Support for Serres® canister system
- Grey angle on the Serres® external canister (connection vacuum hose)
- Vacuum hose with angled connection

With Medi-Vac® canister system

- Angle (connection disposable suction hose)
- Red hose
- Medi-Vac® suction bag
- Connection vacuum hose
- Medi-Vac® external canister
- Support for Medi-Vac® canister system
- Vacuum hose with angled connection

With universal bracket

- Support for canister system
- Vacuum hose with angled connection

The universal bracket is suitable for a secretion canister with a diameter of 11.5 - 12.5 cm.
Do not operate the device without a bacterial filter.

Do not operate the device without a bacterial filter.
Control panel

1. Display of battery status
2. Vacuum controller
3. Vacuum display
4. On/off button
5. Power indicator

Display of the battery status

The following display values are not valid during battery charging.

<table>
<thead>
<tr>
<th>Battery Level</th>
<th>Display</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt; 100 %</td>
<td><img src="battery1.png" alt="" /></td>
</tr>
<tr>
<td>ca. 70 %</td>
<td><img src="battery2.png" alt="" /></td>
</tr>
<tr>
<td>ca. 40 %</td>
<td><img src="battery3.png" alt="" /></td>
</tr>
<tr>
<td>&lt; 10 %</td>
<td><img src="battery4.png" alt="" /></td>
</tr>
</tbody>
</table>

The battery status must be supervised visually. If only the red LED lights up, although the battery has been loaded for a longer period, the battery is defective.

### 3.2 Preparing the device

Prior to first operation peruse the safety notes in chapter „2.0 Hints for your safety“ on page 11.

1. Damaged pump diaphragms due to cold temperatures during transport.
2. After transport in temperatures below -5°C: The device must be acclimatized for up to 6 hours at room temperature before you continue with the next steps.
3. Check the device for any damage in transport.
4. If the device is damaged: Document and report the transport damage. Send in the device to ATMOS (chapter „6.3 Sending in the device“ on page 40).
5. If the device is not damaged: Place it on a safe and even surface.
6. Check the charging accessories for any damage.
7. Replace defective charging accessories immediately.
8. The battery must be fully charged, chapter „3.3 Charging the battery“ on page 17.
9. Remove the canister system from the support.
10. With DDS canister system: Prior to first use, clean the canister system and insert a bacterial filter (chapter „5.0 Cleaning and disinfection“ on page 32 as well as chapter „3.4 Connection and removal of canister system and hoses“ on page 18).
11. Connect the suction hose.
11. Place the canister system upright from above into the support: Chapter „3.4 Connection and removal of canister system and hoses“ on page 18.

12. Wrap the suction hose onto the hose rewind.

13. If you wish to use the device for vacuum mattresses: Check whether a suitable adapter is available for the vacuum mattresses.

### 3.3 Charging the battery

- You can check the battery status by switching on the device. The battery must be fully charged prior to first use.
- Damage to the battery due to deep discharge.

1. Charge the battery at the latest when the red LED on the display of the battery status lights.

2. Only use the enclosed power supply and recharging unit 318.0035.0. Other charging accessories must not be used.

3. Please observe the notes in chapter „6.4 Handling of batteries“ on page 40. During battery recharging full suction performance of the device is still available. If the battery is fully discharged or defective, the device may be operated via the charging accessories.

#### Ambient conditions during charging

- Temperature: -5...+50 °C
- Relative air humidity: 5...95 % without condensation
- Air pressure: 540...1100 hPa

#### Charging with power supply and recharging unit

1. Connect the plug from the power supply and recharging unit to the back of the device 1.

2. Connect the power cable to the power supply and recharging unit.

3. Plug in the power plug of the power supply and recharging unit to the socket.

   » The battery is fully recharged when the top red LED is illuminated.

#### Recharging via the wall and device support

If you have attached the charging accessories to a wall and device support then the device will be charged automatically: Chapter „8.1 Wall and device support“ on page 45.

1. Attach the device to the wall and device support.

   » The battery is fully recharged when the top red LED is illuminated.
3.4 Connection and removal of canister system and hoses

3.4.1 DDS canister system

**WARNING**
Risk of infection by contaminated bacterial filter and canister lid.
Deadly diseases can be transmitted.

- Do not operate the device without a bacterial filter. We recommend you always to store at least one spare bacterial filter.
- Always wear disposable gloves when changing the bacterial filter.
- Prior to each use please check whether the bacterial filter is dry and clean. Replace the bacterial filter with a new filter if it is discoloured, contaminated or oversucked. A bacterial filter may never be reused.
- Exchange the bacterial filter when changing the patient. ATMOS recommends: Replace the bacterial filter after 14 days, even if there is no patient change.

**Removal**

1. Remove the suction hose from the hose rewind and then from the hose guide.
2. Gently unlock the clip 1 of the canister lid from the support of the DDS canister system and lift it upwards:

3. Lift the canister system upwards from the support.
4. Place the canister system on a safe and even surface.
5. Remove the suction hose from the secretion canister.
6. Turn the filter holder anti-clockwise by 90°.
   - The filter holder is difficult to turn because it has to seal the canister lid tightly.

7. Remove the filter holder with bacterial filter from the canister lid.
8. If required: Remove the bacterial filter 7 and the sealing ring 6 from the filter holder 5.

**Risk of infection by overflowing secretion. Deadly diseases can be transmitted.**
9. Hold the secretion canister with one hand and pull it upwards with force.
   - The canister system is open.
10. If required: Press the inner canister lid forward and remove it from the outer canister lid.

11. If required: Remove the float ball from the float ball compartment of the inner canister lid.

**Connection**

When you pour 50-100 ml water or disinfectant into the secretion canister, then it is easier to clean.

1. Press the outer canister lid on the inner canister lid, until it clicks into place.
2. Open the float ball compartment gently and insert the float ball.
3. Gently press the float ball compartment together.
4. Check whether the float ball moves easily and does not fall out of the float ball compartment.
5. Place the secretion canister on a firm surface.
6. Press the canister lid onto the secretion canister. The canister lid cannot be placed in a wrong position.
7. Press the canister lid tightly with both hands as far as it will go onto the secretion canister.
8. Place the sealing ring onto the filter holder.
9. Insert a new bacterial filter onto the filter holder.
10. Insert the filter holder into the canister lid and turn it clockwise until it clicks into place.

11. Connect the suction hose to the canister system.

12. Place the canister system upright into the support and at the same time position the suction hose in the hose guide.

13. Check that the clip 1 of the canister lid is correctly attached to the support of the canister system.

» The vacuum connection from the pump to the canister system is established.

14. Check whether the hose has a kink. If applicable, remove the kink.

15. Insert the suction hose into the hose guide of the device base.

16. Wrap the suction hose onto the hose rewind.

17. If required: Connect a fingertip to the suction hose.

3.4.2 **Serres® canister system**

**WARNING**
Risk of infection by contaminated canister system and hoses.
Deadly diseases can be transmitted.
- Only use Serres® suction bags with integrated bacterial filter.
- Only use sterile packaged parts, when the packaging is undamaged.

**No vacuum or vacuum is too low because of incorrect connection.**
Patient can suffocate.
- Please observe the operating instructions from the manufacturer of the Serres® canister system.

**Removal**
1. Remove the disposable suction hose from the hose guide.
2. Remove the disposable suction hose and the angle 2 from the Serres® suction bag.
3. Close the connection „patient“ at the Serres® suction bag with the green cap 3.
4. Remove the vacuum hose from the Serres® external canister (grey angle 1).
5. Remove the Serres® canister system from the support.
6. If required: Remove the vacuum hose from the device.

**Connection**
1. Connect the vacuum hose to the device.
2. Place the Serres® external canister upright into the support. The scale must be visible.
3. Insert the Serres® suction bag into the Serres® external canister.
4. Connect the vacuum hose to the Serres\textsuperscript{®} outer canister (grey angle 1).

5. Check whether the foil of the Serres\textsuperscript{®} suction bag is fully inserted into the Serres\textsuperscript{®} external canister and the lid tightly fits to the Serres\textsuperscript{®} external canister.

6. Connect the disposable suction hose with the angle 2 to the Serres\textsuperscript{®} suction bag.

7. Close the auxiliary air vent of the fingertip and close the front opening with your thumb.

8. Switch on the device so that the pump builds up a vacuum.

   » The Serres\textsuperscript{®} suction bag develops.

9. Insert the suction hose into the hose guide.

10. Wrap the suction hose onto the hose rewind.


3.4.3 Medi-Vac\textsuperscript{®} canister system

\begin{tabular}{|c|}
\hline
\textbf{WARNING} \\
\hline
\end{tabular}

\textbf{Risk of infection by contaminated canister system and hoses.}
Deadly diseases can be transmitted.

• Only use Medi-Vac\textsuperscript{®} suction bags with integrated bacterial filter.

• Only use sterile packaged parts, when the packaging is undamaged.

\textbf{No vacuum or vacuum is too low because of incorrect connection.}

Patient can suffocate.

• Please observe the operating instructions from the manufacturer of the Medi-Vac\textsuperscript{®} canister system.

\section*{Removal}

1. Remove the disposable suction hose from the hose guide.

2. Remove the disposable suction hose and the angle 2 from the Medi-Vac\textsuperscript{®} suction bag.
3. Close the connection „patient“ at the Medi-Vac® suction bag with the blue cap ➃.
4. Remove the red hose ➀ from the Medi-Vac® suction bag.
5. Close the connection „Vacuum“ at the Medi-Vac® suction bag with the blue cap ➂.
6. Remove the vacuum hose from the red connection ➄ of the Medi-Vac® suction bag.
7. Remove the Medi-Vac® canister system from the support.
8. If required: Remove the vacuum hose from the device.

**Connection**

1. Connect the vacuum hose to the device.

2. Insert the Medi-Vac® suction bag into the Medi-Vac® external canister.
3. Connect the red hose ➀ to the Medi-Vac® suction bag.
4. Place the Medi-Vac® external canister upright into the support.
5. Connect the vacuum hose to the red connection ➄ of the Medi-Vac® suction bag.

6. Check whether the lid tightly fits to the Medi-Vac® external canister.
7. Connect the disposable suction hose ➋ to the Medi-Vac® suction bag.
8. Close the auxiliary air vent of the fingertip and close the front opening with your thumb.
9. Switch on the device so that the pump builds up a vacuum.
   » The Medi-Vac® suction bag develops.
10. Insert the suction hose into the hose guide.

11. Wrap the suction hose onto the hose rewind.
3.5 Support for canister system

3.5.1 DDS canister system

Removal

1. Remove the canister unlocking.

2. Push the support for the DDS canister system backwards up to the middle and take it out of the guides.

Mounting

1. Attach the support for the DDS canister system in the middle on the right side of the device. The bars at the support must be fitted to the two guides at the device.

2. Slide the support for the DDS canister system forward until it is flush with the device. The inlet to the pump must be visible.
3. Attach the canister unlocking.

The canister unlocking is at the same time the connection angle through which the canister system is connected to the pump.

### 3.5.2 Serres® canister system

#### Removal

1. Remove the connection angle.
2. Push the support for the Serres® canister system backwards up to the middle and take it out of the guides.

#### Mounting

1. Attach the support for the Serres® canister system in the middle on the right side of the device. The bars at the support must be fitted to the two guides at the device.
2. Slide the support for the Serres® canister system forward until it is flush with the device. The inlet to the pump must be visible.
3. Connect the vacuum hose with the connection angle.

3.5.3 Medi-Vac® canister system / Universal bracket

Removal
1. Remove the connection angle.
2. Push the support of the canister system backwards up to the middle and take it out of the guides.

Mounting
1. Attach the support of the canister system in the middle on the right side of the device. The bars at the support must be fitted to the two guides at the device.

2. Slide the support of the canister system forward until it is flush with the device. The inlet to the pump must be visible.
3. Connect the vacuum hose with the connection angle.

- The universal bracket is suitable for a secretion canister with a diameter of 11.5 - 12.5 cm.
- Do not operate the device without a bacterial filter.

### 3.6 Hose rewind

#### Removal

Prerequisite: Hose is taken off.
1. Pull the wings outwards so that the hose rewind can be released.
2. Pull the hose rewind from the device.

#### Mounting

Prerequisite: Device base and battery compartment cover are attached.
1. Turn the hose rewind so that the opening points upwards.
2. Attach the hose rewind with force to the support on the left side of the device until it clicks into place.

### 3.7 Device base

#### Removal

Prerequisite:
The following parts are removed:
- Canister system
• Support for canister system
• Hose rewind
• Battery compartment cover
1. Put the device carefully on the front.
2. Remove the device base in the following order A - B - C - D:

Mounting

NOTICE
Incorrectly mounted device base.
Device moves during the operation.
• Attach the device base with particular care, according to the operating instructions.

Requirements:
The following parts are removed:
• Canister system
• Support for canister system
• Hose rewind
• Battery compartment cover
1. Put the device carefully on the front.
2. Take care that the indentations at the device base are fitted to the protruding edges at the device.
3. Attach the device base in the following order D - C - B – A. The arrows show the points at which the device and the device base have to engage with each other.

4. Firmly press on all sides again.
5. Afterwards the following parts can be mounted:
   - Battery compartment cover (chapter „6.5 Battery exchange“ on page 40)
   - Hose rewind (chapter „3.6 Hose rewind“ on page 27)
   - Support for canister system (chapter „3.5 Support for canister system“ on page 24)
   - Canister system (chapter „3.4 Connection and removal of canister system and hoses“ on page 18).
4.0 Operation

**WARNING**

Risk of infection by lack of hygiene or damaged components.
Deadly diseases can be transmitted.
- Please use new consumables and new disposable canister systems or reprocessed DDS canister systems for each patient.
- Prior to each use, please check whether hoses or canister systems are damaged. Replace any damaged parts.

Electric shock from damaged equipment.
Cardiac arrhythmias may be caused.
- Prior to each use, please check whether the device and the recharging accessories are damaged.
- Replace any damaged parts immediately.
- Do not use the device if it is damaged.

Ambient conditions during operation
- Temperature: -5...+50° C
- Relative air humidity: 5...95 % without condensation
- Air pressure: 540...1100 hPa

4.1 Switch on the device

* The device should only be left on as long as you need it. This way you can increase the battery life.
1. Push the on/off button to switch on the device.
   » The pump starts.
   » The LED above the on/off button is illuminated as long as the device is switched on.

4.2 Switch off the device
1. Switch off the device by pressing the on/off button for at least 1 second.

4.3 Vacuum adjustment

**WARNING**

Vacuum is too high.
Patient may be seriously injured.
- Observe the valid guidelines.
- Please select the vacuum according to the patient and the application.

1. Push the on/off button to switch on the device.
   » The pump starts.
2. Turn the vacuum controller for adjusting the required vacuum.
   » The vacuum display shows the current suction performance.
4.4 Suction

**WARNING**
Device failure, if the period of continuous operation is too long.
Patient can suffocate.
- Make sure not to use the device in continuous operation for more than 45 minutes. Otherwise the pump shuts off automatically. In this case let the device cool down for about 1.5 hours.
- Check the status of the battery regularly while you operate the device.

**Risk of infection.**
Deadly diseases can be transmitted.
- Always wear disposable gloves during suction.

**CAUTION**
Risk of injury due to inappropriate material or untrained users.
Injuries in the oral cavity and pharynx of the patient.
- Suction may only be carried out by persons who were medically trained, and were trained in the medical suction.
- Use a suction catheter for tracheal or nasopharyngeal suction.
- If you suck viscous food ingredients in the oral cavity, use the suction hose without suction catheter.

**Connect the suction catheter**
1. Remove the suction hose from the hose rewind.
If you suck viscous food ingredients in the oral cavity:
2. Use the suction hose without suction catheter.
For tracheal or nasopharyngeal suction:
2. Please choose a suction catheter in the appropriate size.
3. Connect the suction hose \(1\) and the suction catheter \(2\) with a fingertip \(4\):

   1. Suction hose
   2. Auxiliary air vent
   3. Suction catheter
   4. Fingertip

**Connect the special suction instruments**
1. Please observe the operating instructions for the suction instruments.
**Suction**

**CAUTION**

Adherence by careless suction.

Injuries in the oral cavity and pharynx of the patient.

- Briefly open the auxiliary air vent, if the suction catheter adheres to the tissue.
- Suck particularly careful in the tracheal area.

1. Push the on /off button to switch on the device.
   » The pump starts.
2. Please select the vacuum according to the patient and the application. Turn the vacuum controller for adjusting the required vacuum.
   » The vacuum display shows the current suction performance.
3. As long as the auxiliary air vent is opened, the device does not suck.
4. Pay attention to the filling level of the canister system.
5. Empty the secretion canister or change the suction bag once it is half full. As soon as the canister system is too full, the float ball seals the intake area. You can then no longer suck with the device.

Make sure that the hose is not kinked during suction. Otherwise the suction at the patient is too low.

If you want to interrupt the suction briefly, you can clamp the suction hose into the opening of the hose rewind.

If secretion has penetrated into the device, please observe chapter „5.6 Oversuction“ on page 38.

**After use**

1. Switch off the device by pressing the on/off button for at least 1 second.
2. Clean the device after every use: Chapter „5.0 Cleaning and disinfection“ on page 32.
5.0 Cleaning and disinfection

We recommend you to document any maintenance work and also any exchange of parts.

⚠️ WARNING
Risk of infection by secretion on the device, accessories and consumables.
Deadly diseases can be transmitted.
- Always wear disposable gloves during any cleaning.
- Clean the device after every use.
- Clean and disinfect the device according to the operating instructions.
- The device must be reprocessed professionally, if secretion has penetrated into the device. Please observe chapter „5.6 Oversuction” on page 38.

5.1 Prepare for cleaning

1. Switch off the device.
2. Remove the recharging accessories from the device.
3. Remove the canister system from the device: Chapter „3.4 Connection and removal of canister system and hoses“ on page 18.
   ⚠️ Risk of infection by overflowing secretion. Deadly diseases can be transmitted.
4. Carefully remove the canister lid / the suction bag.
5. Dispose of the secretion / the suction bag. Please observe the notes in chapter „10.0 Disposal“ on page 48.
6. Dispose of all disposables (e.g. suction catheter, fingertip, single-use suction hose).
   In case you are using the DDS canister system: Dispose of the bacterial filter.
7. Remove the hose rewind.
8. Remove the support for the canister system.

5.2 Cleaning

Please observe the operating instructions of the disinfectant manufacturers. Pay particular attention to the information regarding the concentration of the disinfectants and the material compatibility.

Some disinfectants may stain the parts of the canister lid and silicone hoses. Parts can also stain by autoclaving. However, this has no influence on the properties of the materials.

Please only use disinfectants, which are recommended by ATMOS (chapter „5.4 Recommended disinfectants“ on page 34). The use of other disinfectants may damage the device or the canister system.
**DDS canister system**

Number of reprocessing cycles: max. 50.

1. Disassemble canister lid and filter holder into their individual parts. Chapter „3.4 Connection and removal of canister system and hoses” on page 18.

   ![Diagram of DDS canister system parts](image)

   - Filter holder
   - Sealing ring
   - Inner canister lid
   - Float ball
   - Outer canister lid

2. Rinse the following parts of the DDS canister system with clear water:
   - Secretion canister
   - Inner canister lid
   - Outer canister lid
   - Float ball
   - Filter holder
   - Sealing ring
   - Suction hose
   - Support for canister system

3. Clean the mentioned parts with a brush or a cloth.
4. Disinfect the mentioned parts with a disinfectant which is recommended by ATMOS.
5. Let the individual parts of the canister lid and the filter holder dry.
6. As soon as the individual parts are dry:
7. Insert the new bacterial filter.
8. Reassemble the individual parts of the canister lid and the filter holder.

**Serres®, Medi-Vac® secretion canister system, other canister systems**

- Please observe the instructions in the operating instructions for the canister system.
- Do not operate the device without a bacterial filter. The hydrophobic bacterial and viral filter (REF 443.0738.0) must be used with a secretion canister system which has no integrated bacterial filter.

**Vacuum hose**

After every suction process:

1. Rinse the vacuum hose with clear water for at least 10 seconds.
2. Disinfect the vacuum hose with an instrument disinfectant recommended by ATMOS.

It must be exchanged after each patient or at least once a day.
Device surface

**WARNING**

**Electric shock by liquid in the device.**

- Disconnect the device from the mains power supply prior to cleaning.
- Do not rinse the device under running water and do not immerse it into any liquids.
- Make sure that the cleaning cloth is only damp and not wet.
- Do not autoclave the device.
- Do not immerse the device in disinfectant solution.

1. Clean the entire surface of the device and the hose rewind with a damp cloth.
2. Disinfect the entire surface of the device and the hose rewind with a surface disinfectant.

Power supply and recharging unit

**WARNING**

**Electric shock by liquid in the power supply.**

- Disconnect the power supply and recharging unit from the mains power supply prior to cleaning.
- Do not rinse the power supply and recharging unit under running water and do not immerse it into any liquids.
- Make sure that the cleaning cloth is only damp and not wet.
- Do not autoclave the power supply and recharging unit, do not sterilize it and do not thermally disinfect it.
- Do not immerse the power supply and recharging unit in disinfectant solution.

1. Clean the power supply and recharging unit with a damp cloth. You can use a mild detergent.
2. Disinfect the power supply and recharging unit with a surface disinfectant. Recommended: Terralin® Protect

Wall and device support

1. Clean the wall and device support with a damp cloth.
2. Disinfect the wall and device support with a surface disinfectant.

5.3 After cleaning

**Risk of injury to user and patient by a damaged device.**

1. Check after each cleaning, whether the device is obviously damaged. In case the device is damaged, please send it in to ATMOS.
3. Prepare the device for the next use.

5.4 Recommended disinfectants

If you are using aldehyde and amine-containing disinfectants on the same object, this may cause discoloration.
### 5.4.1 Instrument disinfection

<table>
<thead>
<tr>
<th>Disinfectant</th>
<th>Ingredients</th>
<th>in 100 g</th>
<th>Manufacturer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mucocit®-T</td>
<td>didecyldimethylammonium chloride, alkyl propylene diamine-1.5-bis guanidinium acetate, bis(aminopropyl)laurylamine, laurylpropylen diamine, non-ionic surfactants</td>
<td>3.9 g 4.5 g 2 g 2.8 g</td>
<td>Merz Dental, Lütjenburg</td>
</tr>
<tr>
<td>Gigasept® FF (neu)</td>
<td>succindialdehyde, dimethoxytetrahydrofuran, anionic and non-ionic surfactants, perfumes, methylisothiazolinone</td>
<td>11.9 g 3.2 g</td>
<td>Schülke &amp; Mayr, Norderstedt</td>
</tr>
<tr>
<td>Sekusept® PLUS</td>
<td>glucoprotamín</td>
<td>25 g</td>
<td>Ecolab, Düsseldorf</td>
</tr>
</tbody>
</table>

### 5.4.2 Surface disinfection

#### Coated surfaces

<table>
<thead>
<tr>
<th>Disinfectant</th>
<th>Ingredients</th>
<th>(in 100 g)</th>
<th>Manufacturer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Green &amp; Clean SK</td>
<td>Di alkyl dimethyl ammonium chloride, Alkyl dimethyl ethyl benzyl ammonium chloride, Alkyl dimethyl benzyl ammonium chloride</td>
<td>&lt; 1 g &lt; 1 g &lt; 1 g</td>
<td>Metasys, Rum (Austria)</td>
</tr>
<tr>
<td>Dismozon® pur</td>
<td>magnesium monoperphthalate hexahydrate</td>
<td>80 g</td>
<td>Bode Chemie, Hamburg</td>
</tr>
<tr>
<td>Dismozon® plus</td>
<td>magnesium monoperphthalate hexahydrate</td>
<td>95.8 g</td>
<td>Bode Chemie, Hamburg</td>
</tr>
<tr>
<td>Kohrsolin® FF</td>
<td>glutaral, benzyl-C12-C18-alkyldimethyl-ammonium chloride, didecylammonium chloride</td>
<td>5 g 3 g 3 g</td>
<td>Bode Chemie, Hamburg</td>
</tr>
<tr>
<td>Kohrsolin® extra</td>
<td>(ethylenedioxy)dimethanol, glutaral, didecylammonium chloride</td>
<td>14.1 g 5 g 8 g</td>
<td>Bode Chemie, Hamburg</td>
</tr>
<tr>
<td>Perform®</td>
<td>Potassium peroxymonosulphate</td>
<td>45 g</td>
<td>Schülke &amp; Mayr, Norderstedt</td>
</tr>
<tr>
<td>Terralin® Protect</td>
<td>benzalkonium chloride (benzyl-C12-18 alkylammonium, chloride), 2-phenoxyethanol, aminoaalkylglycine, non-ionic surfactants, perfumes</td>
<td>22 g 17 g 0.9 g</td>
<td>Schülke &amp; Mayr, Norderstedt</td>
</tr>
</tbody>
</table>
### Other surfaces

<table>
<thead>
<tr>
<th>Disinfectant</th>
<th>Ingredients</th>
<th>(in 100 g)</th>
<th>Manufacturer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dismozon® pur</td>
<td>magnesium monoperoxyphthalate hexahydrate</td>
<td>80 g</td>
<td>Bode Chemie, Hamburg</td>
</tr>
<tr>
<td>(Granulate)</td>
<td>End of product 12/2014</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dismozon® plus</td>
<td>magnesium monoperoxyphthalate hexahydrate</td>
<td>95.8 g</td>
<td>Bode Chemie, Hamburg</td>
</tr>
<tr>
<td>(Granulate)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Kohrsolin® FF</td>
<td>glutaral</td>
<td>5 g</td>
<td>Bode Chemie, Hamburg</td>
</tr>
<tr>
<td>(Application concentrate)</td>
<td>benzyl-C12-18-alkyldimethyl-ammonium chlorides</td>
<td>3 g</td>
<td></td>
</tr>
<tr>
<td></td>
<td>didecyldimethylammonium chloride</td>
<td>3 g</td>
<td></td>
</tr>
<tr>
<td>Kohrsolin® extra</td>
<td>(Application concentrate)</td>
<td>14.1 g</td>
<td>Bode Chemie, Hamburg</td>
</tr>
<tr>
<td></td>
<td>(ethylendioxy)dimethanol</td>
<td>5 g</td>
<td></td>
</tr>
<tr>
<td></td>
<td>glutaral</td>
<td>8 g</td>
<td></td>
</tr>
<tr>
<td></td>
<td>didecyldimethylammonium chloride</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mikrobac® forte</td>
<td>benzyl-C12-18-alkyldimethyl-ammonium chlorides</td>
<td>19.9 g</td>
<td>Bode Chemie, Hamburg</td>
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<tr>
<td>(Application concentrate)</td>
<td>N-(3-aminopropyl)-N-dodecylpropane-1,3-diamine</td>
<td>9 g</td>
<td></td>
</tr>
<tr>
<td>Perform®</td>
<td>Potassium peroxymonosulfate</td>
<td>45 g</td>
<td>Schülke &amp; Mayr, Norderstedt</td>
</tr>
<tr>
<td>Terralin® Protect</td>
<td>benzalkonium chloride (benzyl-C12-18 alkyldimethylammonium, chloride)</td>
<td>22 g</td>
<td>Schülke &amp; Mayr, Norderstedt</td>
</tr>
<tr>
<td>(Application concentrate)</td>
<td>2-phenoxyethanol</td>
<td>17 g</td>
<td></td>
</tr>
<tr>
<td></td>
<td>aminoalkylglycine</td>
<td>0.9 g</td>
<td></td>
</tr>
<tr>
<td></td>
<td>non-ionic surfactants, perfumes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SaniCloth® Active</td>
<td>didecyldimethylammonium chloride</td>
<td>&lt; 1 g</td>
<td>Ecolab, Düsseldorf</td>
</tr>
<tr>
<td>Incidin® Active</td>
<td>peracetic acid</td>
<td>&lt; 1 g</td>
<td>Ecolab, Düsseldorf</td>
</tr>
<tr>
<td>Mikrozid® Sensitive Wipes</td>
<td>benzyl-C12-16 alkyl(dimethyl-), chloride</td>
<td>0.26 g</td>
<td>Schülke &amp; Mayr, Norderstedt</td>
</tr>
<tr>
<td></td>
<td>didecyldimethylammonium chloride</td>
<td>0.26 g</td>
<td></td>
</tr>
<tr>
<td></td>
<td>benzyl-C12-14-alkyl [ethylphenyl)methyl] dimethyl-, chlorides</td>
<td>0.26 g</td>
<td></td>
</tr>
<tr>
<td>Gigasept® pearls</td>
<td>Natriumpercarbonat</td>
<td>43 g</td>
<td>Schülke &amp; Mayr, Norderstedt</td>
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<tr>
<td>Suitable for DDS secretion canisters</td>
<td>Tetraacetyl ethylenediamin</td>
<td>22 g</td>
<td></td>
</tr>
</tbody>
</table>
## 5.5 Hygienic plan

### What | How | When | Who | Notices
--- | --- | --- | --- | ---
Parts to be reprocessed | Cleaning | Disinfection | Sterilization | Qualified and trained staff who are familiar with reprocessing

### Surfaces

| Housing | Cleaning and Disinfection | X | X |
| Device base | Cleaning and Disinfection | X | X |
| Hose reel | Cleaning and Disinfection | X | X |
| Wall and device support | Cleaning and Disinfection | X |
| Power supply and recharging unit | Cleaning and Disinfection | X | X |

### Secretion canister system

| DDS secretion canister | Cleaning with a brush; cleaning and disinfection (automatic or manual), sterilization is possible | X | X |
| Outer canister lid (DDS) | Cleaning with a brush; cleaning and disinfection (automatic or manual), sterilization is possible | X | X |
| Inner canister lid (DDS) | Cleaning with a brush; cleaning and disinfection (automatic or manual), sterilization is possible | X | X |
| Float ball (DDS) | Cleaning with a brush; cleaning and disinfection (automatic or manual), sterilization is possible | X | X |
| Filter holder | Cleaning with a brush; cleaning and disinfection (automatic or manual), sterilization is possible | X | X |
| Sealing ring | Cleaning and Disinfection | X | X |
| Bacterial filter | Exchange. If the filter is blocked it must also be exchanged. | X | X |
| Fingertip | Exchange. | X | X |
| Support for canister system | Cleaning and Disinfection | X | X |
| Suction hose (DDS) | When using without suction catheter | X | X |
| Vacuum hose | Cleaning and Disinfection | X | X |

### Recommended disinfectants

- **Surface disinfection for coated surfaces:**
  - Green & Clean SK (ATMOS)
  - Dismoxon® pur (Bode Chemie)
  - Kohrsolin® extra (Bode Chemie)
  - Perform® (Schülke & Mayr)
  - Terralin® Protect (Schülke & Mayr)

### Manual disinfection of instruments:

- **Microutil® T** (Merz Dental)
- **Gigasept® FF neu** (Schülke & Mayr)
- **Sekusept® PLUS** (Ecolab)

For concentrations, contact time, temperature, material compatibility, please see the relevant information from the manufacturer.

### Important information

**Wipe cleaning and disinfection:**

All surfaces have to be wiped with a clean (disposable) wipe which is dampened with disinfectant solution. The entire surface has to be wiped thoroughly and may not be dried afterwards.

**Cleaning and disinfection:**

Various methods, such as cold or hot steam sterilization, are permissible. For concentrations, contact time, temperature, material compatibility, please see the relevant information from the manufacturer.

**Important notice:**

Wrong concentration of disinfectants may lead to material damage!
5.6 Oversuction

If you use the ATMOS C 341 Battery according to instructions, with bacterial filter and float ball, the device cannot be oversucked during normal use. Nevertheless, should secretion penetrate into the interior of the device, the device is oversucked. This can happen, for example, if no bacterial filter is used and the device tips over. Reduced suction power is an indication for an oversucked device. If you suspect that your device might be oversucked, proceed as follows:

⚠️ WARNING

Risk of infection by secretion on and within the device.
Deadly diseases can be transmitted.
- Always wear disposable gloves when touching the oversucked device.
- Clean and disinfect the device.
- Send in the device to ATMOS or an authorized ATMOS service partner, chapter „6.3 Sending in the device“ on page 40.
6.0 Maintenance and service

Maintenance, repairs and period tests may only be carried out by persons who have the appropriate technical knowledge and are familiar with the product. To carry out these measures the person must have the necessary test devices and original spare parts.

ATMOS recommends: Work should be carried out by an authorized ATMOS service partner. This ensures that repairs and testing are carried out professionally, original spare parts are used and warranty claims remain unaffected.

6.1 Period tests

Please comply with the country-specific guidelines regarding regular testing especially for the electrical safety.

ATMOS recommends a test every 24 months.

6.2 Functional check

Perform a function check:

• prior to each use
• after each use or cleaning
• every 4 weeks in case the device is not used
• after every maintenance work, service or repair

 Carry out a function check

1. Please check whether the following parts are damaged or torn:
   - all hoses
   - canister system
2. In case parts are visibly damaged: Please exchange them.
3. Switch on the device.
4. Check whether all LEDs are illuminated.
5. Check the battery status.
6. Connect a fingertip to the suction hose and close the auxiliary air vent.
7. Close the front opening of the fingertip with your thumb.
8. Select the vacuum -0.5 bar.
9. Please check whether the device reaches the vacuum after approx. 20 seconds: The pump switches off and the vacuum display shows -0.5 bar.
10. In case the device does not reach the vacuum within 20 seconds: Please check the device on potential sources of error and remedy the defect: Chapter „7.0 Eliminating errors“ on page 43.
11. You may now use the device or switch it off.
6.3 Sending in the device

1. Remove and properly dispose of consumables.
2. Clean and disinfect the products and accessories according to the operating instructions.
3. Place used accessories with the device.
   ☛ This form is enclosed to each delivery and can be found at www.atmosmed.com.
5. The device must be well padded and packed in suitable packaging.
6. Place the form QD 434 „Delivery complaint / return shipment“ and the respective decontamination certificate in an envelope.
7. Affix the envelope to the outside of the package.
8. Send the product to ATMOS or to your dealer.

6.4 Handling of batteries

Batteries are wearing parts and therefore excluded from the general warranty. There is a function guarantee of 6 months.

Please observe the following notes in order to reach the maximum service life of your battery:

• Exclusively use the lead-acid battery Panasonic LC-R 123R4PG.
• Please observe the operating instructions of the battery manufacturer.
• Prior to first use the battery must be fully charged.
• Battery-run devices should only be stored when they are charged.
• Please fully recharge the battery every 3 months, even if the device is not used.
• Prevent the batteries from direct solar radiation and keep them away from radiators. The perfect storage temperature is between 8 and 15° C.
• Exchange the battery when the remaining battery service life noticeably decreases.
• Batteries are run-down after approx. 400 charging cycles.

6.5 Battery exchange

**NOTICE**

Damage to the electronics due to the use of a third-party battery.

• Exclusively use the lead-acid battery Panasonic LC-R 123R4PG. This battery is included in the scope of delivery and is available at ATMOS. The warranty claim shall not be applicable if non-original spare parts are used.

Requirements: The support for canister system is removed.
1. Switch off the device.
2. Disconnect the device from the power supply.
3. Remove the hose rewind: Chapter „3.6 Hose rewind“ on page 27.
4. Place the device on its back with the control panel facing upwards.
5. Press the battery compartment cover 1 from right a little towards the left.

6. Lift the battery compartment cover slightly and remove it from the upper guide.
7. Remove the red and the black cable from the battery.
8. Put up the device again.
9. Press below on the battery so that it is tilted in the battery compartment.

10. Remove the battery in pulling it upwards.
11. First insert the battery below so that it is tilted in the battery compartment.
12. Press at the top of the battery until it clicks into place in the battery compartment.
13. Place the device on its back with the control panel facing upwards.
14. Connect the red cable to the positive pole on the left.
15. Connect the black cable to the minus pole on the left.

! The cables are mounted in such a way that a mix-up can be excluded.
16. Stow the cables in the battery compartment so that they cannot be damaged by the battery compartment cover.
17. Insert the battery compartment cover in the upper rail.
18. Slide the battery compartment cover to the right until it stops.
19. Press down the battery compartment cover.
20. Slide the battery compartment cover completely to the right.
21. Attach the hose rewind: Chapter „3.6 Hose rewind“ on page 27.
22. Affix the support for the canister system: Chapter „3.4 Connection and removal of canister system and hoses“ on page 18.
23. Perform a function check.
6.6 Exchange release button

- There are 2 springs under the release button. Pay attention that they are not misplaced.
1. Place the screwdriver in the middle of the release button and lift the release button up.

2. Replace with a new release button. Pay attention that the two springs are positioned in the guide of the release button.

3. Press the release button downwards until it engages.
7.0 Eliminating errors

The ATMOS C 341 Battery was subjected to a thorough quality control in the factory. Nevertheless, if a problem may occur you can possibly solve it yourself.

Recharging and battery status

<table>
<thead>
<tr>
<th>Error indication</th>
<th>Possible cause</th>
<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Device cannot be recharged.</td>
<td>The plug from the charging accessories is poorly fitted.</td>
<td>1. Check the connection to the mains supply.</td>
</tr>
<tr>
<td></td>
<td>Charging accessories are defective.</td>
<td>1. Exchange charging accessories.</td>
</tr>
<tr>
<td></td>
<td>Battery is not connected properly.</td>
<td>1. Check the plug connections in the battery compartment.</td>
</tr>
<tr>
<td></td>
<td>Battery temperature too high or too low.</td>
<td>1. After long-term use: Let the device cool down.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. Extreme ambient temperature: Position the unit where appropriate to a cooler or warmer place.</td>
</tr>
<tr>
<td></td>
<td>Defective battery.</td>
<td>1. Exchange the battery.</td>
</tr>
<tr>
<td></td>
<td>Defective fuse or electronics.</td>
<td>1. Send in the device for repair.</td>
</tr>
<tr>
<td>LED on the power supply and recharging unit is not illuminated.</td>
<td>Defective power supply and recharging unit.</td>
<td>1. Exchange the power supply and recharging unit.</td>
</tr>
<tr>
<td></td>
<td>The mains plug is poorly fitted.</td>
<td>1. Check the connection to the mains supply.</td>
</tr>
<tr>
<td>When recharging the battery 100 % cannot be achieved.</td>
<td>Battery service life is exhausted or the battery is defective.</td>
<td>1. Exchange the battery.</td>
</tr>
<tr>
<td>The charging time can take up to 14 hours.</td>
<td>Wrong charging accessories.</td>
<td>1. Only use the provided charging accessories or an original spare part.</td>
</tr>
<tr>
<td>Red LED of the battery status indicator is lit up although the battery is fully charged.</td>
<td>Defective battery.</td>
<td>1. Exchange the battery.</td>
</tr>
<tr>
<td>Battery compartment cover cannot be closed.</td>
<td>Battery is not fitted correctly.</td>
<td>1. Fit the battery correctly.</td>
</tr>
<tr>
<td></td>
<td>Battery compartment cover is installed incorrectly.</td>
<td>1. Mount the battery compartment cover correctly according to operating instructions.</td>
</tr>
</tbody>
</table>
## On and Off switching

<table>
<thead>
<tr>
<th>Error indication</th>
<th>Possible cause</th>
<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Device cannot be switched on or off.</td>
<td>Battery is discharged.</td>
<td>1. Recharge the battery.</td>
</tr>
<tr>
<td></td>
<td>Battery is not connected properly.</td>
<td>1. Check the plug connections in the battery compartment.</td>
</tr>
<tr>
<td></td>
<td>The plug from the charging accessories is poorly fitted.</td>
<td>1. Check the connection to the mains supply.</td>
</tr>
<tr>
<td></td>
<td>Defective fuse or electronics.</td>
<td>1. Send in the device for repair.</td>
</tr>
<tr>
<td>During switch on: Battery LEDs blink once, but the device does not start.</td>
<td>Device was stored outside the operating temperature (battery is in standby).</td>
<td>1. Switch on the device again.</td>
</tr>
<tr>
<td>Pump does not start up.</td>
<td>Vacuum is already built up.</td>
<td>1. Do not switch on the device if the vacuum is already built up.</td>
</tr>
<tr>
<td>Device switches off after 45 min.</td>
<td>Battery is discharged.</td>
<td>1. Recharge the battery.</td>
</tr>
<tr>
<td>Device switches off after &lt; 45 min.</td>
<td>Battery is discharged.</td>
<td>1. Recharge the battery.</td>
</tr>
<tr>
<td></td>
<td>Battery temperature is too high.</td>
<td>1. Let the device cool down or select a lower vacuum.</td>
</tr>
</tbody>
</table>

## Vacuum and suction capacity

<table>
<thead>
<tr>
<th>Error indication</th>
<th>Possible cause</th>
<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vacuum is not built up or cannot be reached.</td>
<td>Battery is discharged or defective.</td>
<td>1. Recharge or exchange the battery.</td>
</tr>
<tr>
<td></td>
<td>Leakages at the hoses or the canister system.</td>
<td>1. Check canister lid and hoses on tight fit. 2. DDS canister system: Firmly insert the bacterial filter and check the sealing ring and the filter holder.</td>
</tr>
<tr>
<td></td>
<td>Fingertip is not closed.</td>
<td>1. Close both openings of the fingertip.</td>
</tr>
<tr>
<td></td>
<td>Liquid has penetrated the device.</td>
<td>1. Send in the device for repair.</td>
</tr>
<tr>
<td></td>
<td>Pump is defective or the device has a leak.</td>
<td>1. Send in the device for repair.</td>
</tr>
<tr>
<td></td>
<td>Low ambient pressure (e.g. high altitude).</td>
<td>Not possible.</td>
</tr>
<tr>
<td>Low suction capacity although the vacuum is reached.</td>
<td>Bacterial filter is clogged.</td>
<td>1. Exchange the bacterial filter.</td>
</tr>
<tr>
<td></td>
<td>Hose is kinked.</td>
<td>1. Check the hoses.</td>
</tr>
<tr>
<td></td>
<td>DDS canister system: Float ball closes the suction area.</td>
<td>1. Check and if needed clean the float ball and the float ball compartment.</td>
</tr>
</tbody>
</table>
### 8.0 Accessories

<table>
<thead>
<tr>
<th>Item</th>
<th>Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>Serres® external canister 1 l</td>
<td>312.0465.0</td>
</tr>
<tr>
<td>Medi-Vac® external canister 1 l</td>
<td>312.0473.0</td>
</tr>
<tr>
<td>DDS canister system 1 l, complete</td>
<td>318.1000.0</td>
</tr>
<tr>
<td>Wall and device support for ATMOS emergency suction devices</td>
<td>318.1250.0</td>
</tr>
<tr>
<td>Retrofit kit DDS canister system</td>
<td>318.1350.0</td>
</tr>
<tr>
<td>Retrofit kit Serres® canister system</td>
<td>318.1450.0</td>
</tr>
<tr>
<td>Retrofit kit Medi-Vac® canister system</td>
<td>318.1650.0</td>
</tr>
</tbody>
</table>

### 8.1 Wall and device support

**Affix the power supply and recharging unit**

Requirements: The velcro fastener 1 is attached to the wall and device support.

1. Damaged cable due to incorrect mounting.
2. Please ensure that the power supply and recharging unit are positioned with the writing towards the wall or the standard rail. Otherwise the cable could be clamped.
3. Affix the parts 2 and 3 and fix the cable 4 with the velcro fastener 1.

**Wall mounting**

1. Only affix the power supply and recharging unit after you have marked the holes to be drilled.
2. Screws are not included in delivery.
3. Only use screws (max. 4 mm), which are suitable for the material of the wall.
4. Position the wall and device support at an easily accessible place.
5. Check whether the wall is smooth and vertical at the mounting position.
6. Hold the wall and device support to the mounting location and align it with a spirit level.
7. Mark the holes to be drilled on the wall.
8. Drill the holes with a, the wall material and the chosen screws corresponding drill.
9. Mount the power supply and recharging unit to the wall and device support. The power supply and recharging unit must not be connected to the mains supply.
10. Screw the wall and device support to the wall with suitable screws.
11. Connect the power supply and recharging unit to the mains supply.
12. Check whether the charging accessory is correctly attached by installing the device.
» The battery will be charged.
Attach to / remove from a standard rail

Requirements: The power supply and recharging unit is mounted.

Mounting

Attaching the device

1. Slide the device from above onto the wall and device support until it clicks into place.

   » If the charging accessory is attached, the battery is charged automatically.

Removing the device

1. Press the release button 1 and pull the device simultaneously vertically upwards.

8.2 Retrofit kit canister system

You may change the canister system. The retrofit kits comprise the canister system as well as the required support for the canister system. The retrofit kits for the single-use canister systems also comprise the vacuum hose.

Retrofitting

1. Remove the existing canister system, chapter „3.4 Connection and removal of canister system and hoses“ on page 18.
2. Remove the existing support for the canister system.
3. Attach the new support for the canister system.
4. Insert the new canister system.
# 9.0 Spare parts and consumables

<table>
<thead>
<tr>
<th>Consumables</th>
<th>REF</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reusable suction hose, Ø 10 mm</td>
<td>318.1012.0</td>
</tr>
<tr>
<td>Fingertip for reusable suction hose, Ø 10 mm, 10 St.</td>
<td>318.1100.0</td>
</tr>
<tr>
<td>Bacterial filter for ATMOS DDS secretion canister, pack of 10 pcs.</td>
<td>340.0054.0</td>
</tr>
<tr>
<td>Suction hose, disposable, Ø 6 mm, L = 1.30 m, 10 pcs.</td>
<td>006.0057.0</td>
</tr>
<tr>
<td>Suction hose, disposable, Ø 6 mm, L = 1.30 m, 50 pcs.</td>
<td>006.0059.0</td>
</tr>
<tr>
<td>Serres® suction bag 1 l, not autoclavable, 36 pcs.</td>
<td>312.0466.0</td>
</tr>
<tr>
<td>Medi-Vac® suction bag 1 l, not autoclavable, 50 pcs.</td>
<td>312.0474.0</td>
</tr>
<tr>
<td>Vacuum hose for single-use canister system</td>
<td>318.1211.0</td>
</tr>
<tr>
<td>Suction catheter Unomedical®, size: CH 12, L = 50 cm, 100 pcs.</td>
<td>000.0294.0</td>
</tr>
<tr>
<td>Suction catheter Unomedical®, size: CH 14, L = 50 cm, 100 pcs.</td>
<td>000.0295.0</td>
</tr>
<tr>
<td>Suction catheter Unomedical®, size: CH 16, L = 50 cm, 100 pcs.</td>
<td>000.0296.0</td>
</tr>
<tr>
<td>Hydrophobic bacterial and viral filter, Ø 8 mm</td>
<td>443.0738.0</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Spare parts</th>
<th>REF</th>
</tr>
</thead>
<tbody>
<tr>
<td>DDS canister system</td>
<td></td>
</tr>
<tr>
<td>DDS secretion canister 1 l</td>
<td>318.1013.0</td>
</tr>
<tr>
<td>DDS outer canister lid</td>
<td>318.1002.0</td>
</tr>
<tr>
<td>DDS inner canister lid</td>
<td>318.1004.0</td>
</tr>
<tr>
<td>Float ball</td>
<td>000.0839.0</td>
</tr>
<tr>
<td>Filter holder</td>
<td>318.1003.0</td>
</tr>
<tr>
<td>Sealing ring</td>
<td>055.0112.0</td>
</tr>
<tr>
<td>Device</td>
<td></td>
</tr>
<tr>
<td>Battery for ATMOS C 341 Battery</td>
<td>318.0001.0</td>
</tr>
<tr>
<td>Battery compartment cover</td>
<td>318.0012.0</td>
</tr>
<tr>
<td>Foam part for battery compartment cover</td>
<td>318.0018.0</td>
</tr>
<tr>
<td>Hose rewind</td>
<td>319.0004.0</td>
</tr>
<tr>
<td>Device base</td>
<td>319.0003.0</td>
</tr>
<tr>
<td>Support for DDS canister system</td>
<td>318.1010.0</td>
</tr>
<tr>
<td>Support for Serres® canister system</td>
<td>318.1210.0</td>
</tr>
<tr>
<td>Support for Medi-Vac® canister system</td>
<td>318.1500.0</td>
</tr>
<tr>
<td>Release button</td>
<td>318.0013.0</td>
</tr>
<tr>
<td>Spring for release button</td>
<td>000.1029.0</td>
</tr>
<tr>
<td>Power supply and recharging unit</td>
<td>318.0035.0</td>
</tr>
<tr>
<td>2 pin power cord</td>
<td>008.0920.0</td>
</tr>
</tbody>
</table>
10.0 Disposal

Packing

1. Please recycle the packing.

Secretion and blood

1. Please dispose of secretion, blood and contaminated parts in line with country-specific regulations.

In the Federal Republic of Germany the „Requirements on the implementation aid for disposal of waste from healthcare institutions“ are valid, a statement of the Federal / State Working Group on Waste.

Canister system

Single-use products may not be reprocessed and may not be reused! Please dispose of disposable products professionally.

The following notes are only applicable for reusable products.

1. Clean and disinfect the reusable products of the canister system.
2. Recycle the disinfected reusable products.

ATMOS C 341 Battery

Do not dispose of the device or the battery in domestic waste.

The ATMOS C 341 Battery does not contain any hazardous goods.

1. Clean and disinfect the device.
2. In Germany: Send in the device to ATMOS or your specialized dealer. They will dispose of the device professionally.
3. In other countries: Dispose of the device professionally and according to country-specific laws and regulations.

In Germany the device is excluded from the Electrical and Electronic Equipment Act (ElektroG) according to the National Register for waste electric equipment because it may be contaminated. Do not dispose of the device in the electrical waste.

Basically, the case is fully recyclable. However, please note the country-specific laws and regulations.
# 11.0 Technical data

## Device

<table>
<thead>
<tr>
<th>Dimensions (W x H x D):</th>
<th>370 x 277 x 146 mm</th>
</tr>
</thead>
<tbody>
<tr>
<td>with DDS canister system</td>
<td>370 x 277 x 146 mm</td>
</tr>
<tr>
<td>with Serres® canister system</td>
<td>370 x 277 x 136 mm</td>
</tr>
<tr>
<td>with Medi-Vac® canister system</td>
<td>370 x 277 x 136 mm</td>
</tr>
<tr>
<td>with universal bracket</td>
<td>370 x 277 x 136 mm</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Weight:</th>
<th>4.6 kg</th>
</tr>
</thead>
<tbody>
<tr>
<td>Device with battery /without canister system and support</td>
<td>1.00 kg</td>
</tr>
<tr>
<td>DDS canister system with support for DDS canister system</td>
<td>0.65 kg</td>
</tr>
<tr>
<td>Serres® canister system with support for Serres® canister system</td>
<td>0.295 kg</td>
</tr>
<tr>
<td>Medi-Vac® canister system with support for Medi-Vac® canister system</td>
<td>0.2 kg</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Operation:</th>
<th>-5° C up to +50° C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperature range</td>
<td>5% to 95% without condensation</td>
</tr>
<tr>
<td>Relative air humidity</td>
<td>540 hPa to 1100 hPa</td>
</tr>
<tr>
<td>Air pressure</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Transport / storage:</th>
<th>-40° C up to 70° C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperature range</td>
<td>5% to 95% without condensation</td>
</tr>
<tr>
<td>Relative air humidity</td>
<td>540 hPa to 1100 hPa</td>
</tr>
<tr>
<td>Air pressure</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Charging:</th>
<th>-5° C up to +50° C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperature range</td>
<td>5% to 95% without condensation</td>
</tr>
<tr>
<td>Relative air humidity</td>
<td>540 hPa to 1100 hPa</td>
</tr>
<tr>
<td>Air pressure</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Maximum operating altitude</th>
<th>5000 m (NN)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Contamination level</td>
<td>Class 1 (fully-sealed housing)</td>
</tr>
<tr>
<td>Overvoltage category</td>
<td>II</td>
</tr>
<tr>
<td>Maximum power consumption</td>
<td>45 W</td>
</tr>
<tr>
<td>Maximum current consumption</td>
<td>3.7 A</td>
</tr>
<tr>
<td>Mains voltage</td>
<td>12 V DC nominal (at least 10 V, max. 15 V) at the charging interface or via the power supply and recharging unit.</td>
</tr>
</tbody>
</table>

### Pump

Vacuum pump (diaphragm pump), 1 head

### Suction capacity at the device inlet (without canister system)

26 l/min ± 4 l/min

### Suction capacity at the inlet of the DDS canister system

23 l/min ± 3 l/min

### Maximum achievable vacuum

0.8 bar* +0.15 bar / -0.06 bar resp. 80 % of the air pressure
<table>
<thead>
<tr>
<th><strong>Vacuum adjustment</strong></th>
<th>Via infinitely variable vacuum controller: -0.1 bar up to -0.8 bar</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Vacuum display</strong></td>
<td>Manometer up to -1 bar at a maximum, accuracy class 2.5 (2.5 %)</td>
</tr>
<tr>
<td><strong>Display</strong></td>
<td>By means of LED on the control panel: On / off, display of battery status, warning (red LED)</td>
</tr>
<tr>
<td><strong>Power cycles (short-term operation)</strong></td>
<td>45 min On, 90 min Off</td>
</tr>
<tr>
<td><strong>Noise level</strong>: Mean sound pressure level in 1 m distance and at -0.8 bar</td>
<td>&lt; 60 dB (A)</td>
</tr>
<tr>
<td><strong>Classification according to EN 60601-1:</strong></td>
<td>Protection class II (during mains and battery operation)</td>
</tr>
<tr>
<td>Protection class against electric shock</td>
<td></td>
</tr>
<tr>
<td>Degree of protection against electric shock</td>
<td></td>
</tr>
<tr>
<td><strong>Degree of protection against</strong></td>
<td>Protection class II (during mains and battery operation)</td>
</tr>
<tr>
<td>Ingress of solid foreign objects</td>
<td></td>
</tr>
<tr>
<td>Penetration of dust</td>
<td></td>
</tr>
<tr>
<td>Ingress of water with harmful effects</td>
<td>IP 34D</td>
</tr>
<tr>
<td><strong>Period tests</strong></td>
<td>Recommended: Testing every 24 months.</td>
</tr>
<tr>
<td><strong>Suspension</strong></td>
<td>Compatible with ATMOS wall and device support</td>
</tr>
<tr>
<td><strong>Classification according to EN ISO 10079-1</strong></td>
<td>High vacuum / high flow</td>
</tr>
<tr>
<td><strong>Product class according to Directive 93/42/EEC</strong></td>
<td>Ila</td>
</tr>
<tr>
<td><strong>UMDNS code</strong></td>
<td>15-016 Suction device, emergency</td>
</tr>
<tr>
<td><strong>GMDN code</strong></td>
<td>36616, Suction unit, transport and emergency</td>
</tr>
</tbody>
</table>

*1 bar = 100 kPa

**Battery**

<table>
<thead>
<tr>
<th><strong>Type</strong></th>
<th>Lead-acid, Panasonic LC-R 123R4PG</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Dimensions (W x H x D)</strong></td>
<td>67 x 134 x 67 mm</td>
</tr>
<tr>
<td><strong>Weight</strong></td>
<td>1.15 kg</td>
</tr>
<tr>
<td><strong>Nominal capacity</strong></td>
<td>3.4 Ah</td>
</tr>
<tr>
<td><strong>Nominal voltage</strong></td>
<td>12 V nominal</td>
</tr>
<tr>
<td><strong>Charging time</strong></td>
<td>Battery status 80 % : 2 h 40 min Battery status 100 %: 14 h Automatic switch-over to trickle charging</td>
</tr>
<tr>
<td><strong>Recharging interval during long-term storage</strong></td>
<td>Every 3 months.</td>
</tr>
<tr>
<td><strong>Battery operating time during continuous operation with fully recharged battery / new battery (&gt;20 l/min, setting -0.8 bar)</strong></td>
<td>23 min at -5° C 40 min at +21° C 40 min at +50° C</td>
</tr>
<tr>
<td><strong>Life cycle</strong></td>
<td>400 recharging cycles within approx. 3 years</td>
</tr>
</tbody>
</table>
### Display

<table>
<thead>
<tr>
<th>Display</th>
<th>Display of battery status during operation and recharging</th>
</tr>
</thead>
</table>

### Typical battery operating life*

<table>
<thead>
<tr>
<th>Pressure</th>
<th>Operating Life</th>
</tr>
</thead>
<tbody>
<tr>
<td>-0.2 bar</td>
<td>40 min</td>
</tr>
<tr>
<td>-0.5 bar</td>
<td>40 min</td>
</tr>
<tr>
<td>-0.8 bar</td>
<td>40 min</td>
</tr>
</tbody>
</table>

* Measured at +21°C, continuous operation, without battery recharging and at free air flow.

### DDS canister system

<table>
<thead>
<tr>
<th>Capacity</th>
<th>1000 ml</th>
</tr>
</thead>
<tbody>
<tr>
<td>Connection reusable suction hose</td>
<td>Ø 10 mm I.D.</td>
</tr>
<tr>
<td>Reusable suction hose: Diameter</td>
<td>Ø 10 mm I.D.</td>
</tr>
<tr>
<td>Length</td>
<td>1300 mm</td>
</tr>
<tr>
<td>Connection to the suction device</td>
<td>Direct connection (without intermediate hose)</td>
</tr>
<tr>
<td>Bacterial filter</td>
<td>Hydrophobic bacterial filter cartridge for use in the secretion canister lid, disposable</td>
</tr>
<tr>
<td>Bacterial filtration efficiency</td>
<td>99.999778%</td>
</tr>
</tbody>
</table>

### Single-use canister systems

<table>
<thead>
<tr>
<th>Capacity</th>
<th>1000 ml</th>
</tr>
</thead>
<tbody>
<tr>
<td>Connection disposable suction hose</td>
<td>Ø 7 mm I.D.</td>
</tr>
<tr>
<td>Disposable suction hose: Diameter</td>
<td>Ø 6 mm I.D.</td>
</tr>
<tr>
<td>Length</td>
<td>1300 mm</td>
</tr>
<tr>
<td>Connection to the suction device</td>
<td>By means of a vacuum hose (intermediate hose)</td>
</tr>
<tr>
<td>Bacterial filter</td>
<td>Integrated in the suction bag</td>
</tr>
</tbody>
</table>

### Power supply and recharging unit

<table>
<thead>
<tr>
<th>Dimensions (W x H x D)</th>
<th>130 x 36 x 60 mm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weight</td>
<td>280 g</td>
</tr>
<tr>
<td>Operation: Temperature range</td>
<td>0°C up to +40°C</td>
</tr>
<tr>
<td>Relative air humidity</td>
<td>10% to 90% without condensation</td>
</tr>
<tr>
<td>Air pressure</td>
<td>700 hPa to 1100 hPa</td>
</tr>
<tr>
<td>Transport / storage: Temperature range</td>
<td>-40°C up to +70°C</td>
</tr>
<tr>
<td>Relative air humidity</td>
<td>10% to 95% without condensation</td>
</tr>
<tr>
<td>Air pressure</td>
<td>700 hPa to 1100 hPa</td>
</tr>
<tr>
<td>Electrical connection</td>
<td>100 V AC to 240 V AC, 50 Hz to 60 Hz</td>
</tr>
<tr>
<td>Maximum current consumption</td>
<td>1.1 A</td>
</tr>
<tr>
<td>Output nominal</td>
<td>13.8 V DC, 3.5 A</td>
</tr>
<tr>
<td>Classification according to EN 60601-1: Protection class against electric shock</td>
<td>Protection class II</td>
</tr>
<tr>
<td>Degree of protection against electric shock</td>
<td>Application part type CF</td>
</tr>
<tr>
<td>Degree of protection against:</td>
<td>IP 40</td>
</tr>
<tr>
<td>------------------------------------------------------------------</td>
<td>------------------</td>
</tr>
<tr>
<td>• Ingress of solid foreign objects</td>
<td></td>
</tr>
<tr>
<td>• Penetration of dust</td>
<td></td>
</tr>
<tr>
<td>• Ingress of water with harmful effects</td>
<td></td>
</tr>
<tr>
<td>Length of output line</td>
<td>1.8 m</td>
</tr>
<tr>
<td>Length of power supply cord</td>
<td>approx. 2 m</td>
</tr>
</tbody>
</table>
12.0 Notes on EMC

- Medical electrical equipment is subject to special precautions with regard to EMC and must be installed acc. to following EMC notes.
- Portable and mobile HF communication facilities can influence medical electrical equipment.
- The use of other accessories, other converters and cables than stated may lead to an increased emission or a reduced interference immunity of the equipment or system.

Guidelines and Manufacturer’s Declaration - Emissions

The ATMOS C 341 Battery is intended for use in the electromagnetic environment specified below. The customer or user of the ATMOS C 341 Battery should ensure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Emissions Test</th>
<th>Compliance</th>
<th>Electromagnetic Environment - Guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF Emissions acc. to CISPR 11</td>
<td>Group 1</td>
<td>The ATMOS C 341 Battery uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.</td>
</tr>
<tr>
<td>Harmonics IEC 61000-3-2</td>
<td>Class A</td>
<td>The ATMOS C 341 Battery is suitable for use in all establishments, including domestic, and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.</td>
</tr>
<tr>
<td>Flicker IEC 61000-3-3</td>
<td>Corresponds</td>
<td></td>
</tr>
</tbody>
</table>

Guidelines and Manufacturer’s Declaration - Immunity for ATMOS C 341 Battery

The ATMOS C 341 Battery is intended for use in the electromagnetic environment specified below. The customer or user of the ATMOS C 341 Battery should ensure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Immunity Test</th>
<th>IEC 60601- Test Level</th>
<th>Compliance Level</th>
<th>Electromagnetic Environment - Guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>ESD acc. to IEC 61000-4-2</td>
<td>± 6 kV Contact</td>
<td>± 6 kV Contact</td>
<td>Floors should be wood, concrete, or ceramics tile. If floors are synthetic, the relative humidity should be at least 30 %.</td>
</tr>
<tr>
<td></td>
<td>± 8 kV Air</td>
<td>± 8 kV Air</td>
<td></td>
</tr>
<tr>
<td>EFT IEC 61000-4-4</td>
<td>± 2 kV Mains</td>
<td>± 2 kV Mains</td>
<td>Mains power quality should be that of a typical commercial or hospital environment.</td>
</tr>
<tr>
<td></td>
<td>± 1 kV I/Os</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Surges IEC 61000-4-5</td>
<td>± 1 kV common-mode</td>
<td>± 1 kV common-mode</td>
<td>Mains power quality should be that of a typical commercial or hospital environment.</td>
</tr>
<tr>
<td></td>
<td>± 2 kV differential mode</td>
<td>± 2 kV differential mode</td>
<td></td>
</tr>
<tr>
<td>Voltage Dips / Dropout IEC 61000-4-11</td>
<td>&lt; 5 % U₁, (&gt; 95 % Dip of the UT for 0.5 Cycle)</td>
<td>&lt; 5 % U₁, (&gt; 95 % Dip of the UT for 0.5 Cycle)</td>
<td>Mains power quality should be that of a typical commercial or hospital environment. If the user of the ATMOS C 341 Battery demands continued function even in case of interruptions of the energy supply, it is recommended to supply the ATMOS C 341 Battery from an uninterruptible current supply or a battery.</td>
</tr>
<tr>
<td></td>
<td>40 % U₁, (60 % Dip of the UT for 5 Cycles)</td>
<td>40 % U₁, (60 % Dip of the UT for 5 Cycles)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>70 % U₁, (30 % Dip of the UT for 25 Cycles)</td>
<td>70 % U₁, (30 % Dip of the UT for 25 Cycles)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>&lt; 5 % U₁, (&gt; 95 % Dip of the UT for 5 s)</td>
<td>&lt; 5 % U₁, (&gt; 95 % Dip of the UT for 5 s)</td>
<td></td>
</tr>
<tr>
<td>Magnetic field at power frequency 50/60 Hz acc. to IEC 61000-4-8</td>
<td>3 A/m</td>
<td>Inapplicable</td>
<td>Power frequency magnetic fields should be that of a typical commercial or hospital environment.</td>
</tr>
</tbody>
</table>

NOTE  U₁ is the mains alternating current prior to application of the test levels.
The ATMOS C 341 Battery is intended for use in the electromagnetic environment specified below. The customer or user of the ATMOS C 341 Battery should ensure that it is used in such an environment.

### Immunity Test

<table>
<thead>
<tr>
<th>Immunity Test</th>
<th>IEC 60601-Test Level</th>
<th>Compliance Level</th>
<th>Electromagnetic Environment - Guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conducted RF IEC 61000-4-6</td>
<td>3 V &lt;sub&gt;eff&lt;/sub&gt; 150 kHz to 80 MHz</td>
<td>10 V</td>
<td>Portable and mobile communications equipment should be separated from the ATMOS C 341 Battery incl. the cables by no less than the distances calculated/listed below.</td>
</tr>
</tbody>
</table>
| Radiated RF IEC 61000-4-3 | 3 V/m 80 MHz to 2.5 GHz | 10 V/m | Recommended distances:  
  - d = 0,35 \sqrt{P}  
  - d = 0,70 \sqrt{P}  
  - 80 MHz to 800 MHz  
  - 800 MHz to 2.5 GHz  
  where "P" is the max. power in watts (W) and D is the recommended separation distance in meters (m). |

Field strengths from fixed transmitters, as determined by an electromagnetic site (a) survey, should be less than the compliance level (b). 

Interference may occur in the vicinity of equipment containing following symbol:

---

**NOTE 1**  By 80 MHz and 800 MHz the higher frequency range applies.  
**NOTE 2**  These guidelines might not be applicable in all cases. The emanation of electromagnetic waves is affected by absorption and reflection of buildings, objects and people. 

- **a** The field strength of stationary transmitters, such as base stations of cellular phones and mobile terrain radio equipment, amateur radio transmitters, cbm broadcast and TV stations cannot be predestined exactly.  
  To determine the electromagnetic environment in regard to stationary transmitters, a study of the location is to be considered. If the measured field strength at the location where the ATMOS C 341 Battery is used exceeds the above compliance level, the ATMOS C 341 Battery is to be observed to verify the intended use. If abnormal performance characteristics are noted, additional measures might be necessary, e. g. a changed arrangement or another location for the device.  

- **b** Over the frequency range of 150 kHz to 80 MHz, field strengths should be lower than 3 V/m.  

The device may not be used directly next to other devices or piled up with other devices. If operation next to or piled with other devices is necessary, please watch the device to check its intended operation in this arrangement.
Recommended separations between portable and mobile RF Communications equipment and the ATMOS C 341 Battery

The ATMOS C 341 Battery is intended for use in electromagnetic environment in which radiated disturbances are controlled. The customer or user of the ATMOS C 341 Battery can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF Communications equipment and the ATMOS C 341 Battery as recommended below, according to the maximum output power of the communications equipment.

<table>
<thead>
<tr>
<th>Nominal output of the transmitter W</th>
<th>Safety distance, depending on transmit-frequency m</th>
</tr>
</thead>
<tbody>
<tr>
<td>150 kHz to 80 MHz</td>
<td>80 MHz to 800 MHz</td>
</tr>
<tr>
<td>d = [0.35] √ P</td>
<td>d = [0.35] √ P</td>
</tr>
<tr>
<td>0.01</td>
<td>0.035</td>
</tr>
<tr>
<td>0.1</td>
<td>0.11</td>
</tr>
<tr>
<td>1</td>
<td>0.35</td>
</tr>
<tr>
<td>10</td>
<td>1.1</td>
</tr>
<tr>
<td>100</td>
<td>3.5</td>
</tr>
</tbody>
</table>

For transmitters for which the maximum nominal output is not indicated in the above table, the recommended safety distance d in meters (m) can be determined using the equation belonging to the respective column whereas P is the maximum nominal output of the transmitter in watts (W) acc. to manufacturer’s specification.

NOTE 1  By 80 MHz and 800 MHz the higher frequency range applies.

NOTE 2  These guidelines might not be applicable in all cases. The emanation of electromagnetic waves is affected by absorption and reflection of buildings, objects and people.